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# Transparent TiO<sub>2</sub> nanotubes supporting silver sulfide for photoelectrochemical water splitting†

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Differences between photoelectrochemical and electrochemical activity were thoroughly investigated for the oxygen evolution reaction mediated by  $Ag_2S$  deposited on two types of ordered titania substrates. Titanium dioxide nanotubes were fabricated by anodization of magnetron sputtered Ti films on ITO-coated glass substrates or directly from Ti foil. Further,  $Ag_2S$  deposition on the nanotubes was carried out using successive ionic layer adsorption and reaction, known as SILAR, with 5, 25, and 45 cycles performed. Two types of nanotubes, one on transparent the other on non-transparent substrates were compared regarding their geometry, structure, optical, and electrochemical properties. It was demonstrated that the composite of  $Ag_2S$  grown on transparent nanotubes exhibits higher catalytic activity compared to  $Ag_2S$  grown on the nanotubes formed on Ti foil. The results showed that transparent nanotubes after modification with  $Ag_2S$  by 25 SILAR cycles exhibit ca. 3 times higher photocurrent under visible light illumination than non-transparent ones treated with the same number of cycles. Furthermore, transparent nanotubes after 45 SILAR cycles of  $Ag_2S$  exhibit enhanced activity towards oxygen evolution reaction with 9.3 mA cm<sup>-2</sup> at 1.1 V vs. Ag/AgCI/0.1 M KCI which is six times higher than titania alone on Ti foil.

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### 1. Introduction

In the world of increasing global energy demand, sustainable and renewable energy sources have become indispensable. As fossil fuel resources dwindle and environmental concerns intensify, scientists are developing innovative solutions to meet the growing energy needs of society. The combustion of carbon-based materials releases greenhouse gases which contribute significantly to negative climate changes and degradation of the environment. Therefore, renewable energy sources such as solar, wind, hydropower, biomass, or geother-

mal offer attractive alternatives without depleting limited resources. Among others, the water splitting process to produce hydrogen used in fuel cell technology is a promising route forward. The water electrolysis process utilizes electrodes coated with a catalyst to perform the reaction. Conventional catalysis is done with noble metals or metal oxides, however, metal sulfides are increasingly being explored as a lower cost alternative.2 Metal sulfides are a class of compounds in which a sulfur (S) anion is combined with a metal (M) cation labelled as  $M_x S_y$  forming sulfides of various stoichiometry.<sup>3</sup> Up to now, various metal sulfides like CdS, 4 Ag<sub>2</sub>S, 5 PbS, 6 NiS<sub>v</sub>, 7 CoS<sub>v</sub>, 8 and FeS<sub>v</sub> have been implemented for oxygen and hydrogen evolution reactions. Typically, sulfur is in the -2 oxidation state and the metal in the +2, +3, or +4 oxidation state. Their unique optical, electrical, and catalytical properties such as bandgap tunability,10 low valence band level,11 quantum size effect,12 facile synthesis methodology, 13 and exposed catalytic sites 3 position them as promising candidates for energy conversion applications. Among others, silver sulfide extends its applicability in many fields especially in photodetectors, 14 solar cells,15 and catalysts for example in the oxygen reduction reaction16 or oxygen evolution reaction (OER).17 Ag2S is an n-type semiconductor with a narrow band gap of 1 eV.18 Furthermore, it exhibits strong absorption from the ultraviolet through the visible spectrum and near-infrared regions.<sup>14</sup> Investigation of the catalytic activity of Ag<sub>2</sub>S towards OER shows that the

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Paper Nanoscale

common practice is to combine silver with another metal to increase the catalytic activity. However, this work aims to explore Ag<sub>2</sub>S alone without transition metal co-catalysts like Ni, <sup>17</sup> Co, <sup>17</sup> or Fe<sup>19</sup> usually used for oxygen evolution processes.

Here, we present the fabrication process where both transparent and non-transparent TiO2 nanotube substrates were modified by successive ionic layer adsorption and reaction (SILAR) of Ag<sub>2</sub>S and thermal treatment. Titanium dioxide nanotubes were chosen as a platform for metal sulfide modification because of their high degree of ordering, larger surface area than planar films, and strong absorption of UV light which can be tuned by the addition of other photosensitizers, thereby leading to an enhancement of charge collection with visible illumination. 20,21 The electrochemical and photoelectrochemical properties of Ag<sub>2</sub>S on transparent TiO<sub>2</sub> nanotubes fabricated on ITO-coated glass were compared with non-transparent nanotubes fabricated on Ti foil, indicating significant changes in the properties of the electrode material depending on the chosen titania support. The electrochemical measurements were carried out in two different electrolytes: 0.5 M Na<sub>2</sub>SO<sub>4</sub> and 0.5 M NaOH, with which the material's (photo)activity toward OER was investigated.

