Real-time tomographic diffraction imaging of catalytic membrane reactors for the oxidative coupling of methane

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Graphical Abstract
Highlights

- Synchrotron X-ray diffraction computed tomography applied to three fixed bed catalytic membrane reactors
- The solid-state evolution of catalysts and membranes is tracked under operating conditions
- A new crystal structure model of BaCo$_{0.4}$Fe$_{0.4}$Zr$_{0.2}$O$_{3-\delta}$ (BCFZ) is suggested and used for diffraction data analysis

Abstract

Catalytic membrane reactors have the potential to render the process of oxidative coupling of methane economically viable. Here, the results from operando XRD-CT studies of three different catalytic membrane reactors, employing
BaCo$_{0.4}$Fe$_{0.4}$Zr$_{0.2}$O$_{3-\delta}$ (BCFZ) and La$_{0.6}$Sr$_{0.4}$Co$_{0.2}$Fe$_{0.8}$O$_{3-\delta}$ (LSCF) perovskite membranes with Mn-Na-W/SiO$_2$ and La-promoted Mn-Na-W/SiO$_2$ catalysts, are presented. It is shown that synchrotron X-ray tomographic diffraction imaging allows the extraction of spatially-resolved diffraction information from the interior of these working catalytic membrane reactors and makes it possible to capture the evolving solid-state chemistry of their components under various operating conditions (i.e. temperature and chemical environment).

**Introduction**

In the past decade, the advances achieved in shale gas extraction, coupled with the continuous global effort to reduce and ban gas flaring, have led to dramatic increases in methane availability [1,2]. Methane is nowadays considered to be an abundant hydrocarbon source and, it being the main component of natural gas, much cheaper than crude oil [3]. These developments have set a strong economic imperative towards the utilisation of methane for the direct production of more precious light olefins and other bulk chemicals [4]. A characteristic example is the direct conversion of methane, under the presence of oxygen, to ethylene which is one of the most important building blocks used in the chemical industry [5]. This approach, known as the oxidative coupling of methane (OCM) reaction, is an alternative to the conventional energy-intensive naphtha steam cracking [6]. Historically, this reaction has been mainly studied using fixed bed reactor configuration but as in the case of many selective oxidation reactions, there is a trade-off between selectivity and yield for the desired product molecule (ethylene).

In the past two decades, there has been considerable effort in exploring alternative reactor designs as a means to reach a stable 30 % yield for C$_2$ molecules (i.e. ethane and ethylene) which is often reported as the target goal for this technology [7]. Among the various reactor types, it has been realised that those incorporating membranes, which act as oxygen distributors to the catalyst bed, hold a lot of potential [8–11]. The OCM reaction typically takes place at high temperatures (i.e. > 750 °C) and for this reason most often perovskite-type materials are employed [12–16]. These have mixed ionic and electronic conducting properties (MIEC materials).
and activate at temperatures similar to the operating ones required for the OCM reaction [17]. The MIEC materials are in the form of dense ceramic membranes allowing only oxygen anionic species to diffuse through them, therefore exhibiting pure permeation selectivity for the desired molecules [18,19]. Several geometries have been developed over the past two decades but the one attracting the most interest is the hollow-fibre geometry due to the very large active surface area (i.e. area-to-volume ratio in the range of 500–9000 m²/m³) [20,21]. The incorporation of dense hollow-fibre perovskite-type membranes in the catalytic processes, especially for the OCM reaction, shows several advantages when compared to conventional reactor configurations [21–26]:

- **Maintaining the oxygen partial pressure (pO₂) low and homogeneously distributing the oxygen over the chemical reactor can lead to an increase in C₂ selectivity by preventing the total oxidation of methane.** The uniform distribution of oxygen mitigates reaction rate and selectivity gradients along the reactor, both of which cannot be avoided in fixed bed reactors.

- **Process Intensification:** The oxygen separation unit (i.e. separation of oxygen from air) is no longer needed which increases the energy efficiency and decreases the cost of the overall process.

- **The formation of NOx is avoided as the membrane is selective only for oxygen.** The risk of having explosive mixtures is also reduced and higher productivity can be achieved compared to fixed bed reactors as the ratio of the reactants can be closer to stoichiometry.

- **The formation of hot spots (common problem in fixed bed reactors) can be mitigated due to the gradual feeding of oxygen along the reactor.**

Two of the most promising MIEC membranes investigated as part of membrane reactors for the OCM reaction are La₀.₆Sr₀.₄Co₀.₈Fe₀.₂O₃₋δ (LSCF) and BaCo₀.₄Fe₀.₄Zr₀.₂O₃₋δ (BCFZ) [27–30]. These membranes have been also coupled with catalyst fixed beds in catalytic membrane reactor configurations. However, these complex integrated reactor systems are conventionally characterised ex situ (i.e. before and after an OCM experiment) with destructive local probing characterisation
techniques (e.g. electron microscopy coupled with elemental analysis) or global probing techniques after sample treatment (e.g. sample grinding for bulk ex situ or in situ powder diffraction). These approaches can provide some information regarding the stability of the various material systems but offer little in terms of understanding their real active state and behaviour under OCM operating conditions or complex chemical environments [25,31–39].

Synchrotron X-ray based techniques have the potential to bridge this operando gap often present in catalytic membrane reactor experiments [40,41]. The brilliant X-rays generated at third generation synchrotrons allow the investigation of multi-component systems under real processing conditions which can help us gain a better understanding of their behaviour under different operating conditions [42–46]. As an example, it is possible to investigate under which conditions the MIEC materials undergo phase transformations, form carbonate layers, or what is the state of the catalyst bed in a membrane reactor compared to a fixed bed one; questions often left unanswered with conventional ex situ characterisation. On the other hand, synchrotron-based X-ray chemical tomography techniques can provide this information and also in a spatially-resolved manner [43-61]. We previously employed X-ray diffraction computed tomography (XRD-CT) to study a BCFZ hollow-fibre membrane combined with a Mn-Na-W/SiO$_2$ catalyst demonstrating for the first time the feasibility of such experiments [66–68]. In this work, we performed a thorough investigation of three catalytic membrane reactors investigating the behaviour of the various reactor components under different operating conditions (i.e. temperature and chemical environment).

