1	Understanding Metal Synergy in Heterodinuclear Catalysts for the
2	Copolymerization of CO ₂ and Epoxides
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9	
10	Abstract
11	The copolymerization of carbon dioxide with epoxides is an industrially relevant means
12	to valorize wastes and improve sustainability in polymer manufacturing, and may also
13	provide an economic benefit to CO2 capture and storage technologies. The efficiency of
14	the process depends upon the catalyst used; previously Zn(II)Mg(II) heterodinuclear
15	catalysts showed good performances at low CO_2 pressures, which has been attributed to
16	synergic interactions between the metals. Here we report a Mg(II)Co(II) catalyst for the
17	production of polyols by copolymerization of CO_2 with cyclohexene oxide that exhibits
18	significantly better activity (turn-over-frequency over 12,000 h^{-1}), high CO ₂ utilization
19	(over 99 %) and high polymer selectivity (over 99 %). Detailed kinetic investigations
20	show a second-order rate law, independent of CO_2 pressure from 1 to 40 bar.
21	Investigations of the synergy between the metal centres showed that epoxide
22	coordination occurs at Mg(II) with reduced transition state entropy, which the carbonate
23	attack step is accelerated at Co(II) through lowering of the transition state enthalpy.
24	

1 Society needs more and better methods to transform CO₂ into products, both to obviate 2 industrial greenhouse gas emissions and to lock-away this recalcitrant molecule into 3 useful products.¹ CO₂ utilization can be an important means to increase product 4 sustainability and better understanding of its chemistry is needed to accelerate 5 technological developments.²⁻⁴ One promising option is its copolymerization with 6 epoxides to yield polycarbonates (for catalysts resulting in perfectly alternating 7 enchainment) or polyether carbonates (for catalysts resulting in less CO₂ uptake) (Fig. 8 1).⁵⁻⁸ Life cycle analysis demonstrates a triple win in terms of greenhouse gas emissions: 9 for every molecule of CO₂ used, two more are saved through replacing the use of the petrochemical (epoxide).⁹ Some polymerization catalysts have also shown good 10 11 compatibility for integration with large-scale CO₂ capture technologies and high tolerance towards common impurities found in gas streams.¹⁰ Given the value of the 12 13 polymeric products, CO₂ copolymerization could provide an economic stimulus for large-14 scale capture and storage technologies. This work focuses on the production of CO₂-15 derived polyols which are a class of polymers showing low molar mass (M_n < 5000 g mol⁻ 16 ¹) but which must be hydroxyl terminated. They show equivalent or better properties 17 than conventional polyether/ester polyols in the manufacturing of rigid and flexible foams, adhesives, elastomers and coatings.¹¹⁻¹⁴ Higher molar mass polycarbonates show 18 19 properties suitable to replace petrochemicals in sectors including packaging, coatings, 20 rigid plastics and medical materials.^{11,15-17} For any application sector, the selection of the 21 polymerization catalyst is central to process productivity and selectivity. This work 22 describes the development and understanding of a new type of highly active 23 heterodinuclear catalyst which exploits metal synergy.

1 Metal synergy is often summoned as the rationale for the high performances and 2 activities of bimetallic catalysts but detailed mechanistic insight and support for the 3 putative cooperative interactions are far less frequently presented.^{18,19} For example, 4 synergic interactions are invoked in mechanisms underpinning large-scale processes 5 such as polymerization, ammonia synthesis, methanol synthesis and Fischer-Tropsch 6 reactions as well as for organic transformations from C-H activations to redox processes 7 but so far detailed understanding of how to design catalysts to exploit or optimize 8 synergy is lacking.²⁰⁻²³ In the field of CO_2 /epoxide copolymerization, we have previously 9 reported a series of dinuclear Zn(II)/Mg(II) catalysts and proposed their superior performances arose from synergic interactions (Fig. 1).²⁴ 10 The nature of the metal 11 combination is clearly important as a series of Zn(II)/M(I/II/III) complexes, where M= Li, 12 Na, K, Ca, Al, Ga and In all showed inferior activity compared to the Zn(II)/Mg(II) 13 catalyst.^{25,26} This finding highlights the need for more detailed understanding and 14 suggests there may be a special role for Mg(II) in this catalysis: it motivated the 15 investigation of other heterodinuclear Mg(II)/M(II) complexes. We reasoned that 16 replacing Zn(II) with Co(II), in combination with Mg(II), might increase carbonate 17 nucleophilicity compared to Zn(II)/Mg(II) analogues and hence accelerate rates. There 18 is less precendent for Co(II) complexes in copolymerization catalysis, although we 19 previously observed that a di-Co(II) catalyst outperforms the di-zinc analogue.^{27,28} In the 20 mechanically related field of cyclic carbonate formation from epoxide/CO₂ coupling, 21 Co(II) complexes are also effective.²⁹⁻³¹ It's important to distinguish the hypothesis that 22 Mg/Co(II) might function synergically from the well-known Co(III) salen catalysts, which 23 show high activity but require co-catalysts and operate by more complex multicomponent mechanisms.³²⁻³⁵ 24

1 **Results and Discussion**

2 Catalyst Synthesis

3 Previous research into heterodinuclear catalysts revealed that the mixed metal 4 complexes are generally thermodynamically more stable than the homodinuclear 5 counterparts.²⁵ Capitalizing on this finding, in this work the **MgCo** heterodinuclear 6 complex was prepared by heating the ligand sequentially with the two metal precursors 7 (Fig. 2a). The **MgCo** complex was isolated as a pink powder in 78 % yield and was 8 characterized by elemental analysis and by MALDI-ToF mass spectrometry which 9 revealed a molecular cation peak [690 Da = $[LMgCo(OAc)]^+$](Supplementary Fig. 1). The 10 infra-red (IR) spectrum showed a series of frequencies, consistent with the molecular 11 structure (Supplementary Fig. 2).

