

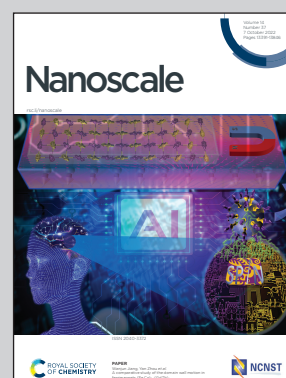


Showcasing research by Chia-Chen Wu and Prof. Yen-Hsun Su from National Cheng Kung University, Taiwan, and Dr Fei Pan from Technische Universität München, Germany.

Machine learning ensures rapid and precise selection of gold sea-urchin-like nanoparticles for desired light-to-plasmon resonance

Genetic algorithm neural network (GANN) is utilized to achieve a synergistically trained model by considering the light-to-plasmon conversion efficiencies and fabrication parameters. The obtained trained model can thus guide us to efficiently gain a rapid and automatic selection of fabrication parameters of the nanoparticles with the anticipated light-to-plasmon resonance.

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## Machine learning ensures rapid and precise selection of gold sea-urchin-like nanoparticles for desired light-to-plasmon resonance

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Sustainable energy strategies, particularly solar-to-hydrogen production, are anticipated to overcome the global reliance on fossil fuels. Thereby, materials enabling the production of green hydrogen from water and sunlight are continuously designed, *e.g.*, ZnO nanostructures coated by gold sea-urchin-like nanoparticles, which employ the light-to-plasmon resonance to realize photoelectrochemical water splitting. But such light-to-plasmon resonance is strongly impacted by the size, the species, and the concentration of the metal nanoparticles coating on the ZnO nanoflower surfaces. Therefore, a precise prediction of the surface plasmon resonance is crucial to achieving an optimized nanoparticle fabrication of the desired light-to-plasmon resonance. To this end, we synthesized a substantial amount of metal (gold) nanoparticles of different sizes and species, which are further coated on ZnO nanoflowers. Subsequently, we utilized a genetic algorithm neural network (GANN) to obtain the synergistically trained model by considering the light-to-plasmon conversion efficiencies and fabrication parameters, such as multiple metal species, precursor concentrations, surfactant concentrations, linker concentrations, and coating times. In addition, we integrated into the model's training the data of nanoparticles due to their inherent complexity, which manifests the light-to-plasmon conversion efficiency far from the coupling state. Therefore, the trained model can guide us to obtain a rapid and automatic selection of fabrication parameters of the nanoparticles with the anticipated light-to-plasmon resonance, which is more efficient than an empirical selection. The capability of the method achieved in this work furthermore demonstrates a successful projection of the light-to-plasmon conversion efficiency and contributes to an efficient selection of the fabrication parameters leading to the anticipated properties.

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## Introduction

Global reliance on fossil fuels has led to an energy crisis and urges a sustainable energy strategy, especially solar-to-hydrogen production, thus stimulating substantial efforts in exploring nanoscale materials.<sup>1–7</sup> In the nano-dimension, materials can alter their characteristics,<sup>8,57–59</sup> such as, the reactivity to light, enabling electron polarization on the metal surface, or facilitating the electron excitation to the conduction band.<sup>9–16</sup> Thereby, nano metals and metal-oxides display prominent functions, especially the light-to-plasmon resonance, which can be utilized to optimize the development of solar cells,

optical sensors, and gas sensors.<sup>17–21</sup> As a typical nano metal-oxide, zinc oxide (ZnO), a conventional n-type semiconductor with a wide bandgap of 3.37 eV at room temperature, has an absorbance peak of 340–370 nm within the ultraviolet range.<sup>22–28</sup> In addition to its excellent conductivity, ZnO has outstanding biological compatibility and low toxicity, unlike most photosensitive materials (CdS, PdS, GaSe, and BiSe). Moreover, ZnO, owing to its hexagonal wurtzite crystallinity, exhibits a variety of anisotropic morphologies associated with the different synthesis methods, *e.g.*, nanorods, nanosheets, nanoflower, and nanowires.<sup>29–32</sup> However, ZnO still presents deficiencies in properties, *i.e.*, the electron-hole recombination rate and the restricted response in the UV region. In this regard, many strategies have been proposed to increase the associated visible light absorption by coating ZnO substrates with various nanometals or narrowing the bandgap of ZnO substrates by introducing defects.<sup>33</sup> These attempts have thus demonstrated that a coating of plasmonic nanoparticles on metal oxides can reduce the electron/hole recombination and

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consequently improves the light-to-electron conversion efficiency to improve the utilization of visible light.<sup>34–36</sup> Moreover, the creation and separation of charge carriers can be enhanced when a Schottky barrier occurs on the heterojunction between metal oxides and noble nanoparticles.<sup>37–40</sup> Therefore, noble nanoparticles are introduced to coat ZnO substrates by applying organic linkers (e.g., MPTMS or similar molecules). However, such coatings cannot subsequently improve the charge transport and thermal endurance of ZnO substrates, although some of their properties can be enhanced by exploiting hazardous organic linkers.<sup>41</sup>