Materials and Methods

Catalyst and Membrane preparation

The 2 wt. % La – 2 wt. % Mn – 1.6 wt. % Na – 3.1 wt. %W/SiO$_2$ and the 2 wt. % Mn – 1.6 wt. % Na – 3.1 wt. % W/SiO$_2$ catalysts were prepared by the sequential incipient wetness impregnation method [69,70]. At first a SiO$_2$ support (commercial Silica gel Davisil 646) was impregnated by an aqueous solution of sodium tungstate dehydrate Na$_2$WO$_4$$\cdot$2H$_2$O and sodium oxalate Na$_2$C$_2$O$_4$ taken in appropriate amounts. The Na-
W/SiO$_2$ was dried at 120 °C for 6 h and was then impregnated by an aqueous solution of manganese (II) acetate tetrahydrate Mn(CH$_3$COO)$_2$$\cdot$4H$_2$O and lanthanum nitrate hexahydrate La(NO$_3$)$_3$$\cdot$6H$_2$O salts, taken in appropriate concentrations. The catalysts were then dried at 120 °C for 6 h and calcined in air at 850 °C for 6 h with a heating rate of 2 °C min$^{-1}$. Each catalyst sample was sieved obtaining a size fraction of 250-500 μm. The catalysts used in this study were kindly provided by the Boreskov Institute of Catalysis.

The ceramic powders used for the membrane production were fabricated by spray pyrolysis by CerPoTech AS (Trondheim, Norway). Aqueous solutions of the respective precursors were prepared and mixed in stoichiometric ratios, before the stable solution was sprayed into a furnace for each material. The as-prepared powders were calcined, ball milled in ethanol with YSZ grinding media and sieved, before a final heat treatment was executed to remove organic residues from powder processing. The BaCo$_{0.4}$Fe$_{0.4}$Zr$_{0.2}$O$_{3-\delta}$ and La$_{0.6}$Sr$_{0.4}$Co$_{0.2}$Fe$_{0.8}$O$_{3-\delta}$ (LSCF) membranes reported here have been manufactured using the spinning and phase inversion methods [71,72]. The starting polymer suspension was prepared from cellulose acetate (CA, Mr ~52000, Fluka), dimethylsulphoxide (DMSO, Synthesis grade, Merck) and de-ionised water that were used as a phase-inversion polymer, solvent and non-solvent additive to the polymer solution, respectively. The ceramic hollow-fibre membranes used in this study were kindly provided by the Flemish Institute for Technological Research (VITO).

**Operando XRD-CT measurements at ID15A, ESRF**

XRD-CT measurements were made at beamline station ID15A of the ESRF using a 92.8 keV monochromatic X-ray beam focused to have a spot size of 25 μm x 25 μm. 2D powder diffraction patterns were collected using a Pilatus3 X CdTe 300K (487 × 619 pixels, pixel size of 172 μm) hybrid photon counting area detector which uses Cadmium Telluride (CdTe) as the semiconducting direct conversion layer. The acquisition time per point was 50 ms. The tomographic measurements were made with 140 translation steps (translation step size of 25 μm) covering 0 – 180 ° angular range, in steps of 1.8 ° (i.e. 100 line scans). The total acquisition time for each XRD-CT dataset was ca. 25 min. The detector calibration was performed using a CeO$_2$ NIST standard. Every 2D diffraction image was converted to a 1D powder diffraction
pattern after applying an appropriate filter (i.e. 10 % trimmed mean filter) to remove outliers using in-house developed MATLAB scripts [73]. The final XRD-CT images (i.e. reconstructed data volume) were reconstructed using the filtered back projection algorithm.

**Reactor Cells**

Three types of catalytic membrane reactors suitable for the oxidative coupling of methane were prepared from hollow-fibre membranes (ca. 2.4 mm Ø and 180 µm wall thickness) packed with a catalyst supported between glass wool. More specifically, the following CMRs were tested: 1) BCFZ membrane with Mn-Na-W/SiO$_2$ catalyst, 2) BCFZ membrane with La-Mn-Na-W/SiO$_2$ catalyst and 3) LSCF membrane with La-Mn-Na-W/SiO$_2$ catalyst. The catalyst loading in each CMR was ca. 30 mg. In every experiment, the catalytic membrane reactor was glued on top of an alumina rod. For the operando XRD-CT experiments, the reactor cell (i.e. the CMR and the alumina rod) was inserted inside a quartz glass tube and a Swagelok union cross was used to allow the use of two different gas streams (see Figures S1 and S2). Dry air and pure gases of CH$_4$ and He were delivered to the reactor by GF40 Brooks mass flow controllers. Air was used at the outer side of the membrane and a mixture of CH$_4$ diluted in He at the inner side of the membrane. The values for the flow rates reported throughout the text correspond to close to NTP (normal temperature and pressure, 20 °C and 1 atm, respectively).

In each experiment, the reactor was mounted into a gas delivery stub, itself mounted to a standard goniometer (to enable alignment). The goniometer was fixed to a rotation stage set upon a translation stage to facilitate the movements required for the CT measurement. Heating was achieved by virtue of two hot air blowers (Cyberstar) heating each side of the catalytic reactor to ensure homogeneous temperature distribution over the CMR [74]. The hot air blowers were mounted on the translation stage and were moving with the reactor during the translation scans, maintaining the hot air blowers-to-reactor distance constant. XRD-CT measurements were performed at nominal temperatures, ambient up to 1000 °C equating to actual temperatures (i.e. after temperature calibration) of ambient and 780 °C with a nominal ramp rate of 10 °C min$^{-1}$. Temperature calibration of the heating system was
performed prior to the OCM experiments (Figure S3). During the operando XRD-CT measurements, the outflow gasses were monitored by mass spectrometry using an Ecosys portable mass spectrometer. The mass spec line was inserted inside the membrane from the top.

