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Depleted uranium catalysts for chlorine production†

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This study demonstrates depleted uranium as a remarkable heterogeneous catalyst for the oxidation of HCl to Cl₂. This reaction comprises a sustainable approach to valorise byproduct HCl streams in the chemical industry. Bulk α -U₃O₈ showed an outstanding stability against chlorination, which is crucial for its durability in catalytic tests. UO₂ and γ -UO₃ transformed into α -U₃O₈ under reaction conditions. Uranium deposition on different carriers by dry impregnation concluded the superiority of zirconia as support. HAADF-STEM investigations revealed that the uranium oxide on the surface of this carrier is present in the form of a film-like nanostructure with a thickness ranging from a monolayer to 1 nm as well as atomic dispersion. The effect of variables (temperature, feed O₂/HCl ratio, metal loading, and Cl₂ cofeeding) on the performance of U₃O₈/ZrO₂ has been studied. The HCl conversion over this catalyst increased with reaction time as a likely consequence of *in situ* re-dispersion of the original uranium phase into atomically dispersed UO_x. As demonstrated by H₂-TPR, the uranium in the generated UO_x phase is more oxidised than in the original U₃O₈. Such a highly dispersed active phase is produced faster in the uncalcined sample. The extraordinary stable Cl₂ production over U₃O₈/ZrO₂ at 773 K for 100 h on stream indicates its potential for application in high-temperature HCl oxidation. Under these conditions, other known catalytic materials suffer from significant deactivation.

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Introduction

Uranium compounds have been used as heterogeneous and homogeneous catalysts. 1,2 Their suitability for redox reactions is related to the wide range of oxidation states that uranium can assume (from II to VI), which in turn derives from the ability of its 5f-electrons to hybridise.3 Specifically for the heterogeneous catalysis field, uranium oxides (mostly U₃O₈) have been recognised since the 1920s for reactions of industrial relevance such as the oxidation of hydrocarbons and the partial oxidation of ethanol.4-6 Later efforts extended the scope of uranium-catalysed transformations to comprise the oxidative destruction of volatile chlorinated hydrocarbons, 7,8 the oxidative coupling of ethylene, acetylene, and acetaldehyde,3 the esterification of formaldehyde,3 and NO_x reduction.9 Relevantly, uranium-based materials were once used in industry for the hydrocracking of shale oil (UO₃/Al₂O₃, UO₃/CoMoO₄)¹⁰ and in the ammoxidation of propylene to acrylonitrile (USb_xO_y) .¹¹⁻¹³

Natural uranium consists of three isotopes, 238 U, 235 U, and 234 U, in the relative abundance of 99.275, 0.720, and 0.005%, respectively. 14 238 U and 234 U are α -ray emitters, while 235 U emits both α - and low-energy γ -rays. Alpha particles are much less

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penetrating than other forms of radiation, thus rendering uranium only a little hazardous (mainly from the γ -rays). Depleted uranium (DU), which is produced as a waste in the uranium enrichment process, is even considerably less radioactive (*ca.* 0.2–0.4% ²³⁵U) and, thus, less harmful. To generate the carbon-neutral energy source, the demand of enriched uranium as a fissile nuclear fuel can be expected to increase,² which represents a strong incentive for the development of novel applications of DU.

The heterogeneously catalysed oxidation of HCl to Cl₂ (Deacon reaction)¹⁵ is an attractive route to recycle chlorine from byproduct HCl streams in the chemical industry, namely in the production of polyurethanes and polycarbonates.¹⁶⁻¹⁸ Two industrial catalysts based on RuO₂, featuring high activity at a relatively low temperature and remarkable stability, have been recently introduced: RuO₂/SiO₂/TiO₂-rutile (by Sumitomo) and RuO₂/SnO₂-Al₂O₃ (by Bayer).¹⁹⁻²⁷ The wide use of ruthenium catalysts for HCl oxidation is hindered by its high and fluctuating market price.¹⁶ This drawback triggered research efforts to develop alternative cost-effective systems. CeO₂-based catalysts represent tangible steps along this direction.^{28,29}

Uranium oxide-based catalysts for HCl oxidation have recently been patented.^{30,31} High single-pass HCl conversion at high temperature and practically negligible active phase loss have been claimed as the key characteristics of these systems. To assess the real potential of uranium-based catalysts for industrial application, further knowledge needs to be gathered. The optimal combination of active phase and support will be

derived only based on a deeper understanding of activity and stability descriptors. The catalyst performance should be then put into perspective with respect to other known catalytic systems and evaluated in an industrially relevant time frame. Herein, we systematically investigated uranium oxides in bulk and supported forms for HCl oxidation. Catalytic tests at ambient pressure in a continuous flow fixed-bed reactor combined with detailed characterisation of the catalysts prior to and after reaction have been applied to gather a solid knowledge of the Deacon chemistry of these materials.

Results and discussion

Bulk uranium oxides

As starting point, the main binary oxides of uranium were considered in this study, namely, UO2, U3O8, and UO3. The X-ray diffractograms of the solids as well as their corresponding structures are displayed in Fig. 1a and b. According to the XRD phase analysis, the catalysts were identified as uranium dioxide (JCPDS 05-0550), α-triuranium octoxide (JCPDS 31-1424), and γ-uranium trioxide (ICPDS 31-1422) with small amounts of β - and α -forms. The crystal structure of UO₂ is of fluorite type with face-centred cubic atomic arrangement. Uranium and oxygen atoms are octaand tetrahedrally coordinated, respectively. 4 α-U₃O₈, one of the two forms (α, β) of this oxide which are stable at ambient temperature,3 crystallises in an orthorhombic structure. All of the uranium atoms are coordinated with oxygen atoms forming pentagonal pyramids.^{1,14} γ-UO₃, the most stable of the seven crystalline phases $(\alpha, \beta, \delta, \epsilon, \gamma, \zeta, \text{ and } \eta)$ of this oxide,³ belongs to the tetragonal crystal system and is characterised by octa- and dodecahedral coordination of uranium to oxygen. All of the three oxides possess a very low total surface area (S_{BET} , Table 1).

