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Assesment of (Mn,Co)₃O₄ powders for possible coating material for SOFC/SOEC interconnects

D. Szymczewska¹, S. Molin², V. Venkatachalam², M. Chen², P. Jasinski¹, P.V. Hendriksen²

- ¹ Faculty of Electronics, Telecommunications and Informatics, Gdansk University of Technology, ul. Narutowicza 11/12, 80-233 Gdansk, Poland
- ² Department of Energy Conversion and Storage, Technical University of Denmark, Frederiksborgvej 399, 4000 Roskilde, Denmark

E-mail: author: sebmo@dtu.dk

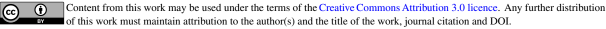
Abstract. In this work (Mn,Co)₃O₄ spinel powders with different Mn:Co ratio (1:1 and 1:2) and from different commercial suppliers are evaluated for possible powder for production of interconnect coatings. Sinterability of the powders is evaluated on pressed pellets sintered in oxidizing and in reducing/oxidizing atmospheres. For selected powder, coatings are then prepared by the electrophoretic deposition method on Crofer 22 APU stainless steel coupons. Effects of dispersant/iodine content and deposition voltage and times are evaluated. Thickness as a function of deposition parameters is described. Results show that with appropriate powder it is possible to produce adherent protective coating with a well-controlled thickness.

1. Introduction

High temperature corrosion of interconnects in Solid Oxide Fuel/Electrolysis stacks is one the most important degradation phenomena for operating stacks [1,2]. Oxides formed on steel interconnects have low electrical conductivity so over time they contribute more and more to the ohmic area specific resistance of the interconnect and thus the full stack [3–5]. Additionally, chromia, formed on the interconnect, can at high pO_2 and in the presence of steam evaporate and poison the oxygen electrode causing more degradation [6–8].

In order to mitigate high temperature corrosion, protective coatings are being developed [9–13]. Coatings can be in general divided into two categories: ones lowering corrosion rates and ones lowering chromia evaporation. For lowering corrosion rates reactive elements (e.g. Y₂O₃, CeO₂) were proven to be very successful whereas for mitigation of chromia evaporation one of the most promising materials reported up to date is the manganese cobalt spinel (Mn,Co)₃O₄ (abbreviated as MCO) [14–17]. It offers good electrical conductivity at working temperatures, good thermal expansion coefficient match to the steel and other cell components. Additionally it is very protective towards chromium diffusion – it blocks evaporation of chromia. One of the most important issues for fabrication of a well performing coating is its deposition method and sintering/thermal treatment.

In the available literature several different methods have been reported: electrophoretic deposition [18], physical vapor deposition [19], plasma spraying [20] and other [21–23]. In this work the powders are evaluated for possible use in the electrophoretic deposition [24,25] method developed in our group [26]. Electrophoretic deposition (EPD) technique is quite fragile in terms of powder requirements and



its high quality. Additionally, as the volatility of powders is not certain due to producer's changes, it seems vital to evaluate different available powders to find possibly the best one and its alternatives.

2. Experimental

2.1. MCO powders.

Five different powders produced by four suppliers were tested. Main difference between powders was the Mn:Co elements ratio, as is shown in Table 1. Ratio 1:1 has been purchased from American Elements and Fuel Cell Materials. Ratio 1:2 from Fuel Cell Materials, Marion Technologies and CerPoTech companies. ID of particular powders will be used throughout the text.

Scanning electron microscopy (SEM) pictures and Energy Dispersive X-ray Spectroscopy (EDS) data of the powders were obtained using Hitachi TM3000 with Energy Dispersive X-ray Spectrophotometer Bruker Quantax 70. X-ray diffractometry (XRD) measurements of the powders were made using X-ray diffractometer Bruker D8.

Table 1. Studied MCO powders.

