# THERMAL DECOMPOSITION OF RHOMBOHEDRAL DOUBLE CARBONATES OF DOLOMITE TYPE

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DTA and TG studies in air were carried out for hydrothermally prepared rhombohedral double carbonates of dolomite type, CaMg(CO<sub>3</sub>)<sub>2</sub>, CaMn(CO<sub>3</sub>)<sub>2</sub>, CdMg(CO<sub>3</sub>)<sub>2</sub>, CdMg(CO<sub>3</sub>)<sub>2</sub>, CdMn(CO<sub>3</sub>)<sub>2</sub> and CdZn(CO<sub>3</sub>)<sub>2</sub>. The solid decomposition products in air have been compared to those obtained under hydrothermal conditions with CO<sub>2</sub> pressure. The dolomite [CaMg(CO<sub>3</sub>)<sub>2</sub>] decomposes in two stages both in air as well as under high CO<sub>2</sub> pressure. The other carbonates studied, follow a single stage decomposition in air and a two stage decomposition under hydrothermal condition. In air, the manganese containing carbonates CaMn(CO<sub>3</sub>)<sub>2</sub> and CdMn(CO<sub>3</sub>)<sub>2</sub>, decompose to form mixed oxides of CaMnO<sub>3</sub> and CdMnO<sub>3</sub> respectively, while CdMg(CO<sub>3</sub>)<sub>2</sub> and CdZn(CO<sub>3</sub>)<sub>2</sub> decompose to their respective two mono oxides.

Keywords: dolomit, rhombohedral double carbonates

### Introduction

Thermal decomposition of dolomite, CaMg(CO<sub>3</sub>)<sub>2</sub>, is apparently straight forward but, has still been studied by several workers because of considerable controversy with respect to the decomposition mechanism of dolomite in air as well as CO<sub>2</sub> atmosphere [1–4]. The studies were reviewed in detail by Otsuka [5] and again re-studied by Engler et al. [6]. The latter used non-isothermal in situ XRD analysis of dolomite decomposition both in air and CO<sub>2</sub>. They proposed a single stage decomposition for dolomite in air indicated by a single asymmetric peak on the DTA curve. But even in this single stage of decomposition, they found the reaction in the initial stage to be:

 $2CaMg(CO_3)_2 \rightarrow CaCO_3 + CaO + 2MgO + 3CO_2$ 

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The CaCO<sub>3</sub> which crystallized along with CaO+MgO in early stages began to decompose at slightly higher temperature during the late stage of the single decomposition peak. In CO<sub>2</sub> atmosphere, two stage decomposition was clear with two endothermic peaks pertaining to reactions;

respectively.

We know now that there are a few more rhombohedral double carbonates which are comparable to dolomite. The natural ones are Kutnahorite  $(CaMn(CO_3)_2)$  and Minrecordite  $(CaZn(CO_3)_2)$ . The synthetic analogues are  $CdMg(CO_3)_2$ ,  $CdMn(CO_3)_2$  and  $CdZn(CO_3)_2$ . Although other combinations of Ca, Cd with transition metals Fe, Ni, Cu, Co are considered theoretically possible, could not hitherto been synthesized. The authors have been studying the hydrothermal decomposition of rhombohedral double carbonates on a P-T plane under  $P_{CO_2}$  of upto 3 kbar in order to determine the equilibrium decomposition curve for these carbonates and use them for retrieval of the thermodynamic data for the double carbonates. The double carbonates so obtained in the hydrothermal studies were used for the decomposition studies in air. In this paper we intend to report the DTA and TG studies in air on the five synthetically prepared dolomite and dolomite type carbonates and compare their decomposition characteristics to those obtained under high  $CO_2$  pressure.

# Experimental

Five synthetic rhombohedral double carbonates namely  $CaMg(CO_3)_2$  (dolomite),  $CaMn(CO_3)_2$  (Kutnahorite),  $CdMg(CO_3)_2$ ,  $CdMn(CO_3)_2$ ,  $CdMn(CO_3)_2$ ,  $CdMn(CO_3)_2$ , are been used for the decomposition studies. The Minrecordite  $CaZn(CO_3)_2$ , could not be synthesized. While the complete phase diagrams for these double carbonates on  $P_{CO_2}$ —T plane will be published elsewhere, we give here the pressures and temperatures of syntheses of these carbonates (Table 1). All syntheses have been done in cold seal Tuttle reactors. The starting materials were the two carbonates (for instance,  $CaCO_3$ + $MgCO_3$  in 1:1 mole ratios for dolomite synthesis) charged in crimped platinum capsules and placed in these reactors. The pressurising gas was dry  $CO_2$ , pumped into the vessels which were externally heated.

The ratio of Ca and Cd and the other combining cations have been determined by measuring the  $d_{104}$  spacing which shows a linear correlation [7] for solid solutions in rhombohedral carbonates. The products which were confirmed by X-ray powder diffraction as double carbonate were used for DTA. The simultaneous

DTA and TG unit Netzsch Gerätebau STA 409 (Pt 10 Rh-Pt thermocouple and alumina sample holder) was used. The heating rate was 10 deg min<sup>-1</sup> and the temperature limit was 1000°C. In each sample after a first run of DTA over the entire temperature range, it was repeated again with a fresh sample in order to arrest the reaction at different stages and identify the products at different stages of heating.

