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Electroless Ni plated nanostructured $TiO₂$ as a photocatalyst for solar hydrogen production†

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Herein, we have demonstrated a facile electroless Ni coated nanostructured TiO₂ photocatalyst for the first time. More significantly the photocatalytic water splitting shows excellent performance for hydrogen production which is hitherto unattempted. The structural study exhibits majorly the anatase phase along with the minor rutile phase of TiO₂. Interestingly, electroless nickel deposited on the TiO₂ nanoparticles of size 20 nm shows a cubic structure with nanometer scale Ni coating (1–2 nm). XPS supports the existence of Ni without any oxygen impurity. The FTIR and Raman studies support the formation of $TiO₂$ phases without any other impurities. The optical study shows a red shift in the band gap due to optimum nickel loading. The emission spectra show variation in the intensity of the peaks with Ni concentration. The vacancy defects are pronounced in lower concentrations of Ni loading which shows the formation of a huge number of charge carriers. The electroless Ni loaded $TiO₂$ has been used as a photocatalyst for water splitting under solar light. The primary results manifest that the hydrogen evolution of electroless Ni plated TiO₂ is 3.5 times higher (1600 μmol g^{−1} h^{−1}) than pristine TiO₂ (470 μmol g^{−1} h^{−1}). As shown in the TEM images, nickel is completely electroless plated on the TiO₂ surface, which accelerates the fast transport of electrons to the surface. It suppresses the electron–hole recombination drastically which is responsible for higher hydrogen evolution using electroless Ni plated TiO₂. The recycling study exhibits a similar amount of hydrogen evolution at similar conditions which shows the stability of the Ni loaded sample. Interestingly, Ni powder loaded $TiO₂$ did not show any hydrogen evolution. Hence, the approach of electroless plating of nickel over the semiconductor surface will have potential as a good photocatalyst for hydrogen evolution. PAPER

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Introduction

The environment and energy have been progressively decisive issues in the last two decades, in connection with the elusive climatic changes emerging worldwide due to the vast use of fossil fuels. Thus, energy sources that are bountiful, regenerative, and free from carbon emissions are in sizable demand. Solar energy is securing a lot of attention as a sustainable and clean energy source. However, for the sake of its practical utilization, solar energy must be reconstructed into other types of energy, such as chemical fuels, electricity, heat, etc. which allow for more effective storage and conveyance. Among chemical fuels, hydrogen $(H₂)$ is quintessentially ascribed to its highest energy capacity of 143 MJ kg^{-1} among all materials over the globe and its emission-free superiority, generating water as the only by-product from combustion energy.¹ Predominantly for the production of hydrogen some cardinal energy systems

are implemented such as thermal, electrical, photonic and biochemical.^{2,3}

Considering the key energy systems, plenty of techniques were employed for the generation of hydrogen, some of them are conventional electrolysis,⁴⁻⁶ high-temperature electrolysis, thermochemical Cu–Cl, hybrid sulphur, Mg–Cl cycles, thermolysis, photoelectrolysis, photoelectrocatalysis, photoelectrochemical cells (PECs), photocatalysis,⁷ hybrid photocatalytic Cu–Cl cycles,⁸ and still search for finding a better technology is going on globally.

Among them, photocatalysis involving the storage of chemical energy in hydrogen molecule enabled by the solar energy is the most cost-effective way of hydrogen production having high efficiency and abundant resource of semiconductor materials such as oxides and sulphides.^{9,10}

Commercialisation of hydrogen production through water splitting has few limitations such as low solar conversion output, fractional solar radiation adsorption range and high rate of charge carrier recombination.^{11,12}

A dependable photocatalyst needs to fulfil important requisites like suitable band gap exhibiting efficient electron–hole separation and transport leading to surface catalytic proton

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reduction as well as water oxidation, and negligeable photocorrosion.¹² Homogeneous photocatalysts possessing these requirements are limited and hence heterogeneous photocatalyst with two or more distinct photocatalysts needs to be employed to overcome the above drawbacks enabling the industrial scale production.¹³

Majority of the semiconductor photocatalysts belong to transition metal cations consisting of metal sulfides, nitrides and oxides having d^0 and d^{10} configurations. Metal sulphide and nitride photocatalysts such as \cos_2 , MoS_2 , Ni_3S_2 , NiCoS_4 , CdS, In_xGa_{1−x}N, GaN, Ta₃N₅ *etc.* are highly prone to photocorrosion and consequently are chemically less stable.¹⁴–¹⁷

Though, metal oxides such as ZnO, WO_3 , Nb_2O_5 , Ag_3PO_4 , $Bi₂WO₆, Ta₂O₅, TiO₂, etc.$ are the most explored materials as photocatalysts, resistance to chemical, thermal and photocorrosion, easy availability, lower production cost and wide absorption range makes $TiO₂$ the material of choice since the time of Honda and Fujishima.^{18,19} Application of TiO₂ in solarlight-driven photocatalysis has been limited owing to its wide bandgap of 3.2 eV which lies in the ultraviolet (UV) region comprising of only 4% of solar spectrum. Paper
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Various approaches have been ventured to overcome the drawbacks associated with $TiO₂$ such as doping (for example with transition metal cations like Cr^{3+} , Fe^{2+} , Ni²⁺ and Rh³⁺, *etc.*)^{20,21} and use of noble metals (Pt,^{22,23} Pd, Rh,^{24,25} Ag and Au^{26,27}) and transition metals like Ni, Cu, Zn, Mn, etc. as cocatalysts.^{11,28,29}

For noble metals, cost is the hindering factor in their practical use as co-catalyst. In recent years, the transition metals like Ni-based (NiO,³⁰ Ni₂P,³¹ NiS,³² and NiSe₂ (ref. 33)), Cu-based $(Cu(OH)_2)^{34}$ CuO³⁵), and Co-based $(COO)^{36}$ Co₃O₄ (ref. 37)) cocatalysts have received considerable attention in the photocatalytic field, attributable to their low-cost, abundant resource, stability, facile preparation and substantial catalytic activity.¹¹