12

13 To allow for proper comparison in catalysis, the homodinuclear analogues, **MgMg** and 14 **CoCo**, were prepared according to published procedures (see Supplementary 15 Information).^{27,36} A range of characterization techniques were used to assess the 16 compounds and to establish the formation of the MgCo catalyst. X-ray photoelectron 17 spectroscopy (XPS) was used to evaluate both the Co oxidation state and to confirm the sample composition (Supplementary Fig. 3). For both **CoCo** and **MgCo**, Co $2p_{1/2}$ and Co 18 19 $2p_{3/2}$ core levels are observed at binding energies (BE) of 780.2 eV and 796.2 eV, 20 respectively. In addition, satellite features at 3-5 eV higher BE are indicative of Co(II) as 21 the oxidation state (Fig. 2b). For **MgCo** it was feasible to compare the Mg 1s and Co $2p_{3/2}$ 22 core levels and to estimate a Mg:Co ratio of 1:1. Using SQUID measurements, the effective 23 magnetic moments (μ_{eff}) per Co(II) centre for **MgCo** and **CoCo** were 4.75 and 4.63 μ_{B} , 24 respectively (Supplementary Fig. 4). Both values are slightly higher than the spin-only 25 value ($\mu_{S0} = 3.87 \ \mu_B$ for S = 3/2) consistent with expected orbital contributions. Magnetic

1 saturation (M_{sat}) was observed at 1.94 and 2.15 µ_B for **MgCo** and **CoCo** (per Co(II) centre), 2 at 2K respectively (Supplementary Fig. 4). These values are consistent with reported 3 octahedral Co(II) centres showing a populated ground-state Kramers doublet.^{37,38} 4 Electrochemical measurements for MgCo, CoCo and MgMg were conducted using cyclic 5 voltammetry (Supplementary Fig. 5). **MgMg** shows a series of ligand centred oxidations 6 at potentials greater than 0.32 V (Supplementary Fig. 5). These processes are assigned to 7 ligand-based oxidations of the phenolate rings and are analogous to previous reports of related ligands.³⁹⁻⁴¹ For **CoCo**, two separate oxidations were observed at E_{pa} = -0.34 V 8 9 (Co(II/III)) and $E_{1/2} = -0.10$ V (Co(II/III)) (Supplementary Fig. 5). The **MgCo** complex shows just one irreversible Co(II/III) oxidation at $E_{pa} = -0.06$ V (Fig 2c). The clear 10 11 differences between the cyclic voltammograms support the formation of MgCo.

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MgCo, **CoCo** and **MgMg** were each tested in the ROCOP of CO₂/cyclohexene oxide (CHO) 13 14 at 1 bar CO₂ pressure (Table 1). The conditions for catalyst testing were optimized and 15 for fast and selective catalysts such as **MgCo**, it is important to establish that rates are 16 independent of stirring speed and that reactions are not under diffusion control 17 (Supplementary Table 2). A series of experiments were conducted in the absence of and with progressively greater quantities of 1,2-cyclohexane diol (CHD) as chain transfer 18 19 agent. These experiments confirm the feasibility of forming high molecular mass 20 poly(cyclohexene carbonate) (PCHC), with the expected bimodal molecular mass 21 distributions (Supplementary Table 2). On the other hand, the catalyst shows 22 outstanding tolerance to protic compounds and functions highly effectively using 20 23 equivalents of 1,2-cyclohexane diol (CHD). Under these conditions it forms exclusively α , 24 ω -hydroxyl telechelic PCHC, i.e. the desired polyol, and does so with a very high TOF 340 25 h⁻¹ (0.1 mol% **MgCo**, 80 °C, 1 bar pressure CO₂). To establish the catalyst remained thermally stable under catalytic conditions, MgCo was heated for 24 hours at 120 °C, in
 dioxane, after which both its ¹H NMR spectrum and performance remained identical
 (Supplementary Fig. 6).

4 Carbon Dioxide and Epoxide Copolymerization

5 The catalyst shows high activity and selectivity at loadings as low as 1:4000 or 0.025 6 mol% (catalyst:epoxide). It also shows excellent CO₂ selectivity (> 99%) across the range 7 of temperatures tested (80-120 °C), with no ether linkages observed by ¹H NMR 8 spectroscopy. Furthermore, all polymerizations resulted only in polymer formation (i.e 9 no cyclic carbonate) and well controlled, monomodal, polymer molar mass distributions. 10 The catalyst functions effectively in the presence of excess chain transfer agent (1:20 11 catalyst:diol) resulting in the formation of polycarbonate polyols (dihydroxyl telechelic 12 polymers). Generally the theoretically predicted molecular mass values and those 13 obtained by GPC analysis showed good agreement (e.g. Table entry 1, M_n theoretical = 14 3,300 g/mol and M_n experimental = 1700 g/mol).

