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## Polymeric bis(triphenylphosphine)iminium chloride as a recyclable catalyst†

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Metal-free catalysts have garnered considerable interest as an environmental and economical alternative to precious metal catalysts. Bis(triphenylphosphine)iminium chloride (PPNCI) has emerged as a prominent choice due to its air and thermal stability and broad reactivity, especially in applications where a bulky cation is needed. The high phosphorus content and synthetic effort required for catalyst synthesis increase environmental impact; the recyclability of PPNCI in catalytic processes remains largely unexplored. The potential development of a polymer-supported PPNCI catalysts therefore desirable to enable this recyclability. In this work, we synthesise polymeric PPNCI (poly(PPNCI)) for the first time. Poly(PPNCI) demonstrates a comparative catalytic reactivity to its small molecule variant when employed as a catalyst in halogen-exchange reactions and CO<sub>2</sub>/epoxide coupling. For the latter the effect of catalyst loading, CO<sub>2</sub> pressure, reaction time and addition of co-catalyst on conversion and selectivity was investigated. Poly(PPNCI) was easily recovered from the crude product by simple precipitation and its catalytic reactivity was well-maintained over three reaction cycles, providing environmental and economic advantages for sustainable reaction development.

## Introduction

Catalysts, often containing transition metals, are essential tools in chemical synthesis.<sup>1-3</sup> Challenges in metal extraction, complex toxicity, and economic cost frustrate the long-term sustainability of their industrial application.<sup>4,5</sup> Consequently, metal-free catalysts have been touted as environmentally friendly, economically viable, and less hazardous alternatives to metal-based catalysts.<sup>6-9</sup> An emerging class of metal-free catalysts are bis(triphenylphosphine)iminium salts,<sup>10-12</sup> thanks to their good solubility, stability at high temperatures and ease of handling.<sup>11,13</sup>

Bis(triphenylphosphine)iminium chloride (PPNCl) has been utilized in carbon dioxide (CO<sub>2</sub>) valorisation<sup>14-18</sup> to affect the coupling of CO<sub>2</sub> and epoxides to form cyclic carbonates as key reagents for the pharmaceutical and fine chemical industry.<sup>19-21</sup> Wing *et al.* demonstrated that both the PPN<sup>+</sup> and Cl<sup>-</sup> play important roles in the reaction mechanism.<sup>22</sup> PPNCl has also been explored as a co-catalyst in the ring-open co-polymerisation of CO<sub>2</sub> and epoxides to form polycarbonates.<sup>23,24</sup> Showing catalyst versatility, PPNCl can also be utilized to facilitate the hydrogenation of CO<sub>2</sub> to yield higher alcohols,<sup>25</sup> and in difficult halogen-exchange reactions to yield

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fluorinated aromatic compounds for pharmaceutical and agrochemical applications (Fig. 1a).<sup>11</sup>

The material-intensive, multi-step synthesis of PPNCl means that catalyst recovery would be important, however, the recycling of PPNCl has thus far not been explored. In recent years,

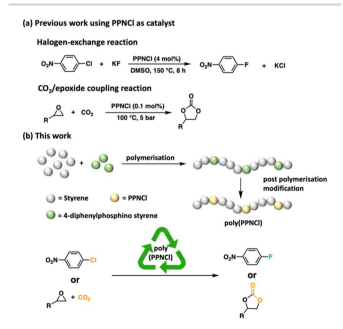


Fig. 1 (a) Previous work using PPNCl as catalysts; (b) poly(PPNCl) as a recyclable catalyst for halogen exchange and  $CO_2$ /epoxide coupling reactions.

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the recyclability of catalysts through incorporation into polymers has gained significant attention. <sup>26-29</sup> Previous work in our group showed that polymer-supported catalysts featuring maingroup elements including phosphorus, can be easily recovered and achieve comparable catalytic efficiencies to their small molecule counterparts. <sup>30-32</sup>

Developing a polymeric variant of PPNCl could generate a catalyst capable of maintaining high activity and new recyclability. In this work, we report for the first time, the preparation of macromolecular **poly(PPNCl)**. We demonstrate the catalytic reactivity of **poly(PPNCl)** in catalyzing both halogen-exchange and CO<sub>2</sub>/epoxide coupling reactions. The catalyst is readily recovered through post-reaction precipitation and serves as an important step in non-metal catalyst recycling (Fig. 1b).