Among these materials, nickel, a low-cost and earth abundant transition metal with a work function of nearly 5 eV has been considered as a potent alternative to noble-metal cocatalysts because it is already employed as a catalyst in many industrial chemical reactions.^{6,7} However, reported hydrogen generation yield of $TiO₂$ –Ni based heterogeneous photocatalytic water splitting systems is very low and there is scope for innovations.³⁸

Liu et al. prepared Ni loaded TiO₂ nanocomposite, with 6 mol% having 10–20 nm size of Ni nanoparticles around the TiO₂ surface using Wet Impregnation Method.³⁹ NiO_x nanoparticles with average size of 24 nm distantly spread over the surface of titania in the 5 wt% NiO_x -loaded TiO₂ nanocomposite.⁴⁰ (9:1) Ni-Zn/TiO₂ in which Ni content is 45 wt% displayed Ni with a diameter over 100 nm, and $TiO₂$ nanoparticles with an averaged diameter of 25 nm. 41 Ni loaded TiO₂ thin films with Ni layer thickness of \sim 50 nm were fabricated by Cheng et al. using the y atomic layer deposition (ALD) technique.⁴² NiO/Ni/TiO₂ sandwich-like heterojunction with Ni layer of ∼0.5 nm grown by the co-precipitation technique over hydrothermally prepared $TiO₂$ nanocables were grown by Ren et al.⁴³ Most of these reports discuss about Ni nanoparticles having the larger diameters or films entirely covering the

semiconductor surfaces which may limit their application as heterogeneous photocatalyst material.

So far, many different methods, including molten-saltmediated process,⁴⁴ precipitation,⁴⁵ solvothermal,^{46,47} sol-gel,⁴⁸ electrodeposition,⁴⁹ and chemical reduction,³⁹ have been reported for synthesis of Ni@TiO₂ systems.⁵⁰ In this context, electroless plating which is generally used for metallization of work-pieces at macro scale, has unexpectedly emerged as a superior wet-chemical nanofabrication alternative.²⁵

This technique, by virtue of its solution chemistry and mechanistic aspects, is capable of creating nanomaterials various shapes such as nanoparticles, nanowires, nanotubes, and ultrathin films, etc., by simply immersing suitable samples into aqueous solutions for defined amounts of time.^{27,51-56}

It offers flexibility in the choice of the substrate to be metallized and is almost independent of its chemical composition, state (solid/liquid/gas), and morphology including gas nano-bubbles, emulsified droplets, biological micro-organisms, microfluidic channels, nanoparticles, and plastics.⁵⁴⁻⁵⁹

In view of this, preparation of $Ni@TiO₂$ photocatalysts was herein accomplished using the in-house developed electroless Ni plating bath and the effect of Ni loading percentage on the photocatalytic hydrogen generation performance has been studied. To the best of our knowledge, preparation of Ni@TiO2 heterogeneous photocatalyst system using electroless method has not been reported. The electroless deposited catalyst has been found to be highly efficient for the hydrogen production.

Materials and methods

Materials

 $TiO₂$ nanopowders were procured from Aldrich (average particle size of 20 nm). For nickel loading commercial electrochemical bath was purchased from Grauer & Weil (India) Limited. Electrochemical commercial bath consisted of cleaning solution, mild etching solution, pre-activator, activator, Ni salt solution and reducing solution. For preparation of 10% sulphuric acid solution, 98% conc. H_2SO_4 procured from SRL chemicals, India was used.

Material preparation

The detailed of bath preparation process has been mentioned in the ESI as Table T1,† all the solutions were prepared as per the data sheet of the manufacturer, Grauer & Weil (India) Limited. Electroless nickel loading process involves different process steps which are carried out under continuous stirring and centrifugation is followed at the end of each process step which are mentioned in the Table T2 in ESI† and Fig. 1 depicts all the processes involved together. In a typical process, $TiO₂$ nanopowder surface was sequentially cleaned using the cleaning solution (maintained at 60 °C for 10 min), mild etching solution (4 min) and acid solution (2 min). At the end of each step, water washing was undertaken followed by centrifugation. Cleaned $TiO₂$ nanopowder surface, was treated with pre-activator solution for 2 min. However, water rinsing was not carried out after this step.

Preactivated $TiO₂$ nanopowder was surface activated by administering it to activator and post activator solutions for 5 min each followed by water rinse. Subsequently, surface activated TiO₂ (Fig. 2) nanopowder was subjected to Ni loading by transferring it to Ni bath maintained at the temperature of 85–90 °C and pH ∼ 5.2, for 15 min. For varying Ni loading, surface activated $TiO₂$ nanopowders were treated with electroless Ni baths of various concentration and the samples, thus prepared, were used for photocatalytic water splitting application (Table 1).

Characterization techniques

Powder X-ray diffraction patterns of the Ni@TiO₂ nanopowders (XRD, Rigaku MiniFlex 600) were recorded using a Cu-Ka radiation source. X-ray photoelectron spectroscopy (XPS) was conducted using PHI Versa Probe XPS. ICP-OES measurements were done using Spectroblue, Amtek. For this study, 200 mg of sample was dissolved in 4 ml con. $HNO₃$ and 1 ml of 30% Hydrogen Peroxide and digested in microwave digestor, diluted up to 50 ml and filtered solution was run through the calibration curve. Raman spectra were measured on a Renishaw InVia Raman microscope equipped with a 532 nm laser. Diffuse reflectance UV-visible absorbance spectra (UV-DRS) were obtained at room temperature in the range of 200–800 nm using

Fig. 2 Ni decoration of $TiO₂$ surface activation followed by electroless deposition method.

the Shimadzu UV-vis-NIR spectrophotometer (Model UV-3600). The photoluminescence emission spectra of prepared powders were carried out by using Shimadzu (RF5301PC) spectrofluorometer at an excitation wavelength 350 nm.

