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Additional Information

'From the Mole to the Molecule': Ruthenium Catalyzed Nitroarene Reduction Studied with 'Bench', High-Throughput and Single Molecule Fluorescence Techniques

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Single molecule fluorescence microscopy techniques are used to complement conventional catalysis and high-throughput experiments in order to gain a complete picture of a model reaction. In these experiments a model nitroarene is reduced to an amine where, upon reduction, a red shift in absorption/emission, as well as increase in emission is observed. The reaction is studied in bulk reaction conditions by NMR spectroscopy and the fluorescence activation makes it possible to also study this reaction at the single molecule level. Fluorescence correlation spectroscopy is a valuable technique in supporting the proposed reaction mechanism and understanding the nature and duration of molecular 'visits' to catalytic sites, where both the starting nitroarene and amine product have an affinity for the catalyst.

15 Introduction

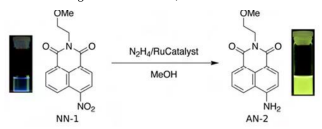
The last decade has seen a dramatic increase in the number and quality of tools available to study catalytic processes. Thus, the conventional organic chemistry or "bench" approach can be combined with high throughput (HT) techniques that enable a combinatorial approach to optimize catalysts and reaction conditions. On the other extreme, single molecule techniques make it possible to visualize how a single molecule arrives at a catalytic site and departs after catalytic conversion; a few examples of this type have been reported, normally using 25 fluorescence changes as the reaction reporter. A

Single molecule, high throughput and bench studies have not been the subject of a single reaction study; yet, this approach, that we describe as "from the mole to the molecule" is feasible today and should provide a unique, intimate understanding of reaction mechanisms; in the future such a combination may become a powerful tool for catalyst design and optimization.

In this study we utilize two new ruthenium supramolecular catalysts based on mesoporous silica⁴ to examine the reduction of nitro-aromatics by hydrazine in what we believe is the first report that combines powerful HT laboratory tools and single molecule fluorescence techniques, to take us from relatively large scale to the single molecule examination of the catalytic reduction of nitroarenes.

Our catalysts, Ru@SBA and Ru@MCM, containing ruthenium 40 on mesoporous MCM-41 or SBA-15 according to TEM images (see Figures S1 and S2 in ESI), have not been tested before in reductive processes, although they had proven valuable in oxidative reactions leading to Wittig chemistry from alcohol precursors. Other ruthenium-based mesoporous catalysts have 45 been used in alkene reductions. We have examined simple

nitroarenes using HT tools as a methodology to identify the best reaction conditions. Selected reactions were also examined at the larger "bench" level to ensure that the conditions selected worked well under classical organic chemistry conditions. Having so achieved this, a fluorogenic nitroaromatic compound bearing the nitronaphthalimide functionality was tested at the bench level and selected for single molecule studies, Scheme 1.



Scheme 1: Reductive conversion of NN-1 to AN-2 yields a strongly fluorescent product that is used as a reporter for fluorescence microscopy.

In the reaction of Scheme 1, reduction leads to an aminonaphthalimide which fluoresces strongly at around 540 nm, a convenient spectral region for single molecule studies. Beyond visualization of the reaction, the work combined with fluorescence correlation spectroscopy (FCS) reveals that molecules generated at the catalytic site spend more time at their nascent place than at randomly selected locations in solution. Thus, our work on two new catalysts bring us from the bench, or 'mole' scale, to the molecule. We can envision a future in which single molecule and computational work will allow us to follow the reverse path, from the molecule, to the mole and scale-up to the factory with catalysts truly designed on the basis of an intimate understanding on how molecules achieve the desired transformation efficiently, rapidly and with high selectivity.

