

Cite this: *Chem. Sci.*, 2019, **10**, 4618 All publication charges for this article have been paid for by the Royal Society of ChemistryReceived 23rd January 2019
Accepted 6th March 2019

DOI: 10.1039/c9sc00385a

rsc.li/chemical-science

Introduction

The alternating copolymerization of carbon dioxide and epoxides yields aliphatic polycarbonates ($\text{CO}_2\text{-PC}$) which contain 30–50 wt% CO_2 .^{1,2} It is a useful means to add-value to a common industrial waste and to reduce greenhouse gas emissions from polymer manufacture.^{3,4} Low molar mass $\text{CO}_2\text{-PC}$ products have properties which allow them to replace petrochemical polyols in the production of polyurethanes.^{5,6} High molar mass products show promising properties for use as elastomers, rigid plastics, scratch resistant coatings and even anti-microbial materials.^{7–9} Future product enhancements and the widespread uptake of these materials still depend upon the development of more highly active and selective catalysts.^{10–15} Heterogeneous catalysts can be highly active but yield much lower CO_2 content in the resulting polymers.¹ On the other hand, homogeneous catalysts can produce perfectly alternating copolymers but remain restricted to a relatively narrow ligand/metal scope.^{10–13} Amongst the most active homogeneous catalysts are $\text{Cr}(\text{III})$, $\text{Co}(\text{III})$ or $\text{Zn}(\text{II})$ complexes coordinated by ligands such as salens,^{13,16–19} phenoxy-amines,^{20–22} and β -diimmines.^{23–28}

^aChemistry Research Laboratory, University of Oxford, Mansfield Road, Oxford, OX1 3TA, UK. E-mail: charlotte.williams@chem.ox.ac.uk

^bSchool of Chemistry, University of Edinburgh, EH9 3FJ, UK

† Electronic supplementary information (ESI) available: All experimental protocols, data for all table entries, spectroscopy and kinetic data. See DOI: 10.1039/c9sc00385a

Heterodinuclear zinc and magnesium catalysts for epoxide/ CO_2 ring opening copolymerizations†

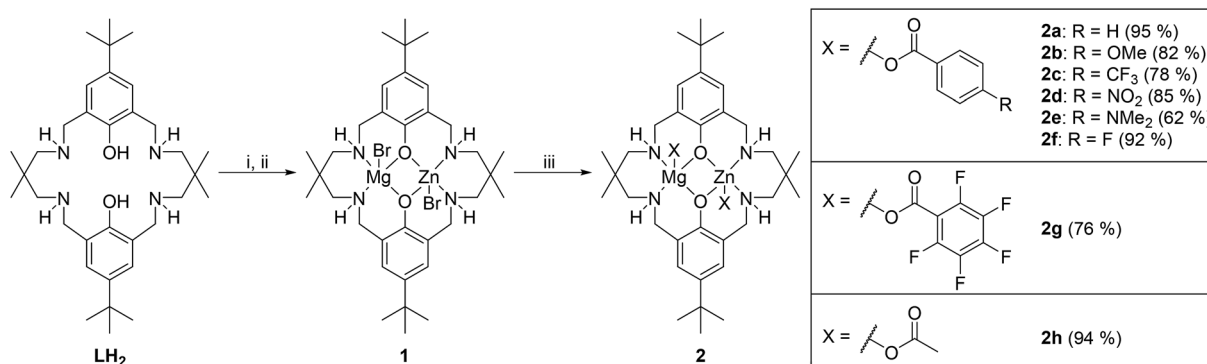
Gemma Trott,^a Jennifer A. Garden ^b and Charlotte K. Williams   ^{*a}

The ring-opening copolymerization of carbon dioxide and epoxides is a useful means to make aliphatic polycarbonates and to add-value to CO_2 . Recently, the first heterodinuclear $\text{Zn}(\text{II})/\text{Mg}(\text{II})$ catalyst showed greater activity than either homodinuclear analogue (*J. Am. Chem. Soc.* **2015**, *137*, 15078–15081). Building from this preliminary finding, here, eight new $\text{Zn}(\text{II})/\text{Mg}(\text{II})$ heterodinuclear catalysts featuring carboxylate co-ligands are prepared and characterized. The best catalysts show very high activities for copolymerization using cyclohexene oxide ($\text{TOF} = 8880 \text{ h}^{-1}$, 20 bar CO_2 , 120 °C, 0.01 mol% catalyst loading) or cyclopentene oxide. All the catalysts are highly active in the low pressure regime and specifically at 1 bar pressure CO_2 . The polymerization kinetics are analysed using *in situ* spectroscopy and aliquot techniques: the rate law is overall second order with a first order dependence in both catalyst and epoxide concentrations and a zero order in carbon dioxide pressure. The pseudo first order rate coefficient values are compared for the catalyst series and differences are primarily attributed to effects on initiation rates. The data are consistent with a chain shuttling mechanistic hypothesis with heterodinuclear complexes showing particular rate enhancements by optimizing distinct roles in the catalytic cycles. The mechanistic hypothesis should underpin future heterodinuclear catalyst design for use both in other (co)polymerization and carbon dioxide utilization reactions.

$\text{Zn}(\text{II})$ or $\text{Mg}(\text{II})$ catalysts are attractive due to the relatively low cost, toxicity, lack of colour and redox activity of these metals.^{20,22} Most research has focussed on zinc complexes and detailed studies indicate many of the best catalysts operate by dinuclear mechanisms.^{15,23–31} For example, optimization of di-zinc β -diiminate catalysts over the last decade has led to very high activities and structure–activity studies indicate that rates depend upon the intermetallic separation and ligand flexibility.^{25–28} In a few cases, it has been shown that replacing the $\text{Zn}(\text{II})$ metal with $\text{Mg}(\text{II})$ in a dinuclear complex increases the activity whilst reducing weight which is particularly desirable for any larger-scale application.^{22,32} Currently the most active catalysts are all homodinuclear complexes, *i.e.* featuring two identical metal centres.^{15,23–31} In 2015, our group reported the first heterodinuclear catalyst, **1**, featuring both $\text{Zn}(\text{II})$ and $\text{Mg}(\text{II})$ metals which are coordinated by a symmetrical macrocycle (Scheme 1).^{33,34} Complex **1** shows significantly higher activity than either the di-zinc or di-magnesium complexes or than mixtures of them.³⁴ Recently some other hetero-catalysts have been reported including a series of zinc tris(lanthanide) complexes some of which show excellent activities at 10 bar pressure.^{35–37}

In general, there are very few reports of the synthesis or properties of well-defined heterodinuclear $\text{Zn}(\text{II})/\text{Mg}(\text{II})$ complexes in the literature, perhaps because these metals are known to be labile, similarly sized and to show equivalent coordination chemistry.^{38–41} In order to properly compare and





Scheme 1 The synthesis of a series of hetero-Zn(II)/Mg(II) catalysts. Reagents: (i) ZnEt₂, THF, 25 °C, 16 h; (ii) MgBr₂, THF, −78 °C → RT, 1 h; (iii) KX, THF, RT, 16 h. Isolated yields in parentheses.

understand the significance of heterodinuclear synergic relationships it is important to use symmetrical ligands but this provides a further difficulty in terms of complex synthesis. In short it requires a means to mono-metallate a symmetrical ancillary ligand.