#### **Experimental** 2.

### 2.1. Reagents

ITO glass substrates (S111, thickness: 1.1 mm, ITO thickness: 100 nm, Ossila), titanium foil (99.7%, thickness: 0.127 mm, Strem), acetone (99.5%, Chempur), ethanol (96%, Chempur), 2-propanol (99.7%, Chempur), ethylene glycol (99.5% Chempur), phosphoric acid (85%, Chempur), ammonium fluoride (98%, Chempur), silver nitrate (POCH), sodium sulfide nonahydrate (98%, Sigma-Aldrich).

#### 2.2. Electrode fabrication

The ITO substrates with a size of  $1.5 \times 2 \text{ cm}^2$  were ultrasonically cleaned in water, ethanol, acetone, and isopropanol for 15 min. The thin Ti interlayer was deposited for 15 seconds using magnetron sputtering at a current of 0.412 A, argon flow of 1 sccm, and pressure of  $2.0 \times 10^{-3}$  Pa (nanoPVD, Moorfield). This initial Ti layer was thermally treated for 3 min at 400 °C on a hot plate (PCE-E6000 Serials). Then, a second round of magnetron sputtering for 30 min was performed to obtain the main Ti layer with a current 0.412 A, argon flow of 2 sccm, and a pressure of  $2.0 \times 10^{-3}$  Pa leading to a film thickness of ca. 400 nm, verified by profilometry (DektakXT, Bruker). Titania nanotubes were fabricated using an anodization process in a two-electrode system, where the Ti on ITO was the anode and a Pt mesh was the cathode. The Ti electrodes were placed in a custom holder where only a defined part of the sample was anodized, namely a circle-shaped area with a diameter of 10 mm (see Fig. 1c). The electrolyte contained 0.27 M NH<sub>4</sub>F/1 M H<sub>3</sub>PO<sub>4</sub>/1 vol% H<sub>2</sub>O/99 vol% ethylene glycol. The anodization was carried out at a temperature of 23 °C controlled with a thermostat and an applied voltage of 40 V for 60 min. Next, the electrodes were thermally treated at 450 °C for 1 h in a

tube furnace in ambient atmosphere. Herein, these electrodes are designated as gTiNT. The first letter "g" indicates a glass substrate was used as a support for the titanium deposition. Alternatively, titania nanotubes were fabricated on Ti foil. First, Ti foil was cut into  $2 \times 3$  cm<sup>2</sup> plates and ultrasonically cleaned in acetone, ethanol, and water for 10 min. The electrolyte contained 0.27 M NH<sub>4</sub>F/15 vol% H<sub>2</sub>O/85 vol% ethylene glycol. The anodization was performed under temperature of 23 °C and voltage 40 V for 1 h. After anodization the samples were annealed in a tube furnace for 1 h at a temperature of 450 °C also in air. These samples are designated fTiNT. The first letter "f" indicating the material was fabricated on foil. Finally, SILAR was carried out both on gTiNT and fTiNT sample sets using silver and sulfur ionic solutions with different numbers of deposition cycles. Before the first SILAR cycle, the electrodes were placed in UV light and ozone for 30 min. Then the modification consisting of 5, 25, and 45 SILAR cycles was performed. The samples were immersed in aqueous 5 mM AgNO3 for 30 min and then in 5 mM Na<sub>2</sub>S for 30 min. The samples were rinsed then with DI water and dried. The procedure was further repeated for the given cycle numbers with the time of the subsequent solution immersion decreased to 2 min. The last step was to anneal in argon for 1 h at the temperature of 250 °C.

### 2.3. Sample characterisation

The morphology of the electrodes was analysed by scanning electron microscopy (SEM) and energy dispersive X-ray spectroscopy (EDX) acquired with a JEOL JSM-F100. Furthermore, Grazing Incidence X-ray Diffraction (Gi-XRD) was used to study the crystallographic structure. Samples were measured in a Panalytical Empyrean, equipped with a Cu lamp working at 4-5 kV 40 mA. The z position of the samples was optimised to faintly detect the most intensive peak of the samples ( $\sim 25^{\circ}$ ) and increments of omega (incident angle) between 0.01 and  $10^{\circ}$  ( $\Omega$ ) were used to collect spectra from 15 to 50° of  $2\theta$ . Samples are represented in a contour map in log scales in y and z. Penetration depth was calculated according to the density of the TiO<sub>2</sub> and the Cu K-alpha (1.5406 Å) wavelength. The optical properties of the electrodes were inspected using an UV-vis spectrophotometer (Lambda 35 PerkinElmer). The spectra were recorded in the wavelength range from 300 to 900 nm.

