Analytical study to determine the optical properties of gold nanoparticles in the visible solar spectrum

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ABSTRACT

In this work the optical properties of the formed gold nanoparticles, that obtained upon reducing the gold(I):6-thioguanosine hydrogel by dimethylamine borane (DMAB) have been studied. The analytical measurements to calculate the optical band gap showed a significant narrowing in the optical band gap value (Eg). Tauc plot was used to estimate the optical band gap (Eg) with the direct and indirect allowed transitions, before and after the reducing process. Narrowing the band gap is very important to increase the efficiency of the semiconductor material as it leads to absorbing in the visible region of the solar spectrum.

1. Introduction

The high emission gases make a great change in the global climate and this was a motivation for the researchers to explore renewable and clean resources of energy supplies to decline the results of consumption the fossil fuel that forms 90% of the total world energy [1]. Several resources are available as alternative resources such as solar cell, wind, geothermal and hydropower, these resources are considered clean compared to the fossil fuel, however, there are some circumstances for each one of these resources [2]. Another energy source which is characterized as a clean, easily stored, and a promising energy source, is the hydrogen (H2) fuel. H2 is mainly produced from petroleum and natural gas, and recently, more attention has been paid to produce the hydrogen, as an alternative fuel resource, from different renewable sources, and the interest in H2 production is increased rapidly. H2 gas can produce from different solid storage resources such as B–N based compounds, and such materials can be considered as a good supplying to a renewable source for sustainable energy in the foreseeable future. Dimethylamine borane (DMAB) is one of B–N compounds that can produce one mole of H2 gas/mole of DMAB in the presence of a suitable catalyst [3, 4, 5, 6]. The mechanism of the photocatalytic method is based on using the solar energy to provide a clean and a high energy in a short time, and this includes absorption photons that lead to excite the components of the reaction, and as a consequence, generates a new species such as radicals or making decomposition the entities of the reaction, and this process normally occurs with the assistance of a photocatalyst [7]. Different materials have been used as a catalyst to generate H2 gas such as metal oxides [8], metal nitride [9], metal sulfides [10], and nanomaterials [11, 12]. Nanomaterials such as gold nanoparticles play a vital role as a catalyst solar cells, for example, by improving the capacity of the photo-absorption and decreasing the effect of the physical properties, via the incorporation of the localized surface plasmon resonance (SPR) in these materials, which leads to induce photo-charging effect by increasing the light absorption [13, 14]. Gold nanoparticles can be excited in the visible region of the solar spectrum due to the electron oscillations that confine in a limited area of the NPs which called as SPR. The size and the distribution size of the AuNPs, that use in the photocatalytic system, has an effect on the position and the width of the SPR band [15]. Gold nanoparticles (AuNPs) are normally used as catalysts by supporting on metal oxides, such as titania and ceria, and it was reported that, the catalytic efficiency of the AuNPs is highly dependent on the properties of the gold such as the morphology and the size [16]. Gold nanoparticles with a smaller size have a highly efficient activity compared to the larger one or to the gold atoms on a bulk metal, and this belongs to the high surface area in the small particles, in addition, to the low coordination number of these small NPs, which make them to be more active, and the reason is attributed mainly to increasing the number of such particles on the surface [17, 18]. Nucleosides and their derivatives are capable of self-assembling and forming different morphology via hydrogen bonding, such as G-quartet, ribbon, nanowires, etc. [19, 20, 21, 22] The multifunctional properties of these compounds have seen growing in their attention due to the important role of these materials in different applications such as nanotechnology, drug delivery, conductivity, and so on [23, 24, 25, 26]. Herein, the mechanism
2.4. Optical band gap ($E_g$) calculation

The optical band gap was obtained by plotting the incident photon energy ($h\nu$) versus the direct and indirect allowed transitions ($\alpha h\nu)^{1/n}$ by using Tauc-Menth plot [28].

2. Experimental

2.1. Preparation gold(I) hydrogel

The sample of Au(I) hydrogel was prepared according to our previous work [23], by equimolar equivalents reaction that occurred between Au(I) ions and 6-thioguanosine nucleoside in aqueous solution. Typically, 1 ml of the hydrogel was prepared by mixing the solution of HAuCl$_4$ (13 mg, 0.033 mmol), formed reduced with two equivalents of 2,2'-thiodiethanol, to the solution of 6-thioguanosine (10 mg, 0.033 mmol).

