Fabrication process parameters significantly affect the perovskite oxygen carriers materials (OCM) performance in chemical looping with oxygen uncoupling (CLOU)

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Abstract
The CLOU performance of the CaTi₄Mn₀.₉₋ₓMg₀.₁O₃ (CMTM) perovskite-type system was investigated, comparing materials produced at laboratory scale with those made at industrial ton scale. The CLOU and conversion performances were studied by a micropacked bed reactor, and crystalline phase structure and homogeneity and bulk density identified as the most important parameters affecting the performance of the OCM. Bulk density is correlated with the sintering temperature, atmosphere and time at sintering temperature, while phase homogeneity is a function of the raw materials chosen, agglomeration method and sintering procedure. Specific challenges are identified in the control of slurry homogeneity and sintering conditions in upscaled production. The degree of sintering affects the chemo-mechanical properties of the material (crushing strength and attrition index), physical properties (specific surface area), and more importantly the crystalline phases formed and their homogeneity: the quantity of “active” crystalline phases present directly determines the thermochemical conversion properties (i.e., CLOU capacity and methane conversion), oxygen transfer capacity and kinetics.

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Graphic abstract
The fabrication parameters of the otherwise same ingredients result in quite different morphology and quality of performance in large scale.

Keywords Calcium manganite · CaMnO₃ · Sintering temperature · CLC · CLOU · TPX · Micropacked bed · In situ XRD · SEM

Introduction
The development of power generation technologies using conventional hydrocarbon fuels with carbon capture and storage (CCS) is a necessary solution to link the contemporary power technologies with environmental demands while all-renewable power technologies emerge. In this context, chemical looping combustion (CLC) is an interesting technology, as CLC allows higher efficiency and lower cost compared to other oxy-combustion technologies. In CLC, oxygen from the air is utilized indirectly in the combustion reactor (fuel reactor) via a solid oxygen carrier material (OCM) that cycles between an air reactor (where OCM takes up oxygen from air) and a fuel reactor (where OCM combusts the fuel). As it intrinsically purifies the used oxygen, it removes the necessity for a stand-alone cryogenic air separation unit.

The OCM is generally a binary, ternary or quaternary transition metal oxide, which utilizes the transition metal redox activity to uptake and release oxygen [1–3]. In the context of fast circulating fluidized bed CLC configuration, the OCM is subject to substantial chemo-mechanical stresses during the redox cycling, through which it must retain a high oxygen capacity, fast redox kinetics and good mechanical properties [4–6]. Furthermore, for industrial application the cost of materials must be low, the component elements abundant, and the active materials and their by-products must be environmentally benign and non-toxic. Considering this long list of demanding requirements, only a handful of materials are proven to pass these criteria.

It has been previously shown that CaMnO₃₋₈ perovskite-type oxides constitute a promising family to be used as OCM. This family of perovskites is especially of interest for large CLC applications, due to low cost and wide abundance of Ca and Mn ores as well as their relatively small environmental impact [7–9]. The redox performance of undoped CaMnO₃, however, deteriorates because of phase separation to CaMn₂O₄ and Ca₂MnO₄ [10, 11]. Although undoped CaMnO₃ perovskite oxide is still considered, it is more common to see doped calcium manganite in publications focusing on perovskite OCMs. On A-site doping, partial substitution of the calcium by strontium, barium and lanthanum [12, 13] is reported, as well as A-site-deficient compounds [14, 15]. More relevant and interesting is the doping of the transition metal on the
B-site, with partial substitution of manganese by magnesium, iron, titanium, or a combination of two or more of them [14–25]. Although it has limited miscibility in the perovskite structure, magnesium has positive catalytic effect on the methane reforming. Doping up to 15% with iron is shown to improve the spontaneous oxygen release kinetics, and widen the operational temperature range [24]. The most important dopant is titanium, as stability of the perovskite structure is shown to be enhanced by substitution of Ti on the B-site [15, 25, 26]. This is due to the higher enthalpy of formation of CaTiO$_3$ compared to CaMnO$_3$ [27], which stabilizes the perovskite structure at reducing conditions of the fuel reactor. CaMn$_{1-x}$Ti$_x$O$_{3-\delta}$ has proven to have high activity with respect to CH$_4$ reduction [28], and introduction of only 12.5% titanium on the B-site improves the spontaneous oxygen release kinetics [29]. Furthermore, it has been shown that it has good chemical stability and shows small dimensional changes upon redox cycling [30]. It is also observed that the re-oxidation is faster for Ti-substituted CaMnO$_3$ at high temperatures [22], and additional Fe doping positively influences the reduction reaction rate and broadened the operation window of the OCM to lower temperatures as well as improve microstructural stability during redox cycling [16]. Although Ti-doping does not assist with the known sulfur intolerance of CaMnO$_3$ [14, 20, 21], remedial technologies for reactivation of the poisoned OCM in situ (in the fuel reactor) are already developed [31].