#### Results and discussion

#### Synthesis of poly(PPNCl)

Macromolecular **poly(PPNCI)** was prepared through postpolymerisation modification of poly(styrene-*co*-diphenylphosphine styrene), **P1**, as shown in Scheme 1. **P1** was prepared by anionic co-polymerisation of styrene and *p*-(diphenylphosphino) styrene using *n*-butyllithium as the initiator in a methodology modified from our previous work on polymeric frustrated Lewis pairs.<sup>30</sup> The styrene co-monomer serves to provide inert spacers in the polymer chains, separating the active catalyst moieties, and improving solubility. The phosphorus groups incorporated into the polymer chains served as anchors for a modified small molecule synthesis.<sup>11</sup> As PPNCl is prepared by the reaction of

Scheme 1 (a) Synthetic procedures of P2 from phosphine-containing copolymer, m:n=95:5 or 90:10; (i) n-butyllithium, toluene, -78 °C, 3 h; (ii)  $C_2Cl_6$ , THF, r.t., 3 h; (b) Synthesis of Ph<sub>3</sub>PNLi from triphenyl-phosphine; (iii) hydroxylamine-O-sulfonic acid, methanol/DCM, r.t., 1 h; (iv) n-butyllithium, THF, initiated at -50 °C, -15 °C 2 h; (c) synthesis of poly(PPNCl) (v) THF, r.t., 3 h.

a dichlorinated phosphine (Ph<sub>3</sub>PCl<sub>2</sub>), accessible in near quantitative conversion, and a lithiated amino(triphenylphosphine) (Ph<sub>3</sub>-PNLi) that forms unavoidable side-products, it is essential to dichlorinate the polymer first. The reaction of P1 with C2Cl6 generates P2 with full conversion and high purity, as shown by NMR spectroscopy (Scheme 1a and Fig. S1†). A broad resonance at 63.2 ppm in the <sup>31</sup>P NMR spectrum is comparable to the chemical shift for Ph<sub>3</sub>PCl<sub>2</sub> (65.0 ppm).<sup>33</sup> Separately, amination of triphenylphosphine afforded amino(triphenyl)phosphonium chloride which was subsequently lithiated to afford lithium (triphenylphosphoranylidene)azanide (Scheme 1b). Finally, the targeted poly(PPNCl) was synthezised by the reaction between P2 and lithium (triphenylphosphoranylidene)azanide (Scheme 1c). The obtained poly(PPNCl) was purified by dialysis to remove small molecule impurities. Analysis by <sup>31</sup>P NMR spectroscopy (Fig. S1†) showed a single resonance at 20.7 ppm, confirming the full conversion of P2 to the desired product. Diffusion order spectroscopy (DOSY) showed a single diffusion coefficient, confirming that styrene and PPNCl moieties were distributed within the same polymer chains (Fig. S3†). This synthetic protocol was modified to prepare catalysts with 5 and 10 mol% PPNCl moieties (polv(PPNCI)-5 and poly(PPNCI)-10 respectively; Table S1†). For further characterisation of poly(PPNCI), thermogravimetry analysis (TGA) and differential scanning calorimetry (DSC) showed that polv(PPNCI)-5 has a lower degradation temperature and glass transition temperature compared to poly(PPNCI)-10 due its lower PPNCl loading in the polymer chains (Fig. S4†). Importantly, the evident absorptions of Fourier-transform infrared spectroscopy (FTIR) in the region between 1100 and 1480 cm<sup>-1</sup> correspond to the vibration of P=N bonds according to the literature,34 further supporting the formation of poly(PPNCI).

#### Halogen-exchange reaction

Halogen-exchange reactions were performed with these novel catalyst systems, **poly(PPNCI)-5** and **10**. The conversion of 1-chloro-4-nitrobenzene (**CNB**) to 1-fluoro-4-nitrobenzene (**FNB**) by **poly(PPNCI)-10** (63%) was surprisingly lower than that of **poly(PPNCI)-5** (80%) (Scheme 2a, Table S2†). While the higher loading offered a higher catalyst concentration, the increased number of PPNCl groups led to poorer solubility of the catalyst, therefore lowering conversion. Similar phenomena have been observed in the literature.<sup>35</sup> The conversion catalysed by **poly(PPNCI)-5** is comparable to that facilitated by PPNCl (81%), suggesting similar efficiency for small and macromolecular systems. With its superior solubility and performance,

(a) 
$$O_2N$$
 CI + KF  $\frac{\text{poly(PPNCI)}}{\text{DMSO, 150°C, 8 h}} O_2N$  F + KCI

(b)  $CI$   $\frac{O}{100°C, 1 \text{ bar, 24 h}} O_2N$ 

Scheme 2 (a) Halogen-exchange reaction from CNB to FNB; (b) coupling reaction of  ${\rm CO_2}$  with GC.