To overcome such bottlenecks, machine learning (ML) methods have emerged as a powerful tool to discover an optimized process for material preparations.<sup>42,43,60</sup> ML as a new tool has shown its uniqueness in materials sciences, e.g., deciphering crystallography<sup>44</sup> and discovering superconducting materials.<sup>45</sup> Moreover, the development of high-performance computing enables an extensive application of artificial neural networks (ANNs) and deep learning. ANNs have been used to construct models to establish the relationship between the physical properties of photonic materials and the associated electromagnetic responses. The established model can subsequently project the electromagnetic responses of a photonic material once their physical parameters are input.<sup>46</sup> However, there are only a few published studies where ML is employed to discover an optimized method to prepare nanoparticles of an anticipated surface plasmon resonance.<sup>42,47</sup>

In this work, we successfully established a model trained by a genetic algorithm artificial network (GANN), a method applied in ML to train models to project the light-to-plasmon resonance. To do so, we empirically synthesized ZnO nanoflowers on the ITO substrates and subsequently coated the gold nanoparticles onto ZnO nanoflowers by ultraviolet treatment.<sup>48</sup> Furthermore, a mixture of sodium citrate, hydroquinone, and HAuCl<sub>4</sub>, is loaded to generate the second growth of gold nanoparticles. The optical and morphological properties of Au/ZnO were subsequently analyzed by UV-visible spectroscopy, quantum efficiency (QE), scanning electron microscopy (SEM), transmission electron microscopy (TEM), and X-ray powder diffraction (XRD). Thereby, we trained the model through GANN by utilizing the parameters of nanoparticle fabrications and the yielded light-to-plasmon resonances. Consequently, we achieved a projection enabling us to select the fabrication parameters for the anticipated light-to-plasmon resonance.

## Results and discussion

### Crystalline characteristics

The crystallography of the fabricated ZnO nanoflowers was analyzed by XRD in Fig. 1. The peak at 34.44° corresponds to the plane (002), indicating that the hexagonal wurtzite structure was preferentially formed in the [001] direction. Furthermore, the measured lattice planes of (100) and (101) suggested the fabricated ZnO nanoflowers have a polycrystalline structure. We could moreover find a peak at 38.1° of low



**Fig. 1** XRD patterns of the Au-decorated ZnO nanoflowers. The symbols depict the positions of the analyzed reflections from the Au, ZnO, and ITO planes. The Au-coated ZnO nanoflowers are synthesized using 10<sup>−4</sup> M chloroauric acid, 0.2 wt% sodium citrate, and 3500 μL of hydroquinone. The phase analysis is executed by comparing to JCPD Standard (No. 75-576).

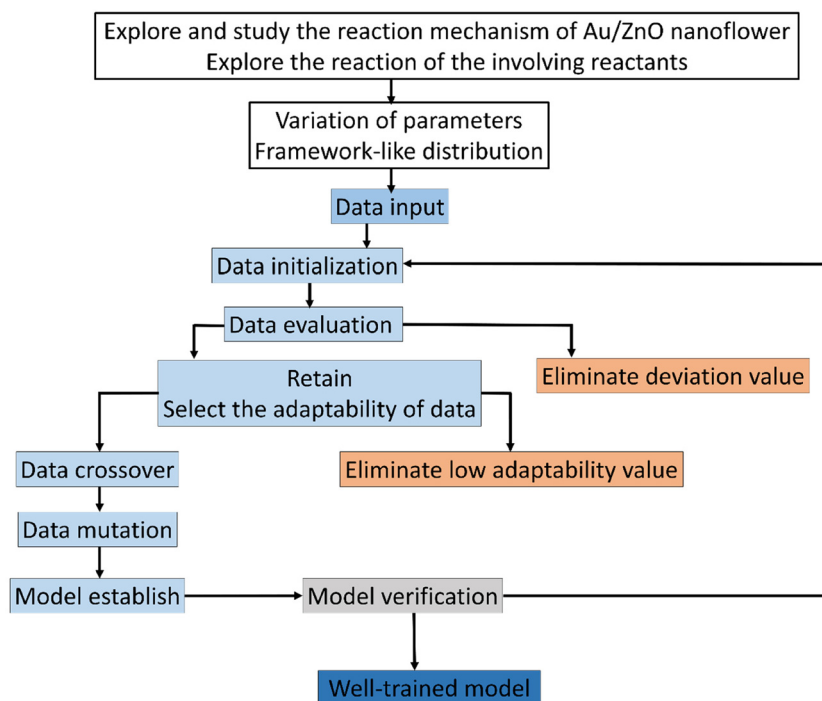
intensity, corresponding to the (111) plane of the metallic gold. The other measured lattice planes, such as (211), (222), (440), (611), (622), and (444), indicated the polycrystalline structure of the ITO thin film.