**Rietveld analysis of the XRD-CT data**

Rietveld analysis was performed using the reconstructed diffraction patterns with the TOPAS software, on a voxel by voxel basis [75]. The results from the Rietveld analysis of the XRD-CT data were imported into MALTAB in order to create the figures presented in this study (e.g. phase distribution maps based on the scale factors, unit cell parameters maps, etc.)[76]. The Rietveld analysis of the XRD-CT data was mainly based on the intensity of the scale factors (unless stated otherwise) and should be treated as a semi-quantitative analysis. In order to have a good starting model, quantitative Rietveld analysis of the summed diffraction pattern of all XRD-CT datasets was performed prior to the Rietveld analysis of the XRD-CT data.

It is important to note that prior to the reconstruction of the XRD-CT images and their analysis several issues needed to be addressed, including self-absorption artefacts and the creation of a crystal structure model for the BCFZ phase. A detailed description of the self-absorption correction strategy used in this work and the BCFZ crystal structure model refined using combined neutron and X-ray powder diffraction data is provided in the Supporting Information (Figures S2-6 and S7-13 respectively).

**Results & Discussion**

**Operando XRD-CT measurements of CMRs during OCM**

**BCFZ membrane with Mn-Na-W/SiO2 catalyst**

A schematic representation of the experimental protocol followed for the OCM experiment with the BCFZ–2 % Mn–1.6 % Na–3.1 % W/SiO2 CMR is shown in Figure 1. XRD-CT measurements were made at nominal temperatures, ambient, 800, 900 and 1000 °C equating to actual temperatures (i.e. after temperature calibration) of ca. ambient, 660, 720 and 780 °C. The temperature of the system was increased with a ramp rate of 5 °C min⁻¹ under the flow of He. More specifically, 30 ml min⁻¹ He flow
rate was used at the inner side of the membrane through the catalyst bed and 30 ml min$^{-1}$ air flow at the outer side of the membrane. After reaching the required temperature for the OCM experiment (i.e. 780 °C), air was used from both sides of the BCFZ membrane as an activation step before switching to OCM conditions (i.e. flow rate of 50 ml min$^{-1}$). Next, different OCM reaction mixtures were used by varying the concentration of CH$_4$ in He (i.e. 5, 20, 50 and 100 % CH$_4$ in He). 3D-XRD-CT was then performed by collecting 11 XRD-CT ‘slices’ at different positions along the reactor (i.e. 25 μm apart) while keeping the reaction mixture the same (i.e. 100 ml min$^{-1}$ of CH$_4$ at the inner side of the membrane through the catalyst bed and 100 ml min$^{-1}$ air at the outer side of the membrane). The system was then cooled to room temperature and a final XRD-CT dataset was collected.

![Figure 1: Schematic representation of the protocol followed during the OCM experiment with the BCFZ – 2%Mn-1.6%Na-3.1%W/SiO$_2$ CMR.](image)

Initially, an XRD-CT scan of the CMR was performed at room temperature. In agreement with our previous study, the main crystalline phases present in the catalyst particles were cristobalite-low, tridymite-low, Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ and Na$_2$WO$_4$ (Figures S14, S15 and Table S3). The phase distribution maps shown in Figure 2 represent the (normalised) values of the scale factors of these crystalline phases as obtained from the Rietveld analysis of the CMR XRD-CT data collected at room temperature. It is seen that the two main crystalline SiO$_2$ phases, cristobalite and tridymite, are homogeneously distributed over the catalyst particles and define well their size and shape. On the other hand, it can be seen that there are certain areas have high concentration of Mn$_2$O$_3$/Mn$_7$SiO$_{12}$. Na$_2$WO$_4$ is present in all catalyst
particles but there are areas of high concentration of this phase too (i.e. hotspots).

Finally, the phase distribution map of BCFZ demonstrates that the membrane wall thickness is fairly uniform (ca. 180-200 μm).

Figure 2: Phase distribution maps created based on the values of the scale factors of the crystalline phases present in the CMR as obtained from the Rietveld analysis of the XRD-CT data collected at room temperature before the OCM experiment. From left to right: BCFZ, cristobalite (alpha - low-T), tridymite (alpha - low-T), Na$_2$WO$_4$ and Mn$_2$O$_3$/Mn$_7$SiO$_{12}$.

The next XRD-CT scan was performed at 660 °C. Careful inspection of the reconstructed XRD-CT data revealed that the high temperature tomographic scans were performed at a seemingly different position compared to the room temperature scan; this is due to thermal expansion/movement of the CMR during heating. It was feasible to identify this vertical change of the probing position after the OCM experiment by comparing the phase distribution maps of cristobalite. As mentioned previously, the phase distribution maps of cristobalite define well the catalyst particles and can be used to correlate different tomographic datasets. As a result, the data of the high temperature XRD-CT scans cannot be directly compared with the room temperature XRD-CT scans. In Figure 3, the results from the high temperature XRD-CT scans are presented which are all at the same position (i.e. XRD-CT cross sections at a constant position of the CMR). The relative change of each phase, as determined from the integrated intensities (scale factors) for each XRD-CT dataset (i.e. integrated over the entire slice), is presented at the right side of Figure 3.

The support of the catalyst particles is seen to evolve at high temperatures which is in full agreement with the results obtained from our previous operando XRD-CT experiments of the catalytic membrane reactor with the same catalyst [68]. The diffraction peaks generated by Na$_2$WO$_4$ have disappeared completely at 660 °C.
although its melting point is 695 °C in 1 bar pressure [77]. The mobile Na+ species are incorporated into the SiO2 framework and lead to the transformation of the remaining amorphous SiO2 to the cristobalite phase. It can be seen though that after the initial spike, the diffraction signal corresponding to the cristobalite phase starts to decrease as it gradually transforms to the tridymite and quartz phases.

Figure 3: Phase distribution maps of Cristobalite (beta - high-T), Tridymite (beta - high-T), Quartz (beta - high-T), Mn2O3/Mn7SiO12 and BaWO4 at high temperatures during the OCM experiment. At the right side, the relative change of each phase, as determined from the integrated intensities for each XRD-CT dataset, is presented.