The reducibility of these materials was studied under a diluted H_2 flow up to 1100 K (Fig. 1c). The reduction profile of UO_2 shows a little H_2 consumption at ca. 880 K. As the XRD

pattern of the reduced sample (Fig. S1 in the ESI†) was unaltered with respect to that of the fresh solid, this feature was attributed to the removal of oxygen species that are known to accommodate in the lattice of the fluorite structure of UO₂ upon exposure to air.¹ For α-U₃O₈, a single and broad peak centred at *ca.* 975 K was evidenced, which is assigned to the reduction of U₃O₈ to UO₂.8 The reduction profile of γ-UO₃ displays a broad signal composed by two main contributions at *ca.* 880 and 963 K, due to the transitions UO₃ \rightarrow U₃O₈ and U₃O₈ \rightarrow UO₂, respectively (Fig. 1c).³² The formation of UO₂ from both α-U₃O₈ and γ-UO₃ was confirmed by XRD (Fig. S1 in the ESI†).

These bulk uranium oxides were tested in the gas-phase oxidation of HCl at $T_{\text{bed}} = 773 \text{ K}$ and $O_2/\text{HCl} = 2 \text{ for } 3 \text{ h}$. The rates of Cl₂ production were stable at ca. 2 mol Cl₂ h⁻¹ mol U⁻¹ for UO_2 and α - U_3O_8 and ca. 3 mol Cl_2 h⁻¹ mol U^{-1} for γ - UO_3 . Normalisation of the rates by S_{BET} of the fresh samples gives the values as 7×10^{-3} , 8×10^{-3} , and 3.4×10^{-3} mol Cl₂ h⁻¹ m⁻² for α -U₃O₈, UO₂, and γ -UO₃, respectively. However, due to transformation of the latter two oxides into the former during reaction (vide infra), rates normalised by the S_{BET} of the used sample are more relevant and lead to a value of 4×10^{-3} mol Cl₂ h⁻¹ m⁻² in all cases. The dependence of the activity of these oxides on temperature was investigated between 673 and 823 K at $O_2/HCl = 2$. The reaction rate scaled linearly with the temperature in the whole range. The apparent activation energy (E_a^{app}) was estimated from the Arrhenius plots at 52, 54, and 40 kJ mol⁻¹ for UO₂, α -U₃O₈, and γ -UO₃, respectively.

The used catalysts were characterised by the same techniques applied to the fresh samples in order to assess possible structural changes upon exposure to reaction conditions. Remarkably, XRD analysis indicated the absence of chlorinated phases in any of the used catalysts. However, we observed the complete conversion of UO_2 and γ - UO_3 into α - U_3O_8 (Fig. 1b). It is suggested that such transformation is due to oxidation by the excess gas-phase O_2 for the former oxide and reduction by feed

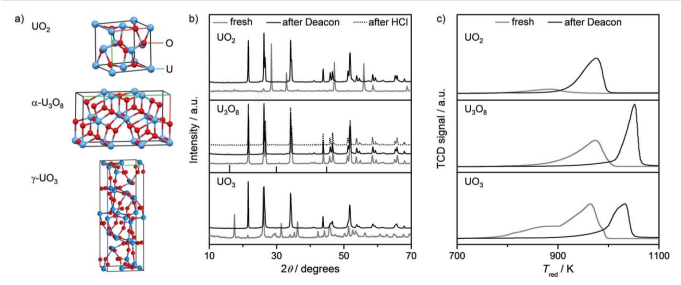


Fig. 1 Structure of the uranium oxides (a) and characterisation results from powder XRD (b) and H_2 -TPR (c) of the samples in fresh form (blue lines), after HCl oxidation at 773 K (black lines), and after HCl treatment at 823 K (dotted lines). Vertical lines at the bottom of the U_3O_8 pattern show the positions of most intense reflections of UCl₄.

Table 1 Characterisation and catalytic data of uranium-based catalysts

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Sample	$\mathrm{U}^a\left(\mathrm{wt\%}\right)$	$S_{\mathrm{BET}} \left(\mathrm{m^2 \ g^{-1}} \right)$	$r^b \pmod{\operatorname{Cl}_2 \operatorname{h}^{-1} \operatorname{mol} \operatorname{U}^{-1}}$	$E_{\rm a}^{\rm app} ({\rm kJ \; mol^{-1}})$
UO ₂	88.1	1	2.2	52
γ-UO ₃	83.1	3	2.9	40
α -U ₃ O ₈	84.8	1	2.0	54
U_3O_8/ZrO_2	9.8	$35 (47)^c$	63.5	50
U ₃ O ₈ /SiO ₂	9.5	136 (193)	45.4	46
U ₃ O ₈ /TiO ₂	9.4	30 (52)	28.6	54
U ₃ O ₈ /Al ₂ O ₃	9.6	131 (191)	27.3	57

^a Determined by ICP-OES. ^b Conditions: W = 0.5 g (bulk oxides) or 0.25 g (supported catalysts), $T_{\text{bed}} = 773$ K, $O_2/\text{HCl} = 2$, $F_{\text{T}} = 166$ cm³ STP min⁻¹, and t = 3 h (bulk oxides) or 1 h (supported catalysts). ^c Surface area of the supports in brackets.