### Decision flow

In order to obtain a well-trained model through GANN for the prediction, a delicate design of experiments is crucial to obtaining high-quality data and thus ensuring the trained model of high accuracy (Fig. 2). In selecting the inputs to train the model, we carefully considered the reaction mechanism, data analysis, and the collected data from previous experiments because a lucid comprehension of the reaction mechanism can lead to an appropriate input of the experimental parameters to train the model. Therefore, the trained model can yield a precise data projection in the range of interpolation and modulation of the activated-plasmon enhancement for the desired condition. In this regard, we systematically arranged the experimental parameters with a carefully designed distribution and could thus decrease the experimental amount. The achieved experimental results alongside the experimental parameters are exploited to initiate the model training with the necessary definitions, e.g., the length of data, the amount of data, the crossover ratio, the mutation ratio, and the calculation functions. However, the data varying beyond the average standard were eliminated to train the model. Subsequently, the residual data were further used and filtered according to their adaptability. However, a careful filtration is necessary as it can mislead the model training and cause mutation. After the preliminary model training, we can establish a premature model, which still has deficiencies to







**Fig. 2** Machine learning framework of a closed learning loop based on GANN. The frame is separated into two sections. White represents the exploration and the parameter setting, and color represents the data input selection, the model establishment, and the feedback mechanism.

match the data. Hence, we need a mutation to process the derived crossover data. In addition, a feedback mechanism is required to guarantee an accurate prediction. Thereafter, we can gain a well-trained model once we accomplish a complete loop as shown in Fig. 2.

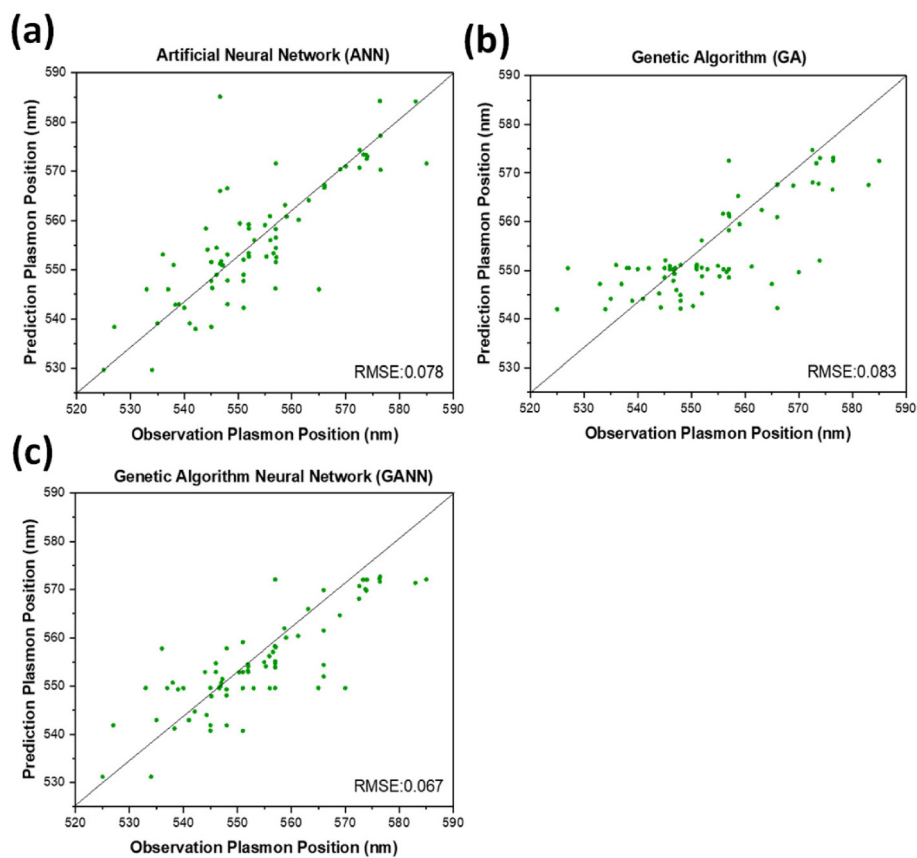
### Model establishment

Three different algorithms, ANN, genetic algorithm (GA), and GANN, are usually applied to address nonlinear questions in conventional programming. We thereby employed these methods to derive a prediction of the position of surface plasmon resonance, which was verified by the empirical investigation (Fig. 2). We additionally defined a regression line ( $Y = X$ ) as an ideal prediction. Namely, the closer the projected data are located to the regression line, the higher accuracy the trained model has. To maintain consistency, we utilized the same sets of parameters in training the model through the three algorithms aforementioned (Fig. 3). The generated results by applying ANN manifested a lower distribution condensation than GA and GANN, suggesting ANN in our study can lead to a more precise prediction (Fig. 3). Such precise prediction relies on the neuromorphic system of ANN, which can optimize the specific weights and generate a better classification. However, due to a lack of evolution, ANN is inefficient in handling complex questions. On the contrary, GA can generate a fitting function for our research by forming a crossover of the input parameters and mutations for multiple solutions. Therefore, a combination of ANN and GA can yield a more efficient model by considering the crossover and mutations.