The multi-national R&D project SUCCESS is globally the largest project so far with rigorous focus on manufacturing and fabrication of oxygen carrier material particles upscaled to industrial scale. The CMT perovskite-type OCM is one of the selected OCM systems. MgO-doped CaMn$_{1-x}$Ti$_x$O$_{3-\delta}$ is one of the materials of focus in SUCCESS [32], selected for upscaling and demonstration. In this study, we analyze the mechanical and thermo-chemical performance of the laboratory-scale and ton-scale OCMs with the nominal composition CaMn$_{0.775}$Ti$_{0.125}$Mg$_{0.1}$O$_3$ fabricated in this project, and try to pinpoint and address some of the remaining challenges that are important to be considered in further upscaling to industrial production scale.

**Experimental**

**Materials synthesis and fabrication**

CaMn$_{0.775}$Ti$_{0.125}$Mg$_{0.1}$O$_3$ (CMTM) is perovskite-type oxides from the CaMnO$_3$ family. Two OCM samples with the same nominal composition were manufactured and studied. The first sample was manufactured at laboratory scale by VITO using a semi-industrial spray-drying method. The second sample was manufactured at the ton scale by Euro Support using an industrial spray-drying technology. The description of manufacturing of both samples is as follows:

**Laboratory-scale batch CMTM**

Stoichiometric quantities of MnO$_2$, Ca(OH)$_2$, MgO and TiO$_2$ were dispersed in deionized water. This suspension was then homogenized using a Netzsch horizontal attrition mill prior to spray drying. During spray drying, the water-based suspension was continuously stirred with a propeller blade mixer while being pumped to the 2-fluid spray-dry nozzle, positioned in the lower cone part of the spray drier (type 6.3-SD, Niro, Denmark). The suspension was atomized into a large number of small droplets in the chamber of the spray drier, which was filled with hot air from the top, thus creating a mixed flow regime. As such, the droplets quickly achieve a spherical shape due to surface tension effects. In addition, the large surface-to-volume ratio of the droplets allows rapid water evaporation, resulting in dry particles which are subsequently separated from the hot air. After spray drying, the fraction within the required particle size range (100–300 µm) was separated from the rest of the spray-dried product by sieving. To obtain oxygen carrier particles with sufficient mechanical strength and with the desired crystalline phases, sintering was performed in air at 1335 °C.

**Ton-scale batch CMTM**

Based on the process developed by VITO described above the spray drying process was upscaled to the ton scale by Euro Support Advanced Material B.V. In order to develop a stable industrial process, the stability of the raw mix slurry needed to be extended. This required substitution of Ca(OH)$_2$ by CaCO$_3$. In order to obtain the desired final particles size with a high yield ratio, a single fluid high-pressure system was used in place of a 2 fluid spray-dry nozzle. This is known to give a sharper particle size distribution especially for bigger spheres as needed for the OCM production. In the high-pressure nozzle, the droplet formation is achieved by a combination of a nozzle plate through which the suspension is pressed via a swirl chamber that creates an additional circular movement in the suspension. The formation of the spheres is a function of the pressure, the nozzle size and the rheological properties of the suspension. Therefore, the change in atomization technology also required changes in the rheological behavior of the suspension in order to get a similar shape for the spheres. The rest of the spray-drying process was identical to the process of VITO. A calcination study of the material produced at Euro Support indicated an optimum firing temperature of 1310 °C set, which was very
close to the temperature optimized by VITO for a different kiln. The full sample was calcined in a stacked saggar configuration.

**Morphochemical characterization**

Samples were characterized before and after the redox cycling. The sieved materials were deposited on conductive adhesive tape and excess material gently blown off. Field-emission gun scanning electron microscopy (FEG-SEM) characterization was performed on a Nova NanoSEM650 (FEI corp.). Images were acquired based on secondary electron (SE) detectors which show topographic contrast and back-scattered electron (BSE) which reflect the local density of the samples (high density induces high brightness). Elemental analysis and mapping was performed using an X-Max50 (Oxford instruments) energy-dispersive spectrometer (EDS) attached to the FEG-SEM instrument.