Table 1 Experimental conditions for synthesis of rhombohedral double carbonates under CO2 pressure

Double	Temperature /	Pressure /	Duration /	d <sub>104</sub>
Carbonates	°C	bar	h	of the
M'M"(CO <sub>3</sub> ) <sub>2</sub>				product
CaMg(CO <sub>3</sub> ) <sub>2</sub>	650	1000	77	2.875
CaMn(CO <sub>3</sub> ) <sub>2</sub>	700	1240	68	2.940
CdMg(CO <sub>3</sub> ) <sub>2</sub>	650	1500	68	2.855
CdMn(CO <sub>3</sub> ) <sub>2</sub>	550	1100	72	2.892
$CdZn(CO_3)_2$	400	2000	68	2.848

#### Results and discussion

Table 2 gives the details of DTA and TG runs for the five rhombohedral double carbonates. The corresponding DTA-TG traces are given in Fig. 1. An examination of these peaks show both single stage and two stage decomposition. though all the runs were done in air. The decomposition stages and the products were also not the same in all these carbonates. Earlier workers [6] while studying the natural dolomite sample with composition Ca<sub>1.0</sub>, Mg<sub>0.9</sub>, Fe<sub>0.1</sub>(CO<sub>3</sub>)<sub>2</sub> reported that the two stage decomposition existed only in the presence of CO2 atmosphere which otherwise was a single stage decomposition. But in our synthetic samples with ordered dolomite structure, the two stages are distinct. The first stage CaCO<sub>3</sub>(ss) forms along with MgO evolving CO<sub>2</sub>. A substitution of upto 10% of Mg was noticed in CaCO<sub>3</sub>. The second stage of decomposition involved the dissociation of CaCO<sub>3</sub> to CaO and CO<sub>2</sub>. In all the rest of the double carbonates studied in air we found a single stage decomposition irrespective of the product, except in the case of CdMn(CO<sub>3</sub>)<sub>2</sub> where a minor second endothermic peak occurs but with negligible weight loss. Both CdMg(CO<sub>3</sub>)<sub>2</sub>, and CdZn(CO<sub>3</sub>)<sub>2</sub>, directly decompose to their two monoxides, but the Mn containing carbonates CaMn(CO<sub>3</sub>)<sub>2</sub>, and CdMn(CO<sub>3</sub>)<sub>2</sub>, result in CaMnO<sub>3</sub> and CdMnO<sub>3</sub> as the final products of decomposition. While in Kutnahorite the CaMnO<sub>3</sub> phase was directly formed as a result of the single decomposition reaction, in the CdMn(CO<sub>3</sub>)<sub>2</sub> the formation of CdMnO<sub>3</sub> was through an intermediate stage where CdO+Cd<sub>2</sub>Mn<sub>3</sub>O<sub>8</sub> crystallized, which on further heating react to form CdMnO3 at higher tempera-

Double	Peak I	ak I	Pea	Peak II	Total	Theor.	
carbonate	Temp./	Weight	Temp. / Weight Temp. / Weight	Weight	weight	weight	Decomposition reactions
$M'M"(CO_3)_2$	ွ	%/ ssol	၁့	loss /% oC loss /% loss /%	%/ ssol	%/ ssol	
CaMg(CO3)2	735	20.8	942	25.5	46.3	47.7	$CaMg(CO_3)_2 \rightarrow CaCO_3(ss) + MgO + CO_2$
							$\rightarrow$ CaO+ MgO+ 2CO <sub>2</sub>
CaMn(CO3)2	862	33.0	ı	1	33.0	38.1	$C_aMn(CO_3)_2 + O_2 \rightarrow C_aMnO_3 + 2CO_2$
CdMg(CO3)2	630	32.3	1	1	32.3	34.2	$CdMg(CO_3)_2 \rightarrow CdO + MgO + 2CO_2$
CdMn(CO3)2	595	24.1	837	1.6	25.7	29.0	$3CdMn(CO_3)_2 + O_2 \rightarrow CdO + Cd_2Mn_3O_8 + 6CO_2$
							$\rightarrow$ 3CdMnO <sub>3</sub> + 6CO <sub>2</sub>
$CdZn(CO_3)_2$	520	29.3	1	1	29.3	29.5	$CdZn(CO_3)_2 \rightarrow CdO + ZnO + 2CO_2$