HCl for the latter. Indeed, treatment of UO_2 and γ - UO_3 in 20 vol % O₂/N₂ at 773 K for 3 h caused the complete transformation of UO_2 into α - U_3O_8 , while it did not affect the state of γ - UO_3 (confirmed by XRD, Fig. S2 in the ESI[†]). All of the H₂-TPR profiles of the uranium oxides after reaction feature a single reduction peak, attributed to the transformation of U₃O₈ into UO₂ (Fig. 1c), in line with the identical bulk composition of the samples after HCl oxidation. The appearance of the peak at higher reduction temperature for used γ -UO₃ and α -U₃O₈ is likely related to certain degree of surface chlorination and/or sintering. With regard to the former, the bulk α -U₃O₈ catalyst after Deacon reaction was calcined in static air at 773 K for 5 h (aimed at removing surface chlorine species) and then measured by H2-TPR. A reduction profile equivalent to that of the fresh α-U₃O₈ sample was obtained (Fig. S3a[†]), which confirmed that the change in reducibility is mainly due to surface chlorination. Further, TEM of α-U₃O₈ in fresh form and after Deacon indicated a slight increase in overall particle size for the latter (Fig. S3b and c†). Since calcination of U₃O₈ after Deacon reproduced the reduction profile of the fresh sample, the effect of sintering on reducibility of U₃O₈ seems to be negligible.

α-U₃O₈ was further assessed under harsher conditions, i.e. at $O_2/HCl = 0.5$ and 0 (without gas-phase O_2) at 823 K for 2 h on stream to evaluate its resistance to bulk chlorination and metal loss. The weight of the reactor before and after the tests remained practically unchanged, suggesting no loss of uranium. Furthermore, the diffractograms of the samples after these treatments indicated the preservation of a pure oxidic phase (Fig. 1b). The endothermic nature of the penetration of Cl atoms to deeper layers (ca. 2 eV) has been already found as a key reason for the robustness of RuO2 against bulk chlorination. 22,28 Similar property could be responsible for the stability of α -U₃O₈ against bulk chlorination. In this line, chlorination of UO2 (which also revealed the absence of any chloride phase upon testing in $O_2/HCl = 0$ at 823 K for 2 h), by Cl_2 to form UCl_4 has been reported highly endothermic ($\Delta G = 148.9 \text{ kJ mol}^{-1}$).³³ Thus, bulk uranium oxide represents an exceptionally stable high-temperature catalyst for HCl oxidation. This finding is particularly striking since CuO, Cr₂O₃, CeO₂, and RuO₂ undergo structural changes at high temperatures. In particular, after testing at 823 K and $O_2/HCl = 0.5$ for 2 h, strong chlorination

was detected (XRD analysis) for the first three oxides, while RuO₂ underwent partial transformation into volatile RuO₄ (*ca.* 20 wt% RuO₂ loss). ¹⁶ It is worth noting, though, that RuO₂ is an outstanding low-temperature (473–673 K) catalyst and is extremely stable under its optimised operating conditions. ^{20,34}

Supported U₃O₈ catalysts

Based on the very promising performance of bulk α -U₃O₈, the next step of the work consisted of finding a suitable support for this uranium-based active phase. Monoclinic ZrO₂, γ -Al₂O₃, SiO₂, and TiO₂-anatase were considered as carriers. The synthesis protocol comprised dry impregnation of these oxides with a uranium precursor (in an amount corresponding to a nominal loading of 10 wt% U), followed by calcination under the same conditions applied for the preparation of bulk α -U₃O₈ (see Experimental section).

The supported U₃O₈ catalysts were screened in HCl oxidation at $O_2/HCl = 2$ in the temperature range of 673-823 K (Fig. 2a). Blank experiments confirmed that the Deacon activity of the pure carriers was negligible under the conditions applied. The HCl conversion displayed a steady increase with the temperature for all supported catalysts, reaching values comprised between 21 and 47% at 823 K. U₃O₈/ZrO₂ was the most active catalyst, followed by U3O8/SiO2 and, finally, U3O8/TiO2 and U₃O₈/Al₂O₃, which were comparably active. With respect to the bulk oxide, only the zirconia- and silica-supported materials offered improved performances (Fig. 2a). Still, as α-U₃O₈ was tested using twice the catalyst amount, a better comparison was drawn on the basis of the reaction rates per mol of U at 773 K. Accordingly, it appeared evident that any of the supports employed determined an activity enhancement, overall leading to 14-30 times higher rates (Table 1). As shown in the same table, the E_a^{app} values (at 723–823 K and $O_2/HCl = 2$) determined from the Arrhenius plots were in the range of 46–57 kJ mol⁻¹ for the supported catalysts, thus being similar to α -U₃O₈. The dependence of the activity on the relative O2 content in the feed was studied over the two most promising catalysts, U₃O₈/ZrO₂ and U₃O₈/SiO₂ (Fig. 2b). In both cases, the HCl conversion increased upon raising the feed O2/HCl ratio and the formal reaction order of O2 was calculated as ca. 0.3. This behaviour is common to the vast majority of Deacon catalysts24,28 and

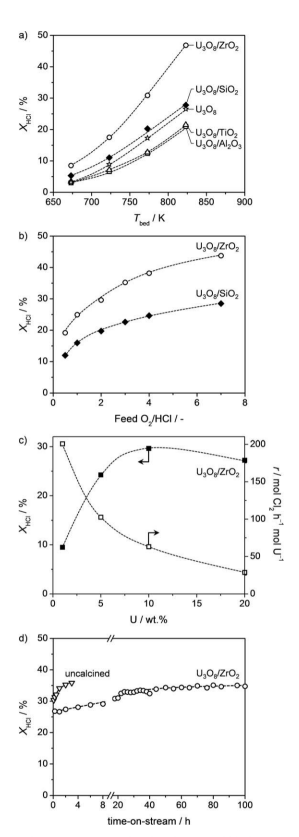


Fig. 2 HCl conversion over U_3O_8 -based catalysts *versus* (a) bed temperature at O_2 /HCl = 2 and (b) O_2 /HCl ratio at 773 K. HCl conversion and U-specific rate over U_3O_8 /Zr O_2 *versus* the uranium loading (c) and HCl conversion over U_3O_8 /Zr O_2 and an as-impregnated zirconia-supported catalyst (uncalcined) *versus* time-onstream (d) at 773 K and O_2 /HCl = 2. Data were acquired after 1 h under each condition for a–c. Other conditions are detailed in the Experimental section.

indicates that catalyst re-oxidation is the limiting step. 27 It is worth noting that the HCl conversion over $\rm U_3O_8/ZrO_2$ remained higher than that of $\rm U_3O_8/SiO_2$ at all $\rm O_2/HCl$ ratios. Overall, the catalytic results indicate that zirconia is the most suitable carrier for uranium oxide.