**Surface area measurement**

The sample’s surface area was measured with a Quantachrome Monosorb instrument. Nitrogen was used as the carrier gas in a single-point BET surface area measurement. Prior to the measurement, the samples were preheated up to 150 °C with the same carrier gas.

**In situ XRD**

The X-ray diffraction (XRD) measurements were performed using a PANalytical Empyrean diffractometer equipped with a CuKα radiation source (λ = 1.541874 Å) and a PIXcel3D solid-state detector. Diffraction patterns were recorded over a 2θ range of 10°–100° in Bragg–Brentano geometry. An Anton Paar XRK 900 in situ high-temperature (25–900 °C) and high-pressure (1–10 bar) reactor cell coupled to an automated gas switching system was used for the in situ XRD experiments. The in situ XRD measurements were performed at 900 °C with a gas flow of 100 mL min⁻¹ through the sample. Initially, the sample was heated up to 900 °C with a heating rate of 5 °C min⁻¹ in synthetic air. The sample was kept at 900 °C in flowing air for 2 h before performing the XRD measurement. The gas was then switched to nitrogen, and the sample was kept in flowing nitrogen for 2 h before measurement. The sample was then cooled down to room temperature in nitrogen. XRD patterns were recorded at room temperature both before and after the experiment.

The crystalline phases present in the diffraction data were identified with reference to the ICDD PDF4+ database and quantitative Rietveld analysis performed using the Bruker DIFFRAC.SUITE Topas v5 software.

**Micropacked bed measurements**

Conversion measurements were performed from 800 to 950 °C with 50 °C incremental steps using a micropacked bed setup (MICROMERITICS AutoChem 2910 TPX apparatus and VG ProLab 1–300 amu mass spectrometer). This reactor has an intersection area of ~ 1 cm². Sieved material in the size range of 125–180 μm was used, with 0.5 ± 0.01 g of sample used in each experiment. The gas flow (30 mL min⁻¹) was successively set from oxidizing (5% O₂ + 5% N₂ in He for 60 min) to inert (He for 30 min) and from oxidizing (30 min) to reducing conditions (10% CH₄ in He for 5 min) with a final inert step (100% He for 10 min). The composition of the outlet gas was followed by mass spectrometry in order to study the CLOU effect and conversion. The gas mixtures used in the experiments are summarized in Table 1.

**Results and discussion**

**Morphochemical structure of the fabricated materials**

As synthesized, the measured surface area for samples was 1 < m² g⁻¹. The laboratory-scale synthesized materials exhibited a density of 1841 kg m⁻³, while the industrial-

### Table 1 Gas mixtures used in the atmospheric experiments

| Step                        | Gas type  | Time at setpoint (min) | H₂ | CH₄ | N₂ | H₂O | He | O₂ |
|-----------------------------|-----------|------------------------|----|-----|----|-----|----|----|
| Spontaneous oxygen release  | 1         | Oxidizing              | 60 | –   | 5  | –   | 90 | 5  |
|                             | 2         | Inert                  | 30 | –   | –  | –   | 100| –  |
| Conversion activity         | 1         | Oxidizing              | 30 | –   | 5  | –   | 75 | 20 |
|                             | 2         | Reducing               | 5  | –   | 10 | –   | 90 | –  |
|                             | 3         | Inert                  | 10 | –   | –  | –   | 100| –  |
scale synthesized materials showed a significantly higher bulk density of 2200 to 2400 kg m\(^{-3}\). Figure 1 shows the SEM–EDS imaging of the laboratory-scale and ton-scale CMTM.

It is seen that the individual particles for both systems are highly spherical and uniform, with agglomerate particle sizes in range 80–180 \(\mu\)m. High-magnification back-scattered electron images confirm that the microstructure and present phases are, however, different.

The fresh laboratory-scale CMTM material is highly spherical and contains minimal doughnut or necking. Higher-resolution images indicate that the grain sizes are in the range of 5 to 10 \(\mu\)m. X-ray diffraction measurements support the SEM–EDS results and suggest that the predominant phase in the sample is a perovskite of approximate stoichiometry of \(\text{CaTi}_{0.15}\text{Mn}_{0.85}\text{O}_3\) (Fig. 2 and Table 2). The magnesium oxide observed by XRD is clearly evident in the EDS element maps (blue areas in Fig. 1), and the elemental analysis confirms that Mg is not dissolved in the perovskite matrix structure. The element analysis also reveals separate Ti-rich and Mn-rich areas, supporting the coexistence of calcium manganate phases with the calcium manganese titanate observed by XRD.