Ruthenium has received a lot of attention as a remarkably active catalyst in both homogeneous^{6, 7} and heterogeneous catalysis.⁸ Ruthenium complexes have been used in metathesis as important catalysts that are both active and selective.7, 9 Recently Ru 5 nanoparticles (RuNP) have been found to be active catalysts for hydrogenation^{8, 10} including asymmetric catalysis. 11 If there is a drawback for using Ru as a catalyst, it is the high cost of the metal; Ru is considered a precious metal and is thus quite expensive. Therefore, to make Ru a viable catalyst it must be re-10 useable, something that is easier to achieve with heterogeneous catalysts. One of the common questions in heterogeneous catalysis is whether the reaction is happening on the surface of the catalyst or as a homogeneous process in solution, as a result of catalyst leaching from the particle surface. 12 One can adjust 15 reaction parameters to gain some insight into the mechanism of the reaction, which is done here as well, but having a direct measure of the retention of reagents and products at the catalytic site is very valuable information in catalysis. Here we use a model reaction, such as shown in scheme 1, to examine catalytic 20 activity of Ru nanoparticles (RuNP) supported on SBA. Using a combination of normal optical microscopy, total internal reflectance fluorescence (TIRF) microscopy and fluorescence lifetime imaging microscopy (FLIM) we are able to image both the catalyst and reaction species and in doing so better elucidate 25 whether or not the reaction is happening on the surface of the catalyst.

Materials and methods

Materials

Tetraethylorthosilicate (TEOS, 98%) was used as silica source and cetyltrimethylammonium bromide (C₁₆TAB, 96%) or pluronic (P123) as structure-directing agents. Aqueous ammonia solution (NH₄OH, 30%) and hydrochloric acid needed for the preparation of the mesoporous MCM-41 and SBA-15 respectively, and aminopropyltriethoxisilane (APTES) were also used in the synthetic protocol to obtain the final mesoporous material. RuCl₃.xH₂O was used as ruthenium source; nitrobenzene, hydrazine monohydrate, the other nitroaromatics and solvents were purchased from Aldrich and used as received without further purification. 1,3-Dimethoxybenzene and 1,3,5-trimethoxybenzene were from Alfa-Aesar. Nitronaphthalimide-1 (NN-1) was prepared according to a procedure described elsewhere.

Catalyst Preparation

The synthesis of MCM-41 or SBA-15 supported RuNPs was reported elsewhere. Briefly, anhydrous toluene (50 mL) was added to dehydrated (473 K for 2 h) MCM-41 or SBA-15 (1.5 g) and this mixture was stirred for 1 h in order to obtain a homogeneous dispersion; then, APTES (0.5 mL) was added and refluxed overnight. The white solids obtained (silica-APTES) were filtered, washed with fresh toluene and acetone and airdried. Then, ruthenium chloride (RuCl_{3.x}H₂O, 9 mg) was added to an aqueous dispersion (110 mL) of the corresponding silica-APTES (1.5 g) and the resulting mixture was stirred for 1 h at room temperature. The grey solids obtained were filtered off and washed (H₂O) to remove unreacted salt. The air-dried new synthesized SRuNPs (2.5 wt % Ru, determined by ICP) were

denoted as Ru@MCM or Ru@SBA, respectively.

