In situ neutron diffraction under high pressure - providing an insight into working catalysts

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Abstract
In the present work the construction and application of a continuous flow cell is presented, from which neutron diffraction data could be obtained during catalytic reactions at high pressure. By coupling an online gas detection system, parallel structure and activity investigations of working catalysts under industrial relevant conditions are possible. The flow cell can be operated with different feed gases in a wide range from room temperature to 330°C. Pressures from ambient up to 60 bar are applicable. An exchangeable sample positioning system makes the flux cell suitable for several different goniometer types on a variety of instrument beam lines. Complementary operational test measurements were carried out monitoring reduction of and methanol synthesis over a Cu/ ZnO/ Al2O3 catalyst at the high-flux powder diffraction beamline D1B at ILL and high-resolution diffraction beamline Echidna at ANSTO.

Keywords: In situ characterization; Heterogeneous catalysis; High-pressure sample environment; Structure-activity correlation

1. Introduction
Nowadays catalyst are considered as dynamic materials whose active centres can be formed or transformed due to the chemical potential of reactants or products under reaction conditions. If such changes are reversible, application of in situ methods is needed to study catalysts in their working state to gain a general understanding of structure-activity relationships. It is especially attractive to bridge the so called ‘pressure gap’ and to go to pressure ranges beyond Ultra-High-Vacuum to ambient pressure regimes. Unfortunately, not many in-situ techniques can be operated at high pressures above ca. 50 bar and allow a direct observation of the working catalyst under realistic chemical potentials as are present in industrial reactors. It often remains questionable, if the properties of model catalysts studied at low pressure can be extrapolated to real catalyst under industrial reaction conditions. Due to their high penetration depth, neutrons allow application of complex sample environment as is needed to study commercial catalyst under industrial reaction conditions, e.g. elevated temperatures and high pressures (up to 60 bar) under strongly reducing gaseous atmospheres like hydrogen/ deuterium-rich synthesis gas. Furthermore neutron diffraction is a powerful tool to study structural and microstructural properties of a catalyst (phase identification, strain, particle size, alloy formation, phase transformations) in operation. A lot of technical effort was made by Turner et al. [1] and Walton et al. [2] to study catalyst or related materials under demanding reaction conditions; but still far away from typical industrial conditions. In this present contribution a reaction setup will be presented, which allows carrying out in situ neutron diffraction studies on various catalyst systems under industrial relevant synthesis conditions.

2. Apparatus design
Aim of the apparatus design was to build a safe reactor, which allows collecting structural data of a working catalyst under industrially relevant conditions with neutron di/r action and a parallel monitoring of the product gas stream by mass spectroscopy to correlate structural and catalytic properties.
The apparatus consists of three basic components: The flow cell including the heated reactor body, the gas supply and the effluent gas analytics.

2.1. Flow cell and reactor body

The operation of a flow cell under high pressure is devoted to strict safety regulations. According to these regulations, a bursting of the cell walls should be excluded at any time of the operation. The most critical point is the balance in between finding a material, which shows a moderate coherent scattering behavior on the one hand and is resistant to high pressures of reactive gases like hydrogen, deuterium or carbon monoxide at elevated temperatures on the other hand. Most common materials used in the nuclear branch like vanadium and zircalloy show hydrogen embrittlement and/ or intragranular corrosion [3], if they are operated under high pressures on a long term scale. Especially under alternating high pressure cycles, which occur in situ conditions a high tensile yield strength of the material must be guaranteed. We have decided to fabricate the cell from thick-walled aluminum-magnesium (AlMg3) offering sufficient pressure and chemical resistance, low absorption and activation and still acceptable coherent scattering. The tubular flux cell was manufactured from a AlMg3 rod with a nominal outside diameter of 20 mm and a tensile yield strength of 283 N mm\(^{-2}\) at room temperature. Using a lathe, the outer diameter was reduced to 19,05 mm and a hole with a diameter of 10 mm was set through, which led to an effective wall thickness of 4,52 mm. Strain calculations by assuming a tensile yield strength of 98 N mm\(^{-2}\) at 300°C and an increased safety factor of 3,6 [4] have shown, that the tube is resistant up to the conditions of 140 bar at 300°C. Static load tests of the setup have been successfully conducted at 90 bar and 100°C for 2 h and the limits for flow operation have been set to 60 bar at 330° C. The total length of the tubular reactor was 150 mm and the catalyst bed bathing in the neutron beam can have a length of up to 70 mm resulting in a volume of up to 5,5 cm\(^3\). To achieve high intensity of the neutron diffraction patterns at short counting times, large sample sizes are required. The cell can be loaded with variable sample amounts from approximately 5g to 20 g. The loaded catalyst bed is fixed by quartz wool plugs, which are inserted from both ends. A thermocouple which is inserted from the top allows measuring the bed temperature in the core of the catalyst bed during the reaction. Reactant feed is injected from the top, product stream flows out at the bottom. Both ends of the tube are supported by cramped stainless steel adapting sleeves to assure a self-tightening seal. By inserting the end of the flux cell into Swagelok\(^{TM}\) stainless steel (SS 316) 3/4 to 1/4 inch reducing unions the cell material forms a tight seal in between the adapting sleeves and the inner mating tape of the reducing unions by its larger thermal expansion coefficient at 250°C.