15 Within the series of catalysts, **MgCo** is clearly the most highly active and increasing the 16 reaction temperature results in a significant rate enhancement from 455 h⁻¹ (80 °C) to 17 1205 h⁻¹ (120 °C) without any compromise in the quality of the polymer produced (Table 18 1). Comparing the activity of all three catalysts under identical conditions reveals that 19 **MgCo** (1205 h⁻¹) is significantly more active than either **CoCo** (712 h⁻¹) or **MgMg** (368 h⁻¹) 20 ¹). It is also >4 times faster than the previously most active heterodinuclear **ZnMg** catalyst 21 featuring the same ancillary ligand (Table 1).⁴² In comparison to other leading catalysts 22 in this field, **MgCo** shows a very high activity (Table 1, Supplementary Fig. 7 shows the 23 structures of the literature catalysts).^{6,7} For example, compared to a recently reported 24 first example of an In(III) catalyst, **MgCo** is thirty times more active.⁴³ It is three times 25 more active than a dizinc complex coordinated by a prolinol derivative,⁴⁴ and approximately twice the activity of a tetranuclear La₃Zn catalyst.⁴⁵ It is also qualitatively
 faster than optimized Co(III) catalysts featuring ligand tethered ionic co-catalysts,
 although direct comparisons may be hindered by the thermal instability of some Co(III)
 complexes.⁴⁶

5 **Polymerization Kinetics and Chain Shuttling Mechanism**

6 To better understand the enhanced activity, the catalytic rate law and rate-determining 7 step (RDS) were investigated. Polymerization conversion vs. time data were acquired 8 using *in-situ* infrared spectroscopy (Supplementary Fig. 8), by analysing the increase in 9 the absorption intensity corresponding to the formation of PCHC. A range of different **MgCo** concentrations were evaluated, from 0.43 – 1.67 mM, using a 3.33 M concentration 10 11 of epoxide (CHO) dissolved in diethyl carbonate (DEC) and by applying 1 bar CO_2 12 pressure, at 100 °C. Plots of epoxide concentration vs. normalized time, using the method 13 described by Bures,⁴⁷ showed the best fits for a first order dependence in catalyst 14 concentration (Fig. 2d). Alternative fits for higher or lower orders in catalyst 15 concentration were poor, substantiating the first order assignment (Supplementary Fig. 16 9). To determine the order in epoxide its concentration was varied from 1– 5 M, in DEC using a concentration of catalyst of 1.67 mM, at 1 bar CO₂ pressure, 100 °C. Semi-17 logarithmic plots of epoxide concentration vs. time (ln[CHO] vs. time) from 5 – 75 % 18 19 epoxide conversion (i.e. an integrated rate approach) showed linear fits indicative of a 20 first order dependence on epoxide concentration (Fig. 2e, Supplementary Fig. 10). The 21 dependence on carbon dioxide pressure was determined from 10 - 40 bar, using a 22 catalyst concentration of 1.67 mM, a CHO concentration of 3.33 M in DEC, at 100 °C (Fig. 2f). Plots of activity (TOF) vs. time, where activity is determined as the initial rate from 5 23 24 - 15 % monomer conversion, showed a zero order dependence on CO₂ pressure. The slight reduction in rate at higher CO₂ pressure is in line with previous observations and
 is probably due to sub-critical CO₂ gas expansion reducing the overall epoxide and
 catalyst concentrations.⁴⁸ Overall the kinetic data are consistent with the homodinuclear
 catalysts,^{42,49,50} and show a rate law dependent to the first order in catalyst and epoxide
 concentration and independent of carbon dioxide pressure (10-40 bar).

6 The rate law is rationalized by a chain shuttling mechanistic hypothesis,^{8,51} whereby the 7 rate limiting step involves metal carbonate attack at the second metal coordinated 8 epoxide. The polymer chain 'shuttles' between the two metals twice per complete cycle 9 of insertions, i.e. it changes the metal at which it is anionically coordinated (X-type ligand) 10 when the epoxide is inserted and again when CO_2 is inserted.⁴⁹ The catalyst features two 11 carboxylate groups but it is proposed that these have different roles in the cycle: one 12 group initiates polymerization but the other remains κ_2 coordinated at the metals.⁴⁹

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This bridging carboxylate ligand is proposed to facilitate chain shuttling by changing its
site of formal anionic coordination to counter-balance charge as the polymer chain moves
(Fig. 3a, Supplementary Fig. 11).⁴⁹

17 To investigate the chain shuttling process in more detail, the temperature dependence of the rate was investigated from 60 – 125 °C, at 1 bar CO₂ pressure (Supplementary Table 18 19 3). The rate of polymerization increases with temperature up to 100 °C, but at higher 20 temperatures polymer (PCHC) formation sharply decreases (Fig. 3b). Concurrently the 21 rate of trans-cyclic carbonate (1825 cm⁻¹)⁵² formation rapidly increases as observed by 22 IR spectroscopy. There is no observable cis-cyclic carbonate (1804 cm⁻¹)⁵² by IR 23 spectroscopy. The loss of selectivity at high temperatures may be rationalized by the 24 decreasing solubility of carbon dioxide (in epoxide), as predicted by Henry's Law.⁵³ As a 25 result, CO₂ insertion becomes rate-limiting and the catalytic resting state shifts from a