The chemical structure of the samples was measured by X-ray photoelectron spectroscopy using a Thermo Scientific<sup>TM</sup> K-Alpha<sup>TM</sup> X-ray photoelectron spectrometer (XPS). The samples were irradiated with Al  $K\alpha$  = 1486.7 eV X-ray radiation. Measurements were carried out under a pressure of 10<sup>-9</sup> to 10<sup>-8</sup> mbar. Survey spectra were registered using a pass energy of 150 eV and a step size of 1 eV. The equipment was calibrated using C 1s (284.5 eV). High-resolution spectra were recorded for the oxygen O 1s, titanium Ti 2p, carbon C 1s, silver Ag 3d, and sulfur S 2p binding energy regions using a pass energy of 20 eV and step size of 0.1 eV.

Raman spectra were recorded by means of a confocal micro-Raman spectrometer (InVia, Renishaw) with sample excitation by an argon ion laser emitting at 514 nm and operating at 10% of its total power from 100 to 1500 cm<sup>-1</sup>. The

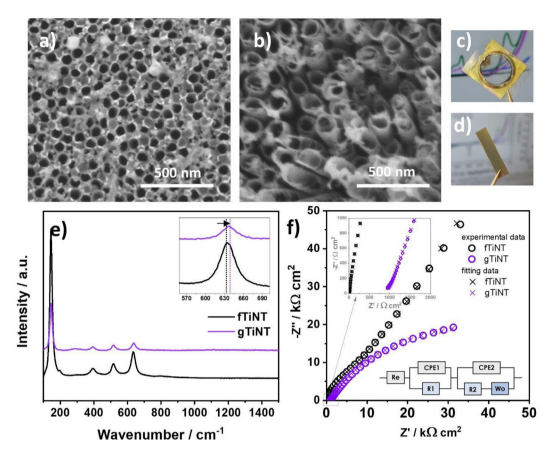


Fig. 1 SEM images of (a) gTiNT, (b) fTiNT, electrode photography of (c) gTiNT, (d) fTiNT, (e) Raman spectra for gTiNT and fTiNT, (f) Nyquist representation of impedance spectra recorded for gTiNT and fTiNT at OCP.

Raman spectra were inspected for gTiNT and fTiNT samples as well as for 5-Ag<sub>2</sub>S/gTiNT, 25-Ag<sub>2</sub>S/gTiNT, 45-Ag<sub>2</sub>S/gTiNT electrodes before electrochemical measurements and after 3 and 10 CVs cycles in 0.5 M Na<sub>2</sub>SO<sub>4</sub> and 0.5 M NaOH solutions.

Electrochemical and photoelectrochemical measurements were carried out in a 3-electrode system, where the TiO<sub>2</sub>-based electrodes were used as the working electrode, Pt mesh as a counter electrode, and Ag/AgCl/0.1 M KCl as a reference electrode. The electrolyte was deaerated with argon, and, during the measurements, argon flow was kept above the electrolyte. The photoelectrochemical measurements were carried out using an AutoLab PGStat 302N potentiostat-galvanostat. Cyclic voltammetry (CV) and linear voltammetry (LV) scans were recorded in 0.5 M Na<sub>2</sub>SO<sub>4</sub> solution in the range from -1 V to +1 V vs. Ag/AgCl/0.1 M KCl. The CVs were performed with a scan rate of 50 mV s<sup>-1</sup>, while the LVs with a scan rate of 10 mV s<sup>-1</sup>. The LV curves were recorded under chopped visible light illumination provided by a xenon lamp (LOT-Quantum Design GmbH) with a cut off filter for wavelengths below 420 nm. The light intensity was calibrated by a silicon reference cell (Rera) to provide 100 mW cm<sup>-2</sup>. The electrochemical measurements were carried out using an AutoLab 204 potentiostat-galvanostat. The CVs measurements were taken in deaerated 0.5 M NaOH solution in the range from -0.4 V to +1.1 V vs. Ag/AgCl/ 0.1 M KCl (see Fig. 7 and 8). The CVs were performed with a

scan rate of 50 mV s<sup>-1</sup>. The stability tests were carried out using both CA and CV techniques. In the case of chronoamperommetry, the current was recorded at +1.0 V (and for comparison at +0.5 V) vs. Ag/AgCl/0.1 M KCl lasted 10 000 s that is ca. 3 hours. Regarding CV, the multicyclic studies (100 cycles) were carried out for 45-Ag2S/gTiNT and the potential was changed from -0.4 to +1.1 V vs. Ag/AgCl/0.1 M KCl. Both results are shown in ESI file as Fig. S7.† The impedance spectra were recorded at OCP that equals +0.1 V vs. Ag/AgCl/0.1 M KCl, in the range from 10 Hz to 0.1 Hz with 10 mV amplitude of AC signal. The fitting procedure was carried out with EIS Analyser program<sup>22</sup> and using the proposed electric equivalent circuit (EQC) that will be discussed later on. The modified Powell algorithm was used with amplitude weighting  $r_a$ :