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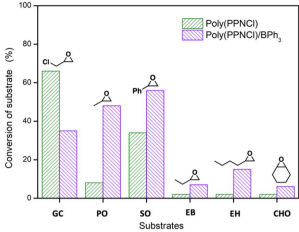


Fig. 2 CO<sub>2</sub>/epoxides coupling reaction to yield cyclic carbonates. Conversions calculated by <sup>1</sup>H NMR spectroscopy. Conversions for GC and PO were taken from the average value of three trials; the rest substrates were conducted once. Reactions were performed in a sealed ampoule on a 12.8 mmol scale at 0.1 mol% poly(PPNCI) with or without 0.1 mol% BPh<sub>3</sub> (0.1 mol% as calculated from the number of expected active functional groups).

poly(PPNCl)-5 was chosen as the preferred catalyst for the rest of this work.

Conversion of CNB with poly(PPNCl)-5 was systematically optimized (Table S2†). Minor increases in conversion by increasing the molar ratio of potassium fluoride (KF) added and when lower concentrations of catalyst were observed (2 mol%, 89%), further improving reaction sustainability. While the reaction worked in DMSO, MeOH and DMF, the optimal solvent was DMSO which correlated with enhanced catalyst solubility. CNB concentration is also important, with both lower (0.5, 1 M) and higher (3, 4 M) concentrations giving decreased conversions. High concentrations of CNB impact catalyst solubility in DMSO, reducing resultant reactivity.

With an optimized system in hand, we investigated whether this substrate scope could be expanded using poly(PPNCI) as the catalyst. Electrophilic aromatic substitution is favoured by the inductive and resonance effects promoted by electronwithdrawing groups (EWGs); previously this reaction was only reported for nitro-substituted aromatics. Displacing the nitro group of CNB with other para-substituted EWGs or metasubstituted electron-donating groups (EDGs) (Fig. S6†) was challenging. Of the substrates tested, only 4-chlorobenzonitrile showed conversion by <sup>1</sup>H and <sup>31</sup>P NMR spectroscopy (Fig. S7†).

#### CO2 and epoxide coupling reactions

The coupling of CO2 and epoxides is an important reaction for CO<sub>2</sub> transformation, as the resultant cyclic carbonates are valuable intermediates for fuel additives,36 polycarbonates,37 and pharmaceuticals.38 In previous reports, the PPNCl catalyst was usually paired with a Lewis acid co-catalyst such as triphenylborane (BPh<sub>3</sub>) to enhance reactivity.<sup>39,40</sup> The catalytic reactivity of poly(PPNCI) for the insertion of CO<sub>2</sub> to epoxides in the absence of a solvent was investigated (Scheme 2b). Glycidyl chloride (GC) was selected as a model substrate, and the results

summarized in Table S3† under standard conditions of 1 bar CO<sub>2</sub> for 24 h with 0.1 mol% poly(PPNCl)-5.

In the absence of poly(PPNCI), no conversion to chloropropene carbonate was observed. With 0.1 mol% (by catalyst moiety) poly(PPNCl) without any borane co-catalyst, 66% conversion was achieved, increasing to 98% at 0.4 mol%. Conversions did not substantially change upon alteration of CO<sub>2</sub> pressure (1-5 bar) (Table S3,† entries 1 and 5-7), suggesting CO<sub>2</sub> is not involved in the rate-determining step. While the rate of epoxide activation is often enhanced through the addition of BPh<sub>3</sub>, the conversion of GC surprisingly decreased to 48% when the Lewis acid is added to the system (Table S3,† entry 9).30 No reaction was observed when employing BPh3 alone. We hypothesize that the coordination of BPh3 to a key intermediate (I in Fig. S9†) would slow the rate of CO<sub>2</sub> insertion and therefore decrease reaction rates. As the reaction works without the addition of an air-sensitive Lewis acid, we explored milder conditions under an air atmosphere (Table S3,† entry 16). However, GC conversion remained unchanged compared to inert atmosphere (Table S3,† entry 2).