Fourier transform infrared (FTIR) spectroscopy measurements were carried out with a Jasco FTIR-6100 spectrometer over a range of 400-4000 cm⁻¹. Field emission scanning electron microscopy (FESEM, Hitachi, Japan with Model S-4800) and field emission transmission electron microscopy (FETEM by JEOL; JEM-2200FS) techniques were used to perform the morphological and microstructural analysis. For typical FETEM analysis, the test sample was carefully prepared by dispersing the sample powder in de-ionized water and drop of the dispersion was then transferred to carbon coated grid. In order to acquire high imaging and spatial resolution with chemical mapping of the material, out Scanning Transmission Electron Microscopy (STEM) in bright field (BF) mode equipped with energy dispersive X-ray spectroscopy (EDS) coupled elemental mapping by using JEOL, JEM2200FS equipment operated at 200 kV with spherical aberration corrector was carried out. The porosity and surface area of the samples were examined using Brunauer–Emmett–Teller analysis (BET: Quantachrome NOVA touch LX1).

Photoelectrochemical study

Incident photon-to-current conversion efficiency (IPCE) for the prepared pristine TiO₂, 0.1% Ni@TiO₂ and 0.5% Ni@TiO₂ nanopowders were measured in the wavelength region of 350– 600 nm (PV Measurement Inc., USA) where a standard tungstenhalogen lamp as monochromatic light source with a broadband bias light for ∼1 sun light intensity was used. The measurements related to electrochemical impedance spectroscopy (EIS) were undertaken on a CH 660C electrochemical analyzer (CH Instruments, Shanghai, China) in a two-electrode configuration where the photoanode comprising the prepared nanopowders and a platinum wire were used as working and counter electrodes respectively. The transport properties of the electrons were measured using electrochemical impedance spectroscopy (EIS) in the frequency range of 10^{-2} to 10^5 Hz at an alternative signal of 10 mV. The study on transient current-time profile for the prepared pristine TiO₂, 0.1% Ni@TiO₂ and 0.5% Ni@TiO₂ nanopowders was carried out using a Kiethley Source Measure Unit (Model 2400) under the illumination of a 1000 W xenon lamp (Oriel 91193). The homogeneity of light intensity over an 8 \times 8 in² area was confirmed by calibrating it under 1 sun light intensity (AM 1.5G, 100 mW cm^{-2}) with a Si solar cell.

Photocatalytic study

Photocatalytic hydrogen generation study of prepared $Ni@TiO₂$ nanopowders was performed using water splitting technique by following previously reported procedure.⁶⁰ For this experiment, in-house designed 100 ml capacity quartz glass reactor was employed which was sealed with the septum for collecting the evolved H_2 with the help of a gas tight syringe. This water splitting mediated evolved hydrogen can then be applied to gas chromatogram (Shimadzu: Model GC 2014) for quantitatively

analysing its production. The main component of this photocatalytic water splitting set up is mercury vapor lamp (400 W) as the light source fitted with continuous water circulation condenser. Such condenser absorbed the IR radiation and minimized the heating effect. For studying the photocatalytic activity of pristine TiO₂ and as synthesized Ni@TiO₂ nanopowders, 20 mg of sample was mixed with 25 ml of 20% methanol–water mixture in the quartz reactor and was kept under continuous stirring. Conventionally, Pt is added as the co-catalyst in a photocatalytic water splitting process. However, in the present case, Ni itself worked as co-catalyst and hence Pt was not added. The dissolved gases, if any, were removed by Ar gas purging (for 2 min). The reactor was kept in photocatalytic reactor set up under mercury vapor lamp irradiation and the evolved/produced hydrogen gas was collected at an interval of 60 min by means of a 5 ml capacity gas fitted syringe and finally hydrogen production was enumerated with the help of fitted standard curve data.

Result and discussion

Structural study by XRD

X-ray diffraction patterns of TiO₂ and Ni@TiO₂ powders are presented in Fig. 3. The diffractograms of all the samples exhibited tetragonal crystal structure with coexistence of anatase and rutile polymorphic phases (JCPDS card no. 02-22- 1272 and 02-22-1276). The ratio of anatase to rutile phase calculated by using Spurr's formula was found to be $81:19.^{61}$ The diffraction peaks present at 2θ values of 25.3, 36.7, 37.8, 38.7, 48.09, 53.74, 55.05, 62.19 and 75.15° correspond to (101), (103), (004), (112), (200), (105), (211), (213) and (215) planes of $TiO₂$ anatase phase respectively. The diffraction peaks observed at 2θ values of 27.39, 36.05, 41.35, 54.18, 56.78, 62.69 and 68.96° are ascribable to (110), (101), (111), (211), (220), (002) and (301) planes of $TiO₂$ rutile phase respectively.