The microstructure and morpho-chemical mapping of fresh ton-scale CMTM samples are also presented in Fig. 1. Although similar in nominal composition, this material differs from laboratory-scale synthesized sample in microstructure and grain sizes. The material’s sphericity is less than that of the laboratory-scale sample, and some necking and doughnut particles are observed. The microstructure is also quite different, and the grain sizes are much smaller. The sample is also inhomogeneous with respect to composition, with Ca-, Mn- and Ti-rich areas evident in the elemental mapping. Despite the relatively
higher density, a lower level of reaction is indicated by the microstructure. It is noted that the mechanical strength is equivalent to that of laboratory-scale batch.

In situ XRD

In situ XRD was performed only on the laboratory-scale CMTM material. This was performed at 900 °C in air, and subsequently in N₂ following an isothermal gas change. The summary of identified phases at room temperature is given in Table 2, with corresponding Rietveld fit to the data shown in Fig. 2. The identified phases and their relative fractions at 900 °C are given in Table 4, with a corresponding Rietveld fit of the data collected in N₂ given in Fig. 3.

As prepared, the sample comprises a major phase of CaMn₁₋ₓTiₓO₃ (orthorhombic perovskite), Ca₂MnO₄ (K₂NiF₄ structure [33]) and CaMn₂O₄ (Marokite [34]),

Table 2 Structure and phase fractions for laboratory-scale CMTM at room temperature before reaction

| Phase | Unit cell 25 °C | Phase fraction 25 °C (mass%) |
|-------|----------------|-----------------------------|
| CaMn₁₋ₓTiₓO₃ (x ≈ 0.1–0.2) | SG = Pbnm(62) | 69.0(4) |
| | a = 5.2872(3)Å |  |
| | b = 5.3127(4)Å |  |
| | c = 7.4729(5)Å |  |
| | vol = 209.91(3) |  |
| CaMn₁₋ₓTiₓO₃ (x ≈ 0.9–1) | SG = Pbnm(62) | 8.2(3) |
| | a = 5.3609(19)Å |  |
| | b = 5.4120(23)Å |  |
| | c = 7.6258(34)Å |  |
| | vol = 221.25(16) |  |
| Ca₂MnO₄ | SG = Fm-3 m(225) | 7.2(3) |
| | a = 5.2117(5)Å |  |
| | c = 23.8989(45)Å |  |
| | vol = 649.1(2) |  |
| CaMn₂O₄ | SG = Pbcm(57) | 12.0(3) |
| | a = 3.1615(4)Å |  |
| | b = 9.9878(13)Å |  |
| | c = 9.6647(12)Å |  |
| | vol = 305.18(7) |  |
| MgO | SG = Fm-3 m(225) | 3.5(2) |

Table 3 Pseudo-cubic lattice parameters for the Mn-rich CaMn₁₋ₓTiₓO₃ phase

| Pseudo-cubic cell parameters at 25 °C | Pseudo-cubic cell parameters at 900 °C in air | Pseudo-cubic cell parameters at 900 °C in N₂ |
|--------------------------------------|----------------------------------------------|---------------------------------------------|
| aₐ = 3.7386 Å                       | aₐ = 3.8031 Å                                | aₐ = 3.8285 Å                              |
| a₈ = 3.7566 Å                       | a₈ = 3.7854 Å                                | a₈ = 3.8063 Å                              |
| a₉ = 3.7365 Å                       | a₉ = 3.7829 Å                                | a₉ = 3.8062 Å                              |

Values are calculated as aₐ = a₉/2√2, a₈ = b₉/2√2, a₉ = c₉/2

higher density, a lower level of reaction is indicated by the microstructure. It is noted that the mechanical strength is equivalent to that of laboratory-scale batch.

Fig. 3 Rietveld fit to the 900 °C/N₂ data for laboratory-scale CMTM sample. Phase peaks positions are indicated for (a) Mn-rich CaMn₁₋ₓTiₓO₃, (b) MgO, (c) Ti-rich CaMn₁₋ₓTiₓO₃, (d) CaMn₂O₄ and (e) Ca₂MnO₄. Inset is a magnification of the 32–42 ° 2θ region with the positions marked of the (absent) diffraction lines which arise from oxygen octahedral tilting.