The kinetics of the GC/CO2 coupling reaction catalysed by poly(PPNCI) was explored (Table S3,† entries 10-15). Conversion of GC increased from 3 h to 48 h, indicating a long-lived catalyst (Fig. S11a†) and first-order kinetics for [GC] (Fig. S11b†). A catalytic mechanism for the GC/poly(PPNCl) system is proposed (Scheme S1†). Switching the substrate to propylene oxide (PO) showed remarkably different results: PO conversion was very low (9%) when using poly(PPNCI)-5 while the introduction of a BPh3 co-catalyst increased conversion to 48% (Table S4†). This reversion of the trends observed for PO suggests a significant change in the mechanism proposed in Scheme S2.† Increased reaction time resulted in a higher conversion (61%) suggesting catalyst death is not the reason for reduced yields. The effect of CO2 pressure on PO conversion gave no significant change in conversion (1-5 bar). PO conversion was enhanced to 71% by increasing catalyst loading to

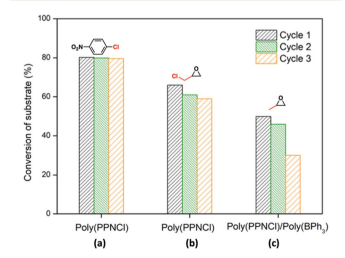


Fig. 3 Reaction conversions over three cycles (a) halogen-exchange, (b) CO<sub>2</sub>/GC coupling, and (c) CO<sub>2</sub>/PO coupling reactions. Conversion calculated by <sup>1</sup>H NMR spectroscopy. Conversions for recycled poly(PPNCI) were obtained from an average value of two trials.

Scheme 3 The selective one-pot, two-step reaction of epoxide conversion to cyclic carbonates. General conditions: the feed ratio of GC: PO: poly(PPNCI):  $BPh_3$  is 1000:1000:4:4.

0.2 mol%. The efficacy of the borane system also allowed us to test the poly(Lewis acid) variant previously developed in our group.<sup>30</sup> Poly(BPh<sub>3</sub>), a co-polymer of styrene and 4-styryl-diphenylborane, gave conversations comparable to poly(PPNCl)/BPh<sub>3</sub> system, demonstrating that both PPNCl and BPh<sub>3</sub> can be switched to recyclable polymeric systems while maintaining catalytic reactivity (Table S4,† entry 10).

Further expanding the substrate scope delineated two clear classes, evaluated by the overall catalytic reactivity of poly(PPNCI) versus poly(PPNCl)/BPh3 under conditions that avoided reaching full conversion in order to probe structure/activity relationships (Fig. 2). Only the most reactive substrate, GC, favoured poly(PPNCI) as a lone catalyst and outperformed all other tested epoxides (66%). Styrene oxide achieved the second highest conversion (34%) using only **poly(PPNCI)**, supporting the theory that the electron-withdrawing nature of the substituents is key to ensuring reactivity. Interestingly, whilst the poly(PPNCl)/BPh3 binary system resulted in a lower conversion of GC than for poly(PPNCI) alone, this trend was reversed for all other substrates screened. As the most reactive substrate, GC can be activated by both poly(PPN<sup>+</sup>) and BPh<sub>3</sub>, the two catalytic mechanisms may compete with each other to stall catalytic turnover. The sterically hindered cyclohexene oxide had the lowest conversion, attributed to its high steric hindrance preventing associative ring-opening.

Importantly, the selectivity of cyclic carbonates is 100% in nearly every case. Under no reaction conditions is polycarbonate and polyether generated in the presence of **poly(PPNCI)** alone. The good leaving group abilities of both poly(PPN<sup>†</sup>) and Cl<sup>-</sup> in the ring-closure step to form chloropropene carbonate likely facilitate this selectivity. For the GC/poly(PPNCI)/BPh<sub>3</sub> system, a secondary product was observed (<sup>1</sup>H NMR: 4.7 ppm, Fig. S17 and S18†), identified as the carbonate anion adduct of BPh<sub>3</sub> following ring opening by Cl<sup>-</sup>. This suggests that chlorine may attack the more substituted carbon to form an off-cycle product.

Finally, the catalytic selectivity and kinetic switch can be used in one-pot stepwise catalytic transformations. **GC** and **PO** could be sequentially converted into their cyclic carbonates as shown in Scheme 3. The epoxide mixture was first treated with **poly(PPNCI)** to yield chloropropene carbonate and propylene carbonate conversions of 97% and 2% respectively. The addition of BPh<sub>3</sub> to the reaction led to the conversion of the unreacted **PO** (74%), evidencing this epoxide selectivity.