1 metal-carbonate to a metal-alkoxide intermediate. Under these conditions, forward 2 polymerization requires carbon dioxide and is limited by its low solubility. In contrast, 3 cyclic carbonate forms from the metal alkoxide species undergoing chain back-biting and 4 this process is independent of CO_2 . Thus, when using low CO_2 pressures, under static 5 conditions, and at high temperatures the reaction conditions favour the formation of 6 mixtures of polymer and cyclic carbonate causing the reaction selectivity to be 7 compromised. To overcome this limitation and to maintain the best rates of 8 polymerization, reactions were conducted at 20 bar CO₂ pressure so as to compensate for 9 high temperature diffusion and solubility limitations (Fig. 3c, Table 2). Under these 10 conditions, the rate of polymerization shows the expected exponential increase from 60 11 - 140 °C and the **MgCo** catalyst achieves an activity of 12,462 h⁻¹, whilst maintaining the highest polymer and carbon dioxide selectivity (> 99%). Under these conditions, the rate 12 13 limiting step remains as metal carbonate attack and the resting state is the metal-14 carbonate intermediate. Since this species does not undergo significant back-biting 15 reactions, the selectivity for polymer is maintained and negligible quantities of cyclic 16 carbonate by-product are formed.

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18 Catalysis Scope and Benchmarking

19 **MgCo** was also tested using a range of other common epoxides, using 0.05 mol% catalyst 20 concentration, 20 bar CO₂ pressure and 100 °C (Supplementary Table 8). Detailed 21 monomer scoping investigations are necessary in future but the preliminary findings 22 show good activity using cyclopentene oxide (TOF = 214 h⁻¹) and vinyl-cyclohexene oxide 23 (TOF = 2900 h⁻¹) (Supplementary Table 8). Poly(cyclopentene carbonate) is of interest in 24 the context of catalysed depolymerization, or chemical recycling, since it shows potential 25 to reform cyclopentene oxide.^{54,55} Vinyl-cyclohexene oxide is of interest for the potential to apply post-functionalization reactions at the alkene as a means to modify the polymer properties and achieve efficient cross-linking.⁵⁶⁻⁶⁰ **MgCo** shows no activity using biobased limonene oxide and, under these conditions, using propene oxide, it catalyzes the formation of propene carbonate (TOF = 5 h⁻¹) (Supplementary Table 8).

5 Objectively, the **MgCo** activity is very high and comparisons both against homodinuclear 6 combinations and other literature catalysts are warranted (Table 2). The most accurate 7 means to compare catalysts is to compare rate coefficients and this is most easily 8 accomplished for the series of complexes featuring the same ancillary ligand and different 9 metal combinations (Table 1).42 Since it's already been established that these 10 homodinuclear (ZnZn, MgMg, CoCo) and heterodinuclear (ZnMg) catalysts show the 11 same rate law, 27, 42, 49, 50 it is possible to properly compare rate coefficients (k_p) and 12 relative rates ($k_{\rm rel}$, against the **ZnZn** benchmark). At 80 °C, the **MgCo** complex shows a 13 relative rate which is 85 times greater than **ZnZn**. At 120 °C, the **MgCo** relative rate is 14 >1000 times greater than the **ZnZn** analogue, 5 times greater than **MgMg** and 15 approximately double the recently reported MgZn. The rate coefficients also confirm 16 that **MgCo** is significantly more active than either of the homodinuclear analogues (**MgMg** 17 or **CoCo**), for example at 120 °C **MgCo** shows double the rate of **CoCo** and four times the 18 rate of MgMg. Polymerization catalysis conducted using an equimolar mixture of MgMg 19 and **CoCo** showed an average rate for the two complexes and the value was three times 20 less than that for **MgCo** (Table 2, entry 8). This experiment underscores the importance 21 of isolation of the pure heterodinuclear complex and provides good evidence of a synergic 22 interaction between the Mg and Co(II) metals. It also shows there is not any appreciable 23 conversion of the homodinuclear complexes to the **MgCo** species under the conditions of 24 the catalysis.^{25,42}

1 Comparisons with literature catalysts, often tested using higher CO₂ pressure regimes, 2 are more complex since the rate laws and associated rate coefficients are rarely reported. 3 A recently reported homogeneous Fe(III) catalyst (see Supplementary Fig. 12 for catalyst 4 structure), in combination with an ammonium chloride co-catalyst, is around twenty 5 times slower than MgCo.⁶¹ Comparing point activity data (TOF values) reveals 6 qualitatively similar, or somewhat higher rates, for **MgCo** compared with the well-known 7 highly active Co(III)-salen/PPNX bicomponent catalyst systems or with trialkyl borane 8 and PPNX catalyst systems.62-64

9 Given the outstanding performance of the catalyst and that the previously most active systems comprised Co(III) complexes, it is relevant to understand whether any 10 11 appreciable Co(II) oxidation occurs during catalysis. It should be noted that such 12 oxidation is not anticipated from the previous Co(III) salen catalytic literature, which has 13 rather shown thermally induced reduction side-reactions to form inactive Co(II) 14 complexes at high-temperatures.⁶⁵ Notwithstanding this prior result, several attempts 15 were made to oxidise the MgCo(II) catalyst but all reactions were unsuccessful and 16 resulted in substantial complex decomposition (Supplementary Fig. 13). Cyclic 17 voltammetry experiments using MgCo(II) confirm the instability of the MgCo(III) 18 intermediate, with no clear reduction potential being observed (Supplementary Fig. 5). 19 In contrast, the **Co(II)Co(II)** analogue shows two clear oxidations (to Co(II)Co(III) and 20 Co(III)Co(III), respectively) and the concomitant two reduction reactions. The findings 21 using **CoCo** confirm that the heterodinuclear complex **MgCo(II)** is expected to be stable 22 with respect to oxidation and that there is not expected to be any substantial formation 23 of MgCo(III) species during catalysis.