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In situ XRD was performed only on the laboratory-scale CMTM material. This was performed at 900 °C in air, and subsequently in N₂ following an isothermal gas change. The summary of identified phases at room temperature is given in Table 2, with corresponding Rietveld fit to the data shown in Fig. 2. The identified phases and their relative fractions at 900 °C are given in Table 4, with a corresponding Rietveld fit of the data collected in N₂ given in Fig. 3.

As prepared, the sample comprises a major phase of CaMn₁₋ₓTiₓO₃ (orthorhombic perovskite), Ca₂MnO₄ (K₂NiF₄ structure [33]) and CaMn₂O₄ (Marokite [34]).
with small amounts of a second orthorhombic perovskite
and MgO also present. The fitted room temperature unit
cell parameters for the major perovskite phase correlate
well to a composition \( x \approx 0.1–0.2 \) in CaMn\(_{1-x}\)TixO\(_3\) [35].

The minor perovskite phase exhibits unit cell dimensions
close to those reported for CaTiO\(_3\) [36] and so is presumed
to be a Ti-rich member of the same solid solution.

On heating to 900 °C in air, CaMn\(_2\)O\(_4\) and Ca\(_2\)MnO\(_4\) are
observed to react into the CaMn\(_{1-x}\)TixO\(_3\) phase. The
orthorhombic distortion of this phase is also reduced with
increasing temperature, as reflected by the pseudo-cubic
unit cell parameters for the room temperature and 900 °C
data (Table 3). Switching atmosphere from air to N\(_2\) at
900 °C induces a marked increase of \( \sim 1.85\% \) in the unit
cell volume for the main CaMn\(_{1-x}\)TixO\(_3\) phase (Table 4).

Though this phase still shows apparent orthorhombicity in
the unit cell parameters, it is also observed that the
diffraction lines arising from the oxygen octahedral tilting
located at \( \sim 37.9^\circ \) 2\( \theta \) (indices (120) and (210)) and
at \( \sim 39.2^\circ \) 2\( \theta \) (indices (121), (211) and (103)) disappear
completely, which suggests that the structure has probably
transformed from orthorhombic to cubic. It is reported by
Taguchi [33] that CaMnO\(_3\) undergoes an orthorhombic to
tetragonal phase transition at 896 °C, with a further tran-
sition from tetragonal to cubic at 913 °C. Though solution
of Ti into CaMnO\(_3\) would be expected to raise these
transition temperatures [36], our observation is in good
qualitative accord with the results of Leonidova et al. [37], who report strong depression of the transition temperatures in pure CaMnO$_3$ with decreasing pO$_2$. A slight shift in the relative phase fractions is also observed on switching from air to N$_2$, with increases in the fractions of Ca$_3$MnO$_4$ and CaMn$_2$O$_4$ and a decrease in the major CaMn$_{1-x}$Ti$_x$O$_3$ fraction. CaMnO$_3$ is known to be unstable in reducing conditions with respect to Ca$_3$MnO$_4$ and CaMn$_2$O$_4$ [10].

**Parameters affecting sintering**

The most important parameters affecting sintering are temperature, time and atmosphere. These parameters were investigated in depth for this system Jing et al. [38] showed that temperature has the greatest effect on the quality of the final product and that after 5–6-h sintering, the effect of time is minimal. It was also shown that for upscaled calcination, an air atmosphere should be used. In Fig. 4, the effect of sintering temperature on the bulk density of a wide range of ton-scale CMTM samples is summarized, and demonstrates the sensitivity of the bulk density to the processing temperature, with only a 100 °C variation in the sintering temperature from 1280 to 1380 °C causing the bulk density to increase from ~1150 kg m$^{-3}$ to nearly 2000 kg m$^{-3}$.

As the firing temperature is closely correlated with crystalline phases formed and their homogeneity, as well as the sample density, it primarily defines most of the performance characteristics of the CMTM OCM: Selectivity and activity toward chemical reactions such as methane conversion are a function of the crystalline phases present and their available surface area, and mechanical properties such as attrition index (through changing the crushing strength) are a function of sample density (Fig. 5). A difficult balance is then sought, which trades density (and thus strength) for increased surface area (and thus increased activity), while ensuring the presence of the correct crystalline phases. To achieve the optimum balance, samples must be processed within a relatively narrow sintering temperature window. The significantly higher density obtained for the ton-scale sample here, despite successful pre-sintering studies to determine the optimum firing temperature, highlights the difficulties with maintaining a homogenous and controlled thermal regime when upsaling.

