

Effect of Equimolar Sodium Borohydride-Ferric Chloride Concentrations on Nano Zero-Valent Iron/Palm Shell Composites for Simultaneous Nanogold Recovery and Hydrogen Generation

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Abstract

Gold-containing waste solutions represent both an environmental liability and a valuable secondary resource, yet few existing technologies integrate nanogold recovery with sustainable hydrogen generation from these streams. In this study, the effect of equimolar sodium borohydride–ferric chloride ($\text{NaBH}_4\text{--FeCl}_3$) concentrations on the synthesis and performance of nanoscale zero-valent iron (nZVI)/palm shell composites was systematically investigated for the simultaneous recovery of nanogold and generation of hydrogen from gold-containing aqueous solutions. The composites were synthesized at different equimolar $\text{NaBH}_4\text{--FeCl}_3$ concentrations (0.5–2.0 M), while maintaining a fixed overall molar ratio, with palm shell biomass employed as a support to suppress particle aggregation and preserve reactive surface area. Nanogold formation was evaluated using UV–Vis spectroscopy via localized surface plasmon resonance, while hydrogen evolution was quantified by a water-displacement method. Surface properties were characterized by BET analysis. Nanogold recovery increased progressively with increasing equimolar precursor concentration, whereas hydrogen production exhibited a non-linear dependence, reaching a maximum of 29.02 mL at 1.5 M, which also corresponded to the highest BET surface area (13.57 m²/g). Further increasing the equimolar $\text{NaBH}_4\text{--FeCl}_3$ concentration to 2.0 M led to surface passivation and diminished reactivity. These results demonstrate that equimolar precursor concentration plays a critical role in governing nZVI/palm shell composite structure and functionality. The optimized composite exhibits strong potential as a multifunctional material for integrated precious metal recovery and green hydrogen production, thereby contributing to sustainable circular resource utilization and clean energy technologies.

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Keywords: Nano zero-valent iron; Equimolar precursor concentrations; Nanogold recovery; Hydrogen generation; Palm shell composites

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1. Introduction

Nanogold, valued for its size-dependent electrical, catalytic, and surface properties, is widely used in electronics, catalysis, sensing, and biomedicine [1,2]. Although electronic waste and

industrial effluents contain abundant gold-bearing precursors [3], the recovery and controlled synthesis of nanogold from these complex matrices remain challenging due to inefficiencies in existing extraction and reduction approaches [4]. Conventional nanogold synthesis methods also often depend on strong chemical reductants, hazardous solvents, and energy-intensive processes, raising both economic and

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environmental concerns [5–9]. In parallel, hydrogen has emerged as a clean and efficient energy carrier critical to achieving global carbon neutrality [10]. Although hydrogen can be produced via routes such as steam methane reforming, electrolysis, and biomass conversion [10,11], most conventional techniques are energy-intensive and generate substantial greenhouse gas emissions [12]. These challenges have intensified the search for cost-effective and eco-friendly nanogold recovery and hydrogen production technologies.

Recently, nanoscale zero-valent iron (nZVI) has attracted increasing attention as a multifunctional material for both nanogold recovery and hydrogen generation from aqueous systems, owing to its high reactivity, large specific surface area, cost effectiveness, and ability to function simultaneously as a reductant and an adsorbent [13]. Its characteristic core–shell structure, featuring a metallic Fe⁰ core and an iron oxide shell, is widely proposed to support complementary reduction and sorption mechanisms. Specifically, the Fe⁰ core generally considered to facilitate electron-transfer reactions, while the oxide shell provides adsorption sites that enhance reactant accessibility and stabilize reaction intermediates [14]. In nanogold recovery, nZVI has been reported to be capable of simultaneously reducing and adsorbing Au³⁺ ions from chloroauric acid (HAuCl₄) solutions to form elemental nanogold (Au⁰), accompanied by oxidation of Fe⁰ to Fe³⁺, as described in Equation (1) [15,16]:

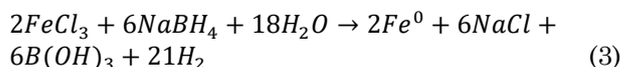


This coupled redox–sorption mechanism is commonly proposed as a simplified explanation for the efficient removal and immobilization of gold species on the nZVI surface without the use of toxic chemical reductants. Studies such as Li et al. [17] have reported high nanogold recovery efficiencies from industrial wastewater using nZVI, demonstrating its potential for sustainable resource reclamation. A similar redox–sorption interplay has also been proposed to contribute to hydrogen generation via water decomposition. In this process, the Fe⁰ core is suggested to donate electrons to water molecules, producing hydrogen gas (H₂), while the oxide shell adsorbs and facilitates the dissociation of water at the solid–liquid interface, enhancing reaction kinetics [18]. The overall reaction is expressed in Equation (2) [18]:



This pathway offers a conceptual route for hydrogen production without emitting greenhouse gases, unlike conventional thermochemical

methods [19]. Although nZVI has been independently employed for nanogold recovery (Equation (1) and hydrogen production (Equation (2)), the integration of both processes within a single system has not been reported yet. The nZVI is typically synthesized through chemical reduction of ferric salts (e.g., FeCl₃) using strong reductants such as sodium borohydride (NaBH₄), as shown in Equation (3) [20].



In this reaction, Fe⁰ represents the reactive metallic core of nZVI responsible for reductive transformations. Theoretically, a NaBH₄:FeCl₃ molar ratio of 3:1 is required for complete reduction of Fe³⁺ to Fe⁰. In practice, NaBH₄ is often used in excess to compensate for its rapid hydrolysis and other competing side reactions, thereby ensuring complete Fe³⁺ reduction [21]. However, excessively high NaBH₄:FeCl₃ ratios may induce uncontrolled hydrogen evolution, over-reduction, and the formation of boron-rich byproducts, which promote particle aggregation and surface passivation [21,22]. While the NaBH₄:FeCl₃ molar ratio governs the stoichiometric balance of the redox reaction, the absolute concentrations of both reagents play a critical role in influencing Fe³⁺ reduction kinetics, as well as nZVI nucleation, particle growth, and colloidal stability [23]. Our previous work identified a NaBH₄:FeCl₃ molar ratio of 4:1, slightly above the stoichiometric 3:1 requirement, as optimal for efficient and controlled nZVI formation [24]. Nevertheless, the influence of absolute reagent concentrations at this fixed optimal ratio remains unexplored. Importantly, varying reagent concentrations while maintaining equimolar NaBH₄ and FeCl₃ solutions and a constant overall 4:1 molar ratio enables isolation of concentration-dependent effects on Fe³⁺ reduction efficiency and nZVI yield, without interference from changes in molar ratio or total reagent volume. Such effects cannot be captured through molar-ratio adjustments alone, which primarily alter reaction stoichiometry rather than the physicochemical environment governing nZVI formation.

Accordingly, this study investigated the effect of varying the absolute concentrations of equimolar NaBH₄-FeCl₃ solutions, combined to maintain an overall NaBH₄:FeCl₃ molar ratio of 4:1, on nZVI yield and its performance in simultaneous nanogold recovery and hydrogen generation from gold-containing aqueous media. Palm shell, an abundant biomass in Malaysia [25], was employed as a support material to fabricate nZVI/palm shell composites, mitigating nZVI aggregation and preserving its reactive surface area. Nanogold recovery was quantified by

UV–Vis spectroscopy based on localized surface plasmon resonance (LSPR) intensity. Hydrogen production was measured via the water-displacement method, while the purity of the generated hydrogen gas was analysed using gas chromatography equipped with a flame ionisation detector (GC-FID). The surface properties and reactivity of the synthesized composites were further evaluated via Brunauer–Emmett–Teller (BET) analysis. Overall, this work provides fundamental insights into the role of reagent concentration in governing nZVI formation and offers enhanced control over nZVI composite reactivity for sustainable metal recovery and green hydrogen production.