#### Catalyst recycling

Polymer-supported catalysts are important additions to sustainable catalyst development as their ready isolation allows for recyclability.<sup>31</sup> We sought to explore the catalytic performance of both **poly(PPNCI)** and **poly(PPNCI)**/poly(BPh $_3$ ) after repeated recovery of the polymer-supported catalysts from crude products (Fig. 3). **Poly(PPNCI)** can be recovered easily from the product by precipitation from n-hexane and dried before reuse in subsequent reactions.

In the halogen-exchange reaction, the addition of *n*-hexane precipitated poly(PPNCI)-5 which was recovered from the crude product by filtration and dried under vacuum prior to weighing between each cycle. Catalytic performance was well-maintained over three reuse cycles (1st cycle: 80%; 2nd: 79%; 3rd: 77%), demonstrating excellent recyclability. For CO2/epoxide coupling, precipitation of the catalyst or catalysts from the crude chloropropene carbonate (poly(PPNCl)-5) and propylene carbonate (poly(PPNCl)-5/poly(BPh3)) reactions was conducted in the reaction vessel. The catalytic reactivity of poly(PPNCl) drops slightly over three reaction cycles of GC/CO2 coupling (1st cycle: 67%; 2nd: 61%; 3rd: 59%). A more precipitous decrease in substrate conversion is observed in the poly(PPNCI)/poly(BPh3) binary catalyst system (1st cycle: 50%; 2nd: 46%; 3rd: 30%) likely due to the mass loss and partial deactivation of airsensitive poly(BPh3) during catalyst recovery.

## Conclusions

In summary, we report the synthesis of **poly(PPNCI)** using post-polymerisation modification. **Poly(PPNCI)** can achieve high conversion at a low feeding ratio in the halogen-exchange reaction of chloroarenes to fluoroarenes, matching catalytic efficiency to the unrecoverable small molecule analogue, PPNCI. In CO<sub>2</sub>/epoxides coupling, the selective formation of cyclic carbonates is catalysed by two mechanisms. Depending on epoxide reactivity, either air-stable **poly(PPNCI)** can be used on its own or with the addition of BPh<sub>3</sub> or poly(BPh<sub>3</sub>) as a co-catalyst. The robust **poly(PPNCI)** can be recovered by simple precipitation and reused in further reactions, with both halogen-exchange and CO<sub>2</sub> coupling of reactive epoxides displaying excellent recyclability. The recovery and reuse of this key metal-free catalyst is an essential step to support sustainable chemical syntheses.

## Data availability

The data supporting this article have been included as part of the ESI.†

#### Author contributions

Ziwei Xu and Meng Wang: investigation, methodology and manuscript writing. Michael P. Shaver: conceptualisation, manuscript writing – review and editing, supervision and funding acquisition.

## Conflicts of interest

There are no conflicts to declare.

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## Notes and references

- 1 J. Magano and J. R. Dunetz, Large-scale applications of transition metal-catalyzed couplings for the synthesis of pharmaceuticals, *Chem. Rev.*, 2011, **111**, 2177–2250.
- 2 N. Saini, A. Malik and S. L. Jain, Light driven chemical fixation and conversion of CO<sub>2</sub> into cyclic carbonates using transition metals: A review on recent advancements, *Coord. Chem. Rev.*, 2024, **502**, 215636.
- 3 X. Wang, Z. Li, Y. Qu, T. Yuan, W. Wang, Y. Wu and Y. Li, Review of metal catalysts for oxygen reduction reaction: from nanoscale engineering to atomic design, *Chem*, 2019, 5, 1486–1511.
- 4 H. Esteves, N. E. Morais Costa, V. K. Tomazett, J. L. S. Milani, R. P. das Chagas and Â. de Fátima, Organocatalysis for the Chemical Fixation of Carbon Dioxide to Synthesise N-Heterocycles, *Curr. Green Chem.*, 2024, **11**, 87–126.
- 5 D. Yang and B. C. Gates, Catalysis by metal organic frameworks: perspective and suggestions for future research, *ACS Catal.*, 2019, **9**, 1779–1798.
- 6 Y. Zhang, Q. Wang, Q. Chen, X. Li, Y. Li, M. Kang, Q. Li and J. Wang, A new boron modified carbon nitride metal-free catalyst for the cycloaddition of CO2 and bisepoxides, *Appl. Catal.*, A, 2024, 119615.
- 7 R. Morodo, D. M. Dumas, J. Zhang, K. H. Lui, P. J. Hurst, R. Bosio, L. M. Campos, N. H. Park, R. M. Waymouth and J. L. Hedrick, Ring-Opening Polymerization of Cyclic Esters and Carbonates with (Thio) urea/Cyclopropenimine Organocatalytic Systems, ACS Macro Lett., 2024, 13, 181–188.
- 8 E. Poursaitidis, P. L. Gkizis, I. Triantafillidi and C. G. Kokotos, Organocatalytic activation of hydrogen peroxide: towards green and sustainable oxidations, *Chem. Sci.*, 2024, **15**, 1177–1203.
- 9 K. A. Ahrendt, C. J. Borths and D. W. MacMillan, New strategies for organic catalysis: the first highly enantioselective organocatalytic Diels—Alder reaction, *J. Am. Chem. Soc.*, 2000, **122**, 4243–4244.
- 10 T. W. Rave, Preparation and properties of bis(phosphoranylidene) ammonium chlorides, *J. Org. Chem.*, 1967, 32, 3461–3466.