24

25 Heterodinuclear Synergy

1 To gain further insight into the metal-metal synergy, the transition state Gibbs free 2 energy (ΔG^{\ddagger}) was determined by an analysis of the temperature dependence of the rate 3 coefficient (Fig. 4a). Experiments were conducted using an initial concentration of 4 epoxide of 3.33 M in DEC, with a catalyst concentration of 1.67 mM, under 20 bar CO₂ 5 pressure and temperatures were varied in 20 °C increments from 60 - 120 °C 6 (Supplementary Fig. 14, Supplementary Tables 4-6). Plots of $\ln(k/T)$ vs. 1/T enabled 7 determine of both the transition state enthalpy (ΔH^{\ddagger}) and entropy (ΔS^{\ddagger}) (Supplementary 8 Fig. 15, Supplementary Table 7). The transition state Gibbs free energy (ΔG^{\ddagger} , T = 373 K) 9 values are 94.5 ± 1.2, 97.3 ± 1.5 and 100.2 ± 1.3 kJ mol⁻¹ for MgCo, CoCo and MgMg, 10 respectively. Over the series of catalysts, the transition state barrier decreases by 5.7 kJ 11 mol⁻¹ which correlates well with the observed 8-fold increase in rate for the 12 heterodinuclear catalyst **MgCo**. The analogous dizinc catalyst (**ZnZn**) shows $\Delta G^{\ddagger} = 107$ kJ 13 mol⁻¹ (T = 373 K)^{49,50} and accordingly **MgCo** is ~85-times faster; these findings clearly 14 demonstrate the benefit of targeting the right metal combinations and synergies. The 15 transition state entropy (ΔS^{\ddagger}) is significantly reduced for catalysts containing Mg(II), with 16 values for **MgMg** and **MgCo** catalysts at -45.4 ± 3.7 and -46.1 ± 3.4 J mol⁻¹, respectively, 17 compared to the **CoCo** analogue, -60.2 ± 4.2 J mol⁻¹. (Fig. 4b, Supplementary Table 7). The 18 entropic benefit of using Mg(II) may arise from its low bond directionality which may 19 increase the degrees of freedom (rotational and/or conformational) associated with 20 epoxide coordination. It could also relate to the higher oxophilicity of magnesium (θ = 0.6) 21 compared to cobalt (θ = 0.4), as quantified in a recent evaluation of oxophilicity values 22 across the periodic table.⁶⁶ The transition state thermodynamic data also show that 23 catalysts containing Co(II) show reduced enthalpy barriers, e.g. $\Delta H^{\ddagger} = 77.3 \pm 1.2$ kJ mol⁻¹ 24 for **MgCo** vs. 83.3 \pm 1.3 kJ mol⁻¹ for **MgMg**, T = 373 K). This finding is in-line with the 25 Co(II)-carbonate being significantly more nucleophilic than its Mg(II) counterpart. Taken

1 together these experimental data can be interpreted as rationalizing the synergy of the 2 **MgCo** complex since it combines the favourable entropy of epoxide coordination at 3 Mg(II), with a highly nucleophilic Co(II)-carbonate (Fig. 4d, Supplementary Fig. 11). The 4 high lability of Co(II)-carbonates was previously observed in literature describing the 5 formation of cyclic carbonates.^{29,30} The ability to replace a Co(II) centre with Mg(II) to 6 accelerate activity is both fundamentally interesting but also practically useful due to its 7 abundance, light-weight and lack of toxicity.⁶⁷ Overall, the enhanced performance for the 8 heterodinuclear MgCo catalyst unambiguously arises from synergic interaction between 9 the metals and kinetic analysis signals that the barrier to the rate limiting step is reduced 10 by using the Mg(II) to carry out epoxide coordination and the Co(II) centre to provide the 11 reactive (nucleophilic) carbonate group to attack and ring-open the epoxide 12 (Supplementary Fig. 11).

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These results provide a new strategy for the design, preparation and understanding of
highly efficient, selective, stable and inexpensive catalysts for CO₂ copolymerization.