The body of the reactor is also made of AlMg3, due to its good heat capacity and corrosion resistance. If the incident neutron beam is poorly collimated, the reactor body shows low activation behavior and shows a good radiation damage resistance [5]. Given that the body is made out of the same material as the flux cell itself, it is practically seamless on the diffraction pattern. Pedestras and sampling base plate are made out of SS316 and fixed with screws via threading. Six heating cartridges are inserted into holes in the reactor body with a total heating power of 600W (2x 150 W, 4 x 75W), enabling heating rates of up to 5 K min\(^{-1}\). Each heating element is equipped with a thermocouple to check its heating behavior for linearity and overheating. Loading of the filled flow cell is performed by removing the frontal heating covers and inserting the cell into the notch. The installed system with a total weight of...
range from 0 to 100 bar. The system pressure was electronically measured with an Endress + Hauser PMP 131 pressure transducer which was connected to a Schwille SPE 670 digital display and linked with a serial cable to a Labview application which allowed automated read-out and data-recording. For additional safety reasons a rupture disk with a specified relief pressure of 85 bar and a check valve was installed between the outlet after the pressure transducer and the reactor inlet, which was able to shut down the gas supply in the case, the flow exceeded 500 Nml min$^{-1}$ (e.g. in case of a rupture). The pressurized product lines can be heated to 150°C-170°C to avoid condensation of products like steam.

### 2.3. Effluent gas analytics

Gas analytics is performed online at the heated product line beyond the back pressure regulator at atmospheric pressure. By switching the gas flow between bypass or reactor cell, the syngas composition or the effluent gas from the reactor can be analyzed (e.g. for calibration). A gas chromatograph or a mass spectrometer can be coupled to the system. We have used the latter on site during the in situ experiments and the former in the laboratory to quantitatively study the system at the same conditions. During the neutron diffraction experiments a Pfeiffer Vacuum ThermoStar Mass spectrometer was used to check the progress of catalyst activation and whether the expected outlet gas composition was reached. Effluent gases could be collected after online analysis in a condenser vessel or released into the venting system of the neutron facility.

### 3. Application example: Methanol synthesis

A prominent example for the importance of in-situ characterisation of structure-activity relationship in heterogeneous catalysis is the methanol synthesis over Cu/ZnO/Al$_2$O$_3$ catalysts. Even though these catalyst (in different compositions) have been used in commercial methanol synthesis for 45 years, the so-called synergy of Cu/ZnO is still under debate in literature. Several models have been introduced [6] which should give a first approach to properties of operating industrial catalyst systems. Some of the observations made on this system like brass formation, dynamical undergoing of morphological changes [7] have been directly obtained only on model catalysts under conditions, where no methanol has been produced. In the present work, the structural properties of the aforementioned industrial catalyst under realistic industrial synthesis conditions is studied using the flow cell reactor system described above. To minimize the effect of incoherent scattering, hydrogen was replaced by deuterium in the reaction gases.
3.1. Catalyst activation in high-flux diffraction

Due to their pyrophoric nature, nano-structured Cu/ZnO/Al2O3 catalysts are handled in their completely oxidized form, i.e. as CuO/ZnO/Al2O3, and the first step of a methanol synthesis experiment is the activation of the catalyst by reduction of the CuO component to metallic Cu:

\[
\text{CuO/ZnO/Al}_2\text{O}_3 + D_2 \rightarrow \text{Cu/ZnO/Al}_2\text{O}_3 + D_2\text{O}
\]  