$$r_{\rm a}(\omega, P_1, ... P_M) = r_{\rm c}^2/(N - M)$$
 (1)

where N is the number of points, M is the number of parameters,  $\omega$  is the angular frequency,  $P_1...P_M$  are parameters. Parameter  $r_c$  is defined as:

$$r_{\rm c}^{\,2} = \sum_{i=1}^{N} \frac{(Z'_i - Z'_{i_{\rm calc}})^2 + (Z''_i - Z''_{i_{\rm calc}})^2}{Z'_i^2 + Z'_{i_{\rm calc}}^2} \tag{2}$$

where i corresponds to the measured values of impedance and  $i_{calc}$ . is attributed to the calculated values; N is the number of points.

# 3. Results

# 3.1 Characterisation of transparent and non-transparent titania nanotubes

3.1.1. Differences between TiO2 nanotubes fabricated on ITO and Ti foil. Titanium dioxide nanotubes exhibit varied morphological, structural, and electrochemical properties. Morphological alterations impact their surface area and therefore their photocatalytic efficiency. Here, electrochemical anodization processing results in formation of aligned nanotubes (see Fig. 1a and b). The outer diameter of TiO<sub>2</sub> nanotubes fabricated on ITO substrates is equal to  $46 \pm 10$  nm, whereas, in the case of TiO2 nanotubes fabricated on Ti foil the diameter reaches 124 ± 10 nm. In both cases, the electrochemical anodization process was carried out at a temperature of 23 °C, applied voltage of 40 V, duration of 1 h, and in a hydrated ammonium fluoride in ethylene glycol electrolyte. It should be noted that anodization of TiO2 nanotubes on various substrates like ITO and Ti foil vields diverse morphologies and growth mechanisms. Ti/ITO substrates promote faster growth compared to Ti foil which can be attributed to a thinner Ti layer. According to Sivaprakash et al., 23 high water content leads to an increased oxide dissolution rate and ion mobility. In our previous work<sup>24</sup> and this work, 15 vol% H<sub>2</sub>O is used for anodization of the Ti foil to obtain aligned TiO2 nanotubes. In the case of Ti on ITO, it can be too high of a concentration which leads to etching of the Ti and ITO surface. Therefore, the electrolyte for the gTiNT contained only 1 vol% H<sub>2</sub>O. As can be seen in Fig. 1c, a transparent circle-shaped area is where nanotubes are present. According to Hankova et al., 25 the anodization process of Ti sputtered on ITO carried out in 5 vol% H2O in ethylene-based electrolyte results in the formation of transparent TiO2 nanotubes. With the usage of 1 vol% H<sub>2</sub>O it was possible to obtain completely non-transparent nanostructures (Fig. 1d). The Raman spectra of the fTiNT and gTiNT are shown in Fig. 1e. Raman bands associated with the anatase phase of TiO<sub>2</sub> <sup>26</sup> are located at 145, 198, 394, 516, and 634 cm<sup>-1</sup>. The slight blue shift from 634 to 637 cm<sup>-1</sup> between the fTiNT and the gTiNT electrode can be correlated with lack of impurities.<sup>27</sup> It should be mentioned that with the foil having 99.7% purity whereas the sputter target used for Ti deposition has a purity of 99.99%, a consequence is that the obtained nanotubes can have a slightly different content of impurities. Electrochemical impedance spectroscopy was used to determine the charge transfer resistance of the fTiNT and gTiNT electrodes (Fig. 1f). The impedance spectra were recorded in 0.5 M Na<sub>2</sub>SO<sub>4</sub> solution at open circuit potential (OCP). The Re(CPE1R1)(CPE2R2Wo) equivalent circuit was proposed that is frequently used in works focused on the electrochemical studies of the titania nanotubes. 28,29 The electrolyte resistance is assigned to Re, while the subsequent EQC parts: (CPE1R1) and (CPE2R2Wo) are introduced to represent two time constants related to the inner and outer porous layer of titania. CPE instead of pure capacitance is used due to the electrode porosity represented by titanium dioxide nanotubes. The impedance of CPE is defined as  $Z = Q^{-1}(i\omega)^{-n}$ , where

energy dispersion denoted as n is considered. The value of n is between 0.54–0.86 which results from the high porosity and is the consequence of the distribution of the relaxation times.  $W_0$  stands for Warburg open element and is assigned to the finite length diffusion with reflective boundary and described by:

$$Z_{W_{\rm o}}(\omega) = \frac{\omega_{\rm or}}{\sqrt{\omega}} (1 - j) \coth[W_{\rm oc} \sqrt{j\omega}] \tag{3}$$

where  $W_{\text{or}}$  stays for Warburg coefficient,  $W_{\text{oc}} = d/D^{0.5}$ . In the case of the studied material,  $W_0$  is related to the charge transport within tubular titania down to the blocking metal oxide support. The quality of fitting was on the level of 10<sup>-4</sup> which suggests a good fit. The calculated values for each element in EQC for nanotubes formed onto Ti foil and ITO/glass substrate are collected in Table S1.† As can be seen, the most significant difference can be observed for resistance elements R1 and R2. In the case of R1, attributed to the outer porous layer, has the smallest value ( $R1 = 41 \Omega \text{ cm}^2$ ) for gTiNT. On the other hand, the value of R2, which represents the resistance of the inner layer, is much lower for fTiNT ( $R2 = 7.1 \Omega \text{ cm}^2$ ). Those differences are related to tubular geometry. According to SEM inspection, the outer diameter of gTiNT is almost 2.7 times lower than fTiNT. Thus, much more nanotubes are present on the same geometric area of glass substrate than on Ti foil and in consequence, the real surface area is much higher for gTiNT. A higher developed area gives higher space charge capacitance Q1 and lower resistance R1 that was obtained for gTiNT. The significantly higher value of resistance R2 for gTiNT, can result from the presence of some amorphous material, as was shown on Gi-XRD, at the barrier layer present between support and titania nanotubes. Regarding values of Q2, the higher one was found for gTiNT also results from the oxide film present at the bottom of nanotubes that could be more thick and porous than in the case of fTiNT.30

The fTiNT and gTiNT electrodes were examined using Gi-XRD (see Fig. 2). This technique confirmed the presence of anatase on both the surface and within the material. The results indicate a minor presence of amorphous titania (indicated in Fig. 2b) below the titanium nanotube surfaces, which is observed only in the ITO substrates and not on the metal foil. Gi-XRD results for all of the samples described in the manuscript are presented in Fig. S2.†

### 3.2. TiO<sub>2</sub> nanotubes modified by Ag<sub>2</sub>S

3.2.1 Morphology. SEM images of  $Ag_2S$ -coated  $TiO_2$  nanotubes fabricated on glass substrates are presented in Fig. 3a–f. The outer diameter of NTs for gTiNT, 5- $Ag_2S$ /gTiNT, 25- $Ag_2S$ /gTiNT, and 45- $Ag_2S$ /gTiNT is equal to 46, 41, 42, and  $42 \pm 10$  nm, respectively. For electrodes with  $Ag_2S$  deposition, the outer diameter become slightly smaller than for pure TiNTs. However, it should be noted that with an increase of SILAR cycles, the regular interior area of the nanotubes starts to disappear. This behaviour is mostly visible in the case of 45- $Ag_2S$ /gTiNT electrode (see Fig. 3f). The number of cycles SILAR is reflected in the colour changes of the electrode surface. Initially pure gTiNT are completely transparent and with an

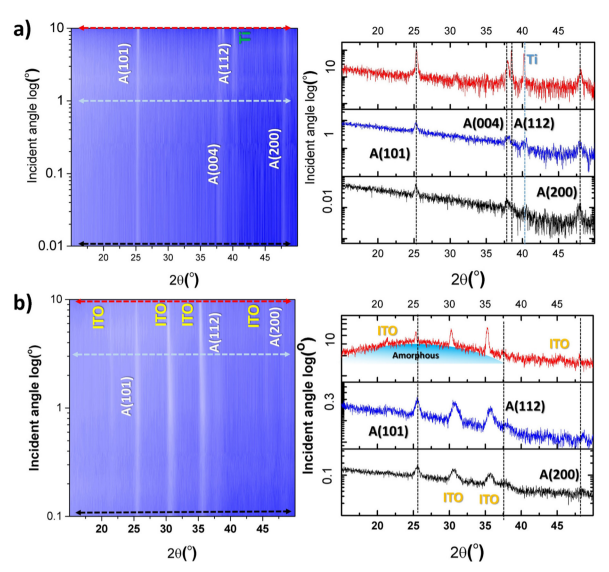


Fig. 2 Gi-XRD for (a) fTiNT and (b) gTiNT. Dashed lines show selected spectra at a given incident angle.

increase of Ag<sub>2</sub>S modification cycles the electrodes become brown colored. SEM images of 5-Ag<sub>2</sub>S/fTiNT, 25-Ag<sub>2</sub>S/fTiNT, 45-Ag<sub>2</sub>S/fTiNT are presented in Fig. S1 in ESI.†

Using EDX analysis, it was confirmed that Ag<sub>2</sub>S is deposited (see Fig. S3 and Table S2†), however due to spectral overlaps and the large interaction volume, the assignment of exact atomic percentages and distributions throughout the tubes is obscured.