Despite their limited synthetic precedent di-Zn(II)/Mg(II) complexes are invoked as active sites in conjugate addition reactions,⁴² as highly reactive agents in Pd(II) catalyzed cross-coupling reactions,⁴⁰ in carbonyl addition reactions³⁹ and in stoichiometric fluoroarene substitution reactions.⁴³ Several metalloenzymes also feature heterodinuclear Zn(II)/Mg(II) cores, for example DNA polymerases, aminopeptidases and alkaline phosphatases, and structural studies have been undertaken to model these active sites.^{44–46} These broader applications for heterodinuclear Zn(II)/Mg(II) complexes underscore the importance of both developing their synthetic chemistry and improving the understanding of reactivity.

Results and discussion

Heterodinuclear complex synthesis

The synthesis of **1** was achieved by the reaction of **LH₂** with, first, diethyl zinc and, second, magnesium bromide (Scheme 1). The reaction improves upon a previously published route in that both steps can now be conducted sequentially and in the same pot by using only THF as the solvent. The improved synthesis allowed the isolation of complex **1** at larger scale (multi-gram scale) and in yields of >90%. By carrying out metathesis reactions using complex **1** a series of 8 new heterodinuclear catalysts featuring benzoate co-ligands were isolated. All the potassium carboxylate metathesis reactions proceeded to high conversions and yielded only the heterodinuclear complex. In a few cases the isolated yields are slightly lower than expected on the basis of the reaction conversion (determined by aliquot analysis) due to differences in the amount of solvent needed to fully remove the potassium bromide by-product.

All the complexes were analysed using NMR spectroscopy, mass spectrometry and elemental analyses (see ESI†). All the ¹H NMR spectra (^d₄-methanol) show 8 distinct methylene

resonances and this lack of symmetry is particularly diagnostic of heterodinuclear complex formation (Fig. S1†). In comparison, the C₂-symmetric di-zinc or di-magnesium complexes show only four methylene resonances.^{20,22,34,47,48} The N–H resonances could not be identified when using ^d₄-methanol as the NMR solvent, presumably due to rapid H/D exchange. The ¹H NMR spectra were also determined in ^d₈-THF and although all the methylene resonances were broadened, due to fluxional processes, the phenolate aromatic signals show two distinctive coupled doublets. In all cases the coupling was confirmed by COSY NMR. These two doublets are an important additional indicator of heterodinuclear complex formation since the homodinuclear analogues show singlet resonances and mixtures of homodinuclear complexes show two resonances but without any coupling (Fig. S2†).⁴⁷ The metathesis reactions do not result in any significant metal re-distribution reactions. To use these complexes in polymerization catalysis it is essential to understand the extent, if any, of aggregation in solution. DOSY NMR spectra were determined for all complexes in THF solution so as to mimic the conditions for copolymerization reactions which are conducted in neat epoxide. In all cases a single diffusion coefficient was observed and, when benchmarked against known zinc complexes, the molar masses are in line with the formation of discrete dinuclear complexes (Fig. S3–S10†).

CO₂/CHO polymerization catalysis

The new complexes were tested under a common set of conditions used previously to evaluate homodinuclear epoxide/CO₂ catalysts: specifically at 0.1 mol% catalyst loading, in neat epoxide, 80 °C and at 1 bar pressure of CO₂.^{22,34} The low pressure polymerizations were all conducted in Schlenk tubes with magnetic stirring. The catalytic activity was assessed by running reactions for fixed time periods and characterizing the extent of epoxide conversion. This allows determination of both the turn-over-number (TON) and turn-over-frequency (TOF). In all cases, polycarbonates were produced with a high proportion of carbonate linkages and showed carbon dioxide uptake >99% (where 100% corresponds to a completely alternating copolymer). Furthermore, in all cases the selectivity for polycarbonate



was very high with no cyclic carbonate by-product detected by either ^1H NMR or IR spectroscopy.

The heterodinuclear catalysts are all significantly more active than homodinuclear analogues featuring the same co-ligands. For example, **2h** has a TOF of 98 h^{-1} whereas the di-zinc complex achieves 18 h^{-1} and di-magnesium 30 h^{-1} , under the same conditions.^{22,34} The activity also differs depending on the nature of the carboxylate co-ligand and even shows reproducible differences with only minor changes to the benzoate co-ligands (Table 1, entries 2–7). The most active catalyst, **2d** (*p*-nitro-benzoate), is ~ 3 times more active than the least active catalyst, **2g** (pentafluorobenzoate).

One of the more active catalysts, **2b**, was also tested for the copolymerization of various other epoxides, including cyclopentene oxide (CPO) at 1 bar pressure of CO_2 (Tables S1 and S2†). For CO_2 /CPO ROCOP showed a slightly lower TOF of 30 h^{-1} compared to a value using CHO of 108 h^{-1} . Other catalysts have also been reported to polymerize CPO more slowly than CHO perhaps due to differences in ring strain between the two epoxides.^{19,49–52} The copolymerization was very selective both for polymer (vs. cyclic carbonate) and carbonate linkage formation (>99%). Under optimized conditions (0.01 mol% **2b**, 20 bar CO_2 , $80\text{ }^\circ\text{C}$), a TOF value of 76 h^{-1} was achieved and poly(cyclopentene carbonate) with molar mass values as high as $42\,000\text{ g mol}^{-1}$ was produced. These activities and molar masses are both at the leading end of values reported for CPO polymerizations.^{19,49–52} **2b** was also an efficient catalyst for ROCOP using vinyl-cyclohexene oxide (TOF = 71 h^{-1} and carbonate selectivity >99%), cyclohexadiene oxide (TOF = 1 h^{-1} , carbonate selectivity >99%) and 3,4-epoxypentahydrofuran (TOF = 14 h^{-1} , carbonate selectivity >99%) (Table S2†). However, it

showed only low activity and selectivity in the ROCOP of PO/ CO_2 (TOF = 8 h^{-1} , carbonate selectivity = 98%, polymer selectivity = 26%) (Table S2†).

Polymerization kinetics

In order to understand the relative activity of the series of complexes detailed analysis of the polymerization kinetics was undertaken using *in situ* IR spectroscopy. This analysis is recommended so as to allow determination of rate coefficients which should allow for more accurate comparison of catalyst performances than point kinetic values (TOF).

The polymerization rate law was determined using **2e** which is one of the most active catalysts of the series. Starting from an idealized rate law dependent upon all reagent concentrations:

$$\text{Rate} = k_p[2\mathbf{e}]^x[\text{CHO}]^y p\text{CO}_2^z$$

Various experiments were conducted to determine the values for x , y and z . Firstly, the order in catalyst concentration (x) was determined by conducting experiments under pseudo first order conditions and applying catalyst concentrations from 1.5–7 mM. In every case *in situ* ATR-IR spectroscopy was used to monitor the reactions and the initial rate coefficient, k_{obs} , determined as the gradient of linear fits to the data over the conversion range 0–20% (see Fig. S43–S47† for each data set and associated fit). The plot of k_{obs} vs. $[2\mathbf{e}]$ shows a linear fit to the data which is consistent with a first order dependence on catalyst concentration (Fig. 1a). Alternatively, plots of $\ln(k_{\text{obs}})$ vs. $\ln[\text{catalyst}]$ show a gradient of 0.95 which is also consistent with a first order dependence on catalyst concentration.

