Solid substrates decorated with Ag nanostructures for the catalytic degradation of methyl orange

Menekse Sakira, M. Serdar Onses⁎

⁎ Corresponding author at: Department of Materials Science and Engineering, Nanotechnology Research Center (ERNAM) Erciyes University, Kayseri 38039, Turkey.
E-mail address: onses@erciyes.edu.tr (M.S. Onses).

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ABSTRACT

There is a strong demand for development of catalytically active solid substrates for heterogeneous catalysis applications. This study reports fully solution-processable and scalable fabrication of solid substrates decorated with Ag nanostructures for the degradation of organic dyes. Ag nanostructures were prepared by direct surface-growth from Pt nanoparticles that were immobilized on Si substrates modified with a layer of end-grafted poly(2-vinylpyridine). The proper choice of the growth conditions and seed-selective growth from Pt nanoparticles were critically important in fabricating Ag nanostructures with high catalytic activity and large surface coverage. The catalytic performance of the presented platform was studied by the reduction of methyl orange by borohydride ions and monitored using UV–visible spectrometry. The substrates exhibited high catalytic activity enabling degradation of 10⁻⁵ M methyl orange solution in less than an hour with an apparent reaction rate constant of 33.5 × 10⁻³ min⁻¹. The substrates can be easily removed from the degradation medium and used multiple times. Our approach presents an effective strategy for waste water management applications avoiding the agglomeration and separation issues of colloidal catalysts and overcoming the need for tedious and costly fabrication of thin films.

Introduction

The growth of the population together with industrialization have resulted in a strong demand for clean water. A source of water pollution consists of organic dye molecules used in textile, paper, leather, plastic, cosmetic, food, and pharmaceutical industries [1–3]. The presence of these hazardous dye molecules in natural water resources poses serious risks to the ecological system and human health. Besides their toxicity and carcinogenicity, these dyes limit penetration of sunlight and reduce the amount of the dissolved oxygen in water [4]. The removal of organic dyes from water is challenging, due to their stable and complex aromatic molecular structure [5]. These dyes, such as methyl orange (MO), and Congo red, are characterized by nitrogen to nitrogen double bounds (N=N), named as azo linkage. The cleavage of azo bond causes to the decoloration of dyes. These dyes can be highly hazardous. MO, for example, is highly toxic, may be fatal if inhaled, swallowed or absorbed though skin [6]. It is known that the reduction of MO is thermodynamically favorable but kinetically unfavorable [7]. This challenge strongly motivates research into classes of materials that can transform these organic dye molecules into environmentally benign forms.

The use of nanomaterials for catalytic degradation of organic dyes presents an effective solution to this challenge. The large surface to volume ratio of nanoparticles (NPs) and high activity of atoms with low coordination number make nanomaterials appealing for heterogeneous catalysis applications [8,9]. Pt NPs are one of the most efficient catalysts due to their stability, the abundance of active edges, and high specific surface area [10]. The limited reserves and high cost of Pt, together with opportunities to synergistically improve their catalytic properties through the charge transfer, lattice strain and tailored surface compositions resulted in investigation of bimetallic nanostructured catalysts prepared by alloying Pt with various noble metals (Au, Ag, Pd) [11–13] and transition metals (Ni, Co, Fe) [14–16]. With its relatively low-cost, unique plasmonic and catalytic properties, Ag is a logical choice [17]. In addition, both Pt and Ag have a face-centered cubic (fcc) crystal structure and their lattice parameters are close to each other, favoring preparation of bimetallic catalysts [18]. The catalytic efficiency of bimetallic nanostructures depends on their morphology, size, composition and chemical structure. As a result, a range of different nanostructured catalysts consisting of Ag and Pt has been reported for...
catalytic degradation of nitrophenol, Congo red, rhodamine B, and methyl violet dyes in water using Ag@Pt core-shell [19], Pt@Ag core-shell [20], hollow Pt/Ag nanocomposite [21], and Ag/Pt bimetallic catalysts [22]. The size, structure and shape of the nanostructures were critically important in the catalytic activity [23,24]. To provide flexibility, reusability and enhanced catalytic activity, these heterogeneous catalysts are often used in colloidal forms either by doping metal oxide crystals such as Al2O3 and SiO2 [25,26] or by attachment on other nanomaterials including graphene oxide layers [27], Si nanowires [28], and nanofibers [29]. Despite the high catalytic activity of colloidal nanomaterials, a number of processing steps, such as centrifugation, filtration, and washing, is necessary for their separation and reuse [30]. Recycled catalysts have significant technological value in the energy, environmental and biomedical fields [31,32]. Additionally, colloidal nanomaterials can easily aggregate resulting in loss of the catalytic activity associated with the decrease in the active surface area.