3.2.2. Optical properties. Fig. 4a and b shows absorbance and reflectance spectra for silver sulfide coated titania nanotubes grown on ITO or Ti foil, respectively. The reflectance spectrum was recorded for fTiNT materials due to the opaque substrate. Both pure TiO2 NTs samples, namely gTiNT and fTiNT, exhibit absorption in the UV region, i.e. below 400 nm, with Raman spectral analysis indicating anatase as the main crystalline phase. However, for gTiNT one can observe fringes that appear from ca. 400 nm. This phenomenon can be explained by photonic nature of the material, i.e. constructive

and destructive interference of reflected light due to the nanostructured titania nanotubes and bilayer structure of the TiNTs on ITO glass making the subtraction of reflectance not feasible.24,31 Nonetheless, morphological features such as nanotube ordering as well as grain size may also affect the optical properties<sup>32</sup> as the optical response depends also on the nanotubes' geometry33 which may explain the reason behind lack of the fringes for fTiNT.34 Deposition of silver sulfide results in disappearance of the fringes for gTiNT and the increase of absorption in the visible range for both gTiNT and fTiNT substrates. This enhanced absorption is attributed to the narrow band gap (1.0 eV) and high absorption coefficient of Ag<sub>2</sub>S.35

As can be seen for fTiNT and gTiNT electrodes, light is absorbed at longer wavelengths (lower energy) than the TiO2 bandgap after coating the NTs with Ag2S, as expected for a material of smaller bandgap. In order to determine the bangap value, we taken into account the values of the expo**Paper** Nanoscale

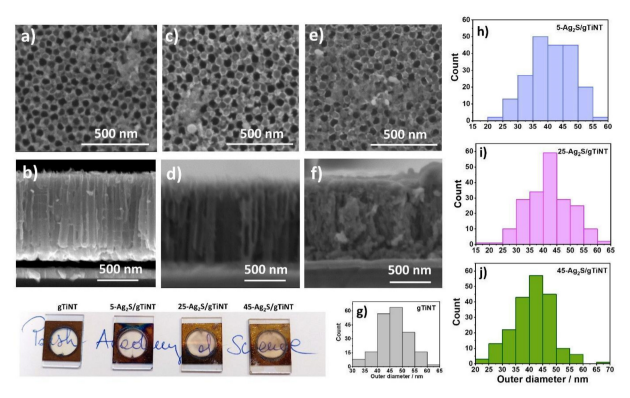


Fig. 3 SEM images of (a and b) 5-Ag<sub>2</sub>S/gTiNT, (c and d) 25-Ag<sub>2</sub>S/gTiNT, (e and f) 45-Ag<sub>2</sub>S/gTiNT, and size distribution of TiO<sub>2</sub> nanotubes outer diameter (g) gTiNT, (h) 5-Ag<sub>2</sub>S/gTiNT, (i) 25-Ag<sub>2</sub>S/gTiNT, (j) 45-Ag<sub>2</sub>S/gTiNT.

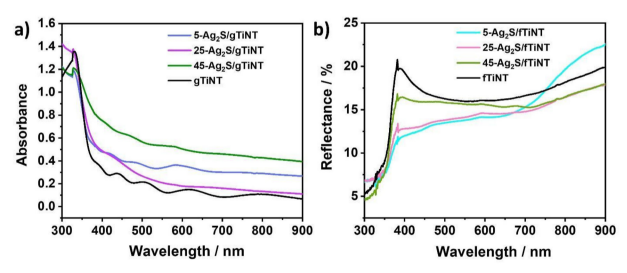


Fig. 4 UV-vis (a) absorbance spectra for Ag<sub>2</sub>S-coated and non-coated gTiNT and (b) reflectance spectra for Ag<sub>2</sub>S-coated and non-coated fTiNT.

nent in Tauc method typical for titania since amount of the TiO<sub>2</sub> nanotubes is much higher comparing to the decorating agent - silver sulphide. As can be seen, the values for fTiNT and gTiNT electrodes hardly differ (2.95 and 2.96 eV, respectively) and are narrower than the one reported for bulk anatase -3.2 eV. However, it should be taken into account that the energy bandgap of the nanomaterial depends on its geometry.36 For both types of substrates, the values of energy

bandgap are lower than for pure titania nanotubes which was expected as the NTs were covered with the material of smaller energy bandgap. For modified gTiNT, the energy bandgap values decrease with increased cycles which is in agreement with the literature data.<sup>37</sup> This may be attributed to the enhanced interfacial chemical interaction and increased coverage of titania with Ag<sub>2</sub>S.<sup>38</sup> This can also explain the fact that although 45-Ag<sub>2</sub>S/gTiNT has a lower energy bandgap than