Catalytic Tests

High Throughput Combinatorial Catalysis Screening. The 60 reactions were tested in plates containing 96 glass shell vials (8 x 40 mm), each with a π -arylene-coated tumble stir bars (Supporting Information, Figure S1). The appropriate amount of catalyst (10-25 mg) was manually weighted into each vial (accuracy of \pm 0.3 mg). Plates for reactions were prepared using a 65 Symyx/Freeslate (Santa Clara, CA) first-generation core module, equipped with a Julabo LH45 temperature control unit. Nitrobenzene solutions in ethanol (0.6 - 1 M) were freshly prepared in 20 mL vials that were also loaded into the sample processor. The robot was programmed to dispense the appropriate 70 substrate solution into each vial (Supporting Information, Figure S2). Next, the plate was cooled down to 15 °C before the solution of hydrazine (6 M) was dispensed in order to avoid gas evolution. The appropriate amount of solvent (ethanol 99%) was programmed to be added to each vial to ensure an equal final volume of 0.5 mL. Finally, the internal standard (1,3,5trimethoxybenzene) was added to every vial (0.030 mL, 0.5 M). Once every substrate solution was dispensed, the plate was sealed under air with a top metal plate consisting of a teflon sheet and an aluminium cover, cushionned by two viton gaskets/sheets. Plates 80 were stirred and heated on a VP Scientific VP710E-2/VP743A-1R microplate tumble stirrer. After that, the complete plate was centrifuged for 15 min in a Genevac EZ-2 Plus and then the vials were daughtered using the Symyx/Freeslate core module, diluted with ethyl acetate and loaded into the GC for analysis. Reactions 85 were analyzed by gas chromatography (Agilent Technologies 6850 GC), equipped with a CTC GC-PAL autosampler; with a HP-1 column (30 m x 0.32 mm x 0.25 µm). The inlet/detector temperatures were set to 250°C, initial oven temperature of 55°C held for 0.5 min, followed by a ramp of 35°C/min up to 120°C, 90 held for 0.5 min and then a second ramp of 45°C/min up to 260°C held for 0.45 min (total runtime of 6 minutes). Retention times were confirmed by comparison with authentic materials when available or by injecting them into GC-MS and compared to the internal library. Yields of the reactions were determined 95 from both consumption of starting material and formation of product. Peak areas were referenced against the internal standard, interpolating from previously made 7-points calibration curves.

Optimized Reduction Conditions. In a typical run, to an ice100 cooled mixture of 25 mg of SBA/Ru (4% mol) and 0.15 mmol of
the nitroaromatic compound in 0.345 mL of EtOH, 0.125 mL of
NH₂NH₂.H₂O 6M (6 eq) and 0.030 mL of internal standard
(1,3,5-trimethoxybenzene in the case of nitrobenzene or 1,3dimethoxybenzene for the other nitroderivatives) (0.5 M) were
105 added. The mixture was stirred under air at 80°C for 21 h. Then it
was centrifuged, daughtered (0.020 mL of the reaction mixture
were diluted up to 1 mL of AcOEt) and analyzed by GC or ¹H
NMR.

Agreement between replicate runs was within $\pm 3\%$.

Catalyst Reusability. To check the reusability of the hybrid materials, they were washed three times with ethyl acetate by consecutive stirring and centrifuging steps. GC analysis of the last aliquot corroborated the absence of reaction products.

Microscopy Measurements

Samples for microscopy were prepared by adding <1mg of prepared catalyst (Ru@SBA) into reaction wells with a cover slip bottom for imaging. 0.4 mL of methanol was added to the catalyst followed by 0.1 mL of a solution of the starting material $_{5}$ (NN-1, 9.2 mM) and 0.010 mL of hydrazine (1.1 x 10^{-3} M). All measurements for single molecule catalysis were performed at room temperature; note that the reaction can be catalyzed even under these very mild conditions. The catalyst with methanol alone was imaged with white light as well as by fluorescence 10 imaging that showed a contrast due to low intensity fluorescence and scattering of the catalyst. After addition of the starting material (NN-1), excitation at 375 nm could be used to image the starting material and to obtain FCS curves of NN-1 while focused on the catalyst, or freely diffusing in solution. After addition of 15 hydrazine, the product (AN-2) could be excited selectively with 440 nm light to image AN-2 in areas at the catalyst and diffusing in solution. When excited by 440 nm (FLIM) or 480 nm (TIRF) light, images were noticeably brighter as the reaction proceeded, as expected with the absorbance increase at longer wavelengths

Fluorescence Activation Reaction and Controls

20 with formation of the product (see Figure 4, vide infra).