In order to rationalise the activity differences, the supported $\rm U_3O_8$ catalysts were characterised in fresh form and after use in the Deacon reaction. The uranium content, as determined by ICP-OES, was close to the nominal value of 10 wt% for all of the catalysts and remained unchanged in the used samples, indicating negligible uranium loss during HCl oxidation. The fresh alumina- and silica-based catalysts featured ca. 4 times larger $S_{\rm BET}$ than the zirconia- and titania-based materials (Table 1). This deviation reflects the difference in surface area of the pure carriers, which was depleted to a similar extent upon uranium incorporation in all cases, likely due to pore blockage. The $S_{\rm BET}$ of the catalysts was also unaltered upon use. Accordingly, the activity trend cannot be explained by differences and/or changes in the active phase content or textural characteristics.

XRD analysis of the fresh materials evidenced the formation of α -U₃O₈ over all supports with exception of titania (Fig. 3). In this latter case, a mixed UTiO₅ phase was detected (JCPDS 49-1397).³⁵ Furthermore, reflections specific to both the anatase and rutile forms of titania were observed, indicating that partial transformation of the carrier structure occurred during the high-temperature thermal activation of the as-impregnated solid. Thus, the loss in the support's surface area during catalyst preparation could be additionally ascribed to phase changes and structural reconstructions for U₃O₈/TiO₂. Based on the much lower intensity of its diffraction lines, the uranium phase is supposed to be present in form of smaller nanostructures on ZrO₂ compared to the other carriers, especially titania. The diffractograms of the samples after reaction revealed the absence of bulk chlorides (Fig. 3), extending the stability of

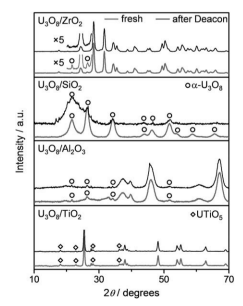


Fig. 3 XRD patterns of supported U_3O_8 samples in fresh form (blue lines) and after Deacon reaction (black lines). Unmarked reflections belong to the corresponding carriers.

 $\alpha\text{-}U_3O_8$ against chlorination also to the supported form. No changes were detected in the patterns of the TiO₂ and Al₂O₃-supported catalysts upon use, while the reflections specific to $\alpha\text{-}U_3O_8$ became less intense for U_3O_8/SiO_2 and disappeared for U_3O_8/ZrO_2 . Since uranium was not lost upon reaction, these alterations might be substantiated by fragmentation of the $\alpha\text{-}U_3O_8$ phase in tinier structures.

In order to further tackle this point and as the XRD analysis hints to differences in the dispersion of the supported active phase as a possible main parameter for determining the activity levels, the two most active catalysts (U₃O₈/ZrO₂ and U₃O₈/SiO₂) were further investigated by electron microscopy (EM, Fig. 4 and 5). For fresh U₃O₈/ZrO₂, aggregates of 20–30 nm sized support grains are visualised by HRTEM (Fig. 4a). However, inspection of surface regions even at higher magnification does not reveal a distinct uranium phase. Thus, based on the significant difference in the atomic numbers of U and Zr ($Z_{\rm U} = 92$ versus $Z_{\rm Zr} =$ 40), HAADF-STEM with Z-contrast was applied as a suitable tool to get information about the distribution of uranium-based phases (Fig. 5). Indeed, the uranium oxide species in the fresh U₃O₈/ZrO₂ are clearly visualised as bright rim or spots (Fig. 5a and b). The presence of uranium in these rims was confirmed by EDXS analysis. Investigation of the surface structure at the edges and on the surface revealed that two types of uranium oxide dispersions are present in the fresh U₃O₈/ZrO₂, namely, (i) a film-like nanostructure with a thickness ranging from a monolayer to 1 nm (Fig. 5a) and (ii) atomically dispersed uranium oxide as identified by bright spots (encircled) on the ZrO₂ support (Fig. 5b). Moreover, analysis of the complete structure of these spots is not possible on the basis of HAADF-STEM and would require more specific methods such as STEM coupled with electron energy loss spectrometer (EELS).36 Nonetheless, based on the studies on identification of single atoms,36 the bright spots seem to be composed of a single uranium atom (likely with some O atoms bound to it) and therefore, in this study they are referred to as atomically dispersed UO_x. Upon exposure to reaction conditions (for 5 h), the catalyst morphology seems to be altered. A film-like nanostructure is less visible and a concentration of bright spots of UO_x appears to be increased (Fig. 5c), suggesting the

transformation into tinier, better dispersed uranium oxide. Thus, uranium oxide on zirconia likely undergoes partial re-dispersion during reaction. This explains the disappearance of the α-U₃O₈ peaks in the XRD pattern of the used sample (Fig. 3). α-U₃O₈ on SiO₂ appears to be carried as nanoparticles of ca. 5 nm in the fresh catalyst (Fig. 4b). Upon use in HCl oxidation, the average particle size was reduced to ca. 2.5 nm (Fig. 4c, inset in b), supporting a certain degree of re-dispersion of the uranium phase. This agrees with the XRD results (Fig. 3). The origin of the active phase re-dispersion phenomenon, apparently common to both the zirconia- and silica-supported catalysts, is not fully understood. It is proposed that disaggregation of the uranium oxide structures might be induced by HCl and Cl2. The latter has been reported to produce such an effect on supported noble metal particles by generation of chlorides which readsorb on the solid carrier and are then reduced by the reaction environment.37,38 In our case, it is possible that uranium oxychloride species (UO_2Cl_2 , melting point = 843 K)³⁹ are formed to some extent. As they are highly unstable and readily re-oxidise under conditions similar to those applied in HCl oxidation,40 uranium will not be lost, but a certain degree of metal migration could be possible. This will ultimately improve the dispersion of the supported phase. Thus, based on the XRD and EM results, the activity differences seem to mainly depend on the uranium oxide dispersion. Still, the possibly different intrinsic activity of the chemical forms of uranium stabilised by the carriers might also play a role.