Table 1 Cyclohexene oxide/ CO_2 copolymerization data for catalysts **1** and **2a–h** at 1 bar CO_2 ^a

#	Cat.	TON ^b	Initiation (min) ^c	TOF (h^{-1}) ^d	CO_2 (%) ^e	Polymer ^f (%)	$k_{\text{obs}} (\times 10^{-5} \text{ s}^{-1})$ ^g	$M_n [D]$ ^h
1	1	247	160	78	>99	>99	3.0	3000 [1.18]
2	2a	407	50	87	>99	>99	2.8	13 300 [1.03]
3	2b [†]	252	40	108	>99	>99	3.5	6400 [1.06]
4	2c	383	30	89	>99	>99	3.5	8700 [1.04]
5	2d [†]	307	20	124	>99	>99	3.8	3500 [1.18]
6	2e [†]	258	27	101	>99	>99	3.3	10 500 [1.03]
7	2f	437	45	83	>99	>99	3.4	4700 [1.13]
8	2g	199	80	43	>99	>99	1.4	10 200 [1.02]
9	2h	438	50	98	>99	>99	3.1	4750 [1.08]
								9980 [1.04]
								4200 [1.14]
								19 000 [1.03]
								8000 [1.18]
								2800 [1.23]
								12 700 [1.04]
								5100 [1.16]

^a Reactions were run using a 1 : 1000 catalyst : CHO, in neat CHO ($[\text{CHO}]_0 = 9.88\text{ M}$), $80\text{ }^\circ\text{C}$, 1 bar pressure of CO_2 , 6 h or where [†] shown for 3 h, stirring rate = 600 rpm. ^b TON = number of moles of cyclohexene oxide consumed/number of moles of catalyst (see ESI for more information).

^c Determined using ATR-IR spectroscopy as the time taken for the on-set of polycarbonate absorptions (Fig. S11–S19). ^d TOF = TON/polymerization time, *i.e.* corrected to account for the initiation period; the average error in TOF is $\pm 4\%$. ^e Determined from the ^1H NMR spectrum by comparing normalised integrals for carbonate (4.65 ppm) and ether (3.45 ppm) resonances (Fig. S29). ^f Determined from the ^1H NMR spectrum by comparing normalised integrals for polycarbonate (4.65 ppm) and cyclic carbonate (4.00 ppm) resonances (Fig. S29).

^g Determined by an initial rates method (see ESI and Fig. S20–S28). ^h Determined by SEC, in THF, calibrated with narrow molar mass polystyrene standards [dispersity values in parentheses] (Fig. S30–S38).



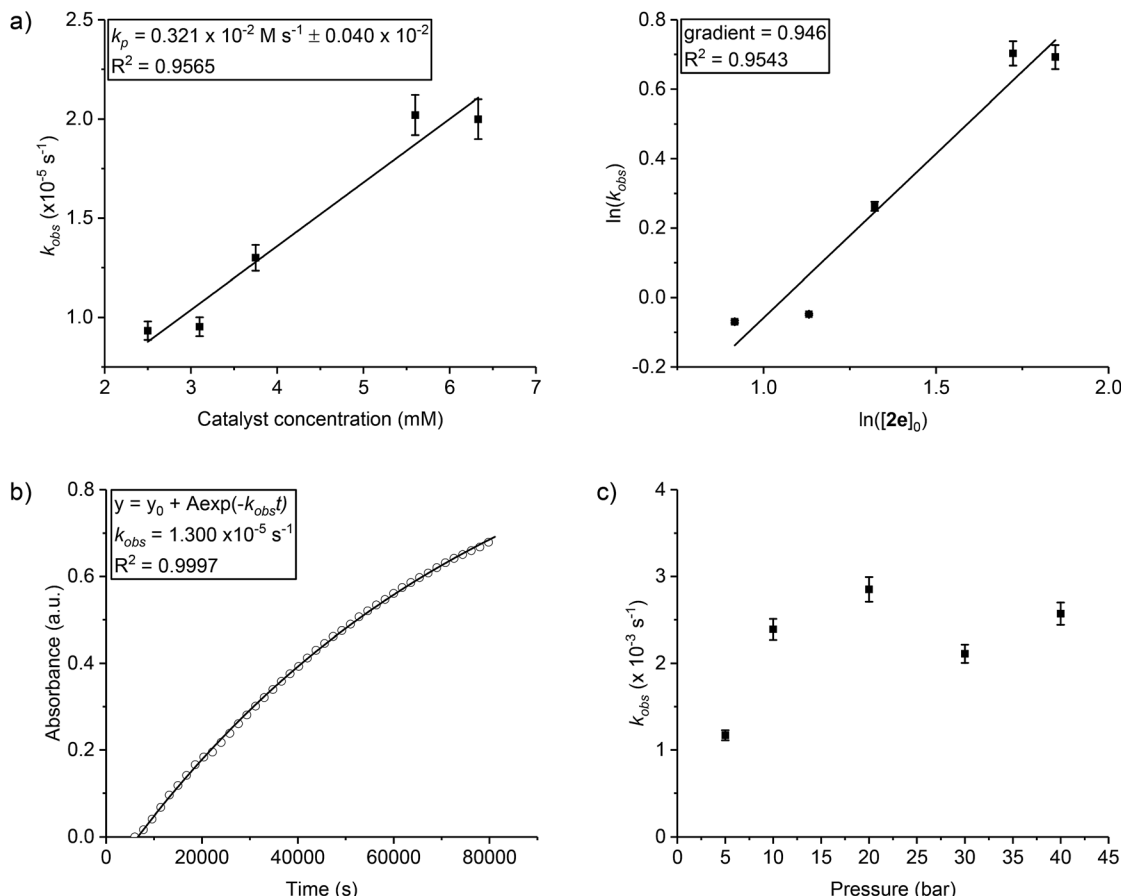


Fig. 1 Reaction kinetic analysis to determine order in (a) catalyst (first order); (b) epoxide, **CHO** (first order); (c) CO_2 pressure (zero order) for catalyst **2e**. Reaction conditions: $[\text{CHO}]_0 = 5 \text{ M}$ in diethyl carbonate, $[\text{2e}]_0 = 5 \text{ mM}$ (for plots b and c), temperature = 80°C (for plots a and b) or 120°C (for plot c), CO_2 pressure = 1 bar (for plots a and b). For clarity only every 30th data point is shown for plot b. 5% error bars shown and further details on data collection methods and analysis are available in the ESI.†

The order in epoxide concentration was determined by an integrated rate law method and involved analysis of the polymerization from 0–65% conversion (after which point viscosity is sufficiently high as to become a limiting factor). Analysis of the absorbance *vs.* time data shows an exponential growth in the absorption intensity assigned to the polycarbonate. Applying an exponential fit to the data demonstrates that the reaction is first order in epoxide concentration and shows a k_{obs} value of $1.3 \times 10^{-5} \text{ s}^{-1}$. It is notable that the rate coefficients obtained using either the initial rates method or the integrated rate law treatment show very good agreement which gives confidence in using the initial rates method to compare catalyst activities (*e.g.* for catalyst **2e** the integrated rate law, $k_{\text{obs}} = 1.30 \times 10^{-5} \text{ s}^{-1}$ *vs.* initial rates method, $k_{\text{obs}} = 1.44 \times 10^{-5} \text{ s}^{-1}$). In order to determine the order in carbon dioxide pressure a series of runs were conducted in a high pressure autoclave equipped with an ATR-IR *in situ* probe. The experiments were conducted at 120°C and, therefore, show overall faster rates than equivalent studies at 80°C . Plots of rate coefficient, k_{obs} , *vs.* CO_2 pressure show a weak correlation between rate and pressure only up to 10 bar and thereafter values are broadly similar (Fig. 1c and S48–S55†). The apparent increment in activity from

5–10 bar is mostly likely due to an experimental limitation: it was observed from the reactor pressure gauge that very rapid carbon dioxide consumption occurred particularly in the early stages of polymerization but the reactor has no automatic refill mechanism to top-up the consumed CO_2 and maintain constant pressure. The relative change in CO_2 concentration as the reaction progresses is therefore much more likely to be significant at 5 bar than at higher pressures. The results suggest that there is a zero order dependence in carbon dioxide pressure over the range 10–40 bar and indicate a zero-order dependence at lower pressures if CO_2 concentration were maintained. The rate law for catalyst **1** was also determined, since it carries a slightly different halide co-ligand: it showed the same second order rate law (Fig. S53–S55†).

