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Towards a better understanding of structure-performance relation in PEMFC fuel cells based on ptychography X-ray nanotomography and scanning small angle X-ray scattering

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Motivation

- Gaining fundamental insights on the structure of advanced composite materials is challenging, but essential for their role in respect to future energy challenges
- This project explores the possibility to combine high resolution imaging with coherent X-rays (Ptychographic X-ray Tomography PXCT) together with SAXS (small angle X-ray scattering) imaging to obtain full scale models on multiple length scales (nano- to millimetre)
- The analysis approach is complemented by additional methods such as X-ray fluorescence performed simultaneously to SAXS imaging

Methods

SAXS@7m det. distance

g/nm

Pilatus 2M

 $\phi_1 \phi_2 =$

 $2\pi^2(\Delta\rho_{SLD})$

Ptychography X-ray Tomography

- PXCT measures far-field diffraction patterns from a sample that is moved across a spatially confined coherent beam in a way that illumination areas overlap. This is carried out for multiple sample orientations
- An iterative phase retrieval algorithm together with a tomography reconstruction is able to reconstruct the electron density (ED) distribution in 3D



Materials and electrochemical performance

Polymer electrolyte membrane fuel cells (PEMFCs) generate electricity by electrochemical reaction that take place in a complex porous material (catalyst layer) with 3 components

Catalyst layer sprayed on Nafion²²¹

Three components:

1.) Support: Carbon (Vulcan XC72R)2.) Binder: Ionomer (Nafion)

[1] http://www.jaist.ac.jp/ms/labs/nagao-www/wp-content/uploads/2018/07/20180704labguide2018.pdf

10 ¹

10⁰

g / nm ⁻¹







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 Samples were measured under cryo-conditions with OMNY^[1] at 6.2keV; 25-40µm sample diameter; 3D half-period resolution of 24-34nm estimated by Fourier shell correlation

2D SAXS & XRF imaging

Isotropic scattering

Catalyst sprayed on

Data merged from

three detectors

Nafion²²¹

- Scanning SAXS measurement at 11.2keV were performed with a local resolution of 10µm, scanning 1x1mm² for all 3 catalysts at two sample-detector distances (2m & 7m)
- X-ray fluorescence was collected simultaneously to complement the scattering data at each position



3.) Active sites: Pt nanoparticle (~3-4nm)



Samples with three different Ionomer/Carbon (I/C) ratios:

	Pt / wt%	lonom. wt%	Carbon wt%	SLD [10 ⁻⁶ Á ⁻²]
I/C 0.2	16.8	13.8	67.4	39.1
I/C 0.54	13.6	30.5	52.9	34.8
I/C 0.95	11.1	43.2	45.7	31.5
ED [e/Å ³]	5.167	0.593 (dry)	0.57-0.63	

0.70.60.50.60.60.50.60.50.60.50.60.50.60.50.60.50.60.50.60.50.60.60.60.50.60.50.6

Flow: 0.2 L/min H_2 and O_2 ; 100%RH; OCV vs 0.3V; 80°C 5cm² cell



10⁻⁸

10-10

10⁻²

 10^{-1}

Input integral or Q-invariant • XRF spectra

- 3 Pt and S line at 2...2.3keV overlap
- Peaks fitted by linear combination of pure materials





Compare and discuss results

XRF maps

- Amplitude histogram for both elements show the expected increase in S with increasing ionomer concentration.
- Pt peaks do not overlap, contrary to expectation, however sample self-absorption correction is still missing which may solve this

Pore size distribution

 Pore size distribution is segmented for all samples from PXCT by threshold segmentation (ED < 0.1 Å⁻³)



Outlook

- SAXS: Full q-range modelling of data, morphological model for micropores and additionally feed models with DFT results from Ar gas adsorption results
- XRF: Correct Pt & S amplitudes by sample selfabsorption
- PXCT: Gradient based segmentation for pores, support and ionomer
- Modelling: Use SAXS, XRF & PXCT results to



 Radius of gyration, assuming spherical shape, matches well with the smallest pore size accumulation of PXCT distributions

Pore volume fraction (PVF)

- PVF for full PXCT is quite different from SAXS results
- SAXS picks up pores from 1...100nm (q-resolution) and agrees with results for small pores < 120nm PVF (PXCT) quite good
- PVF from all pores shows strong increase for I/O 0.2 (good performing) & I/O 0.54 (best performance), missing feature for I/C 0.95 (worst performance) catalyst layer





generate 3D models based on elemental distribution, pore volume fraction and pore size distribution

Summary

- SAXS & PXCT provide data on different length scales, but with a well resolved overlap
 SAXS & XRF maps show homogeneous layers on macroscopic. length scales (µm to mm)
- Element distribution as expected from chemical composition
- Pore volume fraction indicates that large pore network may be responsible for performance decrease of I/C 0.95 (mass diffusion limitations); PXCT ED reconstruction also show dense electron density

Acknowledgement

References

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