No additional peaks belonging to metallic nickel was observed in $Ni@TiO₂$ samples. This may be because of low concentration of nickel as well as electroless deposition which generally gives amorphous coating. Most of the researchers did not get Ni phase in their X-ray diffraction study of $Ni@TiO₂$ and hence validated by XPS.^{40,43,62,63}

X-ray photoelectron spectroscopy study

The confirmation of elemental composition as well as chemical state of 0.1% Ni loaded $TiO₂$ nanopowder was carried out using X-ray photoelectron spectroscopy (XPS). The wide scan survey spectrum validated the presence of Ti, O, and Ni that are the three main elements present in the prepared samples (Fig. 4a). The peaks attributable to Ti $2p_{3/2}$ and Ti $2p_{1/2}$ were found at 458.8 eV and 459.0 eV, and at 464.5 eV and 464.7 eV, respectively (Fig. 4b).^{41,50} The binding energy difference between Ti $2p_{1/2}$ and Ti $2p_{3/2}$ was 5.68 eV, attributable to characteristic Ti–O bond of $TiO₂$.⁵⁰ The O 1s XPS spectra of 0.1% Ni loaded TiO₂ nanopowders was shown in Fig. 4c which revealed their typical peak around 530.1 eV which was assigned to the lattice oxygen of the Ti–O species. Another peak at 532.5 eV related to the hydroxyl groups adsorbed on the sites of surface oxygen vacancies of the Ni@TiO₂ was noticed.

The XPS signal of Ni 2p peak at the binding energy of 851.83 eV (Fig. 4d) is associated with metallic Ni loaded on $TiO₂$

Fig. 3 X-ray diffraction patterns of pristine $TiO₂$ and Ni@TiO₂ powders.

Fig. 4 XPS spectra of 0.1% $Ni@TiO₂$ powder corresponding to (a) survey scan, and deconvoluted high resolution scans corresponding to elements (b) Ti 2p, (c) O 1s and (d) Ni 2p.

surface which is consistent with other available reports.^{40,41} The Ni 2p XPS spectrum exhibited additional peaks at 854.76 and 872.62 eV (Fig. 4d) ascribable to the Ni²⁺ (2p_{3/2} and 2p_{1/2}) of $Ni(OH)₂/NiO$, their respective satellite peaks at 861.54 and 879.63 eV were observed.^{50,64} Thus, coexistence of metallic Ni and $Ni(OH)₂/NiO$, is observed in XPS spectra. This coexistence of metallic Ni and $Ni(OH)_2/NiO$ is mutually beneficial. The defects in TiO₂ create an electron donor *via* captivation of Ni²⁺, generating the Ni-O bonds, which closely fix metallic Ni nanoparticles on the TiO₂ surface.⁵⁰

ICP-OES study

ICP-OES studies of $Ni@TiO₂$ nanopowders are presented in Table 2 which depicts that the observed wt% loading of Ni is consistent with our theoretical calculations carried out while performing the Ni loading experiments.

Raman study

According to the factor group analysis, the six different modes of anatase TiO₂, namely $1\text{A}_{1\text{g}}, 2\text{B}_{1\text{g}}$ and 3E_{g} are the Raman active modes having the following representation $(1A_{1g} + 2B_{1g} + 3E_g)$

	Table 3 Raman vibrational modes of anatase and rutile phases of $TiO2$		
nanoparticles			

and their reported vibrational peak positions are given in Table 3.⁶⁵ Raman spectra of the Ni@TiO₂ are shown in Fig. 5. Raman vibrational modes occurred at frequencies of 142 (E_g), 197 (E_g), 395 (B_{1g}), 516 (A_{1g} + B_{1g}), 637 (E_g) and 793 (first overtone of B_{1g}) cm⁻¹ corroborated the presence of anatase crystalline phase (Fig. 5a). This E_g peak was assigned to the symmetric bending vibration of O–Ti–O in TiO₂ frame. In case of Ni doping, it is reported that vibrational mode at 141 cm⁻¹ of

Fig. 5 Raman spectra of pristine $TiO₂$ and $Ni@TiO₂$ powders for (a) major peaks depiction (inset: enlarged image of peak at 142 cm⁻¹ due to E_q mode) and (b) major peaks in 300 to 850 cm⁻¹ range (inset: deconvoluted peaks).

Ni@TiO₂ samples showed an appreciable shift toward higher frequencies (from 143.2 to 145.0 $\rm cm^{-1})$ by increasing the nickel content in the photocatalyst. However, in our samples nickel loading/decoration rather than doping is carried out, hence, noticeable shift is not observed (inset of Fig. 5a).⁶⁵

Gradual decrease in the intensity of the peaks is noticed with increase in Ni loading concentration. Such quenching may be attributable to the scattering of incident radiation by the Ni on surface of TiO₂ nanoparticles. Ni does not show any peaks in the Raman spectrum as Ni, a face-centred cubic (FCC) metal, has a single atom primitive unit cell and does not show any polarizability change due to the interaction between the electric field of the monochromatic light and dipole moment within the material.⁶⁶

Presence of rutile phase in the sample is confirmed by observed vibrational modes assigned at 142 (B_{1g}), 447 (E_g) and 613 (A_{1g}) cm⁻¹ (Fig. 5b). The shape of the band centred at 637 cm−¹ indicates that there could be a shoulder to this band at the lower frequency side which is further confirmed by peak deconvolution (inset of Fig. 5b) using Lorentzian curve fitting.^{63,67-69} The Raman study clearly reveals the presence of anatase and rutile $TiO₂$ phases.

UV DRS and photoluminescence study

Investigation of the band gap energies of pristine $TiO₂$ and Ni@TiO₂ nanopowders was executed using UV-DRS technique

Fig. 6 (a) UV-DRS spectra (inset: Tauc plot) and (b) PL spectra of pristine TiO₂ and Ni@TiO₂ nanopowders.

and the results are shown in Fig. 6a. When the surface of $TiO₂$ nanopowder is decorated with Ni using electroless coating method, the absorption edges of Ni@TiO₂ samples shift towards longer wavelength and a bathochromic shift (red shift) was observed which is also reported by other investigators.⁷⁰⁻⁷² Red shifted absorption edge was perceived as an effect of interconnectivity between the d electrons of $Ni²⁺$ and the electrons availed from O^{2-} and Ti⁴⁺.