A promising approach to overcome these problems in colloidal catalysts involves the use of solid substrates as supports for the catalytic nanostructures. The direct growth of nanostructures on the surface of solid substrates via vapor-phase deposition methods, such as magnetron sputtering and atomic layer deposition, is a commonly adopted strategy. Miotella et al. synthesized Co NPs embedded B matrix films by pulsed laser deposition (PLD) for hydrogen production by hydrolysis of NaBH4 and NH3BH3. They further assembled Co oxide NPs on thin films fabricated by reactive PLD for the degradation of methylene blue [33]. Henkel et al. prepared TiO2 thin films on quartz and phosphor doped Si substrates by magnetron sputtering for the degradation of methylene blue [34]. Zeng et al. reported substantial improvements in the photocatalytic yields for CO2 reduction to CO through the surface passivation of InP with TiO2 deposited by ALD [35]. These vapor-phase deposition methods allow for fabrication of uniform films of catalytic nanostructures on the surface of the substrates with precise control over the structure and composition of the NPs; however, the need for expensive and complex infrastructure together with limitations in the surface coverage and, throughput challenge their use in practical applications. In a complementary strategy, catalytically active solid substrates can be fabricated by immobilization of colloidal nanostructures or wet-chemistry based surface growth of nanostructures. Catalytically active nanocomposite surfaces were fabricated by in situ synthesis of Pd and Pt NPs inside poly(2-vinylpyridine) (P2VP) brushes [36]. In a recent study, polymer ionic liquids grafted to a glassy carbon electrode, was presented as an efficient electrocatalyst for oxygen reduction reaction [37]. Constantinii et al. reported surface-initiated polymerization followed by in-situ reduction of metal ions to functionalize the inner walls of glass microreactors for the reduction of 4-nitrophenol (4-NP) [38]. Mussel inspired polydopamine is another approach to prepare catalytically active solid substrates through aqueous functionalization followed by the immobilization or growth of metallic NPs [39,40]. A challenge associated with these previous demonstrations is that the surface coverage of NPs was low and there was only one type of nanostructure limiting the processes of electron transfer for development of synergetic catalytically active solid substrates.

Herein we report a scalable, solution-processing based and versatile strategy for fabrication of catalytically active solid substrates. Our strategy relies on surface-growth of Ag nanostructures from Pt NPs that were immobilized on solid substrates functionalized with a layer of end-grafted P2VP. The use of a seed selective reducing agent, hydroquinone, enabled direct surface growth of the Ag nanostructures with high surface coverage on the solid substrates. The catalytic degradation of MO dye by borohydride ions was studied as a model reaction. The catalytic activity of the fabricated substrates was monitored using UV–visible spectrophotometry. The impact of the growth conditions on the size and structure of the nanostructures and their catalytic activity were investigated. The fabricated solid substrates enabled complete degradation of 10−5 M MO solution in less than an hour, showing the promise of the presented platform for catalytic degradation of organic dyes. Our approach presents a simple yet effective strategy avoiding the agglomeration and separation issues of colloidal catalysts and overcoming the need for tedious and costly fabrication of thin films.

**Experimental section**

**Materials**

Si wafers (1 0 0) were purchased from Wafer World Inc. Silver nitrate (AgNO3, ≥99.5%), hydroquinone (C6H4O2, abbreviated as HQ, ≥99%), N, N-dimethylformamide (HCON(CH3)2, abbreviated as DMF, ≥99%), methyl orange (C14H14N3NaO3S, abbreviated as MO), sodium borohydride (NaBH4, ≥98%) were purchased from Sigma-Aldrich. Hydroxyl-terminated poly(2-vinyl pyridine) (Mw = 20.5 kg/mol, polydispersity index = 1.04, abbreviated as P2VP-OH) was purchased from Polymer Source Inc. Citrate-stabilized Pt NPs with an average diameter of 30 nm were purchased from NanoComposix. All aqueous solutions were prepared with purified water.