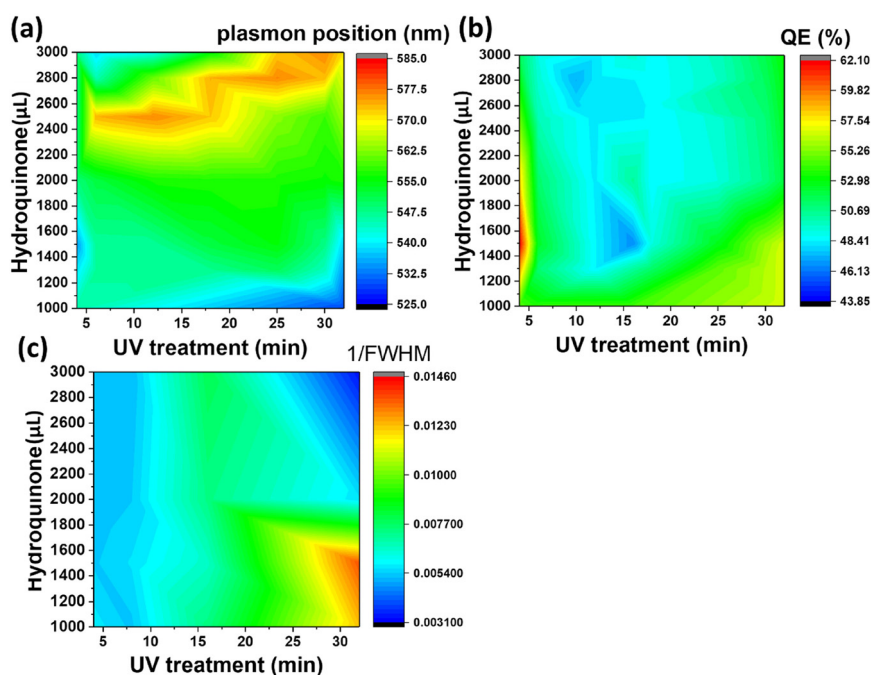
Moreover, this combination supports a feedback mechanism to modify and optimize the network structure, thus realizing deep learning on multiple levels.

After learning cycles, we achieved the prediction model through GANN, and we thereafter obtained the plasmon positions, the quantum efficiencies (QE), and the reciprocals of the full widths at half maximum (FWHMs) (Fig. 4). In training the model, we considered the experimental parameters, *i.e.*, the times of the UV treatment and the hydroquinone amounts, as the dependent variables. On the contrary, we considered the anticipated parameters as independent variables, the positions of the surface plasmon resonance, the QEs, and the FWHMs. Therefore, the positions of the surface plasmon resonance can be projected concerning the times of the UV treatments and the hydroquinone amounts (Fig. 4a), where the color scale bar represents the plasmon shifting of a max wavelength of 585 nm (in red). The other colors in Fig. 4a represent decreasing plasmon positions from orange to blue (min: 525 nm). A larger wavelength of the surface plasmon resonance occurred at a hydroquinone amount in the range of 2200–2600  $\mu\text{L}$  and the gold colloid amount of 7–22  $\mu\text{L}$ . The lower wavelength of the surface plasmon resonance happened with a hydroquinone amount of less than 1100  $\mu\text{L}$  or with a UV treatment over 20 min. Similarly, we obtained a QE prediction in Fig. 4b, where the red color is the maximum efficiency and the blue color is the minimum efficiency. The lower QE occurred when the hydroquinone amount was 1500–1900  $\mu\text{L}$  or 2400–2800  $\mu\text{L}$ . Interestingly, there was a small range displaying an impressive QE up to 66%, where the hydroquinone amount was around





**Fig. 3** Leave-one-out cross-validation results for the three different algorithm models to predict the plasmon positions of the Au-coated ZnO nanoflowers: (a) the artificial neural network (RMSE = 0.078), (b) the genetic algorithm (RMSE = 0.083), and (c) the genetic algorithm neural network (RMSE = 0.067). The  $Y = X$  line is delineated as a representation of an ideal prediction.



**Fig. 4** GANN was utilized to calculate (a) the plasmon positions, (b) the quantum efficiencies, and (c) the reciprocals of FWHMs of the Au-coated ZnO nanoflowers.



1600  $\mu\text{L}$ , and the UV treatment was roughly 4 min. We furthermore performed the FWHM prediction in Fig. 4c, providing information on the morphological peaks. We can thereby find a sharp peak for the UV treatment of roughly 32 min. Moreover, we can smoothen the absorption peak by decreasing the treatment time. This information is very beneficial in designing a material with the anticipated surface plasmon resonance.

### Model validation

In order to validate the model trained by GANN, we cross-checked the data projected by the model, regarding the positions of the surface plasmon resonances, the QEs, and the reciprocals of FWHMs ( $1/\text{FWHM}$ ) (Fig. 5–7). In this regard, the concentration and the volume of the chloroauric acid solution were fixed at  $10^{-4}$  M and 10 mL. We subsequently varied the treatment times of UV from 5 to 32 min, and the hydroquinone amounts from 1000 to 3000  $\mu\text{L}$ . We observed a good correlation between the experimental data (Fig. 5) and the projected data (Fig. 4a): the high wavelength of the surface plasmon resonance appeared with the hydroquinone amount in the range of 2200 to 2600  $\mu\text{L}$ ; the low wavelength appears at

the hydroquinone amount of 1000  $\mu\text{L}$  and the UV treatment over 25 min. We thereby extracted the conditions to synthesize five samples of different wavelengths for the surface plasmon resonance. Additionally, we analyzed the sizes and the morphologies of the synthesized gold nanoparticles, suggesting that the bigger particles appeared at the surface plasmon resonance of a larger wavelength (Fig. 5). We furthermore observed that the thorns became shorter and more difficult to find when the surface plasmon resonance wavelength reached over 554 nm. Moreover, such particles became spherical when the wavelength was larger than 572 nm. Moreover, we noticed that the morphology of gold nanoparticles became random and began to aggregate, with a respective QE of 42% (Fig. 5e). Therefore, we selected  $10^{-4}$  M chloroauric for further validation concerning QEs and  $1/\text{FWHM}$ s (Fig. 6 and 7).