Remarkably, for the pristine $TiO₂$ nanopowder the edge absorbance value was observed to be at 380 nm, which increased significantly to 394, 392 nm for 0.05 and 0.1% Ni@TiO₂ nanopowders and then decreased slightly to 387, 382, 382 and 382 nm, for 0.2, 0.3, 0.4 and 0.5% $Ni@TiO₂$ nanopowders, respectively. Such slight decrease in the absorbance edge at higher Ni loading concentrations may be attributable to excessive interaction d-electrons of $Ni²⁺$ and the sp–d exchanges taking place among the electrons in the conduction band of $TiO₂$.⁷³ Nevertheless, the band gap energy of pristine $TiO₂$ and 0.5, 0.1, 0.2, 0.3, 0.4 and 0.5% $Ni@TiO₂$ nanopowders was computed to be 3.26, 3.16, 3.14, 3.20, 3.23, 3.23 and 3.23 eV, respectively by implementing Tauc plot which revealed slight red shift (inset of Fig. 5a). Paper

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Photoelectronic features of pristine TiO₂ and Ni@TiO₂ was reported using photoluminescence spectra and the range used is 400–800 nm. Wavelength of 350 nm sweetens well to cause the excitation of electrons suitable to understand the recombination of holes and electrons. Numerous concurrent processes occur leading to recombination/reunification of electrons and holes after their excitation by photon which are dominated by electronic stoichiometry and crystal structure.⁷³ In Fig. 6b, the first band observed is at 391 nm (UV region) followed by a low intensity band around 470 nm (visible blue-green emission region) in all the samples. The first band is attributable to the band gap transition or the band edge emission of the host TiO₂. The emission band at 470 nm can be ascribable to self-trapped excitons, oxygen vacancies, surface defects, etc. The blue-green emission band at 470 nm originated from the charge transfer from Ti³⁺ to oxygen anion in $[{\rm TiO}_6]^{8-}$ complex hinting at their intrinsic state nature as against the surface state.⁷²⁻⁷⁴

 $Ni²⁺$ loading concentration on the surface of TiO₂ nanopowder also plays a vital role as quencher of PL peak intensity.⁷⁵ When the surface $TiO₂$ nanopowder is decorated with nickel, it creates an epicentre which is non-radiative in nature and prevents aggressive recombination of e^-/h^+ pair. This phenomenon was also observed in the PL spectra of the present work (Fig. 6b) where decrease in the PL peak intensity was noted with increase of Ni loading percentage responsible for an effective charge separation and an increase of carrier mobility as compared to the pristine $TiO₂$ sample.^{71,75,76} In case of samples prepared with Ni loading of 0.1 and 0.2 wt%, through the intensity of peak due to the band gap transition is decreased little bit as compared to pristine $TiO₂$, the intensity of peak at 470 nm due to self-trapped excitons, oxygen vacancies, surface defects, etc. decreased to a more extent, which may indicate interaction d electrons of Ni^{2+} and the sp–d exchanges dominate the electron–hole decay process taking place among the electrons in the conduction band of $TiO₂$.

FTIR study

FTIR analysis was executed to survey the characteristic vibration bands of $Ni@TiO₂$ catalysts and their spectra in the wavenumber range 400–4000 cm−¹ are presented in Fig. 7. In the range of 400–800 cm−¹ , characteristic vibration bands belonging to metal-oxygen bonds were reported.⁷⁰

From the spectra, absorption band observed in the range of 1000–1700 cm−¹ is designated to water molecules adsorbed on the surface of nanostructure (H–O–H bending: 1639 $\rm cm^{-1})$ and Ti–O–Ti stretching vibration peak at 1116 $\rm cm^{-1}.$

FESEM analysis

The nanoscale morphology and shapes of pristine $TiO₂$ and 0.05–0.5% $Ni@TiO₂$ powders were examined by field emission scanning electron microscopy (Fig. 8). FESEM images of pristine $TiO₂$ nanopowder show nanoparticles having spherical morphology with particle size in the range of 20–30 nm (Fig. 8a and b). After electroless Ni loading on $TiO₂$ nanoparticles, slight increase in the particle size i.e. 35 nm was noted (Fig. 8c–l and ESI Fig. S2†).50,72

Microstructural study by FETEM

FETEM imaging

FETEM images for 0.1% Ni loaded $TiO₂$ nanopowders are shown in Fig. 9. FETEM images at low and intermediate magnification (Fig. 9a and b) display hexagonal and cubical faceted morphologies having sharp-edged geometries. The particles are found to have an average size in the range of 20– 30 nm. Loading of $TiO₂$ sample by Ni was found to occur by two ways as seen from the high magnification images (Fig. 9c-e). Ni nanofilm were found to get intermittently deposited over $TiO₂$ nanoparticles surface (shown by the yellow colour bordered areas in Fig. 9c and d). The width of this film was found to be in the range of 1–2 nm. Secondly, Ni nanoclusters of size around

Fig. 7 FTIR spectra of pristine $TiO₂$ and $Ni@TiO₂$ nanopowders (inset: enlarged region in 1000–4000 cm⁻¹ range).