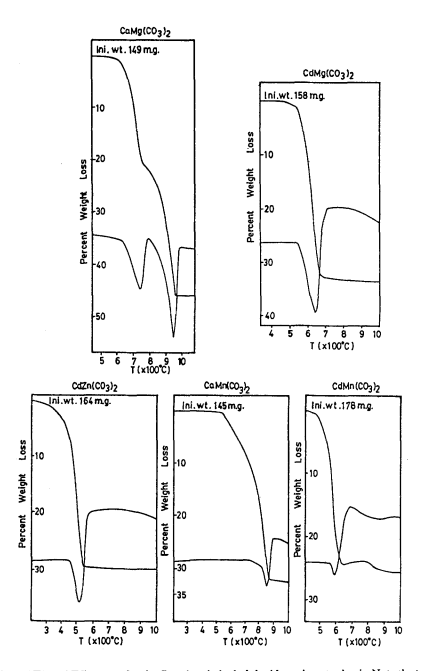


Fig. 1 DTA and TG curves for the five rhombohedral double carbonates in air. Note the two stage decomposition in CaMg(CO<sub>3</sub>)<sub>2</sub>

ture. A small second endothermic peak with minor weight loss (1.6%) observed in the CdMn(CO<sub>3</sub>)<sub>2</sub> decomposition may be due to some non detectable phase remaining but has to be ignored since this could not be reproduced. The theoretical weight loss along with the observed weight loss from TG curves is also given in Table 2. The weight losses agree well except in the Mn containing carbonates, where some complexity exists in the mechanism of decomposition. The formation of CaMnO phase by decomposition of (CaMn)CO<sub>3</sub> under vacuum was reported earlier [8], wherein the entire range of solid solution phases of CaO–MnO were prepared. In order to compare the decomposition in air with those under CO<sub>2</sub> pressure, selected experimental run products at 1 kbar CO<sub>2</sub> pressure are given in (Table 3) along with the temperature of decomposition of double carbonates.

Table 3 Equilibrium decomposition temperatures and pressures of rhombohedral double carbonates under CO<sub>2</sub> pressure of 1000 bar

Double	Pressure	Temperature of	Solid (1st stage)
Carbonates	of CO <sub>2</sub> /	decomposition /	decomposition
	bar	°C	product
CaMg(CO <sub>3</sub> ) <sub>2</sub>	1000	825	CaCO <sub>3</sub> (ss)+ MgO
CaMn(CO <sub>3</sub> ) <sub>2</sub>	1000	730	CaCO <sub>3</sub> + MnO
CdMg(CO <sub>3</sub> ) <sub>2</sub>	1000	715	MgCO <sub>3</sub> + CdO
CdMn(CO <sub>3</sub> ) <sub>2</sub>	1000	600	MnCO <sub>3</sub> + CdO
CdZn(CO <sub>3</sub> ) <sub>2</sub>	1000	420	$CdCO_3 + ZnO$

Note: The products shown in the last column are only the decomposition product, though at equilibrium, the reactant (double carbonate) is also present.

It is interesting to note that the single stage decomposition of dolomite in air reported by earlier workers [6], was not observed with synthetic ordered dolomite presently studied. The decomposition product has been identical in both air and CO<sub>2</sub> for dolomite. But for other rhombohedral carbonates, the presence of CO<sub>2</sub> made a significant difference. The low oxygen fugacity intrinsically generated in steel reactors retained manganese in divalent state and hence the formation of CaMnO<sub>3</sub> and CdMnO<sub>3</sub> as products noticed when their corresponding carbonates decompose in air was not noticed under CO2 pressures. Though the higher stages of decomposition in the carbonates under CO<sub>2</sub> pressure were not investigated because of limitations of instrument tolerance for high temperature, it is quite clear that the distinct stages will appear at different temperatures until all carbonate components decompose to their respective oxides. Thus under CO2 pressure, a single stage decomposition is absent for all dolomite type double carbonates. Secondly mixed oxides such as CaMnO3 and CdMnO3 do not form when the decomposition is under CO<sub>2</sub> pressure. Finally the decomposition of dolomite is in two stages, even in air, unlike the previous report.

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Zusammenfassung — Mittels DTA und TG in Luft werden die hydrothermisch hergestellten rhomboedrischen Doppelkarbonate (Dolomittyp) CaMg(CO<sub>3</sub>)<sub>2</sub>, CaMn(CO<sub>3</sub>)<sub>2</sub>, CdMg(CO<sub>3</sub>)<sub>2</sub>, CdMn(CO<sub>3</sub>)<sub>2</sub> und CdZn(CO<sub>3</sub>)<sub>2</sub> untersucht. Die in Luft erhaltenen festen Zersetzungsprodukte wurden mit denen verglichen, die unter hydrothermischen Bedingungen mit CO<sub>2</sub>-Druck entstehen. Dolomit zersetzt sich sowohl in Luft als auch unter hohem CO<sub>2</sub>-Druck in zwei Schritten. Die übrigen untersuchten Karbonate zersetzen sich in Luft in einem, unter hydrothermischen Bedingungen in zwei Schritten. In Luft zersetzen sich die magnesiumhaltigen Karbonate CaMn(CO<sub>3</sub>)<sub>2</sub> und CdMn(CO<sub>3</sub>)<sub>2</sub> unter Bildung der Mischoxide CaMnO<sub>3</sub> und CdMnO<sub>3</sub>, während aus CdMg(CO<sub>3</sub>)<sub>2</sub> und CdZn(CO<sub>3</sub>)<sub>2</sub> jeweils die entsprechenden beiden Monoxide entstehen.