We performed an FDTD simulation to study the generated electromagnetic field on the surfaces of the gold nanoparticles (Fig. 6). The strength of the electromagnetic field on the gold nanoparticles can consequently impact the resonant reaction of the surfaces.<sup>49,50</sup> Subsequently, we utilized the FDTD models to simulate the particles with similar morphologies to those observed by SEM (Fig. 5a–e). The advantage of FDTD

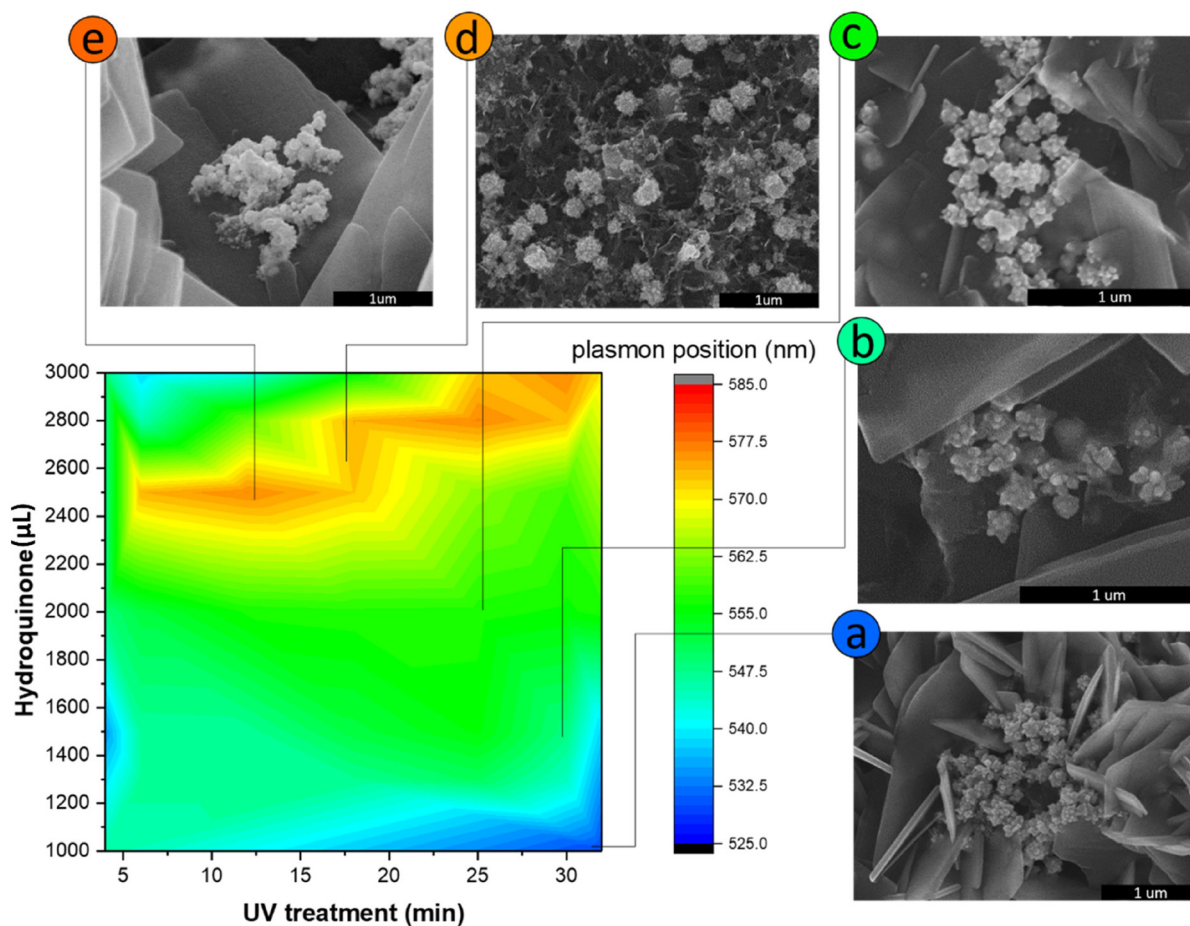


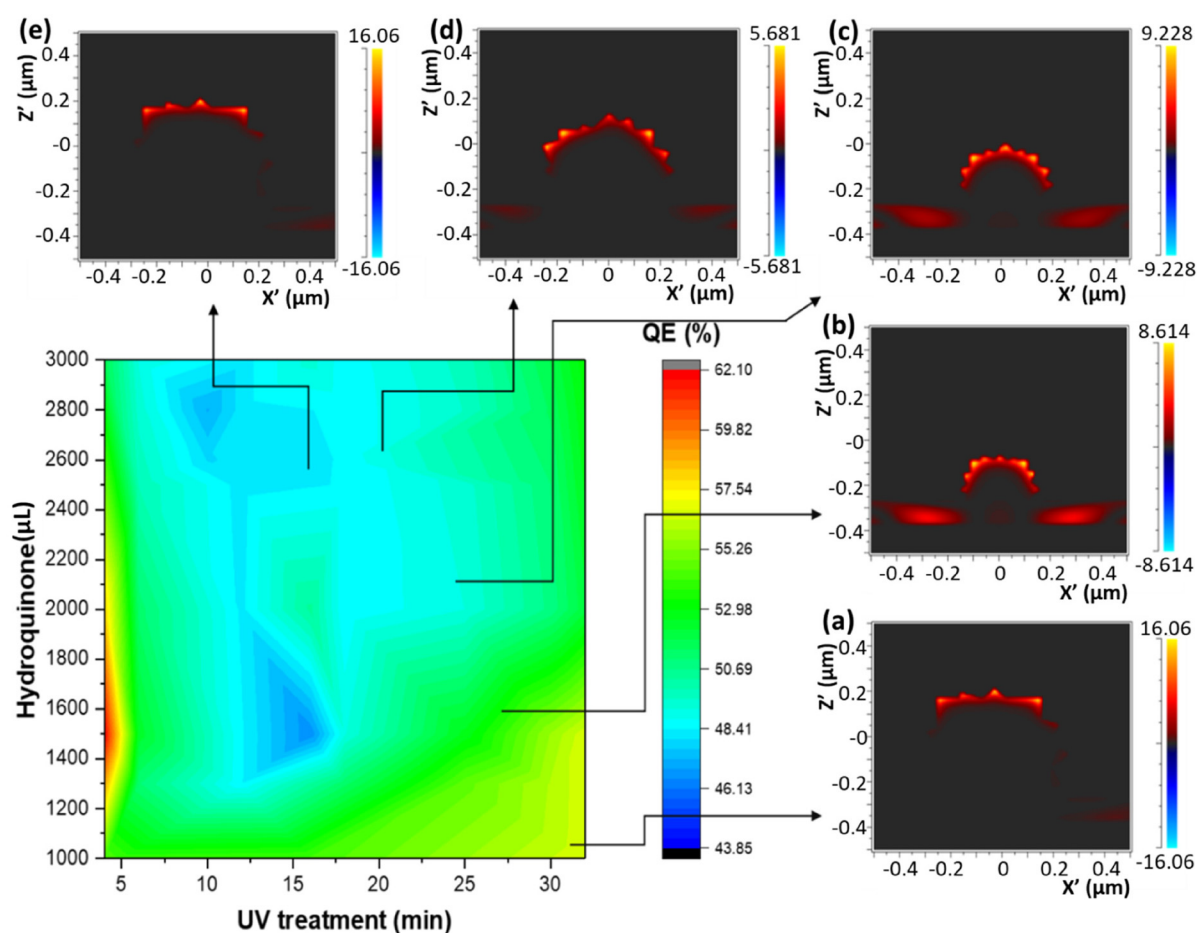
Fig. 5 Configuration between the morphologies and the plasmon positions of the Au-coated ZnO nanoflowers. Size-dependent Au particles were coated on the ZnO nanoflowers under different conditions, displayed from (a) to (e).



models considers the number of thorns, the size of particles, and the thickness of the ZnO. In our modeling, we fixed the thickness of the ZnO thin films to 30 nm and the incident light perpendicular to the ITO thin film. However, in the modeling, we introduced the incident light of two different wavelengths (400 nm and 800 nm) to the Au/ZnO system. Once the incident light of a specific wavelength induces an electron sea on the gold nanoparticle, a surface plasmon resonance can consequently occur (Fig. 6). Thereby, the strongest oscillation can subsequently take place on the surfaces of the gold nanoparticles, attributed to several factors. One of these factors is the interaction between the incident light and the electron sea, leading to the fluctuation in the system of the activated electromagnetic field.<sup>49,51</sup> The fluctuation can further influence the electron coupling with the polar by the surface plasmon polariton. Another factor is the tip-enhancement due to the varying morphologies of the gold nanoparticles.<sup>52</sup>