In view of its potential practical application, the $\rm U_3O_8/ZrO_2$ system was further studied in terms of optimisation of the active phase content as well as durability. Thus, catalysts with U loading comprised between 1 and 20 wt% were prepared and tested at 773 K and $\rm O_2/HCl = 2$ (Fig. 2c). The HCl conversion was found to raise with increasing U contents up to 10 wt%, while a loading of 20 wt% resulted in slightly lower activity. On the contrary, the U specific activity (*i.e.* reaction rate per mol of U) was the highest for the 1 wt% U catalyst and progressively diminished at increased U loadings. Hence, as a compromise between these parameters, a 5–10 wt% U content turns out to be optimal.

The robustness of $U_3O_8(10~\text{wt}\%~\text{U})/\text{Zr}O_2$ in HCl oxidation was tested in a long catalytic run (Fig. 2d). The HCl conversion

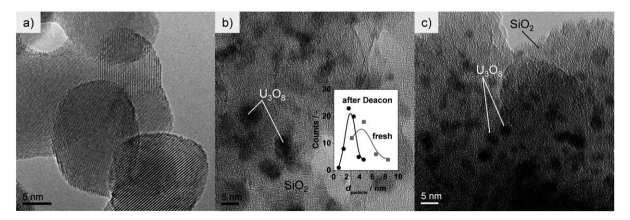


Fig. 4 HRTEM of fresh U_3O_8/ZrO_2 (a), U_3O_8/SiO_2 in fresh form (b) and after Deacon (c). Inset in (b) shows the particle size distribution of the fresh and used U_3O_8/SiO_2 sample.

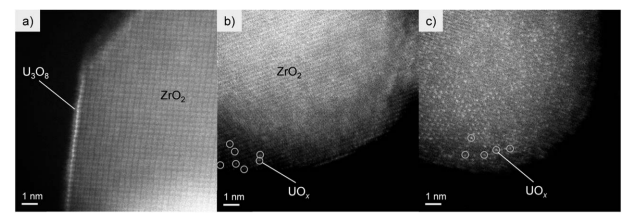


Fig. 5 HAADF-STEM of U₃O₈/ZrO₂ in fresh form (a and b) and after Deacon reaction for 100 h (c). Bright spots (some of which are encircled) in b and c corresponds to atomically dispersed UO_x.

moderately increased from 27 to 35% in the first 85 h on stream, remaining then stable up to a reaction time of ca. 100 h. Overall, this result evidences outstanding longevity, offering bright perspectives for an industrial application of zirconia-supported uranium catalysts in chlorine production. Still, the progressive catalyst activation indicates an alteration of the material's properties upon use. According to the above discussion of the characterisation data, this might originate from an increase in the dispersion of the active phase induced by the exposure to the reaction mixture. To further explore this point, samples after 5, 10, and 100 h on stream were collected and characterised by HAADF-STEM and H2-TPR (Fig. 5 and 6). While an increase in the uranium dispersion to certain extent has been already discussed for the sample after 5 h reaction (vide supra), HAADF-STEM of the sample after 100 h reaction evidenced that the uranium on ZrO2 carrier is mainly present as atomically dispersed UO_x (Fig. 5c). The latter would be characterised by the highest dispersion of uranium oxide. This result provides a direct evidence for the dependence of activity on degree of dispersion.

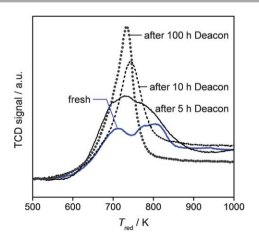


Fig. 6 H_2 -TPR profiles of U_3O_8/ZrO_2 in fresh form and after Deacon reaction for different times.

Additional support was derived from the H2-TPR analysis. The reduction profile of fresh U₃O₈/ZrO₂ features two main peaks at ca. 710 and 800 K (Fig. 6), which could be consistent with the presence of uranium oxide structures of different size (Fig. 5a and b), namely, thin layer (high-temperature signal) and atomic dispersion (low-temperature signal). For the sample collected after 5 h, a broad and more intense reduction peak centred at ca. 730 K with low- (695 K) and high-temperature (775 K) shoulders was visualised, while that taken after 10 h of reaction produced a single, symmetric, and sharper signal with maximum at 740 K. The curve of the catalyst unloaded at the end of the run displays an even narrower and more intense peak, slightly shifted to lower temperature (710 K). The depletion of the high-temperature signals with reaction time and the strengthening of a single peak at lower temperature supports a change in the morphology of α-U₃O₈ phase towards the formation of more uniformly-sized atomically dispersed nanostructures (UOx), in line with the HAADF-STEM results. The latter actually represents the predominant uranium distribution after 100 h on stream (Fig. 5c). Still, considering the modifications in peak position and shape, along with the significant increase in H2 consumption, the presence of more oxidic uranium in UO_x than in the original α -U₃O₈ phase cannot be excluded. Based on the structural equivalence between the zirconia support and β-UO₃ (both monoclinic, the latter having about double cell parameters with respect to the former), it could be possible that α-U₃O₈ undergoes transformation into this oxide during reaction. Although α-U₃O₈ is the most stable bulk oxide under HCl oxidation conditions and γ-UO₃ is converted into it during reaction, it is plausible that, when the incipient uranium oxychloride is oxidised by the O2 excess, the structural matching offered by the support could stabilise β -UO₃ as an oxidation product rather than α-U₃O₈. However, this phase is not detected by XRD owing to its very small size. Thus, from increased H2 uptake and development of atomically dispersed UO_x with reaction time, it can only be suggested that uranium generated in situ as UOx is in higher oxidation state than in the original α -U₃O₈ and the transformation of α -U₃O₈ to UO_x is