25-Ag<sub>2</sub>S/gTiNT, the latter one exhibits better photoelectrochemical activity. Interestingly, contradictory results were obtained for modified fTiNT electrodes, namely the values of energy bandgaps increase with the cycle number. It should be in here mentioned, that apart of the changes in nanotubes diameter, the purity of Ti influences on the wall thickness and it was already reported that it rises with higher purity of layer out of which nanotubes were grown. <sup>39</sup> Similar effect was observed in this work – see Fig. 4(a) and (b).

Therefore, taking into account optical properties, better photoelectrochemical performance of  $Ag_2S$ -coated electrodes under visible light is expected compared to uncoated TiNT. As it will be shown, although 45- $Ag_2S$ /gTiNT absorbs more light than 25- $Ag_2S$ /gTiNT, it does not mean that the sample has the highest photoelectrochemical activity (Table 1).

3.2.3 Chemical structure. The chemical states of titanium Ti 2p (Fig. 5a), oxygen O 1s (b), carbon C 1s (c), silver Ag 3d (d), and sulfur S 2p (e) were determined using X-ray photoelectron spectroscopy analysis. Table S3† presents the percentage contribution and binding energy of electrodes. A detailed interpretation of all recorded spectra was provided based on the peak fitting and deconvolution procedure. The 25-Ag<sub>2</sub>S/ gTiNT and 25-Ag<sub>2</sub>S/fTiNT electrodes were selected for analysis due to their high photoelectrochemical activity as well as the 45-Ag<sub>2</sub>S/gTiNT electrode which exhibits the best activity towards oxygen evolution reaction in alkaline electrolyte. The XPS spectrum of Ti 2p is shown in Fig. 4a with all samples, both on glass and on titanium foil, exhibits a peak doublet. For the pure fTiNT sample, the binding energies are 458.48 and 464.18 eV, corresponding to the spin-orbital splitting of Ti  $2p_{3/2}$  and Ti  $2p_{1/2}$ . These values were assigned to titanium in the +4 oxidation state, indicating a bonding consistent with stoichiometric TiO2.40 Between the glass and foil samples, a slight shift towards higher energies, less than 0.3 eV for the glass sample, has been observed. This result might be correlated with shift observed in Raman spectra due to a lack of impurities (see Fig. 1e). Additionally, it was observed that spectra of samples modified with silver sulfide show slight asymmetry in the peaks. This shift can be correlated with additional modifications of the samples under the influence of UV radiation coupled with ozone generation<sup>41,42</sup> or annealing in argon atmosphere.43 The deconvolution of the O 1s spectrum (Fig. 5b) yielded three peaks at binding energies of 529.98, 531.18, and 532.58 eV. The first peak is, associated with Ti-O bonding from the TiO<sub>2</sub> crystalline lattice. 44 The next maximum corresponds to hydroxyl groups, which occur on the

 $\begin{tabular}{lll} \textbf{Table 1} & Energy & bandgap & values & estimated & for & modified & gTiNT & and \\ fTiNT & samples & & & \\ \end{tabular}$ 

Sample name	Energy bandgap (eV)	Sample name	Energy bandgap (eV)
5-Ag <sub>2</sub> S/gTiNT	2.69	5-Ag <sub>2</sub> S/fTiNT	2.42
25-Ag <sub>2</sub> S/gTiNT	2.65	25-Ag <sub>2</sub> S/fTiNT	2.59
45-Ag <sub>2</sub> S/gTiNT	1.74	45-Ag <sub>2</sub> S/fTiNT	2.9
gTiNT	2.96	fTiNT	2.95