**Fabrication of solid substrates decorated with Ag nanostructures**

**Immobilization of Pt seeds**

Si substrates (1 × 1 cm2) were cleaned in a UV-ozone chamber for 20 min. A film of P2VP-OH was then deposited on the freshly cleaned Si substrate by spin-coating from a 3% solution in DMF at 3000 rpm for 30 s. The substrates were then thermally annealed at 180 °C for 5 min on a hot-plate in a glove box filled with argon. The annealing step was followed by washing with DMF under sonication for three times, 3 min each, and drying with nitrogen. Citrate-stabilized Pt NPs were then immobilized on the P2VP-grafted substrates by spotting a suspension (50 μL/cm2) of the particles for 3 h. The substrates were then washed with water under sonication for 2 min and dried with nitrogen.

**Growth of Ag nanostructures**

The substrates with the immobilized seed particles were immersed into the growth solution including 0.2 mM AgNO3 and 0.4 mM HQ in 50 mL water for varying times (1 h, 4 h, 24 h, 48 h) under constant agitation in a dark environment. Following the growth of the Ag nanostructure on Pt seeds, the substrates were then rinsed with water and dried with nitrogen.

**Characterization**

The morphology and chemical composition of the substrates were studied using scanning electron microscopy (SEM, Zeiss EVO LS10) at 25 keV and energy dispersive X-ray spectroscopy (EDX, Bruker). The X-ray thin film diffraction pattern was recorded with a Rigaku SmartLab diffractometer operating at 40 kV and 30 mA by using Cu Kα radiation source and a scanning rate of 5°/min in the range of 35–70°. UV–visible spectroscopy (Perkin Elmer Lambda 25) was recorded in the wavelength range of 355–575 nm to monitor the catalytic degradation of MO. The thickness of the grafted P2VP layer was measured via an ellipsometer (Gaertner LSE Stokes). The refractive index of P2VP was assumed as 1.595. The compositions of Pt seeds and Ag nanostructures were examined with X-ray photoelectron spectroscopy (XPS, SpecsFlex) using XRTm50 M (UXC1000) source exciting radiation (1486.71 eV). All binding energies were referenced with respect to the C 1s peak at 284.8 eV.

**Catalytic degradation of methyl orange**

The catalytic reduction of MO was selected as a model reaction. The catalytic reduction of MO was carried out in a glass beaker with a volume of 80 mL. In a typical experiment, an aqueous solution of MO with a concentration of 10−5 M and a volume of 35 mL was prepared. The Si substrate with an area of 1 × 1 cm2 decorated with Ag nanostructures were immersed into the reaction mixture (1 g/L) at 100°C, and the reaction was carried out for 30 min. The reaction mixture was centrifuged and filtered, and the absorbance at 465 nm was measured using an UV–visible spectrophotometer. The catalytic activity of the catalysts was calculated using the following equation:

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\text{Catalytic activity} = \frac{\text{Initial absorbance} - \text{Final absorbance}}{\text{Time}}
\]

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was immersed in the MO solution (Fig. S1). The solution was magnetically stirred at room temperature for 30 min to facilitate interaction between the dye molecules and catalytic substrate. The reduction process was initiated by adding 25 mL of freshly prepared 0.033 M NaBH₄ solution. The reaction solution was pipetted into a quartz cell and its absorption spectrum was measured for different reaction times using an UV–visible spectrophotometer. The color of the MO solution gradually changed from yellow to colorless, indicating that the reduction of MO catalyzed by the substrate. The catalytic degradation of MO is schematically illustrated in Fig. S2. The degradation % was calculated according to the following expression:

\[
\text{Degradation} \% = \frac{A_0 - A_t}{A_0} \times 100
\]

where \( A_0 \) is the initial absorbance of MO at 464 nm and \( A_t \) is the absorbance of MO at 464 nm after “t” min. The effect of NaBH₄ was investigated with a control experiment where the degradation was performed in the absence of the catalytic substrate. To test the stability and reusability, the substrates were subjected to 4 consecutive cycles of the catalytic degradation reaction. The catalytic substrates were sonicated in water after each cycle for 15 min to remove the weakly bound dye molecules from the substrate.