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accompanied by an enhancement of the dispersion. Since the presence of some UO_x is evidenced already for the fresh catalyst (Fig. 5b and 6), it could even be possible that a part of the uranium is already stabilised as UO_x during calcination and the atomic dispersion (Fig. 5b) rendering it undetectable by XRD (Fig. 3). An increase of uranium oxide dispersion during HCl oxidation was also evidenced for $\mathrm{U_3O_8/SiO_2}$ (Fig. 3, 4b and c). However, from a similar $\mathrm{H_2}$ consumption of fresh and used catalyst (not shown), it appears that *in situ* oxidation of the uranium phase does not occur on silica. This could be related to a specific property of the carrier and its interaction with the active phase. Thus, it seems that the support determines the degree of redispersion and reoxidation characteristics of the uranium phase. An in-depth understanding of these

Finally, we tested under the same HCl oxidation conditions an as-impregnated catalyst sample with equal U loading (*i.e.* no calcination applied after impregnating the U-precursor). This material reached a similar HCl conversion level (ca. 36%) to U_3O_8/ZrO_2 after only 3 h on stream (Fig. 2d). On the basis of this outcome and of the resemblance of the HAADF-STEM images and XRD pattern of the two catalysts after use (not shown), it is suggested that UO_x can be directly created *in situ* from the uranium precursor and with much faster kinetics. The latter is probably related to the ease of altering an amorphous and unstable deposit rather than a well-crystallised and stable phase.

complex phenomena will require deeper characterisation

studies, which are beyond the scope of this paper.

Comparison with other systems

The performance of U₃O₈/ZrO₂ was contrasted with other known supported HCl oxidation catalysts, namely, RuO2(2 wt% Ru)/SnO₂-Al₂O₃,²⁰ CeO₂(9 wt% Ce)/ZrO₂,²⁹ and CuO(15 wt% Cu)/SiO2 (synthesised by dry impregnation, followed by calcination at 823 K for 10 h). Fig. 7a displays the dependence of the HCl conversion level on temperature for these materials. The equilibrium HCl conversion (dashed line) is reported as a reference. The activity of RuO₂/SnO₂-Al₂O₃ increases with the temperature and reaches a HCl conversion close to the equilibrium value at 673 K. Beyond this temperature the active phase of this catalyst starts to form volatile RuO4.16 This indicates that the optimal high temperature boundary for RuO₂-based catalysts is 673 K. CuO/SiO₂ possesses a volcanoshaped activity profile. A strong deactivation above 723 K is due to huge copper loss in the form of CuCl and CuCl2.41 Differently, U₃O₈/ZrO₂ and CeO₂/ZrO₂ show a steady increase of the HCl conversion with temperature. The difference of activity between these two systems is relatively low (ca. 30 K). Cl₂ co-feeding at comparable initial HCl conversion levels (attained by adjusting T_{bed} , Experimental section) also displays a very similar inhibition of HCl oxidation activity (Fig. 7b). Still, CeO₂/ZrO₂ was observed to undergo bulk chlorination and, thus, deactivation at low O₂ excess.²⁹ Accordingly, U₃O₈/ ZrO2 stands as the most robust catalyst among all and belongs to the category of high-temperature catalysts, similar to CeO₂/ ZrO₂. However, the former offers a superior resistance to bulk chlorination.

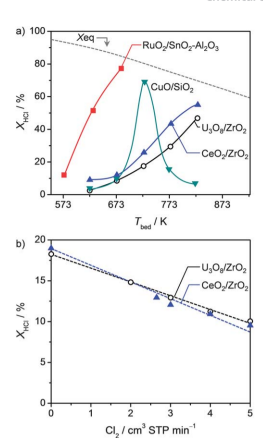


Fig. 7 Steady-state HCl conversion *versus* bed temperature (a) and amount of Cl_2 co-fed (b) at $O_2/HCl=2$. Other conditions are detailed in the Experimental section.

Conclusions

Uranium catalysts have been successfully evaluated for HCl oxidation to Cl2. Extraordinary resistance of bulk uranium oxides against chlorination demonstrates their suitability as a stable active phase for this reaction. While α -U₃O₈ maintains its oxidation state, UO2 and γ-UO3 tend to transform into α-U3O8 under reaction conditions. The support of the uranium phase plays a very important role on its performance. ZrO2 allows depositing of the oxidic uranium phase in the form of film-like nanostructures and atomic dispersion, thus leading to a superior catalyst. U₃O₈/ZrO₂ activates under reaction conditions before reaching a stable performance after ca. 85 h on stream. The catalyst activation is related to in situ re-dispersion and gradual transformation of the original α-U₃O₈ phase into a more oxidic and atomically dispersed UO_x. An uncalcined sample allows faster generation of this highly dispersed UOx. The unique robustness of ZrO₂ supported uranium oxide under the harsh reaction conditions and stable Cl₂ production for more than 100 h on stream justifies its consideration as a hightemperature HCl oxidation catalytic technology. Uranium materials are less sensitive to metal loss and sintering than other known catalysts and are cost-effective since they can be prepared from waste produced in the uranium-enrichment processes.