2. Materials and Method

2.1 Materials

Iron(III) chloride hexahydrate ($\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$, $\geq 99\%$), ethanol ($\text{C}_2\text{H}_5\text{OH}$, 99.7%v/v), and poly(vinyl alcohol) (PVA, $\geq 99\%$, fully hydrolyzed, viscosity 11.6–15.4 cps) were purchased from R&M Chemicals. Sodium borohydride (NaBH_4 , $\geq 98\%$) was obtained from Bellamy Precision, while chloroauric acid ($\text{HAuCl}_4 \cdot 3\text{H}_2\text{O}$, $\geq 99.9\%$) was supplied by Sigma-Aldrich. Deionized water (resistivity 18.2 $\text{M}\Omega \cdot \text{cm}$) was used throughout all experiments. Palm shells were collected from a local palm oil mill, thoroughly washed with deionized water, dried at 80 °C for 24 h, and ground to a fine powder before use. All chemicals were used as received without further purification.

2.2. Fabrication of nZVI/Palm Shell Composites

The nZVI/palm shell composites were fabricated using a modified co-precipitation-reduction method under an inert nitrogen atmosphere, adapted from a previously reported method [26]. The raw palm shell biomass was washed, dried in an oven, ground, and sieved to a powder size of $< 65 \mu\text{m}$. For each experiment, equimolar solutions of FeCl_3 and NaBH_4 with final concentrations of 0.5, 1.0, 1.25, and 1.5 M were prepared by dissolving the corresponding masses of ferric chloride hexahydrate ($\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$) and sodium borohydride (NaBH_4) directly into their respective solvents, while maintaining consistent solution volumes and molar ratios across all conditions. The reagents ($\text{FeCl}_3 \cdot 6\text{H}_2\text{O}$) were premixed with a 4:1 ethanol-water mixture (10 mL ethanol + 2.5 mL deionized water). This solution was mixed with 0.01 g of the palm shell powder in a round-bottom flask and stirred magnetically for 30 minutes to ensure uniform adsorption of iron onto the biomass. Subsequently, 55 mL of the corresponding equimolar concentration NaBH_4 solution was added to the suspension dropwise (~ 20 drops/min)

to the FeCl_3 /palm shell suspension under continuous stirring at 700 rpm. The addition of NaBH_4 initiated an immediate reduction reaction, evidenced by the appearance of a black precipitate upon the first drop. The reaction was allowed to proceed for 1 hour at room temperature to ensure complete reduction. Following the reaction, the black iron nanoparticles were separated from the solution using the centrifuge at 6000 rpm for 7 minutes. Thereafter, the collected precipitate was washed three times with ethanol and dried for 4 hours in a freeze dryer for further use and characterization.

2.3. Preparation of Gold-containing Solutions

A gold-containing solution was prepared to evaluate the nanogold recovery and concurrent hydrogen production performance of the synthesized nZVI/palm shell composites. Specifically, a 0.8 mM HAuCl_4 solution was prepared by appropriately diluting the liquid HAuCl_4 stock in 45 mL of deionized water under gentle stirring at room temperature. Subsequently, 5 mL of 2% (w/w) polyvinyl alcohol (PVA) solution was added as a stabilizing agent and mixed thoroughly. The pH of the gold-containing solution was then adjusted to 2.0 using 1 M HCl or 1 M NaOH, added dropwise with continuous stirring to ensure homogeneity, and the pH was monitored using a calibrated pH meter. Once the desired pH was attained, the solution was used immediately in the nanogold recovery and hydrogen generation experiments with nZVI/palm shell composites.