16 The kinetic and mechanistic findings clearly signal some future directions to improve 17 other catalysts' performances and to design new catalysts for these processes. Most 18 clearly, there is a route to improve existing dinucleating -diiminate and salen catalysts 19 by targeting heterodinuclear complexes, exploiting the thermodynamic stability 20 demonstrated in this work and by using Mg(II) in place of Zn(II)/Co(II/III)/Cr(III).⁶⁸⁻⁷⁰ 21 Such heterodinuclear MgCo(II/III), MgCr(II/III) complexes would have the added benefit 22 of replacing 50% of the expensive, coloured and heavy transition metal with Mg(II). There 23 is also potential to exploit main-group/transition metal synergy in other CO_2 utilizations, 24 for example di-Co(II) complexes are highly active photoredox catalysts for transforming 25 CO_2 into CO_2^{71-73} very recently a Co(II)/Zn(II) complex showed yet higher activity.⁷¹ CO_2 terpolymerizations and switchable catalytic processes are currently limited by low catalyst activities and so better catalysts should allow access to a broader range of CO₂ containing materials and properties.⁷⁴⁻⁷⁸ Beyond the field of CO₂ utilization, there is increasing interest in main group/Co(II) heterodinuclear catalysts, relevant to those explored in this work, for processes including nitrogen activation,⁷⁹⁻⁸¹ electrophilic amination,⁸² CH and CF activation processes.^{83,84}

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8 **Conclusions**

9 A synergic and highly active heterodinuclear MgCo complex for epoxide/ CO_2 copolymerization was synthesized in high yield by a one-pot, thermodynamically 10 11 controlled reaction. It was characterized using a range of techniques, including XPS, mass 12 spectrometry, magnetometry and cyclic voltammetry. The **MgCo** catalyst achieves very 13 high activity at either 1 bar CO_2 pressure (TOF = 1205 h⁻¹) or at 20 bar pressure (TOF = 14 12,400 h⁻¹) and produces perfectly alternating copolymer for CO₂/CHO coupling and was 15 found active for other epoxide monomers including vCHO, CPO and PO (Supplementary 16 Table 8). Detailed kinetic studies reveal the synergy arises because the magnesium centre 17 enhances the transition state entropy, through epoxide coordination, and cobalt reduces 18 the transition state enthalpy, by the greater nucleophilicity of the cobalt-carbonate. This 19 catalyst highlights the potential for heterodinuclear synergy and underscores the 20 importance of metal selection according to its specific role in the cycle. The findings are 21 expected to be broadly applicable to other homodinuclear polymerization and switchable 22 catalysts. Generally, the work provides a rationale and understanding of how to exploit 23 synergic interactions in homogeneous catalysis.

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5

6 **Author Contributions**

- 7 AD and CW conceived the project. AD designed and performed the synthetic experiments.
- 8 SK designed and performed the CV and SQUID experiments. AR designed and performed
- 9 the XPS study. AD and CW prepared the manuscript.

10 **Conflict of Interest**

11 CW is a director of econic technologies.

12 Data Availability

- 13 All the data supporting the findings of this study are available within the paper and its
- 14 supplementary information files or from the corresponding author on request

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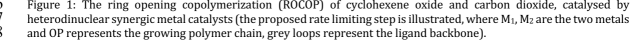
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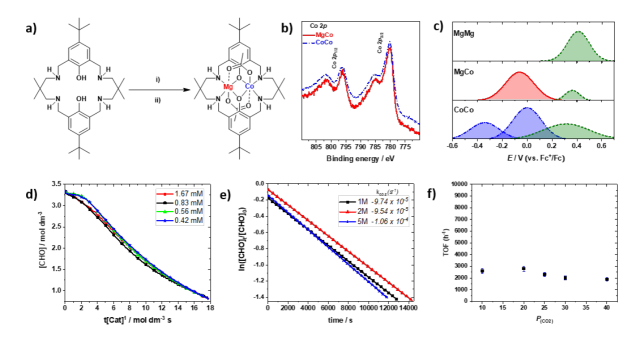


Figure 2: Preparation, characterization and polymerization kinetics of the **MgCo** catalyst. a) Illustration of the route to prepare the heterodinuclear **MgCo** catalyst. Reagents and conditions: i) Mg{N(Si(CH₃)₃)₂}₂, THF, 25 °C, 1 h. ii) Co(OAc)₂, THF, 100 °C, 16 h, 78 %. b) X-ray Photoelectron spectroscopy reveals different binding energies (eV) for the $2p_{1/2}$ and $2p_{3/2}$ orbitals of **MgCo** (—) and **CoCo** (—·—) (for further information see Supplementary Fig. 3). c) Cyclic Voltammetry is used to show different redox potentials (E / V) for **MgMg**, **MgCo** and **CoCo**. The data are obtained vs. Ferrocium⁺/Ferrocene, in THF, 0.1 M [ⁿBu₄N][PF₆] and at 100 mV s⁻¹ (for full cyclic voltammograms see Supplementary Fig. 5). d) Kinetic data using **MgCo** catalyst for the ring-opening copolymerization of cyclohexene oxide and carbon dioxide (CHO/CO₂ ROCOP). The first order dependence on catalyst concentration is determined from the linear plots of [CHO] vs. t[cat]^x, x = 1. e) The order in epoxide concentration is determined from the linear fit to plots of ln[CHO] vs. time. f) The order in carbon dioxide pressure is determined from plots of initial rate (h⁻¹) vs. pressure of CO₂ (bar), with error bars, from duplicate runs, marked in blue.