Overall, the kinetic data indicate a second order rate law and show a first order dependence on both catalyst and epoxide concentration but a zero order dependence on carbon dioxide (10–40 bar). It is proposed that the rate determining step is metal carbonate attack on coordinated epoxide. A feasible mechanistic interpretation for the reaction kinetics is that the polymerization follows the chain shuttling mechanism which is discussed later (Scheme 2).^{29,30,33}

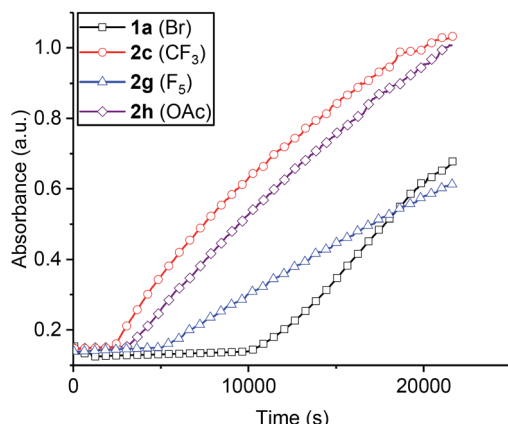
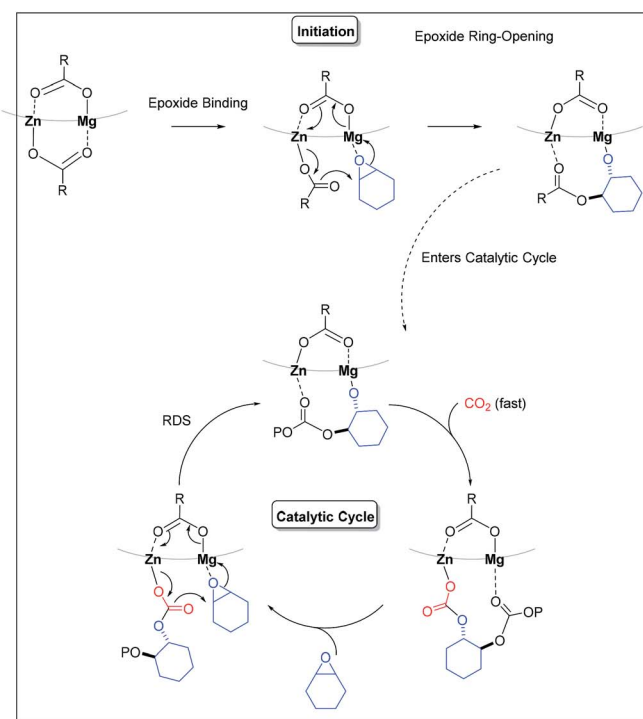


Fig. 2 Raw data plots for selected catalysts (co-ligands) to illustrate the significance of initiation time and propagation rates (the data for all catalysts are shown in the ESI and Fig. S11–S19†). For clarity every 10th data point is shown.

To compare the polymerization behaviour of the series of new complexes, *in situ* reaction analyses and kinetic fits were applied to all 8 catalysts. The initial rates method was used to fit the data and to differentiate between initiation periods and propagation rates for the catalysts (Fig. S20–S28†). All the catalysts showed initiation periods, during which <5% conversion was observed, and the length of these initiation times was variable (Fig. 2, Table 1). This induction period could be

attributed to the dissociation of the bridging carboxylate ligand, providing a coordinative vacancy for epoxide binding, although other processes cannot be excluded such as rates of CO₂ dissolution⁴⁸ and/or epoxide hydrolysis to form diol species.⁵⁴ Following the initiation period, the polymerization conversion *vs.* time data over the range 0–20% conversion was linearly fit, enabling determination of the initial rate coefficient (Table 1). Although the series showed different initiation times, nearly all the values fall within the range $2.8\text{--}3.8 \times 10^{-5} \text{ s}^{-1}$. There is no obvious correlation between electron donating/withdrawing properties of the benzoate co-ligand and propagation rate. The exception is catalyst 2g which features a penta-fluorobenzoate co-ligand and shows a substantially slower polymerization rate. In terms of propagation rates, the largest difference corresponds to the three fold difference between 2g (pentafluorobenzoate) and 2d (nitro-benzoate). Generally, the higher the propagation rate, the shorter the initiation period although there are cases where this correlation breaks down (Table 1, entries 6 and 7).

The similarity between the propagation rate coefficients appears, at first sight, be contrary to the chain shuttling mechanistic hypothesis which suggests that one benzoate ligand remains coordinated during propagation (Scheme 2). There are two alternative interpretations of the activity data: either the chain shuttling mechanism needs fine-tuning to reflect that initiation occurs from both sites (*i.e.* the co-ligand only influences initiation) or most of these benzoate co-ligands show marginal electronic differences and thus there is



Scheme 2 LHS: chain shuttling mechanism proposed for the heterodinuclear catalysts. The ligand framework is shown as a light grey curved line both for clarity and to reflect the finding that the ligand adopts this conformation in solid state structures (with N–H substituents being on the same, convex face of the molecule).²⁹ R is acetate- or (substituted)benzoate-co-ligand and P stands for growing polymer chain. RHS shows the recently reported structure of a related Zn(II)/Mg(II) heterodinuclear complex with THF coordinated at the Mg(II) centre.⁴⁷



no significant moderation of propagation rate with these co-ligands. In support of the second rationale is the observation that the NMR resonances for nearly all the benzoate complexes show very similar chemical shifts, *i.e.* the *para*-substituent on the benzoate ligand does not strongly influence the catalyst electronics. Complex **2g** (pentafluorobenzoate) is, once again, a clear exception as its carbonyl resonance is significantly shifted upfield which indicates a different electronic environment compared to the other benzoates. The most consistent interpretation of the data for complex **2g** is to invoke the chain shuttling mechanism since this allows for the co-ligand to influence both rates of initiation and propagation (Scheme 2). Future research could be directed towards investigating other co-ligands with strongly differing electronic properties. It should be noted, however, that in terms of combining the best catalyst rates with simple process conditions the selection of carboxylate ligands is desirable as they are highly active and air-stable complexes.