Therefore, we used a second-growth method to synthesize the urchin-like gold nanoparticles on the ZnO nanoflower surfaces, yielding many thorns.<sup>53</sup> These thorns of small tip radii manifest strong electric fields, inducing the electron polarization and thus leading to tip-enhancement. Therefore, it is probable to transfer electrons from gold nanoparticles to ZnO semiconductors.

We moreover found the variation of the UV treatment times and the hydroquinone amounts can impact the absorption peaks as well as the reciprocals of FWHMs (Fig. 7). In the configuration diagram, the morphology of the absorption peaks varied slightly, so it was necessary to apply a numerical analysis to investigate the trifling changes. However, the reciprocal FWHMs can directly inspect such variations. Thereby,  $1/\text{FWHMs}$  of the Au/ZnO absorption system showed a resonant coherence due to the two influential factors. The first was the size, the morphology, the distribution, and the stacking of



**Fig. 6** Configuration between the resonance simulation and the plasmon positions of the Au-coated ZnO nanoflowers. The FDTD simulation results are displayed from (a) to (e) concerning the urchin-like gold nanoparticles coated on the ZnO nanoflower surfaces of different sizes and morphologies by varying wavelengths of the incident light. The quantum efficiency (QE) indicates the conversion ratio of the photons of the incident light to the surface plasmon resonance. In order to calculate QE, we collected the scattering light of surface plasmon resonance by integrating the sphere, measured the energy of the scattering light, and then converted the scattering light energy into the quasi-particle number according to the Plank equation ( $E = nh\nu$ ). The quasi-particle number is the surface plasmon resonance number, which serves as a numerator. The power of incident light is also converted into the number of the incident photon by the Plank equation, which acts as a denominator. Thereby, the QE of light-to-plasmon is calculated.