Bench top control reactions were set up as follows: To a mixture of 10 mg of Ru@SBA (7.2% mol) and 10 mg (0.033 mmol) of **NN-1** in 1 mL of ethanol, 0.036 mL of NH₂NH₂.H₂O 6M (6.5 eq) were added. The mixture was stirred under air or H₂ atmosphere at 80°C for 21 h in darkness. Then it was centrifuged, filtered off, concentrated and analyzed by GC and ¹H NMR.

UV Spectroscopy and Fluorescence

UV-vis spectra were recorded on a Cary UV-50 spectrophotometer. Steady state fluorescence spectra were recorded with a Photon Technologies International (PTI) Fluorimeter, equipped with a Xe lamp for excitation, monochromators for both the excitation and emission and a PMT detector for detecting the fluorescence signal. Felix software was used to record and analyze fluorescence spectra.

Advanced Microscopy

TIRF Imaging. Fluorescence imaging was performed with an Olympus FV1000 TIRF (total internal reflection fluorescence) instrument (Olympus, Japan). The instrument is equipped with a 40 488 nm CW laser. The laser beam was collimated and focused through a fiber-coupling unit. A beam splitter cube was used with a dichroic mirror at 488 nm for the excitation, excitation filter centred at 482 nm (18 nm bandpass) and an emission filter centred at 525 nm (45 nm bandpass). An oil immersion TIR (total 45 internal reflection) objective (100×, NA1.45, Olympus, PLAPO) was used to focus both the excitation and emission, and the TIR images were recorded with a Rolera-MGi PLUS high-speed, extremely sensitive Digital EMCCD camera. Several frame TIRF images are presented in the supporting information, where 50 brightest/flashing spots correspond to catalyst where the amine product is generated and has an affinity for the solid. The same camera also collected white light images but using white light excitation from an Olympus TH4-100 powered 100 W halogen

55 *FLIM and FCS*. Fluorescence lifetimes, lifetime images and fluorescence correlation spectroscopy (FCS) curves were

recorded with a Fluorescent Lifetime Imaging System (FLIM, PicoQuant).³ The instrument is equipped with four picosecond pulse diode lasers including the 375 nm and 440 nm lasers used

60 to excite the starting material and product from scheme 1, respectively. The laser beam was collimated and focused through a fiber-coupling unit. The same microscope and oil immersion TIR objective as for TIRF measurements is used for the FLIM studies. Appropriate long pass filters were used to eliminate the excitation from the final FLIM images (400 LP for 375 nm excitation and 450 LP for 440 nm excitation).

FCS measurements were performed by focusing the excitation on regions that contained catalyst as identified by white light, TIRF and FLIM images, and were compared to FCS curves of regions 70 that contained no catalyst (freely diffusing molecules in solution).

Because of the large number of variables that need to be

Results

Catalytic Activity: Reduction of nitroarenes

examined to optimize a green process, such as, temperature, 75 reaction time, percentage of catalyst and support or H₂ source, a combinatorial catalysis screening approach seemed appropriate to optimize the reaction conditions. Initial attempts used nitrobenzene and hydrazine monohydrate or ammonium formate as hydrogen sources, since the use of H₂ and 80 eventually high pressures was ruled out in the development of a green and sustainable process. In the presence of ammonium formate no conversion was observed regardless of other experimental conditions. Efforts were then focused on the optimization of support, mol percent of catalyst, and equivalents 85 of hydrazine. A high-throughput screening approach proved extremely valuable for this process. As shown in Figure 1, the hybrid materials Ru@MCM and Ru@SBA at 1.6-4 mol % in the presence of 1-8 equivalents of hydrazine were tested. At the end of the reaction time, the plate was centrifuged, diluted 90 analyzed by GC. Yields were determined from both consumption of starting material and formation of aniline. As shown in Figure 1, at least 5 equivalents of hydrazine were needed to achieve quantitative conversion of nitrobenzene in the presence of at least 3.2% mol of the catalyst. Higher yields were always obtained 95 when Ru@SBA was employed compared to Ru@MCM at the same mol percent. Next, the effect of reaction time (4 to 24 h) and temperature (40 to 80 °C) were examined. Complete conversion was achieved after 21 h reaction time. Otherwise, several by-products were observed from the GC-MS analysis when temperatures lower than 70°C were used. 14 Nevertheless, increasing the temperature up to 80°C resulted just in the formation of the desired aniline. Overall the best conditions were found using 4 mol % of Ru@SBA in the presence of 5

equivalents of hydrazine monohydrate at 80°C for 21 h under air.