2.2. Kinetic measurements

The data obtained by monitoring time-dependent to indicate the location of the surface plasmon resonance (SPR) for the AuNPs, that formed after reducing the polymer of Au(I):6-thioguanosine hydrogel by using DMAB, were used in this study [27]. The data were carried out by achieving a kinetic study to measure the absorption by using a Cary 100 Bio UV-Vis spectrophotometer with a quartz cuvette at room temperature with a wavelength in the range of 200–600 nm.

2.3. The dehydrogenation reaction of DMAB

A typical sample of Au(I) hydrogel was prepared by drop-casting 2 $\mu$L of the gel into a clean silicon wafer, after leaving to dry in the air, the sample was exposed to DMAB (0.1–0.3 g) in solvent-free medium at room temperature for 1 h. The reaction occurred in a sealed glass Petra-dish [27].

2.4. Optical band gap ($E_g$) calculation

The optical band gap was obtained by plotting the incident photon energy ($h\nu$) versus the direct and indirect allowed transitions ($\alpha h\nu)^{1/n}$ by using Tauc-Menth plot [28].

3. Results and discussion

The optical band gap ($E_g$) of the semiconductor materials should be laid between 1.6-2.5 eV in order to absorb light in the visible region of the solar spectrum, such materials are described as a narrow band gap material, while semiconductor material with wide band gap (~3.2 eV) can only absorb light in the ultraviolet region. Accordingly, to increase the enhancement of conversion the sunlight energy with high efficiency, the absorption range must be extended to involve the regions from visible to the gap between the 5d band and the Fermi level [35, 36]. The most common semiconductor photocatalyst that used in different applications is TiO$_2$, which has a large band gap between 3 eV to 3.2 eV, this titania is normally excited in the UV region and as a consequence, the electron transfers from the valence band (VB) to the conduction band (CB) which leads to form electron-hole pairs, however, the recombination between these electrons and holes makes this semiconductor less efficient, and in order to decrease this recombination, TiO$_2$ is doped with transition metal ions which facilitate transferring the electron to the surface of the photocatalyst and suppressed the electron-hole recombination. Another factor that could lead to increase the efficiency of titania, is to excite this photocatalyst in the visible region, numerous studies have been done to address this approach [37, 38]. This is the first report that has demonstrated the effective role of formed AuNPs, that produced by reducing the Au(I) hydrogel with DMAB, as a photocatalyst in the visible region of the solar spectrum which leads to produce the H$_2$ gas as a product from the dehydrogenation reaction of DMAB.

3.2. Analytical measurements to calculate the optical band gap ($E_g$)

Tauc plot [28] is normally used to indicate the band gap energy of the semiconductor materials. In the direct semiconductor materials, such as ZnO, GaN, etc., the conduction band, CB (which is above the band gap) and the valence band, VB (which is underneath the band gap) have the same momentum. While the value of momentum is different in the indirect semiconductor materials, such as Ge, Si, etc., for both states [29]. Plotting the $(\alpha h\nu)^{1/n}$ versus the $h\nu$ gives the value of the $E_g$ and the straight line can be extrapolated to the baseline of x axis to find the optical band gap. The spectrophotometric measurements that used for achieving...
monitoring time-dependent to indicate the localized surface plasmon resonance (LSPR) were carried out by using Cary 100 Bio UV-Visible Spectrophotometer. The polymer of Au(I):6-thioguanosine hydrogel revealed a strong optical band at 360 nm (3.44 eV), that can be assigned to the charge transfer complex that formed between Au(I) ions and the nucleoside, 6-thioguanosine, via 6S [39], and upon reducing the polymer by using DMAB, a new band revealed at 530 nm (2.33 eV) which can attribute to the formation of the surface plasmon resonance band (SPR) for the formed AuNPs [40]. It is important to note that, the SPR band depends on several factors such as the environment surrounding the nanoparticles; and this factor influences through the optical index, and the higher SPR is related to the higher optical index, another factor is the morphological structure of the nanoparticles; as it was observed that the high symmetric nanoparticles normally have a blue shifted wavelength, and finally the core-shell particles; this factor effects when the size of NPs is larger than 60 nm due to the formation of a large SPR band that leads to shift the dipolar contribution [41]. A strong wavelength peak at 530 nm was observed with the peak formed after 30 min of the reduction with DMAB, and by increasing the reduction time a slight blue shift (~30 nm) was observed in this peak and this occurred as a consequence of the effect of quantum confinement, which confirms that, the strong absorption wavelengths be smaller with smaller NPs and longer with increasing the size, and this leads to produce high and low energy band gaps, respectively [42]. In addition to the emerging a slightly blue shift in the positions of the SPR bands after 30 min of reduction time, noticeably, a gradual decrease in the intensity of the absorption band can, also, be seen after 30 min of the reducing time and this may be attributed to the formation of an agglomeration in the formed NPs which led to decline the