- 11 M. A. Lacour, M. Zablocka, C. Duhayon, J. P. Majoral and M. Taillefer, Efficient Phosphorus Catalysts for the Halogen-Exchange (Halex) Reaction, *Adv. Synth. Catal.*, 2008, **350**, 2677–2682.
- 12 A. Martinsen, J. Songstad, R. Larsson, M. Pouchard, P. Hagenmuller and A. Andresen, Preparation and properties of some bis(triphenylphosphine) iminium salts, [(Ph3P) 2N] X, *Acta Chem. Scand.*, 1977, 31, 645–650.
- 13 J. Ruff and W. J. SCHLIENTZ, Inorg. Synth., 1974, 16, 84.
- 14 M. Scharfenberg, J. Hilf and H. Frey, Functional polycarbonates from carbon dioxide and tailored epoxide monomers: degradable materials and their application potential, Adv. Funct. Mater., 2018, 28, 1704302.
- 15 Y. Xu, L. Lin, M. Xiao, S. Wang, A. T. Smith, L. Sun and Y. Meng, Synthesis and properties of CO<sub>2</sub>-based plastics: Environmentally-friendly, energy-saving and biomedical polymeric materials, *Prog. Polym. Sci.*, 2018, **80**, 163–182.
- 16 H. Zhang and M. W. Grinstaff, Synthesis of atactic and isotactic poly(1, 2-glycerol carbonate) s: degradable polymers for biomedical and pharmaceutical applications, *J. Am. Chem. Soc.*, 2013, 135, 6806–6809.
- 17 L.-Y. Wang, G.-G. Gu, T.-J. Yue, W.-M. Ren and X.-B. Lu, Semiaromatic Poly(thioester) from the Copolymerization of Phthalic Thioanhydride and Epoxide: Synthesis, Structure, and Properties, *Macromolecules*, 2019, 52, 2439–2445.
- 18 J. Huang, Y. Xu, M. Wang and Z. Duan, Copolymerization of propylene oxide and CO<sub>2</sub> catalyzed by dinuclear salcy–CoCl complex, *J. Macromol. Sci.*, *Part A*, 2020, 57, 131–138.
- 19 A. Buonerba, A. De Nisi, A. Grassi, S. Milione, C. Capacchione, S. Vagin and B. Rieger, Novel iron(III) catalyst for the efficient and selective coupling of carbon dioxide and epoxides to form cyclic carbonates, *Catal. Sci. Technol.*, 2015, 5, 118–123.
- 20 A. Sibaouih, P. Ryan, M. Leskelä, B. Rieger and T. Repo, Facile synthesis of cyclic carbonates from CO<sub>2</sub> and epoxides with cobalt(π)/onium salt based catalysts, *Appl. Catal.*, A, 2009, 365, 194–198.
- 21 F. Chen, S. Tao, N. Liu, C. Guo and B. Dai, Hemilabile N-heterocyclic carbene and nitrogen ligands on Fe(II) catalyst for utilization of CO<sub>2</sub> into cyclic carbonate, *Appl. Organomet. Chem.*, 2021, 35, e6099.
- 22 W. N. Sit, S. M. Ng, K. Y. Kwong and C. P. Lau, Coupling reactions of CO<sub>2</sub> with neat epoxides catalyzed by PPN salts to yield cyclic carbonates, *J. Org. Chem.*, 2005, **70**, 8583–8586.
- 23 J. Zhang, L. Wang, S. Liu, X. Kang and Z. Li, A Lewis pair as organocatalyst for one-pot synthesis of block copolymers from a mixture of epoxide, anhydride, and CO<sub>2</sub>, *Macromolecules*, 2021, 54, 763–772.
- 24 C. E. Anderson, S. I. Vagin, W. Xia, H. Jin and B. Rieger, Cobaltoporphyrin-catalyzed CO<sub>2</sub>/epoxide copolymerization: selectivity control by molecular design, *Macromolecules*, 2012, 45, 6840–6849.
- 25 M. Cui, Q. Qian, Z. He, Z. Zhang, J. Ma, T. Wu, G. Yang and B. Han, Bromide promoted hydrogenation of CO<sub>2</sub> to higher alcohols using Ru–Co homogeneous catalyst, *Chem. Sci.*, 2016, 7, 5200–5205.