**Results and discussion**

**Fabrication and structural characterization of the catalytically active substrates**

Our process to prepare catalytically active solid substrates begun with grafting P2VP-OH onto the freshly cleaned Si substrates (Fig. 1a). The grafting was achieved through a condensation reaction between the hydroxyl-terminus of the polymer and surface silanol groups. The washing of the substrates in DMF under sonication was important to remove excess and unreacted polymers from the surface. The thickness of the grafted P2VP layer was 5.6 nm with a grafting density of ~0.16 chains per square nanometer. The grafted P2VP layer provided a covalently bonded, robust interface for the uniform immobilization of seed NPs [41]. The pyridine group is capable of binding to citrate-stabilized metallic NPs through electrostatic interactions [42,43]. We used citrate-stabilized Pt NPs with an average diameter of 30 nm as seeds. The immobilization was performed by drop-casting the suspension of Pt NPs followed by washing in water under sonication. SEM imaging (Fig. 1b) verified the uniform immobilization of Pt NPs with an average surface density of 136 particles/μm². The P2VP grafted substrates with the immobilized Pt NPs were then immersed in the growth solution containing the metal salt and reducing agent. At proper concentrations and ratios of AgNO₃ and HQ, the growth resulted in high density Ag nanostructures over the entire substrate area (Fig. 1c). The surface growth of the Ag nanostructures was achieved via the use of a specific reducing agent, HQ, that only functions in the presence of the seed particles [44]. Control experiments with the P2VP-grafted substrates in the absence of Pt NPs showed that the seed particles are necessary for the growth of Ag nanostructures (Fig. S3).

The concentration of the reducing agent and its ratio to the metal salt was important for the successful growth of the Ag nanostructures. We first fixed the concentration of AgNO₃ at 0.2 mM and varied the concentration of the reducing agent. The growth was performed for 24 h on the P2VP-grafted substrate with the immobilized Pt NPs. At a HQ concentration of 0.4 mM, the entire substrate was covered with the Ag nanostructures with an average size of 511 nm (Figs. 2a, S4a). The further increase of the HQ concentration resulted in the decrease of the size (320 nm) of the Ag nanostructures together with a slight reduction of the surface coverage and increased stacking of the nanostructures (Figs. 2b, S4b). At a fixed HQ concentration of 0.4 mM, the average sizes (Fig. S4c, d) of the Ag nanostructures were 110 nm, 511 nm and 433 nm for AgNO₃ concentrations of 0.1 mM, 0.2 mM and 0.4 mM, respectively. The reduction in the size of the nanostructures at 0.1 mM AgNO₃ could be related to the lack of sufficient metal ions in the solution. HQ has two hydroxyl groups, which get oxidized to reduce silver ions. Our results suggest that a slight excess of HQ is necessary for the high surface coverage of Ag nanostructures that are grown from immobile Pt NPs. These results are in agreement with our previous study on the growth of the Ag nanostructures on solid substrates decorated with Au NPs [44]. The further increase of the HQ concentration (e.g. Fig. 2b) adversely affected the growth process, as reported in the case of colloidal synthesis of gold NPs [45].

The size of the Ag nanostructures was highly dependent on the growth time. Fig. 3 presents SEM images of the Ag nanostructures that were prepared by incubating the substrates with the immobilized Pt NPs in 0.2 mM AgNO₃ and 0.4 mM HQ for different times. The size of Ag nanostructures increased with the growth time. At a short growth time of 1 h, the average size of Ag nanostructures was 208 nm with a low surface coverage (Figs. 3a, S5a). Increasing the growth time to 4 h resulted in a high surface coverage of the Ag nanostructures with a slight increase in their sizes (230 nm) (Figs. 3b, S5b). For a growth time of 24 h, the entire substrate was covered with the Ag nanostructures with an average size of 511 nm (Figs. 3c, S5c). The further increase of the growth time resulted in significantly larger (average size of 1.7 μm) structures, which likely resulted from coalescence of the smaller nanostructures (Figs. 3d, S5d). At such long growth times, detachment of these large particles from the substrate resulted in low surface density of Ag structures. Accordingly, the growth of catalytically active...
The chemical composition and crystal structure of the nanosstructures were characterized by EDX and XRD analysis. The EDX spectrum given in Fig. S6 clearly shows the presence of the characteristic peak (25 keV) of Ag, further verifying the growth of the Ag nanostructures. The peak at ∼2.0 keV corresponds to Pt, which confirms the presence of the seed NPs after the growth process. The ratio of Ag:Pt, as calculated from EDX spectrum, is approximately 4:1. The crystal structure of the substrate decorated with Ag nanostructures was probed with XRD analysis (Fig. 4a). The strong peaks at 38.10°, 44.31° and 64.45° were attributed to diffraction from the (111), (200), and (311) planes of Ag, respectively.