2 nm were also observed over the surface of $TiO₂$ nanoparticles (as indicated by the dotted circle in Fig. 9e). The analysis of lattice fringes in Fig. 9b–e reveal presence of three types of materials. Majority of the particles displayed local single crystalline nature with each particle having a different lattice orientation. For $TiO₂$ nanoparticles, fringes with lattice spacing of 3.50, 2.29 and 1.91 Å belonging to (101), (004), (200) planes, respectively, corresponding to the anatase phase and 3.35 and 1.70 Å with planes for (110) and (211) corroborating the presence of rutile phase were noticed.⁷⁷ The presence of nickel was ascertained by lattice spacing fringes with d-value of 1.98 Å with (111) plane.⁷⁸ The nanocrystalline nature of the samples through selected area electron diffraction (SAED) patterns (Fig. 7f) was evinced by presence of ring like patterns that are not continuous and display regular bright spots.⁷⁷ Confirmation of such morphological features due to Ni loading was carried out by obtaining more such FETEM images as displayed in ESI Fig. S4.† The SAED patterns further substantiated presence of anatase and rutile phases $TiO₂$ when d-values from the corresponding rings were calculated and matched with XRD database. The presence of (101), (103), (004), (200), (105) and (110), (211) planes for anatase and rutile phases of $TiO₂$ respectively and (111) plane for Ni were observed. For sake of comparison, FETEM images of 0.5% Ni@TiO₂ nanopowder sample are shown in ESI Fig. S5[†] which also revealed the loading of TiO₂ nanopowders by Ni. **EXC** Articles. Published on the published on 12/18/2024 3:04:30 PM. This article is the common of the properties are not the interest are

FETEM-STEM-EDS analysis of 0.1 wt% Ni@TiO2 photocatalyst

The semi-quantitative qualification of elements present in the 0.1 wt% $Ni@TiO₂$ sample was ascertained using the energy dispersive X-ray spectroscopy (EDS) analysis in scanning transmission electron microscopy (STEM) mode. Ni was loaded over $TiO₂$ nanoparticles in weight-by-weight manner. The pertinent FETEM-EDS spectrum for 0.1 wt% $Ni@TiO₂$ sample with tabulated quantitative data is presented in Fig. 10a and its inset. The EDS spectrum of 0.1 Ni@TiO₂ catalyst revealed the presence of Ti, O and Ni and disclosed the presence of 65.24 wt% of titanium, 34.68 wt% of oxygen and 0.08 wt% of Ni.

This weight percentage of Ni is close to the calculated value and it is obvious that the XRD pattern did not illustrate any Ni peaks. Thus, EDS and XPS analysis data agreed well with each other (Fig. 4). EDS-elemental mapping images of 0.1 Ni/TiO₂ sample in STEM mode are shown in Fig. 10. The electron image (given in inset of Fig. 10a) and elemental mapping images ascribable to Ti and O (Fig. 10c and d, respectively) correlated well and overlapped with elemental mapping images for Ni also (Fig. 10e) and confirmed the surface loading of Ni over $TiO₂$ nanoparticles. Higher intensity of the colours allocated to Ti and O as compared to Ni was noticed which was tangible as very low amount of nickel was used for the surface loading of $TiO₂$. FETEM images were in tune with FESEM images.

BET surface area analysis

Fig. 11 displayed the N_2 adsorption–desorption isotherms of pristine $TiO₂$ and Ni-TiO₂ composites with varying doses of nickel loading from 0.05 to 0.5 wt%. Both pristine $TiO₂$ and

Fig. 8 FESEM images of (a and b) pristine TiO₂ and Ni@TiO₂ nanopowders with respective Ni wt% loading of (c and d) 0.05, (e and f) 0.1, (g and h) 0.2, (i and j) 0.3 at (k and l) 0.4 at low and high magnification respectively.

Ni@TiO₂ composites revealed type IV nitrogen adsorptiondesorption isotherms possessing type H_3 hysteresis loops. Table 4 summarized the specific surface area, total pore volume and BJH

pore sizes of the as-prepared $TiO₂$ and $Ni@TiO₂$ photocatalyst samples. Till 0.1 wt% Ni loading, the surface area of $Ni@TiO₂$ nanopowders was found to increase to 61.54 m² g⁻¹ from 45.72

Fig. 9 FETEM images of 0.1% Ni@TiO₂ powder at (a) low magnification and their HRTEM images (b-f) the corresponding SAED pattern.

Fig. 10 FETEM-STEM-EDS elemental mapping images of 0.1% Ni@TiO₂ nanopowders: (a) EDS spectra (inset: electron image of selected region and its quantitative EDS data) and its elemental mapping images for (b) elemental mix, (c) Ti, (d) O and (e) Ni.

Fig. 11 BET surface area graphs of pristine $TiO₂$ and Ni@TiO₂ nanopowders (inset: plot of pore diameter vs. pore volume).

 m^2 g⁻¹ for pristine TiO₂. Further increasing the Ni impregnation on TiO₂ surface above 0.1%, gradual reduction in the BET surface area was observed. It is postulated that when Ni loading amount is small (0.05 and 0.1 wt%), occupation of nucleation sites by $Ni²⁺$ species on $TiO₂$ surface leads to generation of Ni dots. As per TEM images the Ni coating is observed on the particles which might be

amorphous after aggregation of Ni dots. At higher concentration electroless nickel plating is intense which might be blocking the surface of $TiO₂$ which is responsible for lowering the surface area.60,79

Photoelectrochemical study

IPCE study. The IPCE results of pristine and Ni@TiO₂ nanoparticles are shown in Fig. 12a. TiO₂ is known for its strong absorption of ultraviolet (UV) light, primarily in the range of 300 to 400 nanometres (nm). Consequently, the IPCE spectrum of pristine $TiO₂$ nanopowder showed high efficiency (with a maximum of 14% at 350 nm) in the UV range, indicating its excellent photoconversion capability in this region. The loading of Ni into $TiO₂$ can enhance its photocatalytic performance by modifying the band structure, improving charge carrier separation and transport, and introducing additional surface states or reaction sites. These effects can lead to an increase in the IPCE, however, 0.1% Ni@TiO₂ nanopowder exhibited better (27.21% at 350 nm) IPCE than 0.5% $Ni@TiO₂$ nanopowder (21.14% at 350 nm). The IPCE spectrum hints about the ability of a semiconductor in converting incident photons into electrical current across a range of wavelengths and improves the understanding of their photocatalytic or photoelectrochemical properties.⁸⁰ Herein, the IPCE results indicated that 0.1% Ni@TiO₂ nanopowder has better charge carrier separation and transport and IPCE was extended to the visible region of the solar spectrum. Published on 2023. The main and the state of the published on 12/18/2024 3:04:30 PM. This article is licensed under a state of published planet is licensed under a state of the state of published under a state of the stat