The results from our previous work on the same CMR system showed that the crystalline Mn2O3/Mn7SiO12 phase decreased with time at high temperatures when a gas mixture of 5% CH4/He is used. However, it is shown in Figure 3 that letting air pass through the catalyst bed leads to a significant increase of the crystalline Mn2O3/Mn7SiO12 phase under OCM operating temperatures (780 °C). This is important as, if Mn2O3/Mn7SiO12 needs to be present under OCM conditions to yield an active and C2+ selective catalyst, then an activation step by using air at high temperatures is necessary; an issue which has been raised before in literature [78]. In the past, the Mn-Na-W/SiO2 catalysts have shown great stability in terms of CH4 conversion but the selectivity towards C2+ molecules has been shown to decrease
with time [79–81]. Therefore, it might be also necessary to switch the inlet gases from reaction mixture to air after many hours of operation at high temperatures as a regeneration step for the catalyst. For example, Hou et al. showed that treating the Mn-Na-W/SiO₂ catalyst with O₂ for 2 h at 800 °C led to the reappearance of Na₂WO₄ and Mn₃O₄/Mn₇SiO₁₂ and the disappearance of MnWO₄ [82].

It should be emphasized though that such (regeneration) treatments may not be needed in CMRs containing Mn-Na-W/SiO₂ catalysts. In the BCFZ – 2 % Mn – 1.6 % Na – 3.1 % W/SiO₂ CMR (Figure 3), the diffraction signal generated by the Mn₃O₄/Mn₇SiO₁₂ phase is seen to decrease under OCM reaction conditions (i.e. gas mixtures of CH₄/He) but the Mn₃O₄/Mn₇SiO₁₂ to MnWO₄ phase transformation was not observed for the duration of the experiment. This is crucial as the catalyst bed was exposed to 100 % CH₄ for almost 10 h and the observed Mn₃O₄/Mn₇SiO₁₂ phase(s) remained present till the end of the OCM experiment. This indicates that the O₂ transferred through the BCFZ ceramic membrane was enough to maintain the Mn₃O₄/Mn₇SiO₁₂ phase(s) and prevent the bulk reduction of the Mn species (from Mn³⁺ to Mn²⁺).

However, another crystalline WO₄²⁻-containing phase appears as a result of the chemical interaction between the catalyst particles and the BCFZ ceramic membrane. This new phase is BaWO₄ and it is seen to form/grow at the inner side of the BCFZ membrane where the catalyst particles are in direct contact with the membrane. As it is shown in Figure 3, BaWO₄ continuously grows during the OCM experiment regardless of the applied chemical environment, proving that this is a purely temperature driven phenomenon.

3D-XRD-CT was also performed under OCM reaction conditions by collecting 11 XRD-CT datasets at different positions along the reactor (i.e. 25 μm apart) at 780 °C (flow rate of 100 ml min⁻¹ of 100 % CH₄ at the inner side of the BCFZ membrane, through the catalyst bed). In Figure 4, the results from the volume rendering of the scale factors data volume (phase distribution volumes) are presented. The scale factors were normalised with respect to the maximum value for each data volume and the colour axes were scaled accordingly in order to improve the contrast of the respective figures (see also Figure S21).
The cristobalite phase distribution volume clearly describes the catalyst particles in terms of both size and shape. This is expected as cristobalite, being the main crystalline phase of the SiO$_2$ catalyst support, is homogeneously distributed (uniform distribution) over the particles. Tridymite is present in most catalyst particles but it can also be seen that the sample regions showing high tridymite concentration also correspond to low cristobalite concentration. Similarly to cristobalite, quartz is seen to be present in all catalyst particles albeit not as evenly distributed.

![Figure 4](image_url)

**Figure 4**: Volume rendering of the normalised scale factors data volume (phase distribution volumes) obtained from the Rietveld analysis of the 3D-XRD-CT data. Top row: Cristobalite (beta - high-T), Tridymite (beta - high-T) and Quartz (beta - high-T). Bottom row: Mn$_2$O$_3$/Mn$_7$SiO$_{12}$, BaWO$_4$ and BCFZ. The values in the colorbar axes have been chosen to achieve the best possible contrast.

The results obtained from the XRD-CT data collected at room temperature before the OCM experiment indicated that Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ is not homogeneously distributed over the catalyst particles but is only present at specific regions with very high concentration, mainly near the surface of the catalyst particles (Figure 2). The Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ phase distribution volume (Figure 4) serves to verify this result and also to show that the Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ distribution does not alter significantly under OCM reaction conditions.
The BCFZ membrane maintained its size and shape for the duration of the OCM experiment and no cracks were formed. This is also implied by the BCFZ phase distribution volume which clearly demonstrates the dense ceramic hollow-fibre design. Finally, the BaWO$_4$ phase distribution volume reveals that there are regions of high concentration of this phase at the inner side of the BCFZ membrane (hotspots of material) which, as it will be shown later, are mainly located at the catalyst-membrane interface. This is attributed to the W containing species, originating from the molten Na$_2$WO$_4$ phase, being both mobile and volatile.

Finally, it was feasible to directly compare the state of the catalyst before, during and after the OCM reaction. As mentioned previously, the CMR slightly moved during temperature ramping but fortunately one of the 3D-XRD-CT scans was at the same position as the one at ambient conditions before the OCM experiment. More importantly, the XRD-CT scan at room temperature after the OCM experiment is also at the same position allowing the direct comparison of these three datasets. The phase distribution maps presented Figure 5 correspond to the respective (normalised) phase scale factors obtained from the Rietveld analysis of the XRD-CT data.
Figure 5: Phase distribution maps of Cristobalite (alpha - low-T), Tridymite (alpha - low-T), Quartz (alpha - low-T), Na$_2$WO$_4$, Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ and BaWO$_4$ at room temperature before the OCM experiment, at high temperatures during the OCM experiment and at room temperature after the OCM experiment. At the right side, the relative change of each phase, as determined from the integrated intensities for each XRD-CT dataset, is presented.

The comparison of these three XRD-CT datasets reveals that by the end of the OCM experiment, cristobalite has been partially consumed in favour of the growth of the tridymite and quartz phases. The Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ phase is also seen to decrease during this OCM experiment but the formation of the MnWO$_4$ phase was not observed. The Na$_2$WO$_4$ phase is seen to reform after cooling to room temperature albeit at a lesser extent (Figures S19-21). This is to be expected as BaWO$_4$ formed and continuously grew during the OCM experiment which was not present in the initial materials (i.e. catalyst and membrane) composing the CMR. The formation of BaWO$_4$ and the redistribution of the WO$_4^{2-}$ species was also verified with complementary ex situ micro-CT and SEM/WDS measurements (Figures S22-23).