Fig. 2. SEM images of the Ag nanostructures grown at varying concentrations of the reducing agent and metal precursor. The growth solutions consisted of (a) 0.2 mM AgNO₃ and 0.4 mM HQ, (b) 0.2 mM AgNO₃ and 0.6 mM HQ, (c) 0.1 mM AgNO₃ and 0.4 mM HQ, and (d) 0.4 mM AgNO₃ and 0.4 mM HQ. The substrate consisted of Pt NPs immobilized on P2VP grafted Si substrates. All growth processes were terminated after 24 h by rinsing with water.

Fig. 3. Influence of the growth time on the size and morphology of Ag nanostructures. SEM images of the Ag nanostructures grown for (a) 1 h, (b) 4 h, (c) 24 h, and (d) 48 h. The growth was performed in a solution containing 0.2 mM AgNO₃ and 0.4 mM HQ.

substrates was performed in 0.2 mM AgNO₃ and 0.4 mM HQ for 24 h, in the rest of the study.

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planes of fcc structured Ag (JCPDS card # 089-3722), respectively [46]. There is an induced strain in the Pt-Ag system [47,48] due to the lattice incompatibility between Pt (3.92 Å) and Ag (4.08 Å). Therefore, Ag ions did not interfere with Pt and grew on the Pt seeds in the crystal structure. The crystal growth is dominated in the (111) plane, since this plane has a low energy [49]. The standard diffraction peaks at 38.69°, 44.97° and 65.49° from the (111), (200), and (220) planes of Pt (JCPDS card # 087-0644) were absent in the XRD pattern of the substrate [46]. The absence of Pt is likely a result of the insufficiently low intensity signals from the Pt NPs underneath the Ag nanostructures [19].

The chemical composition of the nanostructures was further characterized via XPS analysis. The XPS survey scan of the substrates given in Fig. 4b showed the presence of Ag, Pt, C, O and N elements. The characteristic peaks associated with C, O and N likely originated from the growth byproducts and grafted P2VP layer. The predominant peaks centered around 368 eV were referred to the Ag 3d electrons [50]. The Ag 3d peaks confirmed that Ag ions were successfully reduced and deposited onto the Pt seeds. The Pt 4f peaks located around 71.6 eV were also observed in the spectra indicating the presence of Pt seeds [51]. The intensities of the Pt peaks were significantly lower than the peaks of Ag. This result together with the surface-sensitive nature of XPS collecting data from a depth of ~5 nm, further supports that the Ag nanostructures grew on the Pt seeds over the entire substrates with a high surface coverage. To further probe the interaction between Ag and Pt, we analyzed the Pt 4f and Ag 3d regions of the XPS spectra. The Pt 4f XPS spectrum (Fig. 4c) of the Ag nanostructures exhibited two peaks that correspond to Pt 4f7/2 and Pt 4f5/2 electrons at 71.4 eV and 74.9 eV, respectively. In the absence of the growth of Ag nanostructures, Pt seed NPs immobilized on the P2VP grafted substrates displayed peaks centered at 71.8 eV (Pt 4f7/2) and 75.2 eV (Pt 4f5/2). The growth of Ag nanostructures from the Pt seed NPs resulted in a negative shift in the binding energies of Pt. The high-resolution XPS spectrum of the Ag nanostructures over Ag 3d region (Fig. 4d) exhibited the characteristic peaks, which were split into 5/2 and 3/2 states positioned at 368.3 and 374.3 eV, respectively. To compare the electronic structure of Ag nanostructures that were grown on the Pt seeds, we prepared a P2VP grafted substrate and immobilized citrate-stabilized Ag NPs. The Ag 3d5/2 and 3d3/2 peaks were located at 369.0 eV and 375.0 eV for the immobilized Ag NPs. The binding energies of the 3d electrons for the Ag nanostructures have a negative shift of 0.7 eV in comparison with that of Ag NPs. The shifts in the binding energies of both Pt 4f and Ag 3d electrons suggest the modification of the electronic structure of Pt and Ag atoms as a result of the growth of Ag nanostructures from Pt NPs immobilized on the P2VP grafted substrate. This type of modification in the electronic structure was observed for bimetallic catalysts that consist of Au and Pt [52]. The modification of the electronic structure can be related to the transfer of electrons between Ag and Pt. The electronegativity of Ag is lower than Pt, which implies that electrons can be transferred from Ag to Pt. All the structural and spectroscopic characterization confirmed the successful growth of Ag nanostructures on Pt seeds that were immobilized on the grafted P2VP layers.