Transient current-time profile. The transient current-time curves of the pristine TiO₂, 0.1% Ni/TiO₂ and 0.5% Ni/TiO₂ nanopowders are given in Fig. 12b for providing the valuable information about their electrochemical performance. The photocurrent densities of pristine TiO₂, 0.1% Ni@TiO₂ and 0.5% Ni@TiO₂ nanopowders were 15.22, 25.43 and 31.20 μ A cm^{-2} respectively. 0.1% Ni loaded TiO₂ nanopowder showed better photocurrent density than that of pristine and 0.5% Ni loaded TiO₂ nanopowders. The presence of nano Ni loading on TiO2 surfaces introduces additional electrochemical reaction pathways or active sites, leading to an increase in the overall current response in terms of transient current density. In case of 0.1% Ni/TiO₂ nanopowder, the distribution of Ni intermittent films and nanoclusters on the $TiO₂$ surface was relatively sparse, allowing for better access to the active sites by the scavenger

Table 4 BET surface area, total pore volume, BJH pore size, H₂ generation and apparent quantum efficiency (AQE) for pristine TiO₂ and Ni@TiO₂ nanopowders

Samples	BET surface area $(m^2 g^{-1})$	Total pore volume $\rm (cm^3 \ g^{-1})$	BJH pore size (nm)	H_2 generation (µmol g^{-1} h ⁻¹)	AQE^a (%)
Pristine TiO ₂	45.7	0.47359	3.37834	470	3.04
0.05% Ni@TiO ₂	58.3	0.38659	3.36268	987	6.40
0.1% Ni@TiO ₂	61.5	0.2194	3.35446	1600	10.38
0.2% Ni@TiO ₂	52.6	0.39147	3.36362	1141	7.40
0.3% Ni@TiO ₂	49.2	0.42826	3.36568	1106	7.175
0.4% Ni@TiO ₂	47.8	0.43754	3.36374	1103	7.156
0.5% Ni@TiO ₂	46.6	0.45623	3.36992	997	6.468

 a AQE calculated for per gram of sample.

Fig. 12 Photoelectrochemical study of Ni@TiO₂ nanopowders. (a) IPCE vs. wavelength plot, (b) transient current–time profile and (c) Nyquist plot.

solution leading to more efficient charge transfer processes and a good photocurrent response in the transient current–time curve.⁴³ The observed PEC and IPCE results augmented well with the observed PL spectroscopy results.

EIS study. The Nyquist plots of pristine TiO₂, 0.1% Ni@TiO₂ and 0.5% Ni@TiO₂ nanopowders are shown in Fig. 13c. The Nyquist plots generally exhibit a semicircular arc representing the charge transfer resistance at the electrode/electrolyte interface and the efficacy photogenerated electron–hole pairs separation that is associated with the electrochemical reaction

Fig. 13 (a) H₂ generation performance of pristine TiO₂ and Ni@TiO₂ nanopowders and (b) recyclability study for 0.1% Ni@TiO₂ nanopowder.

occurring on the $TiO₂$ surface and a linear plot associated with the diffusion region.^{43,80} Ni loading over TiO₂ surface was found to decrease the charge transfer resistance $(R_{\rm ct})$ in comparison to pristine $TiO₂$ nanopowders as indicated by a smaller semicircular arc of the Nyquist plot in Fig. 12C. The lower charge transfer resistance indicated improved charge transfer kinetics and enhanced electrochemical activity in Ni-loaded TiO₂. 0.1% $Ni@TiO₂$ nanopowder showed the lowest charge transfer resistance due to favourable electronic interactions at the Ni– $TiO₂$ interface, optimal active sites, and band structure modification. These factors contributed to efficient charge transfer and improved electrochemical performance. In nutshell, the photo-electrochemical and allied studies revealed that 0.1% Ni@TiO₂ nanopowders exhibited the least charge transfer resistance and optimal photocurrent density as well as IPCE values revealing that it is equipped with a good charge transfer interface and larger number of active sites.

Photocatalytic water splitting. The hydrogen generation ability of all in situ Ni decorated $TiO₂$ nanopowders based heterogeneous photocatalytic system was revealed to be at least more than double than that of pristine $TiO₂$ nanopowder (Table 4). It was observed that up to 0.1 wt% Ni loading, increase in the hydrogen generation rate was observed which was further reduced upon higher Ni loading. The maximum hydrogen generation rate of ~1600 µmol g^{-1} h⁻¹ was achieved for 0.1% Ni@TiO₂, which is \sim 3.5 times higher than that of pristine TiO₂ nanopowder (470 µmol g^{-1} h^{−1}) (Fig. 13a). The optimized AQE was observed to be 10.38% at 450 nm incident light wavelength for 0.1% $Ni@TiO₂$ photocatalyst, attributable to efficient Ni co-catalysts on the surface activated TiO₂ nanoparticle facets leading to smoother charge separation (Table 4).⁸¹ The formulae used in the calculation of AQE are given in ESI, Section S1.†