Powder source	Molecular formula	Mn:Co ratio	ID
American Elements	Mn _{1.5} Co _{1.5} O ₄	1:1	AE
Fuel Cell Materials	$Mn_{1.5}Co_{1.5}O_4$	1:1	FCM11
Fuel Cell Materials	$MnCo_2O_4$	1:2	FCM12
Marion Technologies	MnCo ₂ O ₄	1:2	MT
CerPoTech	$MnCo_2O_4$	1:2	CPT

2.2. Sinterability.

From AE, FCM11, FCM12 and MT powders pellets were prepared and sintered in four different conditions. Temperatures of 900°C and 1000°C were used. For each temperature two conditions were tested (listed in Table 2). First sintering procedure consisted of only one step – sintering in air. Second procedure had two steps – in the beginning a reducing atmosphere (H₂/N₂) atmosphere was used. After furnace cooling reduced pellets were subjected to sintering in air as a second step. This kind of sintering, first in reducing and then in oxidizing atmosphere is often used for the spinel powder and is termed "redox" sintering.

Table 2. Sintering conditions.

Temperature [°C]	Cooling/heating rate [°C/h]	Sintering time [h]	Atmosphere
900	120	2	Air
1000	120	2	Air
900	120	2	H_2/N_2
	120	2	Air
1000	120	2	H_2/N_2
	120	2	Air

Sintered pellets were measured geometrically to calculate shrinkage and the density of each pellet. Information about density was received also from Archimedes measurements. SEM and EDX



examinations of pellets polished cross-sections were made using Hitachi TM3000 with Energy Dispersive X-ray Spectrophotometer Quantax70.

2.3. Electrophoretic deposition of American Elements powder.

For EPD only American Elements powder was used. For substrates for the deposition, stainless steel coupons were prepared. These were cut from a 0.3 mm thick Crofer 22 APU steel sheet and were subsequently cleaned in ethanol. For the deposition, ethanol and isopropanol in volume ratio 1:1 were mixed as solvents for each suspensions. Three different suspensions were made with varying I_2 content. I_2 amount was different in each suspension: 0.1 wt.%, 0.2 wt.% and 0.3 wt.% of I_2 was used (wt.% in relation to powder weight). Powder content in the suspension was 2 wt.%. Influence of deposition time on layer thickness was checked. Layers were deposited for 1 min, 1.5 min and 2 min. A voltage of 60 V was used for deposition of this set. Additionally, the influence of the voltage during the deposition was also examinated. For this purpose solution with 0.2 wt.% I_2 was used and 2 min deposition time was chosen. Deposition voltages of 30 V, 40 V, 60 V, 80 V and 120 V were evaluated.

Deposited layers were sintered in two steps (redox sintering): 900°C in H₂/N₂ atmosphere for 2 hours, and after furnace cooling again at 900°C for 2 hours in air. Sintered layers were examined using Hitachi TM3000 with Energy Dispersive X-ray Spectrophotometer Quantax70. XRD measurements were made using X-ray diffractometer Bruker D8.

3. Results

In the first step the powder morphology and composition was checked by means of SEM and EDS analysis. This was done in order to ensure powder composition and quality.

SEM analysis of loose powder (attached to a carbon tape) is presented in Figure 1. All powders have fine, less than 1 micrometer grains. Powders produced by CerPoTech, coming from a spray drying process have a core-shell structure and to be suitable for the electrophoretic deposition would require an additional grinding/milling step. Therefore this powder has been excluded from further (sintering) analyses.

As provided by the manufacturers, Mn:Co ratio in the powders should be either 1:1 or 1:2 depending on the composition. All tested powders roughly fall in the expected range with a small and constant deviation, that all powders are slightly more Co rich then the original ratio. About 5-10 at.% enrichment in Co content is measured for all powders.

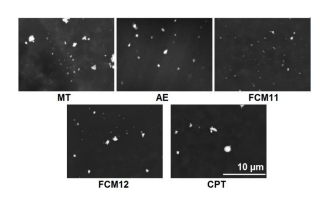


Figure 1. MCO powders

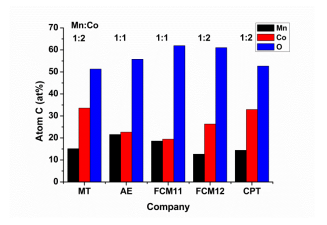
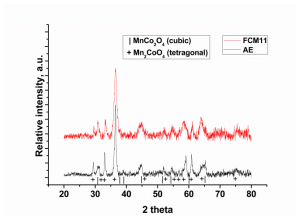


Figure 2. Atomic composition of different asdelivered powders as determined by EDS analysis.