Catalytic conditions

Next, the influences of the reaction conditions on activity and selectivity were explored. These studies were conducted using three of the best performing catalysts (**2b**, **c**, **e**) at 1 bar (Table 2). Increasing the reaction temperature from 80 to 120 °C resulted in a significant increase in the catalytic activity and TOF values reached $\sim 400 \text{ h}^{-1}$ for all three catalysts. Even at this elevated reaction temperature there was no evidence for any by-product formation and high proportions of carbonate linkages were observed. Using high temperature conditions and an autoclave to ensure sufficient carbon dioxide concentration (20 bar) allowed the loading of catalyst **2c** to be reduced significantly and resulted in a very high TOF value of 8830 h^{-1} . The catalyst even retained acceptable activity at a loading as low as 1 : 20 000 (**2c** : CHO).

Polymerization control

The catalysts also showed good polymerization control, as assessed by linear increases in polymer molar mass vs. conversion (Fig. 3b). The polycarbonates generally showed bimodal molar mass distributions, albeit with narrow dispersity in each distribution. Such bimodality has frequently been

observed previously in the literature and has been extensively studied.^{4,22,53,54} Bimodality commonly results because of a side-reaction between residual water and cyclohexene oxide to generate 1,2-cyclohexanediol (CHD).⁵⁵ Since all chains propagate at the same rate those initiated from mono-functional groups (*i.e.* catalyst carboxylates) show approximately half the molar mass of chains initiated from di-functional groups (*i.e.* the diol).^{21,54–56} MALDI-ToF analysis of the chain end groups substantiates this hypothesis since two series of chains are observed due to carboxylate and hydroxyl telechelic chains, respectively. When reactions were conducted with the addition of 10 equivalents of CHD or water, monomodal molar mass distributions were observed with narrow dispersity. The MALDI-ToF spectra, under these conditions, showed only chains end-capped with hydroxyl groups (Fig. 3a). The excess chain transfer agent reduces slightly the activity and productivity of the catalyst, in particular when using excess water, possibly due to accelerated catalyst decomposition pathways (Table 3).

A significant benefit of this series of compounds is their air-stability and to demonstrate how this could be exploited a reaction was conducted using **2b**, dissolved in CHO, which was stirred in air for 1 h prior to addition of CO₂. The polymerization was successful and resulted in good activity and selectivity values (Table 3). The resulting polymer showed a monomodal molar mass distribution without requiring any addition of chain transfer agent.

Discussion

Overall this series of catalysts shows high activities and selectivity and performs successfully in the low CO₂ pressure regime. In order to provide context to the activity and selectivity values these new heterodinuclear catalysts are compared against some of the most successful literature catalysts (Fig. 4). The best Zn(II)/Mg(II) catalyst (**2d**, TOF = 124 h^{-1}) is significantly more active than either the di-Zn(II) or di-Mg(II) analogues (18 h^{-1} and 30 h^{-1} , respectively, under equivalent conditions).³³ The activity is also higher than previously reported di-Zn(II) catalysts coordinated by a non-macrocyclic ligand (TOF < 3 h^{-1}).^{57,58} The activity matches that of proline ligated di-Zn(II) catalysts (TOF $\sim 149 \text{ h}^{-1}$)⁵⁹ but is lower than optimized Co(III)-salen catalysts (TOF = 263 h^{-1}).⁶⁰ One advantage of these Zn(II)/Mg(II)

Table 2 Copolymerization reaction conditions^a

Cat.	Catalyst (mol%)	Temp. (°C)	Pressure (bar)	PCHC ^b (%)	TON ^b	TOF (h ⁻¹) ^b	M _n [D] ^b
2b	0.1	120	1	>99	435	377	12 280 [1.04] 5340 [1.13]
2c	0.1	120	1	>99	466	419	14 490 [1.06] 5930 [1.15]
2e	0.1	120	1	>99	645	430	21 760 [1.04] 9090 [1.15]
2c	0.01	120	20	>99	4415	8830	44 400 [1.04] 21 200 [1.05]
2c	0.005	120	20	>99	5435	1359	54 380 [1.04] 26 550 [1.04]

^a Reactions were carried out in a Parr high pressure vessel with an impeller at 20 bar. ^b See Table 1 and ESI for all data (Fig. S56–S60).



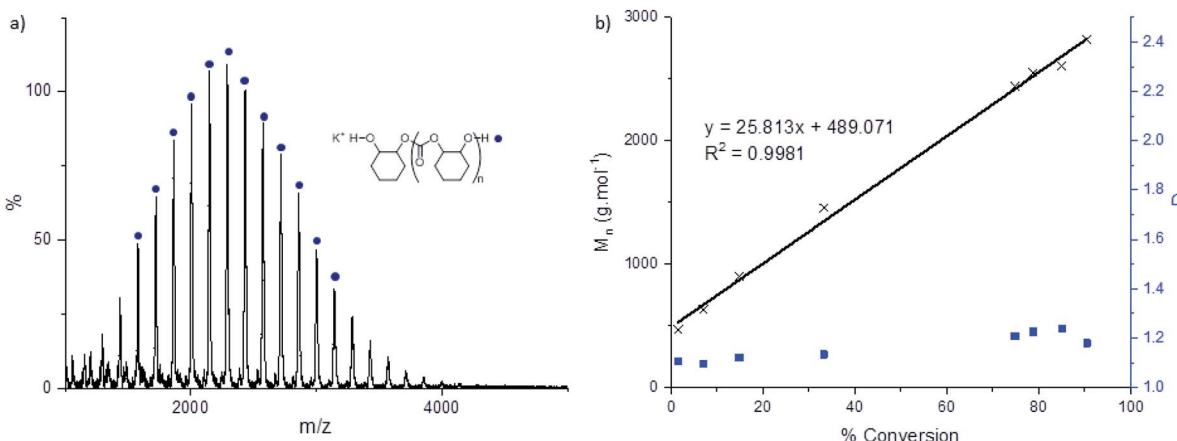


Fig. 3 (a) MALDI-ToF mass spectrum of poly(cyclohexene carbonate) formed from catalyst **2c** (0.1 mol%) with 10 equivalents of added CHD. (b) Plot of the polycarbonate M_n (black crosses) and D (blue squares) versus % CHO conversion. Reaction conditions: **2b**/CHO/CHD = 1/1000/10, 80 °C, 5 M in toluene (Fig. S65–S72†).

heterodinuclear catalysts is that very high CO_2 and polymer selectivity is maintained even at elevated temperatures which contrasts with simple Co(III) salen catalyst systems that are well-known to be thermally unstable with respect to cobalt reduction and catalyst death. For the Zn(II)/Mg(II) complexes, the best activity value reaches 654 h^{-1} which is at the upper end of values for the low pressure regime.^{10,14,15}

At higher CO_2 pressures (20 bar), catalyst **2c** showed an activity of 8800 h^{-1} . This value significantly exceeds the performance of tetranuclear Zn(II)/Ln(III) catalysts ($\text{TOF} = 200\text{--}300 \text{ h}^{-1}$, 10 bar).³⁶ It is also higher than a series of new organocatalysts, formed *in situ* by reaction between boranes and ionic salts ($\text{TOF} = 600 \text{ h}^{-1}$, 10 bar),⁶¹ and higher than for optimized Cr(III)-salen catalyst systems (1153 h^{-1}).⁶² Indeed, the activity is only surpassed by a tethered di-zinc(II) β -diiminate catalyst ($\text{TOF} = 155\,000 \text{ h}^{-1}$), but its selectivity for CO_2 is significantly higher (>99% for **2c** vs. 80% for di-zinc catalyst).²⁵ The activity values obtained using cyclopentene oxide/ CO_2 are also high compared to other catalysts.^{19,49–52}