2.4. Recovery of Nanogold and Hydrogen Gas Collection

The procedure for simultaneous nanogold recovery and hydrogen generation was conducted following modified protocols reported in the literature [27,28]. Figure 1 schematically depicts the experimental setup used. All reactions were performed in a 50 mL three-neck round-bottom flask immersed in a thermostatic water bath maintained at 37 °C. The flask was equipped with a thermometer and connected via rubber tubing to a pneumatic trough for hydrogen collection using the water displacement method, with an inverted 50 mL water-filled burette serving as the gas-measuring device. Before each experiment, the gold-containing solution in the flask was purged with high-purity nitrogen (99.99 %) for 20 minutes at 80 mL/min to remove dissolved oxygen and establish an inert atmosphere [27,29]. Nitrogen was introduced through a submerged inlet needle while gently stirring the solution to enhance deaeration. After purging, the nitrogen inlet was closed and the system was immediately sealed to form a gas-tight setup consisting of the reaction

flask, connecting tubing, and inverted burette. A slight positive nitrogen headspace was maintained to minimize oxygen ingress, while the water in the inverted burette served as a liquid seal to prevent back-diffusion of air. The system was allowed to stabilize until the burette water level remained constant, confirming the absence of leaks, and establishing a stable baseline. The reaction was then initiated by introducing 8 g/L of nZVI/palm shell composites synthesised using different equimolar NaBH_4 and FeCl_3 concentrations (0.5-2 M) and a fixed amount of palm shell powder, as described in Section 2.2. The suspension was stirred at 300 rpm and allowed to react for 24 hours, during which hydrogen evolution was continuously collected and analysed via gas chromatography (GC-FID, Agilent Technologies) with nitrogen gas as a carrier gas to ascertain the hydrogen concentration in the gas mixture. After 1 minute of reaction, aliquots were withdrawn, and the magnetic nZVI/palm shell composites were separated using a magnet. The resulting supernatant containing nanogold colloids was filtered, rinsed, and analysed by UV-Vis spectroscopy to detect the LSPR peak in the 530 nm - 550 nm range.

3. Results and Discussion

This study examined how varying equimolar NaBH_4 - FeCl_3 concentrations (0.5–2.0 M) influence the dual performance of nZVI/palm shell composites in simultaneous nanogold recovery and hydrogen generation. Although both processes started concurrently, nanogold

reduction dominated initially due to its higher redox potential [17], while hydrogen generation occurred gradually over 24 hours, as discussed in the following sections with supporting UV-Vis and BET analyses.

3.1. Effect on Nanogold Recovery

Figure 2 shows the UV-Vis spectra of nanogold generated using nZVI/palm shell composites synthesised at varying equimolar NaBH_4 - FeCl_3 concentrations (0.5–2.0 M). All spectra exhibit a well-defined localized surface plasmon resonance (LSPR) peak centered around

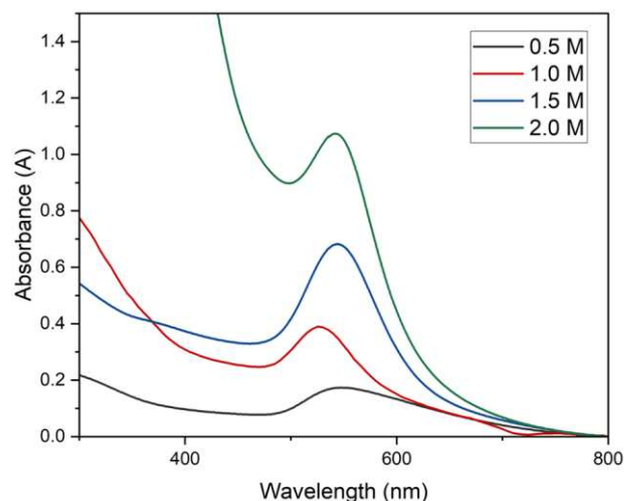


Figure 2. UV-Vis absorption spectra of nanogold recovered using nZVI/palm shell composites synthesized at varying equimolar NaBH_4 - FeCl_3 concentrations.