Experimental

Materials

ZrO₂-monoclinic (Saint-Gobain NorPro, 99.8%), γ-Al₂O₃ (Alfa Aesar, catalyst support, 43855), SiO₂ (ABCR, 99%), and TiO₂anatase (Aldrich, nanopowder, 99.7%) were calcined at 773 K (10 K min⁻¹) for 5 h prior to their use. The starting uranium compounds UO2 and UO2(NO3)2·6H2O (International Bio-Analytical Industries) derive from depleted uranium sources and were used as received. The most important precaution for the safe handling of uranium compounds is to avoid their access to the body, through direct contact with the skin and/or inhalation, and dispersal in the environment. In the present case, personal protective equipment such as impervious gloves, boots, and an apron were worn to prevent skin contact. U3O8 and UO3 were prepared by thermal decomposition of UO₂(NO₃)₂·6H₂O in static air following existing protocols.8,42 U3O8 was obtained by two-step calcination of UO₂(NO₃)₂·6H₂O. The uranyl nitrate was treated at 573 K (5 K min⁻¹) for 1 h and then, without intermediate cooling, at 1073 K (5 K min⁻¹) for another 3 h. UO₃ was synthesised by calcination of the uranyl nitrate at 723 K (5 K min⁻¹) for 3 h. Supported catalysts were prepared by dry impregnation of the carriers with an aqueous solution of uranyl nitrate (nominal 1-20 wt% U), followed by drying at 338 K for 12 h and calcination, according to the same protocol applied for the synthesis of bulk U₃O₈. Unless stated otherwise, the supported catalysts, denoted as U₃O₈/support, contain 10 wt% U.

Characterisation techniques

Powder X-ray diffraction (XRD) was measured in a PANalytical X'Pert PRO-MPD diffractometer. Data were recorded in the 10-70° 2θ range with an angular step size of 0.017° and a counting time of 0.26 s per step. N₂ sorption at 77 K was performed in a Quantachrome Quadrasorb-SI gas adsorption analyser. Prior to the measurement, the samples were evacuated at 473 K for 12 h. Temperature-programmed reduction with hydrogen (H2-TPR) was measured in a Thermo TPDRO 1100 unit. The samples were loaded in a quartz micro-reactor (11 mm i.d.), pre-treated in He $(20 \text{ cm}^3 \text{ STP min}^{-1})$ at 473 K for 30 min, and cooled to 323 K in He. The analysis was carried out in 5 vol% H_2/N_2 (20 cm³ STP min⁻¹), ramping the temperature from 323 to 1173 K at 10 K min⁻¹. Highresolution transmission electron microscopy (HRTEM) measurements were undertaken on a FEI Tecnai F30 microscope (field emission gun), operated at 300 kV. High-angle annular dark field scanning transmission electron microscopy (HAADF-STEM) investigations were performed on an aberration-corrected Hitachi HD-2700CS microscope, operated at 200 kV and equipped with an energy-dispersive X-ray spectrometer (EDXS, EDAX) for elemental analysis. The incorporated probe correction system (CEOS) enables a resolution of below 0.1 nm to be achieved.43

Catalytic tests

The gas-phase oxidation of hydrogen chloride was studied at ambient pressure in a continuous-flow set up⁴⁴ composed of mass-flow controllers to feed HCl (Messer, purity 2.8, anhydrous), O₂ (Pan Gas, purity 5.0), Cl₂ (Messer, purity 2.8,

anhydrous), and N2 (Pan Gas, purity 5.0), a home-made electrically heated oven hosting a 8 mm i.d. quartz micro-reactor, and a Mettler Toledo G20 Compact Titrator for quantitative Cl₂ analysis in the product stream. The catalysts were loaded in the tubular reactor and pre-treated in N2 at 673 K for 30 min. Thereafter, steady-state experiments at variable bed temperatures ($T_{\text{bed}} = 673-823 \text{ K}$), inlet O_2/HCl ratios (0.5-7), and catalyst amounts (W = 0.25 or 0.5 g for supported or bulk catalysts, respectively) were carried out. The inlet HCl concentration and total volumetric flow (F_T) were fixed at 10 vol% and 166 cm³ STP min⁻¹, respectively. The O₂/HCl dependence was measured by increasing the O₂ content in the inlet mixture from 5 to 70 vol% with N₂ as balance gas. The influence of Cl₂ co-feeding on the rate of HCl oxidation was studied by introducing variable amounts (2-5 cm³ STP min⁻¹) of Cl₂ to the inlet feed with $O_2/HCl = 2$ at 733 K and 703 K over U_3O_8/ZrO_2 and CeO_2/ZrO_2 , respectively. Used samples were collected for characterisation after rapidly cooling down the reactor to room temperature in N₂ flow. The percentage of HCl conversion was determined as $X_{\rm HCl} = (2 \times \text{mole Cl}_2 \text{ at the reactor outlet/mole HCl at the})$ reactor inlet) \times 100.

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Notes and references

- 1 S. T. Taylor, in *Metal Oxide Catalysis*, Wiley-VCH, Weinheim, 2009, ch. 13, p. 539.
- 2 A. R. Fox, S. C. Bart, K. Meyer and C. C. Cummins, *Nature*, 2008, 455, 341.
- 3 H. Idriss, Surf. Sci. Rep., 2010, 65, 67.
- 4 W. J. Hale, US1 595 299, assigned to Dow Chemical Company, 1926.
- 5 A. E. Craver, US1 636 854, assigned to Barrett Company, 1927.
- 6 Z. R. Ismagilov, S. V. Kuntsevich, N. V. Shikina, V. V. Kuznetsov, M. A. Kerzhentsev, V. A. Ushakov, V. A. Rogov, A. I. Boronin and V. I. Zaikovsky, *Catal. Today*, 2010, 157, 217.
- 7 G. J. Hutchings, C. S. Heneghan, I. D. Hudson and S. H. Taylor, *Nature*, 1996, 384, 341.
- 8 S. H. Taylor and S. R. O'Leary, Appl. Catal., B, 2000, 25, 137.
- 9 T. Campbell, M. A. Newton, V. Boyd, D. F. Lee and J. Evans, *J. Phys. Chem. B*, 2005, **109**, 2885.
- 10 P. L. Cottingham and L. K. Barker, *Ind. Eng. Chem. Prod. Res. Dev.*, 1973, 12, 41.
- 11 J. L. Callahan and B. Gertisser, US3 198 750, assigned to The Standard Oil Company, 1965.
- 12 J. L. Callahan and B. Gertisser, US3 308 151, assigned to The Standard Oil Company, 1967.