After the SEM/EDS analysis, X-ray diffractometry was performed to check phase purity of the powders. Although no impurities (no foreign elements) were detected by the EDS analysis, still it is

interesting to see phases present. Spinel with a nominal cation ratio of 1:1 is at room temperatures a mixture of a cubic MnCo₂O₄ and tetragonal Mn₂CoO₄ phases. This has been confirmed by XRD analysis of powders FCM11 and AE shown in Figure 3. Spectra are quite noisy with broad peaks which might indicate a small crystalline size and/or only partial crystallization of the powders that might require further thermal processing upon heating of the spinel powder, only single cubic phase remain at 800°C. On the other hand, XRD analysis of CPT, FCM12 and MT (Figure 4) powders revealed only a single phase of cubic spinel. In all cases peaks are narrower. No impurity phases were found in any of powders.



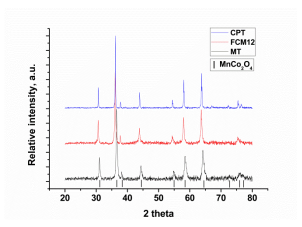


Figure 3. XRD spectra of powders with Mn:Co ratio of 1:1: AE and FCM11.

Figure 4. XRD spectra of powders with Mn:Co ratio of 1:2: CPT, MT and FCM12.

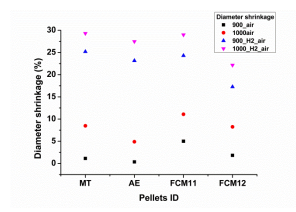
3.1. Sinterability

After ensuring powders quality, its sinterability was evaluated. Pellets were pressed from MT, AE, FCM11 and FCM12 powders. Pellets were then sintered in 4 different conditions (two temperatures and two sets of atmospheres). As a measure of sinterability, linear shrinkage, calculated density (based on pellet geometrical shrinkage) and measured density (by Archimedes principle) were used. Linear shrinkage of the pellets is presented in Figure 5. Geometrical measurement can be influenced by several factors: cracks or voids inside, parts of the pellet cracked away etc, so more reliable are the values from the Archimedes measurement and these are presented in Figure 6.

For sintering in air, there is a clear difference between the temperatures of 900°C and 1000°C. MT and AE powders have shown little sintering at 900°C and FCM 11 and FCM12 powders have shown visibly higher shrinkage (~5 %). For sintering at 1000°C, all powders sintered better than at 900°C with shrinkages in the range 5-12 %. Powders FCM11 and FCM12 again have shown superior shrinkage.

Sintering in reducing and then oxidizing atmosphere provided samples with much lower porosities. Even at 900°C samples are less porous then samples sintered in air at 1000°C. After sintering with the redox procedure, density of MT, AE and FCM11 pellets were very similar. Shrinkage was ~25 %. Only pellet prepared from the FCM12 powder reported density and shrinkage lower than expected both for 900°C and 1000°C.





900_air 1000air 900_H2_air 1000_H2_air Density (g/cc) мт FCM11 FCM12 ΑE Pellets ID

Figure 5. Measured shrinkage of pellets prepared from different powders and sintered at different conditions.

Figure 6. Measured density (by Archimedes method) of pellets prepared from different powders and sintered at different conditions.

Results of shrinkage and density analysis of sintered pellets were further compared with SEM images of pellets cross sections. Cross section images from MT, AE, FCM11 and FCM12 pellets sintered in all 4 conditions are shown in Figure 7. For samples sintered only in air, there is a visible difference in porosities of samples sintered at 900°C and 1000°C. Also grain size increase is noticed, especially from both FCM powders.