The spatially-resolved XRD patterns obtained from the XRD-CT data were not of high enough quality (i.e. in terms of number of observation points, angular resolution and signal-to-noise ratio) to allow for the refinement of structural parameters (i.e. thermal parameters and occupancy of the various atoms present in BCFZ). It was though possible to extract the scale factor and lattice parameter of BCFZ in a spatially-resolved manner and the results are presented in Figure 6. As mentioned previously, the BCFZ membrane retained its shape and size for the duration of the OCM experiment. However, as it is shown in Figure 6, Co is seen to gradually leave the BCFZ unit cell and be present as the crystalline CoO phase. It is still present as a minor phase/traces in the CMR XRD-CT data collected at room temperature after the OCM experiment but it raises questions regarding the long-term chemical stability of the BCFZ membrane under OCM reaction conditions and reducing chemical environments in general.
Figure 6: First and second row: Phase distribution maps of BCFZ and CoO. Third row: Maps corresponding to the BCFZ unit cell parameter during the OCM experiment.

BCFZ membrane with La-Mn-Na-W/SiO₂ catalyst

As mentioned previously, the same experimental protocol was followed for all the operando CMR XRD-CT experiments (Figure 1). Initially, an XRD-CT scan of the BCFZ – 2 % La-2 % Mn-1.6 % Na-3.1 % W/SiO₂ CMR was performed at ambient conditions. The phase distribution maps of the various crystalline phases, as obtained from the Rietveld analysis (scale factors) of the XRD-CT data, are presented in Figure S28. The main difference between the La-promoted and the unpromoted catalyst lies in the distribution of the Mn₂O₃/Mn₇SiO₁₂ phase as on the former it is seen to be homogeneously distributed.

The next three XRD-CT scans were performed at 660, 720 and 780 °C respectively. As in the case of the previous CMR experiment, inspection of the reconstructed XRD-CT data (cristobalite XRD-CT images) revealed that the high temperature XRD-CT scans were performed at a different position compared to the room temperature scan due to thermal expansion/movement of the CMR during heating. As a result, the XRD-CT data collected at high temperatures cannot/should not be directly compared with the XRD-CT data acquired at room temperature. The results from the Rietveld analysis of the high temperature XRD-CT data collected at the same position (i.e. XRD-CT cross sections at a constant position of the CMR) are presented in Figure 7. These phase distribution maps correspond to the values of the corresponding (normalized) scale factors, as obtained from the Rietveld analysis of the XRD-CT data. The relative change of each phase, as determined from the integrated intensities (i.e.
integrated over the entire slice) for each XRD-CT dataset, is presented at the right side of Figure 7.

**Figure 7**: Phase distribution maps of Cristobalite (beta - high-T), Tridymite (beta - high-T), Quartz (beta - high-T), Mn₂O₃/Mn₇SiO₁₂, NaLa(WO₄)₂ and BaWO₄ at high temperatures during the OCM experiment. At the right side, the relative change of each phase, as determined from the integrated intensities for each XRD-CT dataset, is presented.

There is an evolution of SiO₂ support at high temperatures; this phenomenon was thoroughly discussed previously and it is attributed to the presence of mobile Na+ species. The unexpected high concentrations of cristobalite and tridymite at 660 °C (compared to the other high temperature XRD-CT datasets) is explained by the fact that the CMR did not fully stabilize before reaching 780 °C. This can be seen after inspecting the cristobalite phase distribution maps presented in Figure 7 which suggest that the XRD-CT scan performed at 660 and 720 °C probed different positions of the CMR (i.e. with respect to all the XRD-CT data collected at 780 °C). The results obtained from the XRD-CT scans performed at the same position are though consistent with that which has been observed previously; namely that there is an
initial increase in the presence of cristobalite at 780 °C but that this is gradually consumed under OCM operating conditions while the tridymite and quartz phases increase as a function of time. The Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ phase is seen to increase in the presence of air which is also consistent with was observed in the BCFZ – 2% Mn-1.6% Na-3.1% W/SiO$_2$ CMR. The concentration of the Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ phase is though seen to decrease with time under OCM reaction conditions, especially when CH$_4$ rich (i.e. > 20% CH$_4$/He) reaction mixtures are used.

It can be seen that BaWO$_4$ is forming and growing only at specific regions (hotspots in Figure 7) where the catalyst particles are in direct contact with the BCFZ membrane. Only after for several hours of operation at high temperatures (each XRD-CT lasted approximately 45 min.) a uniform “ring” (layer) is observed at the inner BCFZ membrane wall. This delay could be attributed to the presence of the La species in the catalyst particles and specifically the NaLa(WO$_4$)$_2$ phase. It is indicated that the presence of this phase slows down the formation and growth of BaWO$_4$ compared to the unpromoted catalyst, by partially stabilising the mobile and volatile Na$^+$/WO$_4$$^{2-}$ species. However, the slow decomposition of this newly formed phase at high temperatures was also observed. The delayed effect this phase had though suggests the solution to the formation of BaWO$_4$ phase lies on chemically trapping the volatile Na$^+$/WO$_4$$^{2-}$ species.

As in the previous CMR experiment, 3D-XRD-CT was performed under OCM reaction conditions by collecting 10 XRD-CT cross sections at different positions along the reactor (i.e. 25 μm apart) at 780 °C (flow rate of 100 ml min$^{-1}$ of 100% CH$_4$ at the inner side of the BCFZ membrane, through the catalyst bed). The phase distribution maps presented in Figure S29 correspond to the scale factors (normalised to the maximum value) as obtained from the Rietveld analysis of the 3D-XRD-CT data. As mentioned previously, these XRD-CT data should not be directly compared as they correspond to different sample volumes. If the solid-state changes taking place in the CMR were only a function of position (spatial dependence only), then the observation points shown on the right side of Figure S29 should not follow any specific trend. These plots correspond to the summed intensity of the various scale factors (normalised to the maximum value) as obtained from the Rietveld analysis of the 3D-
XRD-CT data. However, for some crystalline phases there is a clear trend (even linear in some cases) indicating that some of the solid-state changes can be tracked as a function of time (temporal dependence). For example, it can be seen that cristobalite decreases with time while tridymite and quartz increase. More importantly though, there is a continuous growth of the NaLa(WO₄)₂ and BaWO₄ phases. This indicates that the formation/growth of NaLa(WO₄)₂ is not completely prevented by the formation/growth of BaWO₄. This is a key result as it implies that if the catalyst is thermally treated (e.g. extended calcination periods) prior to its insertion in the CMR in order to convert the Na₂WO₄ to NaLa(WO₄)₂, then the formation/growth of BaWO₄ could be potentially suppressed.