**Micropacked bed reactor**

In order to study CLOU, redox performance and differences in the reaction rates of the two CMTM samples, pulsed and continuous analyses are performed in a micro-packed bed reactor. The CLOU performance and sufficient CLOU capacity are essential for ensuring full combustion.
For the CLOU experiment, the oxygen uptake and release experiment is measured, using gas switching sequences of 60 min of oxidizing oxygen uptake (5% N₂ and 5% O₂ in He) followed by 32 min of inert atmosphere (100% He), and finally 30 min re-oxidation (5% N₂ and 5% O₂ in He). The results for both samples are summarized in Fig. 6. The left part of this figure shows the CLOU effect (oxygen release) under He inert condition, while the right part shows the re-oxidation (oxygen uptake). Considering the equal amounts of O₂ and N₂ in the oxidizing gas prior to and after the inert step, the difference between the O₂ and N₂ graphs corresponds to the oxygen CLOU capacity of the sample. Performing both oxygen release and uptake steps allows calculation of the samples’ CLOU effect in two ways, from both released oxygen and from uptaken oxygen. It is clear from Fig. 6 that the N₂ concentration in the gas has a constant dependency versus time, regardless of temperature or the OCM used in the reactor, and as N₂ does not participate in any chemical reaction, the relaxation and response graphs for N₂ are only dependent on physical and geometrical characteristics of the reactor. Therefore, at any time, the N₂ concentration can be used as a reference.

**Fig. 6** Oxygen release in He CLOU (left) and oxygen uptake during oxidation (right) for laboratory-scale (top) and ton-scale (bottom) samples
By applying this reference, via subtraction of N<sub>2</sub> Fig. 7 is obtained. The CLOU capacity for the materials based on reduction was found to fit well with the CLOU capacity measured from the OCM’s re-oxidation. From Fig. 7, it is clear that the oxidation which is much quicker and where background noise is not a part of the subtraction is the best way to establish the CLOU capacity.

As the nominal composition of both samples is equal, it is clear that the better homogeneity and lower density of the laboratory-scale sample result in better CLOU performance than for the ton-scale sample.

The CLOU capacity estimated from fixed bed oxygen release and oxygen absorption processes was found to be in the same range with a deviation varying between 1 and 8%.

Figure 8 shows the CLOU capacity plotted versus temperature for oxygen release over a period of 30 min at different temperatures (800–950 °C). The CLOU capacity decreases with decreasing temperature, and the laboratory-scale material has the highest CLOU capacity at all the temperatures of study. This is due to an increased fraction of CMT perovskite structure in this sample. The ton-scale sample exhibits much lower CLOU capacity compared to its nominal value, due to its lack of complete intermixing and unreactive CaO left over in the structure.

Experiments using CH<sub>4</sub> as reductant were also performed on both samples. Reduction was performed under 10% CH<sub>4</sub> in He followed by oxidation under 5% N<sub>2</sub> and 5% O<sub>2</sub> in He. The results for this are shown for both
samples in Fig. 9. This figure focuses only on the reduction part and shows the concentration of CO₂ in outlet gases during reduction with methane. This confirms the lower CLOU capacity for the ton-scale sample, as the CH₄ conversion is also lower compared to the laboratory-scale one. This difference is particularly pronounced at lower temperatures where the bulk oxygen diffusion is significantly lower, and cannot balance for the smaller specific surface area of this sample.

**Conclusions**

The formed crystalline phases, their homogeneity and achieved bulk density are identified as the most important parameter affecting the performance of the OCM. Bulk density is directly and significantly correlated with the sintering temperature, under the applied conditions. In other words, most of the important parameters defining the performance of the perovskite OCM are decided by the sintering temperature (if time and atmosphere are kept constant). These parameters include crushing strength, attrition index, specific surface area, CLOU capacity and gas conversion. One of the great challenges in upscaling of the perovskite OCM systems is to achieve a homogeneous sintering temperature at desired values, and the fact that the final product’s performance is greatly and sensitively affected by this parameter. Production of granulates with high ratios of crystalline perovskite phases in large scale needs further optimization in order to better homogenize formed crystalline phases, most importantly through optimization of large-scale calcination procedure.
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