Catalytic activity

Degradation of methyl orange and kinetics of reaction

The catalytic activity of the substrates decorated with Ag nanostructures was investigated using the reduction of MO in the presence of
NaBH₄. These experiments were performed by immersing the substrates into a solution of MO. The degradation process started with the addition of the reducing agent, NaBH₄. Fig. 5a presents UV–visible absorption spectra of the MO solution taken at different times (0, 5, 10, 20, 30, 40, 50, and 60 min). Before the degradation process, MO exhibited the maximum absorbance in the visible region at a wavelength of 464 nm associated with the azo linkage [53,54]. This absorbance was taken as the basis for monitoring the degradation process in the rest of the study and the degradation percentage was calculated using Eq. (1). When the catalytically active substrate and the freshly prepared reducing agent were added to the dye solution, the absorption at 464 nm decreased as the time progressed. The degradation rate increased rapidly on the Ag nanostructures following the first 10 min, perhaps related with the activation of the surface and diffusion of the dye and reducing agent to the catalytically active sites. The large surface area, high surface coverage, and irregular shape of the Ag nanostructures likely contribute to this behavior. The absence of such an abrupt increase in the degradation rate for the solid substrates decorated solely with the Pt NPs can be related to the low degradation rate together with accessibility of the surface of the NPs due to large particle-to-particle distances. At the end of 60 min, 83% of the initial MO was degraded. We performed control experiments to understand the origin of this catalytic activity (Fig. 5b). In the first set of experiments, the degradation process was monitored in the absence of any substrate (Fig. S7a). The degradation proceeded slowly and only 18% of the initial dye was degraded at the end of 60 min when there was no substrate in the solution. The second control experiment was performed using just the Pt seed NPs immobilized on the P2VP grafted substrate (Fig. S7b). The degradation percentage was as high as 22% at the end of 60 min. Despite a slight increase in the efficiency with the presence of the immobilized Pt NPs, the degradation is much more limited in comparison with the Ag nanostructures grown on the Pt seeds. This result clearly demonstrated the high catalytic activity of the fabricated substrates. Three factors contribute to this high catalytic activity: i) The total catalytically active surface area of the Ag nanostructures is considerably larger than the surface area of the Pt seeds, due to the high coverage of the former structure and the reduced active area of the Pt NPs associated with the interaction with the end-grafted P2VP layer. ii) The work function of Ag NPs is lower than the Pt NPs [55]. iii) The processes of electron transfer between Pt NPs and Ag nanostructures can synergistically improve the catalytic activity [56].

The catalytic activity of the substrate depended on the extent of Ag growth on the immobilized Pt NPs. Fig. 5c compares the degradation of MO using substrates prepared by growth of Ag for 1 h, 4 h, 24 h, and 48 h. The degradation was highest in the case of Ag nanostructures grown for 4 h. After 60 min of degradation, for example, 83%, 74%, 56%, and 46% of the initial MO were degraded for substrates with the Ag growth times of 24 h, 48 h, 4 h, and 1 h, respectively (Fig. S8). Considering these results with the SEM images of these structures (Fig. 3) suggest that both the size of Ag nanostructures and their surface coverage are important for the catalytic activity. At long growth times, the large size of Ag structures approaching the micrometer length scale likely decreased the catalytic activity of the substrate. The short growth times, on the other hand, were not sufficient for the growth of Ag nanostructures over the entire substrate.

Kinetics and mechanism of degradation of methyl orange We investigated the kinetics of the catalytic degradation reaction on the substrates decorated with Ag nanostructures. In the presence of excess of the reducing agent, the rate of the degradation of organic dye molecules is commonly [57] expressed by pseudo-first order kinetics, which is derived from the Langmuir-Hinshelwood mechanism [58]. The rate of the reaction expression is given by Eq. (2) where Cᵣ refers to the concentration of MO at time t. The concentration of MO can be approximated by the absorbance, Aᵣ, at 464 nm. The linearized form of the expression is obtained by integrating the expression and given by Eq. (3), where A₀ and Aₜ are the initial and final absorbance of MO at 464 nm, respectively [7].

\[
\frac{dC_r}{dt} = \frac{dA_r}{dt} = -k_{app}A_t
\]  

Eq. (2)
ln \left( \frac{C_t}{C_0} \right) = ln \left( \frac{A_t}{A_0} \right) = -k_{app}t \quad (3)