To study the phase evolution during reduction a commercial CuO/ZnO/Al2O3 catalyst the beamline D1B at ILL in Grenoble was used. While requesting the highest available neutron flux for a sufficient time resolution of the experiment, a focusing highly ordered pyrolitic graphite monochromator was used, to get a wavelength of \(\lambda=2,52\text{Å} \) which lead to an effective flux of \(6,5\times10^9 \text{ n cm}^{-2} \text{ s}^{-1} \) at the sample [8]. By setting the 3 He/Xe position-sensitive detector to a take-off angle of 45,06° the angular range up to 124,86° 2θ (corresponding Q-range 1,91Å^{-1} to 4,2Å^{-1} ) was covered. The reduction of 6g catalyst was carried out with a feed stream of 100 Nml min^{-1} pure D2, bed temperature was ramped from 28°C at 1 K min^{-1} to 250°C at ambient pressure. Effluent gas composition was tracked by mass spectrometry from mass 0 to 50 in analog scan mode; the probing cycle was 11 second per spectrum. During the reduction procedure 250 patterns were acquired with an acquisition time of 5 min per pattern. The patterns were normalized to monitor counting rate. Afterwards the intensities of the CuO(11-1) at Q= 2,4925Å^{-1} and Cu(111) at Q= 3,0122Å^{-1} peaks were fitted using a pseudo-Voigt peak shape function. After normalizing them to the highest intensity they were plotted on an absolute timescale. The normalized integrated intensities of the Cu(111) and CuO(11-1) peaks are correlated to the catalyst bed temperature and the effluent gas composition in Fig. 3.

3.2. Working catalyst in high-resolution diffraction

High-resolution diraction was performed on ECHIDNA at ANSTO by using a Ge(335) monochromator at a angle of 70°, giving a highly collimated beam at a wavelength of \(\lambda=1,622\text{Å} \). A large array of 128 position sensitive 3 He detectors allowed to cover a angular range of \(4°<2\theta<164° \) which corresponds to a Q-range of 0,28Å^{-1} to 7,7Å^{-1} [9][10]. The reduction of the catalyst was carried out on-site at the diffraction experiment under the same conditions as in the highflux experiment. After reaching the temperature plateau at 250°C, the feed is switched to syngas consisting of \(\text{D}_2\), \(\text{CO}_2\), CO and Ar as internal standard and the flow cell is pressurized by the back pressure regulator with a rate of 0,78 bar min^{-1}. The methanol synthesis reaction, formally according to

\[
\text{CO}_2 + 3 \text{D}_2 \rightarrow \text{CD}_3\text{OD} + \text{D}_2\text{O}
\]  

was conducted in thermodynamic equilibrium at 250°C and 60 bar. After reaching stable operating conditions the effluent gas composition is monitored by mass spectrometry. At stable equilibrium composition of the effluent gas, diffraction patterns with 1 or 2 hours acquisition time were recorded. The observed intensities in the monitor- and efficiency normalized patterns were evaluated by multiple peak-fitting to account for sample and cell-material contributions.

4. Results

With increasing temperature the reduction of the CuO-containing precursor phase is initiated around 115°C and finishes at 180°C. Metallic Cu appears in the diffraction pattern at 164°C. \(\text{D}_2\) consumption starts around 100°C and ends in a regime, were \(\text{CuO}\) is completely reduced. Additionally to the moderate angular resolution of D1B, the poor crystallinity of all component in the nano-structured catalysts, in particular of \(\text{ZnO}\) and \(\text{Al}_2\text{O}_3\) contributes to a relatively high background, which leads to a larger uncertainy of the low-intensity and broad peaks of \(\text{CuO}\) compared to the more crystalline metallic Cu.
conversion of D$_2$ was reached within the maximum of D$_2$O desorption in coincidence with metallic Cu evolution, which is in good accordance with literature [11]. No indication of intermediate formation of crystalline Cu$_2$O has been observed in the diffraction patterns, but the asymmetric shape of the curves in Fig. 3 and the inception at a normalized intensity > 0.5 suggest the presence of a undetected, probably amorphous intermediate, which may be a form of Cu(I)-oxide. Fig. 4 shows a diffraction pattern of a commercial CuO/ZnO/Al$_2$O$_3$ catalyst in operation under 250°C and 60 bar pressure. Strong diffraction peaks from the cell material can be seen and are highlighted with green profiles. The peaks of the catalytically active copper phase of the catalyst (red profile) can be clearly distinguished and indexed. The scattered intensities are fitted by a pseudo-Voigt peak shape function for further evaluation. Such patterns can serve as a starting point to investigate the structural answer of a working catalyst to variation of the reaction conditions and to correlate such changes to catalytic performance. More detailed results of the methanol synthesis catalysts will be published elsewhere.

5. Discussion

A flow cell for in-situ neutron diffraction during continuous catalytic experiments under high pressure was designed and constructed. It was successfully tested for catalyst activation and methanol synthesis over CuO/ZnO/Al$_2$O$_3$ under equilibrium conditions at 250°C and 60 bar while obtaining structural information of the catalyst. Online effluent gas analytics allows direct correlation of structural properties with catalytic activity. Although there is a strong scattering signal from the flow cell wall material, the structural signature of the investigated catalytic system is strong enough to give detailed results concerning the crystal- and microstructure of a catalyst under industrially relevant reaction conditions.

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References