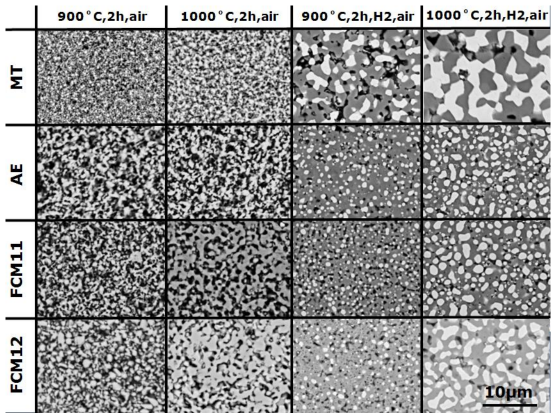


Figure 7. SEM cross sections of polished pellets prepared from different powders and sintered at different conditions.



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SEM images of samples sintered using the "redox" procedure show an interesting microstructure consisting of two phases (bright and dark phases). Apparently, the darker phase is the spinel, whereas the brighter phase is a still reduce powder. After sintering in reducing conditions a mix of Co and MnO is formed that upon reoxidation should transform back to the oxide. However, due to kinetic limitations and porous structure of the samples, reoxidation did not occur fast enough. Grain size in MT and FCM12 powders (both powders with a pure cubic phase composition) was higher than for the other two powders with a mixed composition. It seems that at 900°C there is less metallic phase left after oxidation than at 1000°C. This might also indicate that the reduction conditions were not sufficient to fully reduce the powder. To fully oxidize samples longer exposure times would be needed. Due to importance of the sintering step and possible influence of the sintering atmosphere, temperature and time these studies will be continued in the coming future.

3.2. EPD of American Elements powder

3.2.1. Different I_2 contents and deposition time

In order to deposit powders via electrophoretic deposition, particles in the suspension must be given some electric charge. For this purpose in this study I₂ is added to the suspension. Different amount of I₂ has to be added to powders with different surface area etc. For the AE powder amounts of 0.1 wt.%, 0.2 wt.% and 0.3 wt.% were tested. Samples were deposited at 1 min, 1.5 min and 2 min. After deposition coatings were sintered using a "redox" procedure at 900°C. SEM of the as-sintered coatings are shown in Figure 8.

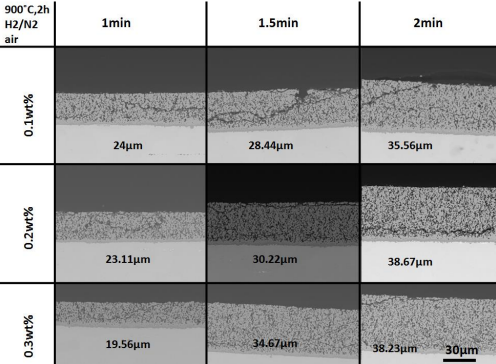


Figure 8. SEM polished cross sections of EPD coatings prepared from AE powder with different iodine content and varying deposition time. Thickness values measured from SEM pictures.

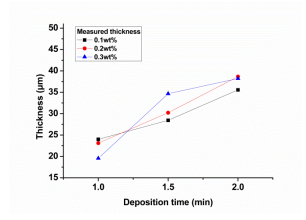
For coatings prepared on steel, there seems to be no unreacted reduced phase present after the oxidation stage of the sintering. It might be due to a relatively low thickness of the coating (~30 µm instead of few mm for pellets) which would facilitate gas exchange due to diffusion.



Thickness of the produced coatings, has been evaluated from the SEM images (Figure 9) and additionally, just after the deposition from the weight gain of sample coupons (Figure 10). For this a 30% porosity was included into calculations. Thickness calculations from the weight gain seems to be representative for real coating thickness and therefore can be used as an assessment of sample quality without the need to destroy the sample.