The results from the volume rendering of the scale factors data volume (phase distribution volumes) are presented in Figure 8. The scale factors were normalised with respect to the maximum value for each data volume and the colour axes were scaled accordingly in order to improve the contrast of the respective figures. As expected, the various crystalline SiO₂ phases are present in all catalyst particles. As expected, in contrast to the unpromoted catalyst (Figure 4), the Mn₂O₃/Mn₇SiO₁₂ phase in the La-promoted catalyst is seen to be homogeneously distributed over all catalyst particles probed during the 3D-XRD-CT scan. The BaWO₄ phase is present in all XRD-CT datasets but it can also be observed that there are regions of high BaWO₄ concentration (hotspots). These hotspots are located at the interface between the catalyst particles and the inner BCFZ membrane wall.
**Figure 8:** Volume rendering of the normalised scale factors data volume (phase distribution volumes) obtained from the Rietveld analysis of the 3D-XRD-CT data. Top row: Cristobalite (beta - high-T), Tridymite (beta - high-T) and Quartz (beta - high-T). Bottom row: Mn₂O₃/Mn₇SiO₁₂, BaWO₄, BCFZ and NaLa(WO₄)₂. The values in the colorbar axes have been chosen to achieve the best possible contrast.

In contrast to the previous CMR experiment, the XRD-CT data acquired at room temperature after the OCM experiment did not probe the same sample volume as the XRD-CT data before the experiment (different position in the CMR). This means that these two datasets should not be directly compared in terms of quantitative analysis but can provide a good indication of the solid-state changes that took place in the CMR during the OCM experiment.

As in the case of the previous CMR, the BCFZ structure was also added to the Rietveld refinement and the results are presented in Figure S30. The maps corresponding to the BCFZ unit cell parameter are homogeneous and change only as a function of temperature. It should also be pointed out that shape and size of the BCFZ membrane did not change during the OCM experiment and no cracks were formed. However, the formation and growth of CoO is observed again showing that the results obtained from the previous CMR experiment are directly reproducible (Figure 6). Whilst the CoO is only a minor phase present in the diffraction patterns but it suggests there may be issues regarding the long-term chemical stability of BCFZ under OCM operating conditions.

In Figure 9, the phase distribution maps of various crystalline phases from the XRD-CT datasets are presented. These datasets correspond to the following XRD-CT scans: room temperature before the OCM experiment, at 780 °C under 100 % CH₄ flow (inner side of BCFZ membrane) and room temperature scan after the OCM experiment. It should be noted that the high temperature XRD-CT scan was performed at the same CMR position as the room temperature XRD-CT scan before the OCM experiment.
Figure 9: Phase distribution maps of Cristobalite (alpha - low-T), Tridymite (alpha - low-T), Quartz (alpha - low-T), Na$_2$WO$_4$, Mn$_2$O$_3$/Mn$_7$SiO$_{12}$, NaLa(WO$_4$)$_2$ and BaWO$_4$ at room temperature before the OCM experiment, at high temperatures during the OCM experiment and at room temperature after the OCM experiment. At the right side, the relative change of each phase, as determined from the integrated intensities for each XRD-CT dataset, is presented.

It can be seen that the evolution of the SiO$_2$ support is consistent with what has been observed in the previous CMR experiment (i.e. consumption of cristobalite and consequent growth of tridymite and quartz). As it is also shown in Figure S32, Na$_2$WO$_4$ reforms upon cooling to room temperature after the OCM experiment. Of course, the concentration of Na$_2$WO$_4$ is lower compared to the fresh catalyst as both NaLa(WO$_4$)$_2$ and BaWO$_4$ formed/grew during the operation of the CMR. The fact that the NaLa(WO$_4$)$_2$ phase was seen not only to form but also to increase at high temperatures in this CMR (Figure S32 and Figure 9) suggests that the NaLa(WO$_4$)$_2$ is chemically stable under these conditions. This means that not only the La species can
stabilise the mobile WO$_4^{2-}$ species but it is also implied that if NaLa(WO$_4$)$_2$ is the only Na-W containing phase in the fresh catalyst (no Na$_2$WO$_4$) then the catalyst may be stable the formation/growth of BaWO$_4$ can be suppressed and maybe even completely prevented.

**LSCF membrane with La-Mn-Na-W/SiO$_2$ catalyst**

The last CMR experiment was performed using a La$_{0.6}$Sr$_{0.4}$Co$_{0.2}$Fe$_{0.8}$O$_3$ (LSCF) membrane and the 2 % La-2 % Mn-1.6 % Na-3.1 % W/SiO$_2$ catalyst. The same experimental protocol was followed as in the other CMR experiments (Figure 1). Unfortunately, the quality of the LSCF – 2 % La-2 % Mn-1.6 % Na-3.1 % W/SiO$_2$ CMR data was significantly inferior compared to the other CMR data. More specifically, the raw diffraction data suffered from saturation (spottiness). This is clearly shown in Figure S33 where a summed 2D diffraction pattern derived from a line scan of the LSCF CMR XRD-CT data collected at room temperature is shown. Apart from the presence of large grains in the LSCF membrane that led to the formation of spots (single-crystal diffraction) on top of the powder diffraction data, the LSCF – La-Mn-Na-W/SiO$_2$ XRD-CT data severely suffered from self-absorption problems too (Figure S34).

The phase distribution maps of the various crystalline phases initially present in the CMR are shown in Figure S35. As in the previous two CMR experiments, the thermal expansion of the LSCF membrane resulted in probing different sample volumes at high temperatures compared to the room temperature XRD-CT scan. The results from the high temperature XRD-CT data, collected at the same position, are presented in Figure 11. The phase distribution maps shown in Figure 11 correspond to the values of the corresponding scale factors, normalised with respect to the maximum value, as obtained from the Rietveld analysis of these XRD-CT datasets (Figures S32-34). The relative change of each phase is plotted on the right side of Figure 11.