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- 27 Y. Chauvin, D. Commereuc and F. Dawans, Polymer supported catalysts, *Prog. Polym. Sci.*, 1977, 5, 95–226.
- 28 B. Chen and F. Jäkle, Boron-Nitrogen Lewis Pairs in the Assembly of Supramolecular Macrocycles, Molecular Cages, Polymers, and 3D Materials, *Angew. Chem.*, 2024, 136, e202313379.
- 29 U. Yolsal, T. A. Horton, M. Wang and M. P. Shaver, Cyclic ether triggers for polymeric frustrated Lewis pair gels, *J. Am. Chem. Soc.*, 2021, 143, 12980–12984.
- 30 M. Wang, F. Nudelman, R. R. Matthes and M. P. Shaver, Frustrated Lewis pair polymers as responsive self-healing gels, *J. Am. Chem. Soc.*, 2017, **139**, 14232–14236.
- 31 T. A. Horton, M. Wang and M. P. Shaver, Polymeric frustrated Lewis pairs in CO<sub>2</sub>/cyclic ether coupling catalysis, *Chem. Sci.*, 2022, **13**, 3845–3850.
- 32 M. Wang, M. Shanmugam, E. J. McInnes and M. P. Shaver, Light-Induced Polymeric Frustrated Radical Pairs as Building Blocks for Materials and Photocatalysts, *J. Am. Chem. Soc.*, 2023, **145**, 24294–24301.
- 33 Q. Yin, Y. Ye, G. Tang and Y.-F. Zhao, Studies on the structure behavior of triphenyldichlorophosphorane in different solvents, *Spectrochim. Acta, Part A*, 2006, **63**, 192–195.

- 34 C. S. Davis and R. J. Richardson, Organophosphorus compounds, *Experimental and clinical neurotoxicology*, 1980, vol. 1.
- 35 S. L. Regen, Triphase catalysis. Kinetics of cyanide displacement on 1-bromooctane, *J. Am. Chem. Soc.*, 1976, **98**, 6270–6274.
- 36 T. Chang, H. Jing, L. Jin and W. Qiu, Quaternary onium tribromide catalyzed cyclic carbonate synthesis from carbon dioxide and epoxides, *J. Mol. Catal. A: Chem.*, 2007, **264**, 241–247.
- 37 J. Sun, S.-i. Fujita and M. Arai, Development in the green synthesis of cyclic carbonate from carbon dioxide using ionic liquids, *J. Organomet. Chem.*, 2005, 690, 3490–3497.
- 38 X.-B. Lu, Y.-J. Zhang, B. Liang, X. Li and H. Wang, Chemical fixation of carbon dioxide to cyclic carbonates under extremely mild conditions with highly active bifunctional catalysts, *J. Mol. Catal. A: Chem.*, 2004, **210**, 31–34.
- 39 K. A. Andrea and F. M. Kerton, Triarylborane-catalyzed formation of cyclic organic carbonates and polycarbonates, *ACS Catal.*, 2019, **9**, 1799–1809.
- 40 K. A. Andrea and F. M. Kerton, Functionalized polycarbonates via triphenylborane catalyzed polymerization-hydrosilylation, *RSC Adv.*, 2019, 9, 26542–26546.