photocatalytic of AuNPs as a result of decrease both the surface area and the possibility to access into the active sites [3, 27]. Figure 2 (a) shows a kinetic study for monitoring time-dependent using UV-Vis spectroscopy of the formed AuNPs, the data display that the optimum time was to be 30 min. Also, the data reveal the location of the surface plasmon resonance (SPR) that formed after reducing Au(I) hydrogel at 530 nm, while Figure 2 (b) reveals that the dehydrogenation reaction of DMAB, that involves the formation of AuNPs and generates H2 gas, follows the first-order kinetics, plotting the relation of ln A0/At versus the reducing time yielded the rate constant (k), for this reaction which was equal to 0.0036 min−1. A0 and At are the absorbanse values at t = 0 and the optimum time (30 min), respectively.

3.3. The relationship between the band tail and sub band gaps

Tauc plot was used to determine the band gap energy (Eg) for the Au(I) hydrogel before and after reducing with DMAB within 70 min at room temperature, by plotting the energy of light (hv/eV) versus (αhν)2 (eV⋅cm−2) & (αhν)0.5 (eV0.5⋅cm−0.5), for the direct and indirect allowed transitions, respectively. Extrapolating the straight line to the x axis produced the value of Eg. The data present in Figure 3 (a & b) reveals clearly the difference in the optical band gap energy values before and after reducing the sample, respectively, the reason can be assigned to the effect of AuNPs that formed after reducing the sample of Au(I) hydrogel with the DMAB. Before reducing the sample of the gel, the value of the optical band gap was 3.15 eV, and the optical absorption of the sample of the gel was 3.44 eV, however, upon reducing the sample of the gel with the DMAB, the value of Eg decreased to 2.45 eV, in addition, a new optical absorption band was formed at 2.33 eV corresponding to the surface plasmon resonance (SPR). It was reported that, the VB energy of the 5d band in AuNPs is 7.88 eV and the energy of the CB band of 6sp is 4.37 eV [33], and this means, the reduction potential of H+/H2 is less positive than the bottom of 6sp and the oxidation potential of DMAB is less negative than the top of 5d, and in accordance with, it seems that the locations of these bands are responsible for transferring the electrons and they clarify the ambiguity about the important role of AuNPs in the dehydrogenation reaction of DMAB. In addition, the presence of AuNPs can lead to broadening the valence band (VB) which results in narrowing the band gap. On the other hand, AuNPs can induce the formation of a strong band tail, however, such band tail was not found in the absence of AuNPs before the reducing process, as demonstrated in the direct and indirect allowed transitions in Figure 3 before reducing (a & c) and after reducing (b & d), respectively [14, 34, 43, 44]. It is worthy to note, the difference in the value of the band gap between the direct and the indirect allowed transitions for both the reactions, before and after, reduction. Direct transitions take place by interactions of photon-electron, while indirect transitions involve the incorporation of phonons in addition to the photons and electrons, and the type of the transition could be depending on the selection rules of quantum
mechanics in addition to the occupy and empty states of the electrons in
the top of the valence band or conduction band, respectively. Normally,
the direct transitions are combined with the emission or the absorption of
the photons, while the probability of photon emission with indirect band
gap semiconductor transitions is much lower because the wavenumber of
the electrons (and therefore the momentum) is equal for direct transitions
and its change for indirect transitions [45, 46]. The onset shape of the
direct electronic transitions seems as a step, while that of the indirect band
gap reveals with much gradual, and this could be assigned to the rela-
tionship between the shape of the transitions and their symmetries of the

Figure 3. Tauc plot for determining the optical band gap (Eg) for Au(I) hydrogel with direct allowed transitions before and after reducing (a, b) and with indirect
allowed transitions, before and after reducing (c, d), respectively. The data obtained with the optimum time (30 min).