Polymerization mechanism

The polymerization rate law is second order and shows a first order dependence on catalyst and epoxide concentration but appears largely independent of CO_2 pressure. The polymerization kinetics are interpreted using a chain shuttling mechanism

(Scheme 2).^{29,33} According to the mechanism, with each monomer insertion step the growing polymer chain moves metal. The rate law indicates the rate determining step is metal carbonate attack on an epoxide molecule coordinated at the second metal. DFT studies indicate that once the epoxide is ring-opened the polymer chain shuttles to the metal originally coordinating the epoxide.²⁹ To balance charge, the carboxylate co-ligand also changes its anionic coordination site to the opposite metal. The fast step in the catalytic cycle is proposed to be carbon dioxide insertion and DFT studies have indicated this step also involves migration of the metal-alkoxide growing polymer chain end from one metal to the other.²⁹ Once again, the carboxylate co-ligand changes its site of anionic coordination to the opposite metal. The proposed mechanism requires that the polymer chain ‘shuttles’ between the two metal centres twice per complete cycle of insertions and that one of the benzoate co-ligands remains coordinated throughout the polymerization and serves to counter-balance charge and polymer chain movement on the opposite face of the catalyst. The results of the structure-kinetic study presented here show only a weak correlation between propagation rate and benzoate co-ligands but this appears most likely due to the limited electronic differences between the benzoates. In the case of the pentafluorobenzoate co-ligand there is NMR spectroscopic evidence for a stronger electronic influence and concomitant evidence of a significant

Table 3 Cyclohexene oxide/ CO_2 copolymerizations with Chain Transfer Agents (CTA)^a

Entry	Catalyst	CTA (eq.)	CO_2 ^b (%)	Polymer ^b (%)	TON ^b	TOF (h ⁻¹) ^b	$M_n [D]^b$
1	2c	None	>99	>99	634	106	7000 [1.04] 3400 [1.07]
2	2c	H_2O (10)	>99	>99	274	46	600 [1.14]
3	2c	CHD (10) ^c	>99	>99	440	73	1700 [1.10]
4	2b ^d	None	>99	>99	559	93	2870 [1.13]

^a Reactions were run using a catalyst : CHO molar ratio of 1 : 1000 at 80 °C under 1 bar pressure of CO_2 for 6 hours. ^b See Table 1 and ESI for all data (Fig. S61–S64). ^c CHD = 1,2-cyclohexanediol. ^d Catalyst and CHO were stirred in air for 1 h at 25 °C before purging the system with CO_2 and heating to 80 °C.

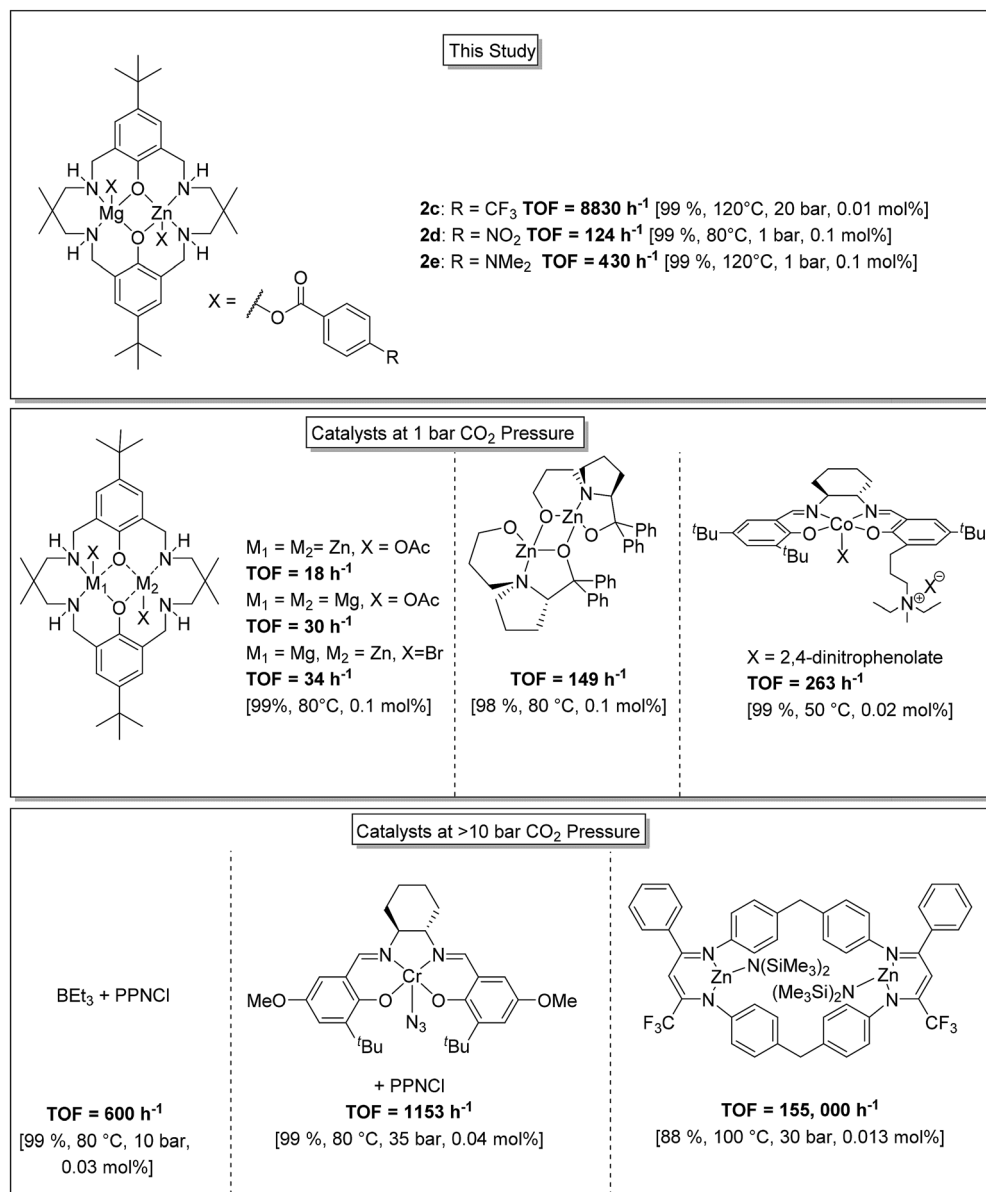


Fig. 4 Illustration of the structures, activity and selectivity for some of the highest performing catalysts reported for CO₂/CHO ROCOP.^{22,25,33,34,59,61}

difference in propagation rate. This data can be best interpreted according to the chain shuttling process since the benzoate remains coordinated during propagation and can influence the propagation rate.