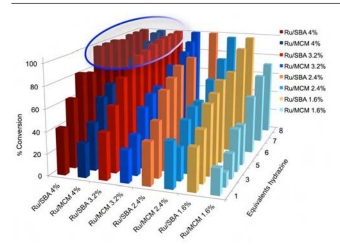


Figure 1. Conversion of nitrobenzene as a function of employed catalyst, at various mol percentages and different equivalents of hydrazine monohydrate. The oval indicates the region of near-quantitative 5 conversion as selected from the HT data.

Chemoselectivity of the reduction was also investigated (results are summarized on Table 1). Excellent selectivities were obtained for the reduction of the nitroarenes with substituents such as -OCH₃, -CN (entries 2-3). However, the reduction of 3-10 nitrostyrene (entry 4) resulted in the formation 3-aminostyrene together with 3-ethylaniline (1.6:1). The use of lower temperatures (60°C) and lower equivalence of hydrazine monohydrate did not increase the selectivity towards the nitro group. The already optimized reduction conditions were then 15 applied to the "more substituted" naphthalimide NN-1 (entry 5). Moreover, the reusability of hybrid materials was tested in the reduction of nitrobenzene under the optimized reaction conditions. Results are summarized in Figure 2. The two catalysts, Ru@MCM and Ru@SBA, exhibited a 20 recyclability; in fact, interestingly higher yields were obtained after the second reuse of Ru@MCM. This is commonly attributed to catalyst activation, in other words, generating reactive sites while under reaction conditions. Results obtained using Ru@HMS are also included for comparison.⁴

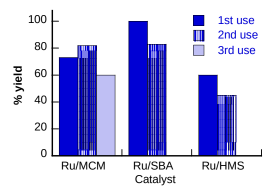


Figure 2. Percentage of the reusability of the materials studied for the reduction of nitrobenzene under the optimized reaction conditions.

Table 1.30 Catalytic reduction of different nitroarenes⁶

#	Substrate and product	Yield (%)
1	\sim NO ₂ \rightarrow	>99
2	MeO NO_2 MeO NH_2	60
3	NC NC NC NC NC NC NC NC	>99
4	NO ₂ NH ₂ NH ₂ NH ₂	>99
5	OMe OMe ONO NO NO NH2	>99

^aNitroarene (0.15 mmol), Ru@SBA (4 mol %), hydrazine monohydrate (5 equivalents) in EtOH (99%) at 80 °C for 21 h under air.

Bench Top Results: transition to single molecule studies

In order to use single molecule techniques and to move from the mole to the molecule, a material for reduction was required such that the reaction produces a fluorescent product different from the starting material. The nitro compound NN-1 provided the requirements needed with a red shift and increase in emission quantum yield upon reduction to AN-2. The reaction was performed with the best catalyst (Ru@SBA) from the high-throughput experiments which required elevated temperatures (80°C) to reach completion. For this reason, ethanol was used for benchtop reactions to reach higher temperatures, whereas methanol was sufficient as a solvent for the room temperature microscopy experiments.