Figure 4. Determination of the band gap energy values (Eg) via direct allowed transitions using the Tauc plot for Au(I) hydrogel, that reduced by DMAB during a
period of 0–70 min at room temperature, revealed narrowing in the band gap energy from 3.15 eV (before reducing) to 2.45–2.7 eV (after reducing). The inset shows
the appearance of sub-gap bands in the range of 2.03–2.34 eV for the formed AuNPs after reducing within 70 min.
energy and the shape of the band gaps. This phenomena could indicate the nature of the compound if it has amorphous or crystalline properties [47]. Materials with direct allowed transitions can absorb only photon energy without disturbing by adsorbing any phonon, and for this reason they have a potential for application in optoelectronic devices [48]. Also, the presence of indirect band gap with a smaller value compared to that for a direct band gap in the same compound leads to produce an efficient catalyst for achieving the excitation in the solar light and this can be attributed to the availability of sufficient phonons that help in the indirect transition of electrons from the valence band to the conduction band [49].

3.4. The effect of sub-gap bands on the efficiency of the photocatalyst

The appearance of a long tail in the fundamental band gap, which seems to be shifted to the red area of the energy level, can easily refer to the presence of a sub band-gaps [50, 51]. This finding was consistent with the data collected from the kinetic study to monitor the progress in the reduction of Au(I) xerogel at room temperature within a period of 70 min, as shown in Figure 4. The data display with more clearly emerging long bands tails that extending to the red area of the light energy level associated with the reducing time during the intervals from 20 min to 70 min, while such band tail has not been seen in the sample before starting the reduction process (pink colour line corresponding with interval l0 min in Figure 4). The inset in Figure 4 reveals the formation of the sub-gap bands covering the area from 2.03 eV to 2.34 eV [52]. In accordance with, the interval of energy between two levels will be \( E_3 < E_5, E_2 < E_5 \), etc. The significant sub band gap can be clearly observed in Figure 4. The emerging of sub-band-gaps are very interesting in the materials that can narrow the band gap via direct allowed transitions as such properties can enhance the solar cell efficiency [53].

The variation in the band gap values after reducing the sample at different intervals time refers to the shift that occurred in the conduction and valence bands by the effect of AuNPs [14]. The schematic in Figure 5 shows the changing that occurred in the value of the band gap energy for the Au(I) hydrogel upon reducing with DMAB, as this value was narrowed from 3.15 eV to 2.45-2.7 eV due to the effect of AuNPs that formed after reducing the sample of the gel, also, the scheme shows the energy levels of the sub-band gaps and the broadening of the VB band that forms and lead to suppress the recombination of the electron-hole pairs and as a consequence increasing the catalytic efficiency of AuNPs.

4. Conclusions

In summary, the work principle of AuNPs as a high efficiency photocatalyst in dehydrogenation reaction of the DMAB to generate \( \text{H}_2 \) gas has been presented in this work. The photocatalytic process occurred in the visible light at room temperature in solvent-free medium. The formed AuNPs, that produced by reducing the Au(I) hydrogel by DMAB, play a vital role in the dehydrogenation reaction of the DMAB owing to the relativistic effect property in the gold, which helps in the formation of an interband as a result of transferring the electrons from 5d to 6sp. This band is responsible for the catalytic efficiency of the AuNPs in this reaction. In addition, the data that obtained by using Tauc plot revealed an important narrowing in the band gap energy after reducing the polymer of the gel with the DMAB, and this can be assigned to the effect of AuNPs which leads to broadening the valence band. Also, the presence of AuNPs suppresses the recombination between the electron-hole pairs which leads to increase the efficiency of AuNPs in the production of \( \text{H}_2 \) gas by transferring the electrons onto the surface of AuNPs then move to reduce the hydride ions. Interestingly, the reaction was very fast and the reduction occurred within 1h. It is worth to note that, the data showed formation of sub-band gaps after reducing the hydrogel, and formation of such sub band-gaps are very important in the development of the energies of solar cells.

Declarations

Author contribution statement

Lamia L. G. Al-mahamad: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

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Data availability statement

Data included in article/supplementary material/referenced in article.

Declaration of interests statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

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