The chain shuttling mechanism allows for the key roles in catalysis to be differentiated and may also rationalize the improved performances of heterodinuclear catalysts *vs.* homodinuclear counterparts. Since the rate limiting step is likely metal epoxide attack by the second metal carbonate group, an obvious question is: which metal coordinates the epoxide and which the carbonate? A detailed DFT study was conducted comparing the rate limiting steps for di-zinc, di-magnesium and a Zn/Mg complex (see ESI† for detailed discussion and results). For the heterodinuclear complex the propagation

cycles were examined for either zinc-epoxide or magnesium-epoxide coordination pathways. The DFT results substantiated the mechanistic hypothesis by showing a substantially lower barrier for epoxide molecule coordination at Mg(II) compared to Zn(II) [with the epoxide coordinated intermediates showing relative stabilities of -7.7 kcal mol⁻¹ (Mg(II) coordination) *vs.* -1.4 kcal mol⁻¹, (Zn(II) coordination), respectively]. All the catalysts showed transition state barriers, for the rate limiting step, that were not significantly different, within the error limits for DFT. Although this finding does not allow DFT to provide any further mechanistic insight, it is consistent with the observed differences in rates. Very recently a solid state structure was obtained for another heterodinuclear complex (analogue of complex 1) obtained by growing crystals from

a THF solution (Scheme 2b).⁴⁷ The structure shows the two different metals and has a molecule of THF coordinated at the Mg(II) centre.

Accordingly the chain shuttling mechanistic hypothesis can be fine-tuned for Zn(II)/Mg(II) complexes. The polymer chain is alternately coordinated by the two metal centres. In the rate limiting step the zinc centre provides the carbonate group which attacks the magnesium coordinated epoxide molecule. This results in the formation of a magnesium coordinated alkoxide group which undergoes fast CO₂ insertion and forms a zinc carbonate intermediate. As the Zn(II)/Mg(II) complexes show significantly greater rates than di-zinc or di-magnesium analogues it is also clear that the proximity of the magnesium centre stabilizes the zinc carbonate group. This could occur through electronic communication through the phenolate oxygen atom which bridges the two metal centres and provides a means to control electron density and hence reactivity at the metals.^{63,64} The mechanistic findings from these hetero-Zn(II)/Mg(II) catalysts are expected to be more broadly applicable to other dinuclear CO₂ copolymerization catalysts. A clear recommendation is to prepare Zn(II)/Mg(II) complexes of dinucleating bis(β-diiminate) ligands which would be expected to show even higher activity than the best dizinc(II) complexes. The chain shuttling pathway is also expected to apply to other polymerization processes and can be exploited to enhance rates of epoxide/anhydride alternating polymerizations and epoxide/heterocumulene processes. Generally there is intense interest and motivation to activate carbon dioxide in chemistry and the concepts illustrated in this work of reagent shuttling and activation between two dissimilar metal sites as a means to increase catalytic performances warrant investigation for other CO₂ utilization processes.

Conclusions

The successful isolation of 8 new heterodinuclear Zn(II)/Mg(II) complexes results in highly active catalysts for CO₂/epoxide alternating copolymerization. All the new complexes show very good performances both in terms of activity (TOF = 8800 h⁻¹) and selectivity (>99% CO₂ uptake over the entire temperature range). The catalysts are amongst the most active reported both at low CO₂ pressure (1 bar) and in the higher pressure regime (>10 bar). The polymerization kinetics, analysed using *in situ* IR spectroscopy, showed an overall second order rate law. The rates were first order dependent on catalyst and epoxide concentrations but independent of carbon dioxide pressure (1–40 bar). Polymerization control was high in all cases and the reaction conditions were controlled to allow the preparation of polycarbonate polyols which are relevant to polyurethane manufacture. Overall, these heterodinuclear catalysts warrant further investigation both as catalysts for polymerizations and for a range of organic transformations, such as conjugate additions, fluorinations and even hydrolysis reactions. In terms of CO₂/epoxide alternating copolymerization catalysis there is still significant scope to moderate both the metals, ligands and co-ligands so as to optimize performances. Heterodinuclear complexes of Zn(II)/Mg(II) using other dinucleating ancillary

ligands should be explored as a means to increase activity and maintain high overall selectivity and control.

Conflicts of interest

Charlotte K. Williams is a director of Econic Technologies.

Acknowledgements

The EPSRC (EP/L017393/1; EP/K014668/1) and Climate KIC (Project EnCO₂re) are acknowledged for research funding.

Notes and references

- 1 J. Artz, T. E. Muller, K. Thenert, J. Kleinekorte, R. Meys, A. Sternberg, A. Bardow and W. Leitner, *Chem. Rev.*, 2018, **118**, 434–504.
- 2 Y. Zhu, C. Romain and C. K. Williams, *Nature*, 2016, **540**, 354–362.
- 3 N. v. d. Assen and A. Bardow, *Green Chem.*, 2014, **16**, 3272–3280.
- 4 A. M. Chapman, C. Keyworth, M. R. Kember, A. J. J. Lennox and C. K. Williams, *ACS Catal.*, 2015, **5**, 1581–1588.
- 5 J. Langanke, A. Wolf, J. Hofmann, K. Bohm, M. A. Subhani, T. E. Muller, W. Leitner and C. Gurtler, *Green Chem.*, 2014, **16**, 1865–1870.
- 6 S. H. Lee, A. Cyriac, J. Y. Jeon and B. Y. Lee, *Polym. Chem.*, 2012, **3**, 1215–1220.
- 7 T. Stoesser, C. Li, J. Unruangsri, P. K. Saini, R. J. Sablong, M. A. R. Meier, C. K. Williams and C. Koning, *Polym. Chem.*, 2017, **8**, 6099–6105.
- 8 O. Hauenstein, M. Reiter, S. Agarwal, B. Rieger and A. Greiner, *Green Chem.*, 2016, **18**, 760–770.
- 9 M. A. Subhani, B. Kohler, C. Gurtler, W. Leitner and T. E. Muller, *Angew. Chem., Int. Ed.*, 2016, **55**, 5591–5596.
- 10 Y. Y. Wang and D. J. Daresbourg, *Coord. Chem. Rev.*, 2018, **372**, 85–100.
- 11 A. W. Kleij, M. North and A. Urakawa, *ChemSusChem*, 2017, **10**, 1036–1038.
- 12 M. I. Childers, J. M. Longo, N. J. Van Zee, A. M. LaPointe and G. W. Coates, *Chem. Rev.*, 2014, **114**, 8129–8152.
- 13 X.-B. Lu and D. J. Daresbourg, *Chem. Soc. Rev.*, 2012, **41**, 1462–1484.
- 14 G. Trott, P. K. Saini and C. K. Williams, *Philos. Trans. R. Soc., A*, 2016, **374**, 19.
- 15 J. A. Garden, *Green Mater.*, 2017, **5**, 103–108.
- 16 D. J. Daresbourg and S. J. Wilson, *Green Chem.*, 2012, **14**, 2665–2671.
- 17 S. J. Na, S. Sujith, A. Cyriac, B. E. Kim, J. Yoo, Y. K. Kang, S. J. Han, C. Lee and B. Y. Lee, *Inorg. Chem.*, 2009, **48**, 10455–10465.
- 18 Y. Liu, W.-M. Ren, J. Liu and X.-B. Lu, *Angew. Chem., Int. Ed.*, 2013, **52**, 11594–11598.
- 19 Y. Liu, W.-M. Ren, K.-K. He and X.-B. Lu, *Nat. Commun.*, 2014, **5**, 5687–5694.
- 20 M. R. Kember, P. D. Knight, P. T. R. Reung and C. K. Williams, *Angew. Chem., Int. Ed.*, 2009, **48**, 931–933.