In electroless coating method, semiconductor surface was first cleaned, activated and then metallic loading was undertaken which might be helpful for uniform decoration of Ni on TiO₂. It may be noted that Ni nanoparticles generated on TiO₂ are not just adhered to the surface by physical forces, but chemically through sp–d exchanges. Additionally, presence of nickel obliterated necessity of Pt, a commonly used co-catalyst in water splitting reactions which makes the process economical. Recyclability study of 0.1% $Ni@TiO₂$ photocatalyst was also carried out and the results are shown in Fig. 13b. The decrease in the photocatalytic activity from 6391 µmol g^{-1} to 5795, 4878, 4748 and 4728 µmol g^{-1} in the 2nd, 3rd, 4th and 5th cycles correspond to loss of 90.67%, 76.32%, 74.29% and 73.97% respectively than the initial photocatalytic activity. However, these losses are attributed to the material loss during the processing and washing of the catalysts after first cycle. The 20, 18 mg of catalysts are used for $1st$, $2nd$ respectively and for $3rd$ cycle onwards 16 mg of catalyst was used. It showed almost similar hydrogen evolution for three cycles, ultimately revealing the good stability of the photocatalyst. FETEM images of recycled 0.1% $Ni/TiO₂$ nanopowder photocatalyst are given in ESI Fig. S6† which also revealed the stability of the photocatalyst by the presence of Ni over $TiO₂$ nanoparticles. RSC Advances α -1500 μ med ¹ ¹ was achieved for 0.1% Right(2), which is produced view High Place Paper Macched Creative CD₁ and the machine of the positive CD₁ published on 12/18/2024 2024 3:04:30 PM. This art

Mechanism of photocatalytic H_2 generation of prepared $Ni@TiO₂$ nanopowders. Photocatalysis process can be divided onto homogeneous and heterogeneous based on the use of only catalyst and additional co-catalyst respectively. The schematic representing the mechanism of photocatalytic H_2 generation in heterogeneous scheme was presented in Fig. 14. In the watersplitting reaction, the photogenerated electrons and holes are

Fig. 14 Mechanism of photocatalytic H_2 generation of prepared Ni@TiO₂ nanopowders.

produced when $Ni@TiO₂$ nanopowders were irradiated by light.^{9,11} During this process, the migration of photoexcited electrons and holes to the surface of $TiO₂$ occurs, that are responsible for H^+ reduction. For H_2 evolution, the more negative CB potential of $TiO₂$ than the reduction potential of water leads to generation of a swift pathway for photogenerated species for migration towards reaction sites present on its surface, which in turn, reduces the probability of recombination. The further enhancement of photocatalytic efficacy for Ni@TiO₂ nanopowders than pristine TiO₂ nanopowder might be due to the effective electron accepter property of Ni and the Schottky barrier formation between these two materials.

In photocatalytic H_2 generation with methanol as sacrificial agent, photogenerated electrons are used to combine with protons in water to generate H_2 , while the methanol molecules get oxidized by capturing excess photogenerated holes through
a series of reactions finally yielding three hydrogen molecules as
given in eqn (1).
 $CH_3OH + H_2O \frac{h\nu / N i@TiO_2}{\sim} CO_2 + 3H_2$ (1) a series of reactions finally yielding three hydrogen molecules as given in eqn (1).

$$
CH3OH + H2O \xrightarrow{hv/Ni@TiO2} CO2 + 3H2
$$
 (1)

Ni loaded on $TiO₂$ as co-catalysts, absorbs photogenerated electrons to promote the separation and transfer of interface charge carriers, but also provide efficient proton reduction centres.⁸²

It may be noted that the CB level of TiO₂ is -0.50 V above the $H₂$ evolution potentials and that of Ni is just -0.23 V and this potential gradient helps in transformation of photoexcited electrons from CB of TiO₂ to Ni nanoparticles which subsequently react with H^+ ions generated by oxidation of water to form H_2 gas.

Beyond the optimized loading, the decreased in the photocatalytic activity was found. In this context, it may be noted the photocatalytic hydrogen generation results corroborate well with BET and PL spectroscopy results. BET surface area was found to increase for Ni decorated $TiO₂$ nanopowders than pristine TiO₂ up to 1 wt% Ni loading and then decreased gradually. Similarly, quenching in PL intensity peak and the peak due to presence of defects was noted (Fig. 5b). Both these phenomena were attributed to the density of non-radiative epicentres (causing the increase in Ni particle size as well as defect centres) which, in turn, are responsible for recombination of electron-hole pairs and ultimately reflects on the photocatalytic activity.

Conclusion

Nickel-decorated $TiO₂$ photocatalytic system proficient of responding to visible light was successfully synthesized by autocatalytic deposition (electroless) method. In this study, nickel loading wt% increased on the $TiO₂$ nanopowder surface as a function of Ni concentration in electroless bath. The existence of metallic Ni (in the form of nanocluster and intermittent films) that was uniformly infused onto the $TiO₂$ nanopowder surface was observed in the FETEM images which was further confirmed by EDS (and allied elemental mapping images) and

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XPS spectroscopy techniques. $Ni²⁺$ loading concentration on the surface of TiO₂ nanopowder also plays a vital role as quencher of PL peak intensity and as an effective charge separator. The further enhancement of photocatalytic efficacy for $Ni@TiO₂$ nanopowders than pristine $TiO₂$ nanopowder might be due to the effective electron accepter property of Ni and the Schottky barrier formation between these two materials. 0.1% Ni@TiO₂ nanopowder exhibited the optimized hydrogen generation rate of ~1600 μmol g⁻¹ h⁻¹ which is ~3.5 times higher as compared to pristine TiO $_2$ nanopowder (470 µmol $\rm g^{-1}$ $\rm h^{-1})$ and the photocatalyst exhibited fairly good repeatable behaviour. In nutshell, electroless-plating is an efficient technique to decorate $TiO₂$ nanopowder surface using nickel which can not only serve as an efficient and cost-effective heterogeneous photocatalyst but in other applications like gas sensors. **Paper**
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Author contributions

The experiments have been planned by B. B. K., S. B. R., S. K. K., S. J. and P. N. B. performed the experiment. The data were analysed by P. N. B., M. D. S. and S. S. A. All the authors participated in writing the manuscript under the lead of B. B. K.

Conflicts of interest

There are no conflicts to declare.

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