For the amount of dispersant of 0.1 wt.% and 0.2 wt.% the thickness vs. deposition time is very similar. Whereas for 0.3 wt.% for 1 minute coating thickness is lower than anticipated. One possible reason is that it has been too much of I₂ and some of it reacted at the electrodes and then the amount was reduced and for longer times results were similar for all studied concentrations. It seems from the images, that for higher iodine content, produced coating is less cracked and of higher quality. For 0.3 wt.% of iodine almost no cracks are visible. Looking at the results, using 0.2 wt.% or 0.3 wt.% with the AE powders seems sufficient to ensure homogenous coating.

By controlling the time of deposition coatings with thicknesses ranging from 20 µm to 40 µm might be produced.



0.1wt% 0.2wt% 45 40 Thickness (µm) 35 30 25 20 1.0 1.5 2.0 Deposition time (min)

Figure 9. Thickness of the coatings as measured from the SEM cross section images.

Figure 10. Thickness of the coatings as measured from weight gain after the EPD deposition.

After evaluation of coatings fabrication parameters, cross sections were analysed in more details by SEM/EDS analysis to check chromium diffusion and oxide layer thickness formed on steel during the "redox" sintering procedure. Samples analysed were samples prepared with 0.1, 0.2, 0.3 wt.% iodine and the deposition time was 1 minutes. Results of the analysis are presented in Figure 11. For all samples 3 regions of interest were analysed. "1" is the alloy, analysed just to ensure proper calibration of the EDS. "2" is the interface of the alloy and the coating, where the oxide (Cr₂O₃ and (Mn,Cr)₃O₄ spinel should occur). "3" is the bulk of the coating, far away from the interface. The composition of points number "1" is for all samples quite similar and close to the expected composition (between 21 and 24 wt.% as given by the producer of steel). At the interfaces of all samples, Cr rich phase is detected. Due to spatial limitation of the EDS detector it is not possible to clearly distinct the chromium oxide from the bulk of the alloy. Also Mn,Co spinel is detected in point "2". In points "3", major elements present are Mn and Co in the 1:1 ratio as expected for the AE powder. For all samples also Fe and Cr are detected in the coating. Up to 6 wt.% of Cr is present in the coating after sintering. This amount seems quite high. One possibility is that Fe and Cr diffused into the coating during sintering. However, the time was quite short, in total 4 hours at 900°C so maybe metallographic preparation of the samples was not careful enough and some alloy particles were embedded in the coating during polishing. Certainly this will be studied further.

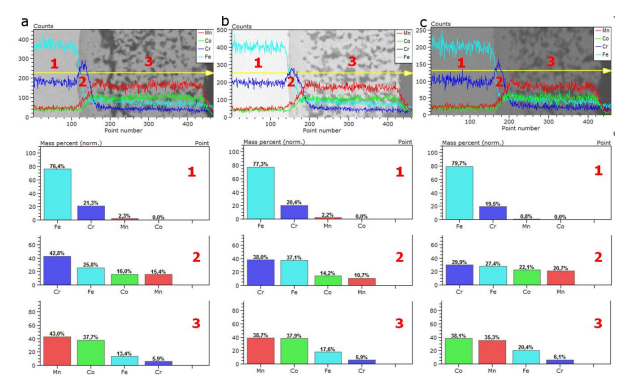


Figure 11. SEM and EDS line profiles of Mn, Co, Cr and Fe on the sintered coatings. Samples with a) 0.1 wt%, b) 0.2wt%, c) 0.3wt% of I₂.

4. Conclusions

In this study several commercial spinel powders were evaluated for their phase purity, composition and sinterability. Four different sintering conditions were used. Results show that sintering using a "redox" procedure results in much denser pellets. For pellets, reduction and/or reoxidation step seems to be kinetically/diffusion limited so that large amount of the reduced phase remain in the pellet after "redox" sintering. This is not present in the coatings prepared on stainless steels.

Electrophoretic deposition method has been evaluated for producing coatings with different dispersant content. For iodine content >0.2 wt.% crack free and homogenous layers were prepared. By finding suitable solvent, dispersant content and deposition parameters (voltage, time) protective coatings for steel interconnects can be produced reproducibly by the EPD method.

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