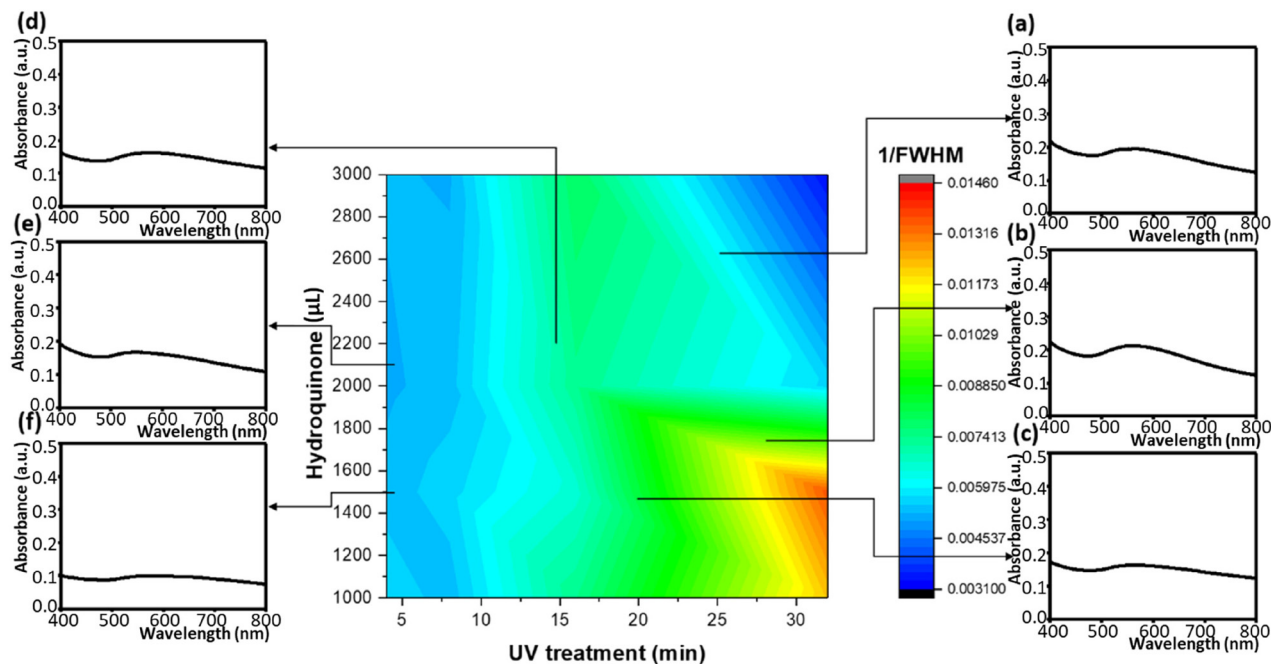


Fig. 7 Configuration between the absorption spectra and the reciprocals of the FWHMs of the Au-coated ZnO nanoflowers. The absorption peaks of various morphologies at different conditions are displayed from (a) to (f).

gold nanoparticles on ZnO nanoflowers. The second was whether the incident light could induce the electron sea on the gold nanoparticles or the tip-enhancement, dependent on

the profile of the gold nanoparticles, the shape of thorns, the density of thorns, and the distribution of thorns.<sup>54</sup> Overall, most of the above effects were directly related to the charge distribution and indirectly associated with the structure of the gold nanoparticles. A combination of material development and machine learning coupled with a genetic algorithm neural network can precisely project the material characteristics and optimize the fabrication process of the material composite. Moreover, the developed projection model can effectively and accurately simulate the empirical results.

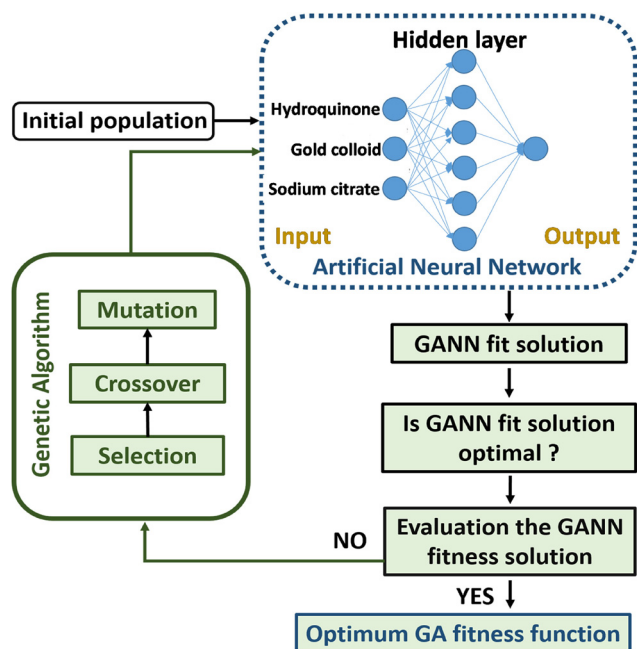


Fig. 8 Flow chart of the empirical parameters in the artificial neural network coupling with the genetic algorithm of the Au-coated ZnO nanoflowers. The framework consists of two sections: the blue section (the structure of the artificial neural network) and the green section (the genetic algorithm process).