21 M. R. Kember, A. J. P. White and C. K. Williams, *Macromolecules*, 2010, **43**, 2291–2298.

22 M. R. Kember and C. K. Williams, *J. Am. Chem. Soc.*, 2012, **134**, 15676–15679.

23 S. D. Allen, D. R. Moore, E. B. Lobkovsky and G. W. Coates, *J. Am. Chem. Soc.*, 2002, **124**, 14284–14285.

24 M. Cheng, E. B. Lobkovsky and G. W. Coates, *J. Am. Chem. Soc.*, 1998, **120**, 11018–11019.

25 S. Kissling, M. W. Lehenmeier, P. T. Altenbuchner, A. Kronast, M. Reiter, P. Deglmann, U. B. Seemann and B. Rieger, *Chem. Commun.*, 2015, **51**, 4579–4582.

26 D. R. Moore, M. Cheng, E. B. Lobkovsky and G. W. Coates, *J. Am. Chem. Soc.*, 2003, **125**, 11911–11924.

27 M. Reiter, S. Vagin, A. Kronast, C. Jandl and B. Rieger, *Chem. Sci.*, 2017, **8**, 1876–1882.

28 S. Kissling, P. T. Altenbuchner, M. W. Lehenmeier, E. Herdtweck, P. Deglmann, U. B. Seemann and B. Rieger, *Chem.-Eur. J.*, 2015, **21**, 8148–8157.

29 A. Buchard, F. Jutz, M. R. Kember, A. J. P. White, H. S. Rzepa and C. K. Williams, *Macromolecules*, 2012, **45**, 6781–6795.

30 F. Jutz, A. Buchard, M. R. Kember, S. B. Fredrickson and C. K. Williams, *J. Am. Chem. Soc.*, 2011, **133**, 17395–17405.

31 C. Romain, A. Thevenon, P. K. Saini and C. K. Williams, in *Carbon Dioxide and Organometallics*, ed. X. B. Lu, 2016, vol. 53, pp. 101–141.

32 S. Ghosh, D. Pahovnik, U. Kragl and E. Mejia, *Macromolecules*, 2018, **51**, 846–852.

33 P. K. Saini, C. Romain and C. K. Williams, *Chem. Commun.*, 2014, **50**, 4164–4167.

34 J. A. Garden, P. K. Saini and C. K. Williams, *J. Am. Chem. Soc.*, 2015, **137**, 15078–15081.

35 J. Qin, B. Xu, Y. Zhang, D. Yuan and Y. Yao, *Green Chem.*, 2016, **18**, 4270–4275.

36 H. Nagae, R. Aoki, S.-n. Akutagawa, J. Kleemann, R. Tagawa, T. Schindler, G. Choi, T. P. Spaniol, H. Tsurugi, J. Okuda and K. Mashima, *Angew. Chem., Int. Ed.*, 2018, **57**, 2492–2496.

37 J. A. Garden, A. J. P. White and C. K. Williams, *Dalton Trans.*, 2017, **46**, 2532–2541.

38 R. E. Mulvey and S. D. Robertson, in *Organometallic Compounds*, ed. Z. Xi, Springer-Verlag Berlin, Berlin, 2014, vol. 47, pp. 129–158.

39 D. R. Armstrong, W. Clegg, P. Garcia-Alvarez, A. R. Kennedy, M. D. McCall, L. Russo and E. Hevia, *Chem.-Eur. J.*, 2011, **17**, 8333–8341.

40 T. D. Bluemke, W. Clegg, P. Garcia-Alvarez, A. R. Kennedy, K. Koszinowski, M. D. McCall, L. Russo and E. Hevia, *Chem. Sci.*, 2014, **5**, 3552–3562.

41 M. Westerhausen, *Dalton Trans.*, 2006, 4755–4768.

42 B. M. Trost and M. J. Bartlett, *Acc. Chem. Res.*, 2015, **48**, 688–701.

43 C. Bakewell, B. J. Ward, A. J. P. White and M. R. Crimmin, *Chem. Sci.*, 2018, **9**, 2348–2356.

44 P. S. Freemont, J. M. Friedman, L. S. Beese, M. R. Sanderson and T. A. Steitz, *Proc. Natl. Acad. Sci. U. S. A.*, 1988, **85**, 8924–8928.

45 S. K. Burley, P. R. David, A. Taylor and W. N. Lipscomb, *Proc. Natl. Acad. Sci. U. S. A.*, 1990, **87**, 6878–6882.

46 T. Tanase, J. W. Yun and S. J. Lippard, *Inorg. Chem.*, 1996, **35**, 3585–3594.

47 A. C. Deacy, C. B. Durr, J. A. Garden, A. J. P. White and C. K. Williams, *Inorg. Chem.*, 2018, **57**(24), 15575–15583.

48 C. Romain, J. A. Garden, G. Trott, A. Buchard, A. J. P. White and C. K. Williams, *Chem.-Eur. J.*, 2017, **23**, 7367–7376.

49 Y. Z. Hua, X. C. Yang, M. M. Liu, X. X. Song, M. C. Wang and J. B. Chang, *Macromolecules*, 2015, **48**, 1651–1657.

50 D. J. Daresbourg, W.-C. Chung and S. J. Wilson, *ACS Catal.*, 2013, **3**, 3050–3057.

51 D. J. Daresbourg, S.-H. Wei, A. D. Yeung and W. C. Ellis, *Macromolecules*, 2013, **46**, 5850–5855.

52 L. S. Huang, C. Y. Tsai, H. J. Chuang and B. T. Ko, *Inorg. Chem.*, 2017, **56**, 6141–6151.

53 M. R. Kember, J. Copley, A. Buchard and C. K. Williams, *Polym. Chem.*, 2012, **3**, 1196–1201.

54 G.-P. Wu and D. J. Daresbourg, *Macromolecules*, 2016, **49**, 807–814.

55 N. Yi, J. Unruangsri, J. Shaw and C. K. Williams, *Faraday Discuss.*, 2015, **183**, 67–82.

56 W. J. van Meerendonk, R. Duchateau, C. E. Koning and G.-J. M. Gruter, *Macromolecules*, 2005, **38**, 7306–7313.

57 Y. Xiao, Z. Wang and K. Ding, *Chem.-Eur. J.*, 2005, **11**, 3668–3678.

58 Y. Xiao, Z. Wang and K. Ding, *Macromolecules*, 2006, **39**, 128–137.

59 M. Schutze, S. Dechert and F. Meyer, *Chem.-Eur. J.*, 2017, **23**, 16472–16475.

60 W.-M. Ren, X. Zhang, Y. Liu, J.-F. Li, H. Wang and X.-B. Lu, *Macromolecules*, 2010, **43**, 1396.

61 D. Zhang, S. K. Boopathi, N. Hadjichristidis, Y. Gnanou and X. Feng, *J. Am. Chem. Soc.*, 2016, **138**, 11117–11120.

62 D. J. Daresbourg, R. M. Mackiewicz and D. R. Billodeaux, *Organometallics*, 2005, **24**, 144.

63 L. Mandal and S. Mohanta, *Dalton Trans.*, 2014, **43**, 15737–15751.

64 S. R. Bayly, E. R. Humphrey, H. de Chair, C. G. Paredes, Z. R. Bell, J. C. Jeffery, J. A. McCleverty, M. D. Ward, F. Totti, D. Gatteschi, S. Courric, R. S. C. Barry, B. R. Steele and C. G. Serettas, *J. Chem. Soc., Dalton Trans.*, 2001, 1401–1414.