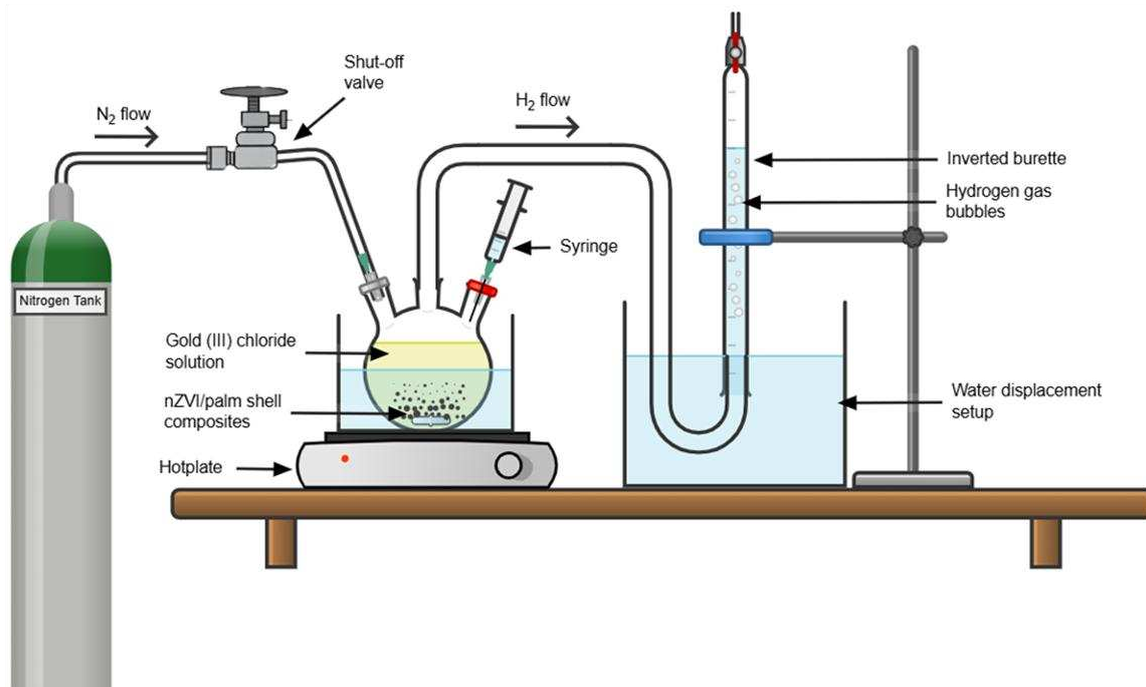


Figure 1. Experimental setup for simultaneous nanogold recovery and hydrogen generation using nZVI/palm shell composites.

530 nm, confirming the successful reduction of Au^{3+} to Au^0 by nZVI. The presence of a single, symmetrical LSPR peak indicates that the resulting AuNPs were predominantly spherical with a relatively narrow size distribution, consistent with typical spherical AuNPs absorbing in the 520–550 nm region [30,31]. The visible colour transition of the solution from light yellow to purple-red further corroborates AuNPs formation. Notably, the intensity of the LSPR peak increased systematically with precursor concentration, suggesting that the amount of nanogold produced is highly sensitive to the NaBH_4 – FeCl_3 molarity used in composite synthesis.

This concentration-dependent behaviour is further quantified in Figure 3, which plots the average absorbance (used as a proxy for nanogold yield) as a function of the equimolar NaBH_4 – FeCl_3 concentration. Absorbance values rose nearly linearly from 0.2227 A at 0.5 M to 1.0175 A at 2.0 M, demonstrating that higher precursor concentrations result in significantly greater nanogold recovery. This enhancement is likely associated with the increased availability of reactive Fe^0 sites generated in the composites at higher molarities, which may facilitate more rapid and extensive reduction of Au^{3+} to Au^0 . The favourable redox potential gradient between $\text{Fe}^0/\text{Fe}^{3+}$ (+0.33 V) and $\text{Au}^{3+}/\text{Au}^0$ (+1.50 V) further promotes efficient electron transfer [16,17], accelerating nanogold nucleation. As Fe^0 availability increases, a greater proportion of Au^{3+} ions undergo reduction, leading to higher LSPR intensities and greater nanogold formation. This trend is consistent with previous findings, such as those reported in a prior study [16], which also demonstrated improved Au^{3+} reduction efficiency with increasing iron precursor concentrations.