- Edge Article
- 13 R. K. Grasselli and J. L. Callahan, *J. Catal.*, 1969, **14**, 93.
- 14 I. Grenthe, J. Drozdzynski, T. Fujino, E. C. Buck, T. E. Albrecht-Schmitt, S. F. Wolf, in *The Chemistry of the Actinide and Transactinide Elements*, Springer, Dordrecht, 2006, vol. 1, ch. 5, p. 253.
- 15 H. Deacon, US85 370, assigned to Gaskell, Deacon and Co., 1868.
- 16 J. Pérez-Ramírez, C. Mondelli, T. Schmidt, O. F.-K. Schlüter, A. Wolf, L. Mleczko and T. Dreier, *Energy Environ. Sci.*, 2011, 4, 4786.
- 17 K. Seki, Catal. Surv. Asia, 2010, 14, 168.
- 18 H. Over, J. Phys. Chem. C, 2012, 116, 6779.
- 19 A. Wolf, L. Mleczko, O. F. Schlüter and S. Schubert, EP2026905, assigned to Bayer MaterialScience, 2006.
- 20 C. Mondelli, A. P. Amrute, F. Krumeich, T. Schmidt and J. Pérez-Ramírez, *ChemCatChem*, 2011, 3, 657.
- 21 D. Crihan, M. Knapp, S. Zweidinger, E. Lundgren, C. J. Weststrate, J. N. Andersen, A. P. Seitsonen and H. Over, *Angew. Chem., Int. Ed.*, 2008, 47, 2131.
- 22 N. López, J. Gómez-Segura, R. P. Marín and J. Pérez-Ramírez, J. Catal., 2008, 255, 29.
- 23 M. A. G. Hevia, A. P. Amrute, T. Schmidt and J. Pérez-Ramírez, J. Catal., 2010, 276, 141.
- 24 D. Teschner, R. Farra, L.-D. Yao, R. Schlögl, H. Soerijanto, R. Schomaecker, T. Schmidt, L. Szentmiklósi, A. P. Amrute, C. Mondelli, J. Pérez-Ramírez, G. Novell-Leruth and N. López, J. Catal., 2012, 285, 273.
- 25 F. Studt, F. Abild-Pedersen, H. A. Hansen, I. C. Man, J. Rossmeisl and T. Bligaard, *ChemCatChem*, 2010, 2, 98.
- 26 T. Hibi, H. Nishida and H. Abekawa, US5 871 707, assigned to Sumitomo Chemical Company, 1999.
- 27 D. Teschner, G. Novell-Leruth, R. Farra, A. Knop-Gericke, R. Schlögl, L. Szentmiklósi, M. G. Hevia, H. Soerijanto, R. Schomäcker, J. Pérez-Ramírez and N. López, *Nat. Chem.*, 2012, 4, 739.
- 28 A. P. Amrute, C. Mondelli, M. Moser, G. Novell-Leruth, N. López, D. Rosenthal, R. Farra, M. E. Schuster,

- D. Teschner, T. Schmidt and J. Perez-Ramírez, *J. Catal.*, 2012, 286, 287.
- 29 M. Moser, C. Mondelli, T. Schmidt, F. Girgsdies, M. E. Schuster, R. Farra, L. Szentmiklósi, D. Teschner and J. Pérez-Ramírez, *Appl. Catal.*, *B*, 2013, **132–133**, 123.
- 30 A. Wolf, L. Mleczko, S. Schubert and O. F. Schlüter, US20100202959, assigned to Bayer Technology Services, 2010.
- 31 A. Wolf, L. Mleczko, O. F. Schlüter and S. Schubert, US20110180419, assigned to Bayer Technology Services, 2011.
- 32 K. J. Notz and M. G. Mendel, *J. Inorg. Nucl. Chem.*, 1960, **14**, 55.
- 33 Y.-S. Yang, Y.-H. Kang and H.-K. Lee, *Mater. Chem. Phys.*, 1997, **50**, 243.
- 34 A. P. Amrute, C. Mondelli, T. Schmidt, R. Hauert and J. Pérez-Ramírez, *ChemCatChem*, 2013, 5, 748.
- 35 R. H. Marsuall and H. R. Hoekstra, *J. Inorg. Nucl. Chem.*, 1965, 27, 1947.
- 36 C. Colliex, A. Gloter, K. March, C. Mory, O. Stéphan, K. Suenaga and M. Tencé, *Ultramicroscopy*, 2012, 123, 80.
- 37 K. Foger and H. Jaeger, J. Catal., 1985, 92, 64.
- 38 P. Birke, S. Engels, K. Becker and H. D. Neubauer, *Chem. Technol.*, 1980, **32**, 253.
- 39 T. Subramanian, B. P. Reddy, P. Venkatesh, R. Kandan, T. Vandarkuzhali, K. Nagarajan and P. R. V. Rao, in Pyrochemical Separations: Workshop Proceeding, Avignon, France, OECD Publishing, Paris, 2000, p. 190.
- 40 T. Sato, S. Shiota, S. Ikoma and F. Ozawa, *J. Appl. Chem. Biotechnol.*, 1977, 27, 275.
- 41 C. Mondelli, A. P. Amrute, T. Schmidt and J. Pérez-Ramírez, *Chem. Commun.*, 2011, 47, 7173.
- 42 V. J. Wheeler, R. M. Dell and E. Wait, *J. Inorg. Nucl. Chem.*, 1964, 26, 1829.
- 43 H. Inada, L. Wu, J. Wall, D. Su and Y. Zhu, *J. Electron Microsc. Tech.*, 2009, **58**, 111.
- 44 A. P. Amrute, C. Mondelli and J. Pérez-Ramírez, *Catal. Sci. Technol.*, 2012, 2, 2057.