The SBA supported RuNPs (Ru@SBA) catalyzed reduction of NN-1 (Scheme 1) was first performed in bulk experiments (together with several control reactions—see Table 2) and analyzed by ¹H NMR and UV-vis absorbance and fluorescence spectroscopy. The absorbance and emission spectra of the reaction are shown in Figure 3 (bottom), where a red-shift in both the absorbance and emission were observed. The product was identified by ¹H NMR as AN-2 and it displays a 7x increase in maximum emission counts, see SI. ¹³The shift in absorbance and increase in quantum yield of emission allowed for easy confirmation that the reaction had occurred.

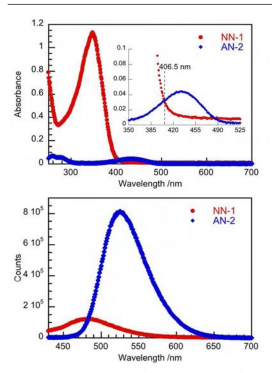


Figure 3. Absorbance spectra of **NN-1** and **AN-2** (after reaction), top, as well as (bottom) emission spectra of for both compounds (with matched 5 absorbance and excitation at 406.5 nm – see SI).

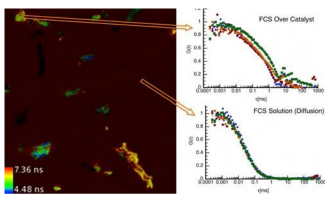
¹H NMR spectroscopy was used to identify products and to quantify the conversion of the reaction shown in scheme 1 (see ESI). The first entry in Table 2 shows that the reaction only goes to completion within 21 h when catalyzed by Ru@SBA. A minor conversion (6 %) can be observed when the hydrazine reducing agent is replaced by a H₂ filled balloon (entry 4). While complete conversion of the starting material is observed without Ru@SBA in the same amount of time (entry 6), no starting material remained unreacted but only 20 % conversion to AN-2 could be achieved.

Table 2. Data from the bench top reactions converting **NN-1** to **AN-2.** All reactions at 80° C for 21h.

Entry	Conditions	Hydrazine monohydrate (Equivalents)	Conversion (% AN-2)
1 ^b	Ru@SBA (7.2 mol%)	6.5	100
2	H ₂ atmosphere	0	0
3	SBA (no RuNPs) H ₂ atmosphere	0	0
4	Ru@SBA (7.2 mol%) H ₂ atmosphere	0	≈ 6
5 ^b	Ru@SBA (7.2 mol%)	0	0
6 ^b	No Catalyst or H ₂	6.5	20°
7 ^b	SBA	6.5	20°

^aDetermined by ¹H NMR. ^b runs performed under air. ^c100% conversion of starting material but only, 20% **AN-2**.

Microscopy. The ¹H NMR product analysis shows that the Ru supported catalysts increase the reaction rate, as well as the fact that addition of the less efficient H₂ gas instead of hydrazine as a reducing agent, works most effectively with the Ru catalyst 25 present. Both results suggest that the reaction occurs at the surface, or at the very least is activated by the catalyst. In order to further establish if the reaction occurs on the catalyst, single molecule fluorescence is used to follow the variations in diffusion of the NN-1 and AN-2 (Scheme 1), in solution as compared with 30 the catalyst. When a laser is focused on a spot, fluorescence correlation spectroscopy (FCS) can be used as a measure of the average time molecules spends within that focused spot. Therefore, by comparing FCS plots when the laser is focused on the catalyst or if it is focused in solution (after some AN-2 35 product has accummulated), gives different results if there are some binding affinities for the surface. 15 With a 440 nm laser, the products of the reaction can be selectively excited (Figure 3). In this way the image in Figure 4 was recorded, the brightest spots in the FLIM images corresponding to catalyst (confirmed by 40 white light and TIRF images similar to those in the SI where the starting material imaging is shown). Representative FCS traces for catalyst and solution spots are given in Figure 4 as indicated.