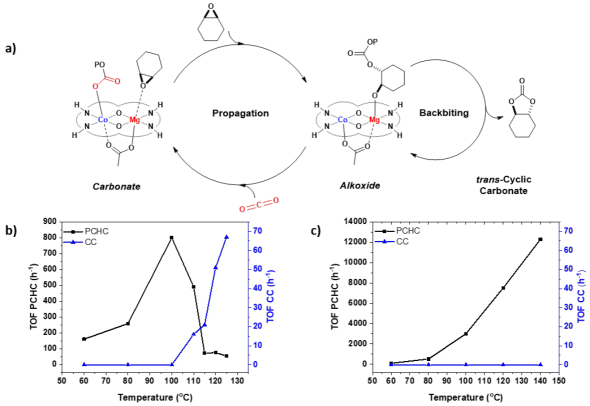


Figure 3: The Chain Shuttling Mechanism for the copolymerization (CHO/CO₂ ROCOP) using **MgCo**. a) Illustration of the Chain Shuttling propagation mechanism, showing the formation of the polycarbonate (PCHC) and side-reactions which occur at higher temperature (low carbon dioxide pressure) to form trans-cyclic carbonate (CC). b) and c) Show the relationship between the catalyst activity (h^{-1}) and temperature (°C) for the formation of polymer (PCHC \blacksquare) and cyclic carbonate (CC \blacktriangle) at 1 bar pressure of carbon dioxide (b) and the same data but determined at 20 bar CO₂ pressure(c).

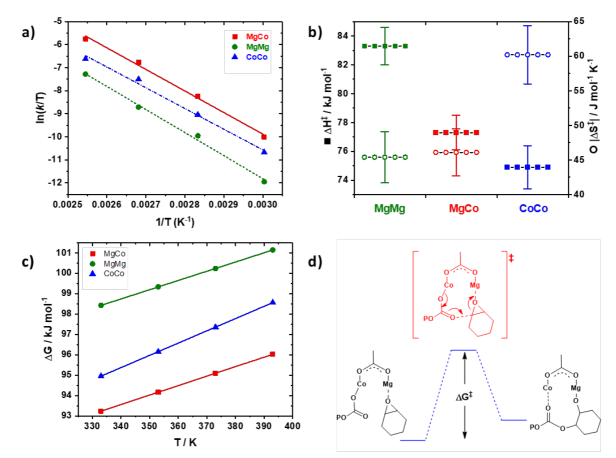


Figure 4: Data providing insight into the factors governing heterodinuclear synergy in polymerization catalysis using kinetic data to compare the MgCo heterodinuclear catalyst with homodinuclear analogues MgMg and CoCo. a) Van't Hoff plots of $\ln(k/T)$ vs. 1/T (K⁻¹) for MgMg (\bullet) CoCo (\blacktriangle) and MgCo (\blacksquare) over the temperature range 60 – 120 °C, under 20 bar CO₂ pressure. b) The kinetic data allow determination of the transition state enthalpy values, ΔH^{\ddagger} (\blacksquare , with errors ± 1.3), and entropy values, ΔS^{\ddagger} (**O**, with errors ± 3.7), for **MgMg**, **CoCo** and **MgCo**. c) The plot shows the variation in the overall transition state Gibbs Free Energy (ΔG[‡]) vs. temperature (K) for MgMg (•, ±1.3) CoCo (•, ±1.5) and MgCo (•, ±1.2). The errors are determined using Least Squares Fitting Analysis. d) Illustrates for the rate determining step occurring in the Chain Shuttling Mechanism with the transition state Gibbs Free Energy (ΔG^{\ddagger}) marked (Supplementary 10 Fig. 11 illustrates the reactions occurring during initiation, propagation and termination).

List of Tables:

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15 Table 1: Shows data for the ring-opening copolymerization (ROCOP) of carbon dioxide/cyclohexene oxide (CO₂/CHO)

16 at 1 bar CO₂, using the heterodinuclear catalyst MgCo and compared with homodinuclear catalysts, CoCo and MgMg.^a

#	Catalyst	Т	Time	CO ₂ ^b : Polym.	TON ^d	TOF	M_{n} [Đ] ^f
		(°C)	(min)	(%) ^c		(h-1)e	(kg mol ⁻¹)
1	MgCo	80	60	> 99: > 99	455	455 (±15)	1.7 [1.13]
2	MgCo	100	40	> 99: > 99	465	699 (±24)	1.6 [1.15]
3	MgCo	120	25	> 99 : 99	502	1205 (±41)	2.1 [1.24]
4	СоСо	120	60	> 99 : 96	712	712 (±24)	2.5 [1.20]
5	MgMg	120	60	> 99 : > 99	368	368 (±13)	1.8 [1.16]

6	MgZn <i>^{g, 42}</i>	80	360	> 99 : > 99	438	98	12.7[1.04] 5.1 [1.16]
7	LIn(O ^t Bu) ^{g, 43}	80	2880	> 99 : 95	350	15	3.4 [1.32]
8	$L_2 Z n_2 {}^{g, 44}$	80	1440	> 99: 98	1684	149	2.7 [1.28]
9	LZn₃Ce(OAc) ₃ ^{<i>h, 45</i>}	100	180	> 99 :> 99	900	300	15.0[1.20]
10	LCo(X) ^{<i>i</i>, 46}	50	300	> 99 : > 99	1315	263	48.2[1.12]