## Conclusion

In this work, we coated the urchin-like gold nanoparticles onto ZnO nanoflower surfaces and successfully established a precise prediction model trained by GANN. There were four factors significantly impacting the precision of the trained model. Moreover, these four factors can be observed from the decrease of RMSE and the convergence of the tested and trained data. The first was the amount of the hidden layer, separating data into different districts in order to clarify the whole database. The second was the neural number in the hidden layer. An increased number of neurons can enlarge the complexity of the training process. Furthermore, the number of neurons was double the number of the independent variables. The third was the mating ratio, the adjustment of which was crucial to establishing a well-training model. Moreover, an appropriate mating ratio can optimize the solution. The last was the learning cycle, whose prolongation can reduce the RMSE and yield a better projection. Consequently, the utiliz-





ation of the model obtained by machine learning can finally generate predictable results in line with an empirical fabrication, facilitating a tuning of the wavelength of the surface plasmon resonance and the quantum efficiencies. We can thereby produce gold nanoparticles of the desired light-to-plasmon resonance, guiding the design of high-performance water-splitting cells.

## Materials and methods

### Materials

Zinc acetate ( $\text{Zn}(\text{CH}_3\text{COO})_2 \cdot 2\text{H}_2\text{O}$ ), chloroauric acid ( $\text{HAuCl}_4$ ), sodium citrate ( $\text{Na}_3\text{C}_6\text{H}_5\text{O}_7 \cdot 2\text{H}_2\text{O}$ ), and hydroquinone ( $\text{C}_6\text{H}_4(\text{OH})_2$ ) were purchased from Sigma Aldrich. The applied chemical reagents were of analytical grade. Deionized water (DI) was used throughout the experiment.

### Preparation of pure ZnO

The ITO substrate was cleaned with ethanol and acetone by ultrasonication for 10 minutes, rinsed with DI water, and dried with  $\text{N}_2$ . The precursors of the ZnO seed layer were 0.05 M zinc acetate in 0.02 M hexamethylenetetramine (HMTA). After coating the precursors on the ITO substrates, samples were then annealed at 350 °C for 10 minutes. Subsequently, the samples were immersed in the precursor solution of ZnO nanorods at 95 °C for 3 hours, which were prepared by dissolving the 0.02M zinc acetate solution in HMTA (0.02 M, 0.01 M, and 0.005 M). Afterward, the samples were annealed again at 350 °C for 30 minutes. The annealed samples were then immersed into the precursors of ZnO nanoflowers containing 0.15 M zinc acetate in methanol at 60 °C for a given time (16, 18, 20, 22, and 24 hours).

### Preparation of Au-modified ZnO

ZnO nanoflower substrates were immersed in a  $10^{-4}$  M  $\text{HAuCl}_4$  solution, and then an ultraviolet (UV) light was applied for an appropriate time (4 min, 8 min, 16 min, and 32 min) to obtain the reduced  $\text{Au}^{\text{III}}$  on the ZnO nanoflower surfaces for the generation of spherical gold nanoparticles. The samples were afterward rinsed twice with DI water. The 0.2 wt% sodium citrate solution in 3 mL of hydroquinone was subsequently added to the samples, and  $\text{HAuCl}_4$  was then added.

### Characterization

The surface morphology of the gold sea-urchin-like nanoparticles was determined by field emission scanning electron microscopy (FESEM, Hitachi). The transmission electron microscopy (TEM) images were recorded with a JEOL 3010 TEM (Netherlands, Philips).<sup>55,56</sup> The X-ray powder diffraction (XRD) patterns were obtained from a D8 Advance diffractometer with Cu K alpha radiation (wavelength = 0.111 nm). The absorption spectra of the samples were measured using a Lambda 950 UV-vis spectrophotometer (PerkinElmer, New York, US).

### Machine learning

Predictable results in materials science are the sought-after research goals that can guide controllable properties, characteristics, and functions of the fabricated materials. Artificial neural networks are well-known predictive models containing functional algorithms for different situations and complicated problems. In addition, a genetic algorithm neural network (GANN) has been successfully applied to model and tune the ample variables for practical processes. Such machine-learning studies were conducted with a commercial software “Super PCNeuron”.

A genetic algorithm neural network (GANN) consists of two main calculation parts: a genetic algorithm for optimal process and an artificial neural network for establishing a non-linear model. The genetic algorithm contains selection, crossover, and mutation, each based on the natural reproduction rule. An artificial neural network is based on the operation of the human brain containing neurons and synapses. Thereby data can be better classified and evaluated.

The network contains a single hidden layer and 6 neurons as shown in Fig. 8. The input data comprise experiment parameters: various UV treatment times and hydroquinone amounts. After a matrix calculation, the accuracy of the trained model is inspected. If the trained model shows a low accuracy or does not fit, the model will be refined until the GA process is optimized.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

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