45 Figure 4. FLIM image acquired with 440 nm laser excitation during reduction of NN-1, where the bright spots correspond to catalyst (left). FCS curves when focused on catalyst spots (top right) and solution spots (bottom right). Solution spots measured after some accumulation of the product AN-2.

spot and a solution spot are given in Figure 5. The catalyst spots and a solution spot are given in Figure 5. The catalyst spots consistently show a longer τ , indicating that the products spend more time on the catalyst than freely diffusing through solution. Similar images and FCS traces were recorded for the starting material with 375 nm excitation (see SI). A similar longer τ is observed for the starting material at the catalyst as compared to in solution, indicating some binding to the catalyst for the starting material as well. These results are consistent with the reaction being catalyzed at the surface through binding since both the reagents and products have an affinity for the surface. These experiments cannot distinguish between residence at the external or internal surface of a mesoporous material.

the laser focal area increases from about 25 µs to about 1 ms when the latter corresponds to a nascent product at the catalytic site. The former (25 µs residence) corresponds to **AN-2** in solution, added as a control and not a reaction product.

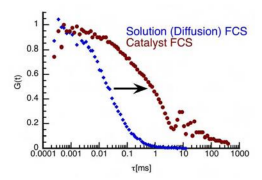


Figure 5. Representative FCS curves with 440 nm excitation when excitation/image acquisition is focused on either a catalyst (red) or 5 solution (blue) spot.

Videos showing reaction triggered blinking are available as Supporting Information. They reveal multiple catalytic sites within a single particle and repeating spots for blinking; this is characteristic of reactions occurring at specific catalytic sites, presumed to be ruthenium-rich spots although diffraction limitations prevent us from visualizing these. Visual analysis of peaks and valleys in enlarged images such as those in Figure 6, suggest 'hot' catalytic spots on SBA are located roughly 0.5 to 2 um apart. The data was recorded at a rate of 10 frames per second 15 for 50 seconds and is displayed in a cumulative way in Figure 6, where panels A and B display the accumulated data for frames 1-250 (25 s) and 251-500 (next 25 s), respectively. While not identical both panels show the same pattern and visually appear to have similar intensity. The difference between panels A and B 20 is shown in panel C at the same vertical scale as A and B and expanded 5× in panel D. Importantly, the peaks in panel D, while minute compared with A and B are all positive revealing that the rate decreases slightly as the reaction progresses, not surprisingly as reagents are being consumed.

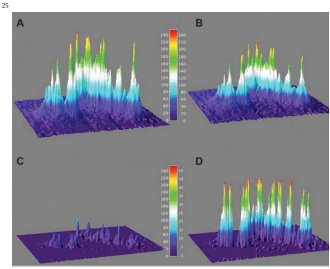


Figure 6: 3D plots of accumulated emission intensity over a catalyst surface over 25 s (A) the following 25 s (B), a difference spectrum of (A) minus (B) shown in (C) and a rescaled image of (C) at $5 \times$ the scale of (A), (B) and (C) shown in panel (D). The area displayed is $15.75 \times 22.5 \,\mu m$.

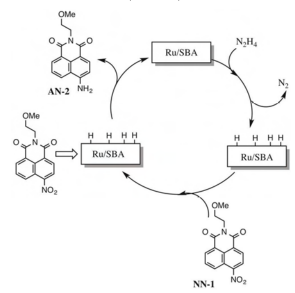
Discussion

A common question for heterogeneous catalysis in liquid systems is whether or not the reaction is happening at the surface of the catalyst or in solution? In other words, do heterogeneous catalysts promote homogeneous catalysis through ion leaching