As expected, the cristobalite phase decreases with time at the high temperatures applied for the OCM experiment while tridymite is seen to be fairly stable and quartz significantly increases. As shown previously, the Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ phase is seen to
increase when air is used as the inlet gas mixture through the catalyst bed but then decreases with time under OCM reaction conditions.

It can be seen that in this CMR, although Ba is absent in the membrane material, there is chemical interaction between the catalyst particles and the LSCF membrane which leads to the formation and growth of a new phase, identified as SrWO₄. However, as in the case the second experiment (and in contrast to the first one), the SrWO₄ is primarily seen to form and grow only at specific regions (hotspots in Figure 11) where the catalyst particles are in direct contact with the LSCF membrane. This phenomenon, as described previously, was attributed to the presence of the NaLa(WO₄)₂ phase which partially stabilises the volatile Na⁺/WO₄²⁻ species. It can be seen that the SrWO₄ phase increases with time at high temperatures, eventually forming a uniform layer at the inner LSCF membrane wall, as in the case of the BaWO₄ phase in the BCFZ CMRs. This is the key result of this experiment as it proves that the main drawback of the three CMRs is related to the unstable WO₄²⁻ species present at the catalyst at high temperatures.
Figure 11: Phase distribution maps of LSCF, Cristobalite (beta - high-T), Tridymite (beta - high-T), Quartz (beta - high-T), Mn$_2$O$_3$/Mn$_7$SiO$_{12}$, NaLa(WO$_4$)$_2$ and SrWO$_4$ at high temperatures during the OCM experiment. At the right side, the relative change of each phase, as determined from the integrated intensities for each XRD-CT dataset, is presented.

As in the previous two CMR experiments, 3D-XRD-CT was also performed at 780 °C while maintaining the OCM reaction mixtures constant (100 ml min$^{-1}$ of 100 % CH$_4$ at the inner side of the membrane through the catalyst bed and 100 ml min$^{-1}$ air at the outer side of the LSCF membrane). The results from the Rietveld refinement of the 3D-XRD-CT data are presented in Figure S39. It can be seen that there is a continuous consumption of cristobalite and growth of quartz which is in full agreement with what has been observed in the previous experiments. It should also be noted that the SrWO$_4$ phase (Figure S39) is also seen to continuously grow as a function of time which is identical to the behaviour of the BaWO$_4$ phase in the BCFZ CMRs.

The results from the volume rendering of the 3D-XRD-CT data are presented in Figure 12. As expected, the three crystalline SiO$_2$ phases (cristobalite, tridymite and quartz) are homogeneously distributed over the catalyst particles. Similar is the case for the
Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ phase which is in full agreement with the results obtained from BCFZ CMR containing the same catalyst. The NaLa(WO$_4$)$_2$ phase is seen to be present in most catalyst particles indicating that the formation/growth of the SrWO$_4$ phase only delays but does not prevent its evolution.

It should be noted though that no cracks formed in the LSCF membrane and no other La, Sr, Co or Fe containing phases were observed during the OCM experiment (apart from the SrWO$_4$ phase). This is in contrast to what was observed in the BCFZ CMRs where the Co was seen to leave the BCFZ unit cell and form CoO. This result indicates that the LSCF membrane is more chemically stable under OCM reaction conditions compared to the BCFZ membrane.

![Figure 12: Volume rendering of normalised the scale factors data volume (phase distribution volumes) obtained from the Rietveld analysis of the 3D-XRD-CT data. Top row: Cristobalite (beta - high-T), Tridymite (beta - high-T) and Quartz (beta - high-T). Bottom row: Mn$_2$O$_3$/Mn$_7$SiO$_{12}$, SrWO$_4$, LSCF and NaLa(WO$_4$)$_2$. The values in the colorbar axes have been chosen to achieve the best possible contrast. Finally, the results from the Rietveld analysis of three XRD-CT datasets collected at the same position are presented in Figure 13. These datasets correspond to XRD-CT data collected at room temperature before the OCM experiment, at 780 °C under 100 % CH$_4$ flow and at room temperature after the OCM experiment. In accordance to what was previously observed, cristobalite is seen to have been partially consumed by the end of the OCM experiment while tridymite and quartz have grown. The Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ phase is seen to decrease with time under OCM reaction conditions which in agreement with what has been previously observed (behaviour
of the La-promoted catalyst under OCM reaction conditions). The Na$_2$WO$_4$ is seen to be present at the room temperature XRD-CT scan after the OCM experiment. However, its concentration is significantly lower as both SrWO$_4$ and NaLa(WO$_4$)$_2$ formed and grew during the operation of the CMR (Figure S42).

**Figure 13**: Phase distribution maps of Cristobalite (alpha - low-T), Tridymite (alpha - low-T), Quartz (alpha - low-T), Na$_2$WO$_4$, Mn$_2$O$_3$/Mn$_7$SiO$_{12}$, NaLa(WO$_4$)$_2$ and SrWO$_4$ at room temperature before the OCM experiment, at high temperatures during the OCM experiment and at room temperature after the OCM experiment. At the right side, the relative change of each phase, as determined from the integrated intensities for each XRD-CT 'slice', is presented.
Summary and Conclusions

Catalytic membrane reactors have the potential to render the process of oxidative coupling of methane to produce ethylene economically viable. For example, two of the highest values for C\textsubscript{2} yield (>35\%) ever reported in literature have been achieved using catalytic membrane reactors [9,83]. However, a number of questions regarding the true state of the active catalyst/membrane and their stability over time remain unanswered. These integrated reactor systems are typically characterised ex situ with electron microscopy techniques (SEM/TEM/EDX) and bulk powder diffraction.

Herein, three different catalytic membrane reactor systems have been studied, for the first time, under real OCM reaction conditions with XRD-CT. The operando XRD-CT measurements allowed to spatially-resolve the pertinent changes in the chemical/physical state within all working CMRs. Despite the challenging nature of these experiments it has been shown that spatially-resolved diffraction information can be extracted from the interior of working catalytic membrane reactors. Indeed, the state of two different catalysts was captured prior to, under and after OCM operando conditions.