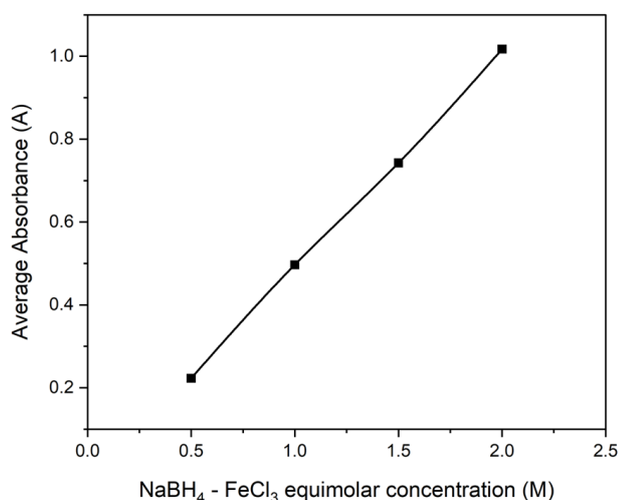


Figure 3. Average UV-Vis absorbance (proxy of nanogold yield) versus equimolar NaBH_4 – FeCl_3 concentration for nZVI/palm shell composite synthesis.

3.2. Effect on Hydrogen Volume

In this system, hydrogen generation occurs as nZVI within the nZVI/palm shell composites undergoes corrosion in aqueous environments, releasing electrons that reduce protons (H^+) or water molecules to produce hydrogen gas. The quantity and reactivity of the nZVI formed, both governed by the NaBH_4 and FeCl_3 concentrations used during composite synthesis, play a central role in determining the overall hydrogen output. As shown in Figure 4, the hydrogen volume exhibits a distinctly non-linear dependence on the equimolar NaBH_4 – FeCl_3 concentrations, contrasting with the more linear trend observed for nanogold recovery profile (Figure 3).

At lower concentrations (0.5–1.0 M), hydrogen volume remained relatively stable (19.08–19.88 mL). Hydrogen generation increased substantially at 1.5 M, reaching 29.02 mL, before dropping sharply to 12.20 mL at 2.0 M. These results contrast with the continuous upward trend observed for nanogold recovery (Figure 3). The divergence arises from the different kinetics of the two redox processes: Au^{3+} reduction is rapid and dominates the early reaction period, whereas corrosion-driven hydrogen evolution proceeds more slowly over 24 hours. At 1.5 M, sufficient Fe^0 is available not only to fully reduce Au^{3+} but also to sustain long-term hydrogen evolution, resulting in the highest hydrogen volume. However, beyond 2.0 M, excessive precursor amounts produce larger quantities of nZVI, which accelerates Au^{3+} reduction but simultaneously leads to the rapid consumption of reactive Fe^0 . Based on the observed reduction in hydrogen volume at higher concentrations, despite increasing nZVI availability, we infer that passivating layers of

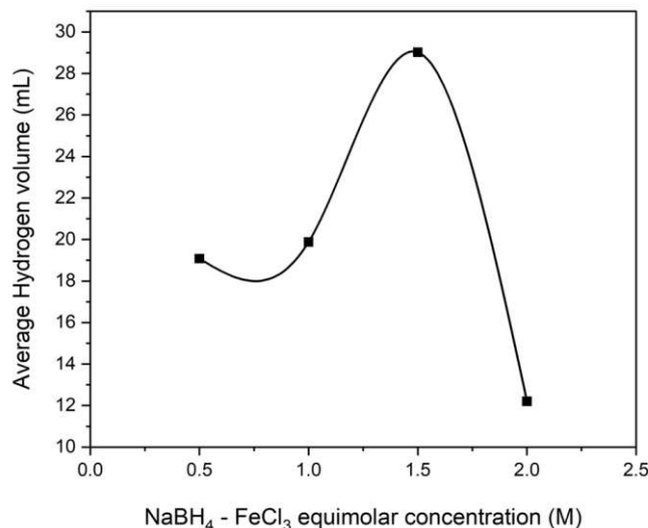


Figure 4. Average hydrogen volume versus equimolar NaBH_4 – FeCl_3 concentration for nZVI/palm shell composite synthesis.