5 promote homogeneous catalysis through ion leaching mechanisms?¹²

Table 2 summarizes the results for the reaction in scheme 1 (selected for single molecule work) as well as control reactions. From the table of results, it is clear that the reaction is catalyzed 40 by the RuNP supported on SBA as it is the only entry with complete conversion of NN-1 to AN-2. Gas evolution is observed upon addition of hydrazine to solutions of catalyst, presumably due to H₂ evolution. Therefore, it was thought possible that this reaction could occur by H2 reaction with NN-1 in solution and 45 that the catalyst merely facilitates the oxidation of hydrazine to produce H₂. For this reason, the reaction was tested with saturated solutions of H₂ gas rather than hydrazine as the reducing agent (entries 2, 3 and 4 in Table 2, respectively). With the catalyst present and under a H₂ atmosphere, only 6 % conversion was 50 observed as compared with 100 % conversion when hydrazine was used. This indicates that the catalyst has a more important role than the mere release of H₂ from hydrazine to the reaction solution. In fact, this is consistent with the idea that hydrazine activation occurs at the catalyst surface.

ss Microscopy studies, and in particular FCS curves, can be used to examine the binding of both the starting material and products of the reaction with 375 nm and 440 nm excitation, respectively. The results indicate that both the starting material and products have an affinity for the catalyst, spending more time bound to the surface than for normal diffusing in solution. Since the FCS curves eventually drop to zero in all cases, this indicates that, not only are they binding to the catalyst, but the binding appears reversible, as expected for an efficient catalytic process. If the binding was irreversible, residual fluorescence would remain constant over time. From these results and the NN-1 conversion studies, we propose a mechanism for this reaction that involves the RuNP activation of hydrazine as well as surface catalyzed reduction of NN-1 to AN-2 (Scheme 2).



70 Scheme 2. Proposed mechanism for the Ru@SBA catalyzed reduction of nitro compounds to amines, illustrated for the example of NN-1.

The microscopy studies reveal that the spots where catalysis occurs keep repeating themselves, and that the overal shape of the active area corresponds to that of the support particle, ^{12, 16} thus demonstrating that catalysis occurs at the catalyst as opposed to 5 in solution as the result of catalyst leaching. ¹⁷ Further, comparison of sequential groups of frames show that there is a slight decrese in rate as the reaction proceeds (Figure 6); beyond the obvious slow down due to reagent gradual depletion, there may be a small decrease of catalytic actitvity, as suggested also

by the data in Figure 2. Further, FCS studies reveal that once the reaction occurs, the product stays at the catalyst site for about 1 ms; clearly, strategies that reduce this unproductive product retention could enhance the overall catalytic efficiency. The mechanism proposed involves reduction by activated hydrogen generated on the catalyst surface; simply making H₂ available does not lead to comparable reduction levels.

Conclusions

RuNPs highly dispersed into the channels of SBA have given excellent yields when catalyzing the reduction of nitroarenes under air at 80°C, using hydrazine monohydrate as the reducing agent. The experimental conditions were optimized based on a high-throughput screening analysis. From high throughput screening of reactions and bench top reduction of **NN-1** to **AN-2**, to single molecule microscopy experiments we examine

25 to single molecule microscopy experiments, we examine reduction reaction 'from the mole to the molecule'. The rationale for this approach is to elucidate a complete picture of the mechanism of this reaction performing the single molecule spectroscopy for systems that can be tested at the large scale level and where the catalyst and the catalytic conditions have been optimized.

In order to be able to use single molecule microscopy techniques, we use a model reaction with fluorescence activation to study the activity of SBA supported RuNP as a catalyst for the reduction of nitro compounds to amines. The reduction results in a red-shift in absorption/emission as well as a fluorescence quantum yield increase. Single molecule fluorescence microscopy techniques results are consistent with the proposed reversible binding of the reaction species with the catalyst and with the proposed mechanism of hydrazine activation, as well as reduction of the nitro compound on the catalyst surface. Overall, this example not only shows the selective activity of this RuNP catalyst towards reduction of nitro compounds to amines, but also provides an example of how advance microscopy, in particular FCS and

45 TIRF, can be advantageous in establishing catalytic mechanisms. We anticipate that in the near future it may be possible to use the reverse approach "from the molecule to the mole", combined with recent advances in computational catalysis 18 will become a powerful tool in the design, optimization and scale-up of catalytic 50 processes.

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