^aCopolymerization conditions: 0.05 mol % cat loading, 20 equiv. 1,2-cyclohexane diol, 1 bar CO₂, CHO neat (9.99 M). ^bExpressed as a percentage of CO₂ uptake vs the theoretical maximum (100 %), determined by comparison of the relative integrals of the ¹H NMR resonances due to carbonate (δ 4.65 ppm) and ether (δ 3.45 ppm) linkages in the polymer backbone. ^cExpressed as a percentage of polymer formation vs. the theoretical maximum (100 %), determined by comparison of the relative integrals of the ¹H NMR resonances due to carbonate (δ 4.65 ppm) and ether (δ 3.45 ppm) linkages in the polymer backbone. ^cExpressed as a percentage of polymer formation vs. the theoretical maximum (100 %), determined by comparison of the relative integrals of the ¹H NMR proton resonances due to polymer (4.65 ppm), cis-cyclic carbonate (4.68 ppm) and trans-cyclic carbonate (4.00 ppm). ^dTurnover number (TON) = number of moles of cyclohexene oxide consumed / number of moles of catalyst. ^cTurnover frequency (ToF) = TON / Time (h). ^TDetermined by SEC, in THF, using narrow-M_n polystyrene standards as the calibrant; dispersity is given in brackets.^g These literature catalysts were tested at 0.1 mol % cat, 1 bar CO₂ for more details see references.⁴²⁻⁴⁴ h This literature catalysts was tested at 0.05 mol % cat, 1 bar CO₂.⁴⁶ For the chemical structures of all the literature catalysts see Supplementary Fig. 7.

9 10 Table 2: 5 11 and com

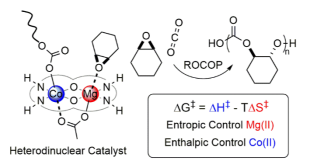
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Table 2: Shows the data for the ROCOP of CO_2/CHO under 20 bar pressure (CO_2) and variable temperatures using **MgCo** and compared with homodinuclear analogues and other high-performance catalysts from the literature.^a

		Т	Doly	TOF	$l_{\rm r}$ (v.1.03)	
#	Cat.		Poly.	TOF	$k_p({ m x10^3})$	$k_{ m rel}{}^{ m e}$
		(°C)	(%) ^b	(h-1)c	(dm ³ mol ⁻¹ s ⁻¹) ^d	MIEI
1	MgCo	60	> 99	80 (±2)	15.1 (±0.5)	14
2	MgCo	80	> 99	510 (±15)	93.1 (±2.8)	85
3	MgCo	100	> 99	3200 (±96)	428.9 (±12.9)	390
4	MgCo	120	> 99	7200 (±216)	1231.2 (±36.9)	1120
5	MgCo	140	> 99	12460 (±374)	1784.4 (±53.5)	1622
6	MgMg	120	> 99	1060 (±11)	269.7 (±8.1)	245
7	СоСо	120	> 99	4200 (±126)	559.2 (±16.8)	508
8	CoCo:MgMg (50:50)	120	> 99	2400 (±72)	405.0 (±12.2)	368
9	ZnZn	80	92	20	11.0	1
10	MgZn <i>f</i> ,42	120	> 99	2400	514.0	467
11	BEt ₃ /PPNCl g, 62	80	> 99	600	-	-
12	CrSalen, PPNCl h,63	80	> 99	1153	-	-
13	LCo ₂ X, PPNX ^{i, 64}	25	> 99	1356	-	-
14	LCo₂X <i>i</i> , 64	25	> 99	200	-	-
15	LFeCl, ⁿ Bu ₄ NCl ^{j,61}	80	> 99	400	0.0056	5

^aCopolymerization conditions: cat : CHO 0.05 mol %, 20 eq. CHD, 20 bar CO₂. All entries report >99 CO₂ selectivity vs. theoretical maximum (100 %), determined by comparison of the relative integrals of the ¹H NMR resonances due to carbonate (δ 4.65 ppm) and ether (δ 3.45 ppm) linkages in the polymer backbone δ 4.65 ppm) and ether (δ 3.45 ppm) linkages in the polymer backbone. Entries 1-8 gone to full conversion (>90 %, >1800 TON) forming polycarbonate polyols of molecular weight 1-3 kg mol⁻¹. ^bExpressed as a percentage of polymer formation vs. the theoretical maximum (100 %), determined by comparison of the relative integrals of the ¹H NMR proton resonances due to polymer (4.65 ppm), cis-cyclic carbonate (4.68 ppm) and trans-cyclic carbonate (4.00 ppm). ^cTurnover frequency (TOF) = TON / Time (h) (measured between 5-20 % conversion). ^dk_p = k_{obs} / [cat]¹, k_{obs} determined from the gradient of ln([CHO]₁/[CHO]₀) vs. time plot [cat] = 1.67 mM, over the conversion range 5-75 %. ^eRate constant relative to **ZnZn** at 80 °C, k_{rel} = k/0.0011. ¹0.1 mol% cat, 20 bar CO₂. ¹0.1 mol% cat, 0.1mol% ^eRu₄N, 10 bar CO₂. For structures of literature catalysts see Supplementary Fig. 12.

Graphic Abstract/TOC



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- 2 \quad The copolymerization of CO_2 with epoxides is an attractive approach for valorizing waste
- 3 products and improving sustainability in polymer manufacturing. Now, a heterodinuclear
- 4 Mg(II)Co(II) complex has been show to act as a highly active and selective catalyst for this 5 reaction at low CO₂ pressure. The synergy between the two metals was investigated using
- 6 polymerization kinetics.