Fig. 6a presents the plot of ln(A_t/A_0) against time for different types of substrates. A reasonably good linear relationship was obtained in the plot, showing that the degradation follows a pseudo first-order kinetics. The apparent reaction rate constant, k_{app}, was derived from the slope of the linear fit to the plot. The apparent reaction rate constant for the Ag nanostructures was 33.5 \times 10^{-3} \text{ min}^{-1}, which was roughly ten times higher than the constant in the case of just Pt seeds (k_{app} = 3.4 \times 10^{-3} \text{ min}^{-1}) and degradation in the absence of a substrate (k_{app} = 2.29 \times 10^{-3} \text{ min}^{-1}). This result further highlights the significant enhancement of the degradation with the use of the substrates decorated with Ag nanostructures. The reaction rate constant of our platform is higher than the recent reports based on the green Ag NPs [4], AgCl nanowires decorated with Au NPs [59] and organic semiconductor film@Au [60].

The degradation of MO on the substrates decorated with Ag nanostructures is likely to proceed through the mechanism depicted in Fig. 6b. The catalytic degradation process relies on the transfer of electrons from the donor (NaBH_4) to the acceptor (MO). The simultaneous adsorption of both the dye and BH_4^- ions on the surface of the nanostructures is the first step for the degradation. The Ag nanostructures serve as electron relay systems facilitating the electron transfer between the electrophilic dyes and nucleophilic BH_4^- ions [61]. The processes of electron transfer results in the gain of electrons by MO, leading to the reduced form of the dye molecule. Previous studies suggested that the electron transfer from BH_4^- ions to the dye molecules mediated by the metallic nanostructures in water results in formation of BO_3^{3-} [62]. Such electron transfer can further proceed and result in more reduced forms of the dye molecule [63]. The high catalytic activity of our platform can be related to the large surface area to volume ratio, high surface coverage and irregular shape of the Ag nanostructures, which facilitate electron transfer and enable overcoming the kinetic barrier for the degradation reaction. Since the reaction occurs on the surface, increasing the area enhances the rate of the degradation. The processes of electron transfer between Ag and Pt suggested by XPS analysis can further contribute to this catalytic activity [50]. The interaction of functional groups [64] presented by the end-grafted P2VP with noble metals can also play a role in the catalytic activity of the presented platform.

Reusability of catalytic substrates

Inherent advantages of the catalytically active solid substrates are the ease of separation from the reaction medium and ability for the repeated use of the catalyst. Together with the efficiency of the catalytically active surfaces, the stability of the material is highly critical for practical waste-treatment applications. To investigate the stability of the solid substrates decorated with Ag nanostructures, we performed four consecutive cycles of degradation on the same substrate. The substrate was simply removed from the reaction medium without any additional processing. The substrate was washed under sonication to remove the bound dye molecules, prior to the following cycle. Fig. 7 presents the degradation percentage of MO after each cycle. The substrate retained most of its catalytic activity with a slight decrease in the degradation efficiency with the increasing number of cycles (Fig. S9). The degradation, for example was reduced by 3.5% and 12% after the second and fourth cycles with respect to the first cycle, respectively. This reduction in the degradation efficiency is perhaps associated with...
the incomplete removal of the bound dye molecules from the surface, resulting in the decay of the catalytically active sites. The possibility of destruction and removal of the nanostructures are excluded because SEM imaging of the substrates following four cycles showed the retention of the Ag nanostructures (Fig. S10). These results further verify the easy separation of the catalyst from the reaction medium without any additional processing steps and ability of reusing the fabricated catalytically active solid substrates.

Conclusions

In conclusion, this study demonstrated the promise of solid substrates decorated with Ag nanostructures for the catalytic degradation of organic dyes. The versatility and scalability of our strategy emerge from the all-solution processing-based fabrication of catalytically active solid substrates. The conditions for growth of Ag on Pt NP's immobilized on end-grafted 2PVP determine the size, surface coverage and catalytic activity of the nanostructures. The seed-mediated growth of Ag nanostructures on the immobilized Pt NPs plays a key role in the high degradation performance of our platform by presenting large areas of catalytically active surfaces. The easy removal of the solid substrates from the degradation medium together with the ability to reuse the substrates are important advantages for practical applications. The presented results motivate research in other classes of materials for fabrication of catalytically active solid substrates decorated with nanostructures.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.rinp.2018.12.084.

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