The three CMRs investigated under OCM operating conditions were a BCFZ – 2 % Mn-1.6 % Na-3.1 % W/SiO\textsubscript{2} CMR, a BCFZ – 2 % La-2 % Mn-1.6 % Na-3.1 % W/SiO\textsubscript{2} CMR and an LSCF – 2 % La-2 % Mn-1.6 % Na-3.1 % W/SiO\textsubscript{2} CMR. The three CMRs were operated under atmospheric pressure but future experiments could also investigate the impact of high pressure in the evolving chemistry of the various reactor components. It should be emphasized that no carbonate phases were observed under any imposed chemical environments (even under 100 % CH\textsubscript{4}) for any CMR. It should also be pointed out that all three CMRs were captured in their active state where CO\textsubscript{2} is continuously produced as a side product of the OCM reaction. This is a very important result as it demonstrates that both BCFZ and LSCF are chemically stable under OCM reaction conditions and the produced CO\textsubscript{2} is not able to significantly damage the membranes; this is typically done through the formation of phases such as BaCO\textsubscript{3} and SrCO\textsubscript{3}.

In all experiments, the Na\textsubscript{2}WO\textsubscript{4} phase, present in the fresh catalysts, is seen to melt before reaching the required high temperatures for the OCM reaction (780 °C) and
coincident to this event the formation of a new phase is observed. In the BCFZ CMRs, this phase was identified as BaWO$_4$ while in the LSCF CMR the new phase was identified as SrWO$_4$. These new phases are seen to initially form at the interface between the catalyst particles and the inner side of the membranes. The mobility of W-containing species was independently confirmed by micro-CT and ex situ SEM/WDS measurements. These phases are seen to continuously grow under the high temperatures used for the OCM reaction and regardless of the chemical environment present (reaction mixtures used) indicating that their formation and subsequent growth is a purely temperature driven phenomenon. The growth of stable the BaWO$_4$ phase in the BCFZ CMRs and the SrWO$_4$ phase in the LSCF CMR is extremely important as:

1. loss of the W-containing species is expected to have an impact on the stability of the catalyst (loss of one of the main catalyst components),

2. it might lead to the deactivation of the membranes (BCFZ/LSCF) after many hours of operation by forming an inner layer at the membrane and therefore blocking the flow of oxygen to the catalyst bed.

At this point it should be emphasized that in the La-promoted catalysts, the loss of WO$_4^{2-}$ species and the formation/growth of BaWO$_4$/SrWO$_4$ did not prevent the formation of NaLa(WO$_4$)$_2$ in the catalyst particles. In fact, the NaLa(WO$_4$)$_2$ phase was seen to grow with time under OCM operating conditions (although with a delay compared to the respective fixed bed reactor experiments). This is important as the formation and subsequent growth of NaLa(WO$_4$)$_2$ suggests that if the catalyst if pre-treated before it is inserted in the CMR (e.g. extensive catalyst calcination periods), then the formation/growth of the BaWO$_4$/SrWO$_4$ can be suppressed if not completely prevented during the operation of the CMRs.

Regarding the catalyst behaviour, the SiO$_2$ support of both catalysts is also seen to evolve with time under OCM operating conditions. More specifically, cristobalite is seen to gradually transform to tridymite and then to quartz at high temperatures (780 °C). This phenomenon was attributed to the presence of mobile Na+ species and their incorporation to the SiO$_2$ framework. The Mn$_2$O$_3$/Mn$_7$SiO$_{12}$ phase remains
present for the duration of the OCM experiments. The transformation of the \(\text{Mn}_2\text{O}_3/\text{Mn}_7\text{SiO}_{12}\) phase to the \(\text{MnWO}_4\) phase was not observed in the Mn-Na-W/SiO\(_2\) catalyst. This result suggests that the bulk reduction of the Mn species (Mn\(^{3+}\) to Mn\(^{2+}\)) was prevented in the CMRs. This observation becomes more important when it is taken into account that the 100 % CH\(_4\) was used as the inlet gas to the catalyst bed for more than 8 h at 780 °C. This result implies that the oxygen permeation flux through the dense ceramic membranes was enough to prevent the bulk reduction of the Mn species present in the catalysts.

Last but not least, a crystal structure model of was created using high resolution neutron powder diffraction data collected at the ISIS neutron source. The model provided an opportunity to extract more information on the nature of the BCFZ membrane’s performance in operation. The BCFZ structure was added to the Rietveld refinement model and it was shown that it was possible to create BCFZ phase distribution and lattice parameter maps. BCFZ has also been shown to be a promising electrode material for fuel cell applications and now a structure model is available for the fuel cell community. We expect that the structure model will be used in future studies to gain a better understanding of the performance of this material under different operating conditions (e.g. crystallographic properties as a function of temperature, chemical environment or potential).

Author Contributions

The OCM experiments were conceived by A.V., V.M., S.D.M.J. and A.M.B. M.D.M. and A.V. wrote the macros for the data acquisition and the scripts for the sinogram data pre-processing. A.V., D.M., P.S., S.D.M.J., S.W.P. and A.M.B. performed the OCM experiments. M.D.M. was responsible for ID15A instrumentation and setup at the ESRF. Data processing and analysis was performed by A.V. with assistance from M.D.M., S.D.M.J., D.M. and A.M.B. A.V. and S.D.M.J. developed the self-absorption correction algorithms. V.M. was responsible for the hollow-fibre dense ceramic membrane preparation and for the Scanning Electron Microscopy (SEM)/Energy Dispersive X-ray Spectrometry (EDS) sample preparation and measurement. P.C. was responsible for the BCFZ XAFS data analysis. C.L.B. was responsible the ISIS neutron powder diffraction data collection. The analysis of the BCFZ neutron powder
Diffraction data was performed by A.V. with assistance from D.M., C.L.B., S.D.M.J. and A.M.B. A.V., D.M., S.D.M.J. and A.M.B. are responsible for writing the manuscript with feedback given by all contributors. A.M.B. directed the research.

Declaration of interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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