iron oxides/hydroxides may build on the nanoparticle surfaces, limiting electron transfer and suppressing long-term hydrogen production. Although no direct surface characterisation of passivation layers was conducted out in this study, this interpretation is compatible with the observed kinetic behaviour and with mechanistic models reported in the literature. Similar behaviour has been reported in the literature [12], where reduced long-term hydrogen generation was observed at higher nZVI dosages. Comparable trends were also reported in other studies [32,33], which attributed that hydrogen evolution decreased exponentially over time, consistent with progressive surface passivation by oxide/hydroxide layers that block access to the underlying Fe⁰ and impede electron transfer, thereby reducing overall reactivity.

Although nanogold recovery continued to increase up to 2 M (Figure 3), hydrogen generation peaked at 1.5 M (Figure 4). This difference shows a fundamental trade-off in the system: increasing equimolar NaBH₄-FeCl₃ concentration speeds up the reduction of Au³⁺ ions, but it also slows down the long-term corrosion needed for hydrogen evolution. The reaction condition of 1.5 M therefore represents an optimal balance where both nanogold recovery and hydrogen production are jointly maximized, confirming the synergistic dual-function nature of the nZVI/palm shell composites.

3.3 Hydrogen Purity and Gas Composition

The evolved gas mixture was characterised by GC-FID to determine hydrogen purity and quantify the composition of gaseous products generated from the nZVI/palm shell system. Table 1 shows the concentrations of the gaseous species detected in the collected samples. As summarised in Table 1, hydrogen accounted for 99.88 mol% of the total detected gas, indicating that the nZVI/palm shell system predominantly produces hydrogen with negligible interference from other gaseous species. Only trace amounts of CO₂ (0.01 mol%) and CO (0.03 mol%) were detected, confirming minimal side reactions following correction for the blank response. This level of hydrogen purity is consistent with prior results from nZVI-based hydrogen production systems. For instance, hydrogen produced from a Bi/Fe⁰

Table 1. Concentration of gases collected from nZVI/palm shell composites.

Total gas concentration (%mole)	Gas concentration (%mole)		
	H ₂	CO ₂	CO
14.28	99.88	0.01	0.03

water system was reported to contain negligible levels of common gaseous impurities such as CO₂ and CH₄, highlighting its potential as a clean energy carrier [18]. Similarly, hydrogen formation arising from nZVI dissolution has been shown to correlate with minimal CO₂ generation under anaerobic conditions, indicating that nZVI itself can serve as a direct source of hydrogen in biomass-based systems [27]. Compared to previous findings, the present system has a high hydrogen purity, showing that the palm shell-supported nZVI promotes selective iron corrosion while inhibiting competing side reactions. This indicates that the nZVI/palm shell composites can produce high-purity hydrogen with little gaseous byproducts, highlighting their potential for clean and sustainable energy applications.

3.4. BET Surface Area and Pore Size Analysis of nZVI/Palm Shell Composites

The specific surface area and pore size distribution of the nZVI/palm shell composites were evaluated using BET analysis, with the results shown in Figure 5. The composites exhibited the highest BET surface area (13.57 m²/g) at an equimolar NaBH₄-FeCl₃ concentration of 1.5 M, while the largest pore size (217.77 Å) was observed at a concentration of 2.0 M. The increase in specific surface area from 8.69 m²/g (0.5 M) to 13.57 m²/g (1.5 M) is significant, as increased surface area enhances adsorptive interactions and redox activity of Fe⁰ nanoparticles. This is consistent with the improved nanogold recovery and hydrogen production observed at 1.5 M (Figures 3 and 4), implying that this concentration generated an optimum composite microstructure with a high density of active Fe⁰ sites.

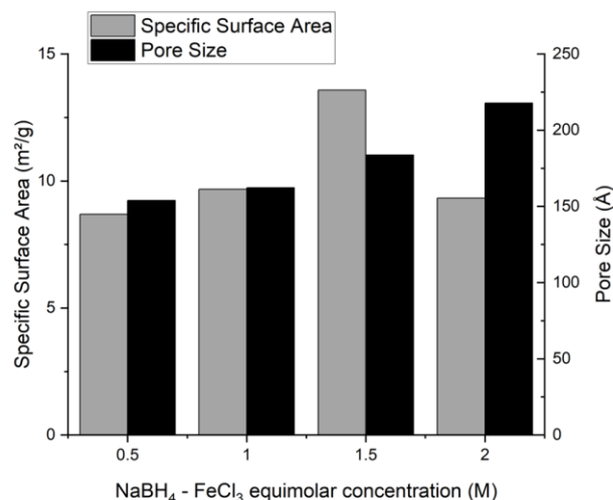


Figure 5. Specific surface area and pore size of nZVI/palm shell composites synthesized at varying equimolar NaBH₄-FeCl₃ concentrations.

At 2.0 M, however, although the pore size was the highest, the specific surface area decreased to 9.33 m²/g. This suggests that excessively high NaBH₄ and FeCl₃ concentrations may cause pore coalescence or partial structural collapse, weakening the internal pore network. Under these conditions, the accelerated nucleation and growth of nZVI can produce larger agglomerates that combine and block nearby pores within the composites [34]. Although no direct microscopic or spectroscopic evidence of pore collapse or surface passivation was determined in this study, the reduction in BET surface area, together with the decrease in hydrogen evolution at 2.0 M (Figure 4), provides indirect evidence that structural densification and surface deactivation may occur at higher precursor concentrations. These structural alterations may limit prolonged redox activity by reducing the accessibility of Fe⁰ surface sites and impeding effective electron transfer [35]. Similarly, the structural instability in nZVI composites at high precursor concentrations has been reported to result in poor textural properties and the formation of a passivating oxide layer that inhibits electron transfer and increased aggregation [34,36].

Overall, the BET results highlight those textural properties play a crucial role in determining Au³⁺ reduction and hydrogen generation efficiency. The nZVI/palm shell composites synthesized at 1.5 M exhibited a well-developed mesoporous structure that balances surface area for rapid nanogold recovery and structural stability for long-term hydrogen generation. These findings emphasize the importance of NaBH₄ and FeCl₃ concentration optimization to engineer surface characteristics that enhance multifunctional nanocomposite performance. Future research should utilize direct surface characterization techniques [36], and explore kinetics and thermodynamics of the sequential redox reactions between Fe⁰, Au³⁺, and water to identify rate-limiting steps and optimize overall performance, similar to other nanocomposite systems [37].

4. Conclusion

This study demonstrated the successful synthesis of palm shell-derived nZVI composites for simultaneous nanogold recovery and hydrogen generation from HAuCl₄ solutions. By varying equimolar NaBH₄-FeCl₃ concentrations, 1.5 M was identified as the optimal condition, producing composites with the highest BET surface area and a stable mesoporous structure. This balance of reactivity and structural integrity enabled rapid Au³⁺ reduction and sustained hydrogen evolution, with 1.5 M serving as the “sweet spot” that maximizes both nanogold recovery and hydrogen

production. Using palm shell biomass as the support further enhanced the economic and environmental sustainability of the material. Overall, the findings highlight the potential of nZVI/palm shell composites as a multifunctional platform for wastewater treatment, precious metal recovery, and hydrogen-based energy applications.

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CRedit Author Statement

Author Contributions: P. N. S. Nordin: Conceptualization, Methodology, Investigation, Resources, Writing, Review and Editing, Visualization; A. S. N. Helmy: Methodology, Investigation, Resources; C. J. C. Derek: Resources, Funding acquisition; M. F. Rajuli: Resources, Funding acquisition; S. H. Chang: Validation, Review and Editing, Supervision, Project administration. All authors have read and agreed to the published version of the manuscript.

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