sample's surface, 45 or oxygen vacancies and excess electron density in Ti.46 The last peak can be assigned to carboxyl groups. 47 For the C 1s spectrum shown in Fig. 5c, three peaks were assigned with binding energy values of 284.58, 286.08, and 288.38 eV. The C 1s peak at approximately 284.58 eV is typically attributed to residual C in the precursor organic compound, which has adsorbed onto the surface of the photocatalyst, specifically to C-C bonds. The subsequent peaks at 286.08 and 288.38 eV are assigned, respectively, to C-O and O-C=O bonds, specifically to carbonate species present from electrolyte. 48,49 Next, deconvolution of the Ag 3d spectrum (Fig. 5d) was performed, assigning a doublet of peaks. The first maximum at 367.38 eV is associated with 3d<sub>5/2</sub>, and the second, with a value of 373.38 eV, originating from 3d<sub>3/2</sub>. These maxima indicate the presence of the Ag<sup>+</sup> valence state in the form of Ag<sub>2</sub>S. <sup>50,51</sup> Additionally, a shift of the maximum can be observed for the sample made on the foil, but smaller than 0.3 eV. Lastly, deconvolution was performed for the S 2p spectrum (Fig. 5e). A doublet was detected, indicating sulfur bonding with only one element. In the observed spectra, two maxima were found: 160.38 and 161.58 eV, representing the spin-orbit pair 2p<sub>3/2</sub> and 2p<sub>1/2</sub> in metal sulfides, providing evidence of Ag<sub>2</sub>S 50 though with the low sulfur signal for the fTiNT samples, more quantitative analysis is not possible. As the number of SILAR cycles increases, the percentage of sulfur increases from 0.84 at% to 1.75 at%, and the percentage of silver from 1.54 at% to 2.07 at% for samples on glass substrates.

### 3.2.4 Electrochemical and photoelectrochemical activity

Electrochemical activity in 0.5 M Na<sub>2</sub>SO<sub>4</sub>. The electrochemical activity of Ag<sub>2</sub>S-coated gTiNT and fTiNT electrodes first was tested using cyclic voltammetry in 0.5 M Na<sub>2</sub>SO<sub>4</sub>. The results are presented in the Fig. 6. When the electrode is immersed in the solution, OH<sup>-</sup> groups adsorb on Ag<sub>2</sub>S. During the anodic potential scan silver oxides are formed on the surface ((A1), (A2) and (A3)) leading to the release of sulfur ions, whereas, during the reverse scan silver oxides are reduced ((C2) and (C1)):<sup>52</sup>

$$2[Ag(OH)_2]_{aq}^- \to Ag_2O + H_2O + 2OH^-$$
 (A1)

$$Ag_2O + 2OH^- \rightarrow 2AgO + H_2O + 2e^- \tag{A2}$$

$$2AgO + 2OH^{-} \rightarrow Ag_{2}O_{3} + H_{2}O + 2e^{-}$$
 (A3)

$$2AgO + H_2O + 2e^- \rightarrow Ag_2O + 2OH^- \tag{C2} \label{eq:C2}$$

$$Ag_2O + H_2O + 2e^- \rightarrow 2Ag + 2OH^-. \tag{C1} \label{eq:C1}$$

The 5-Ag<sub>2</sub>S-coated gTiNT and fTiNT electrodes are not characterized by any significant anodic or cathodic peaks from the above list of proposed reactions (see Fig. 6a and b). However, as the number of cycles increases, an anodic peak ascribed as (A1) at ca.-0.6 V vs. Ag/AgCl/0.1 M KCl assigned to oxidation of Ag to Ag<sub>2</sub>O is characterized by higher current density values. The reduction peak assigned as (C1) at ca.-0.9 V behaves analogously. Considering the (A2) and (A3) peaks, which are the most visible for the 25-Ag<sub>2</sub>S/gTiNT electrode,

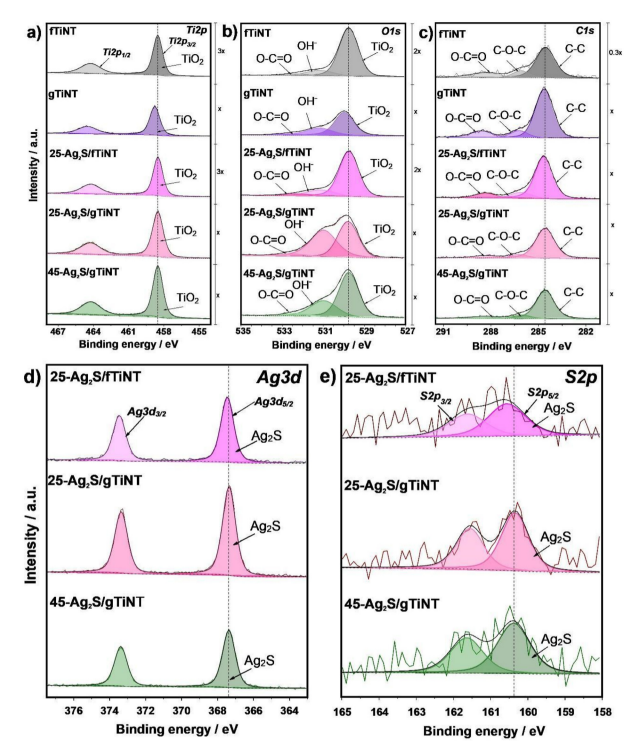


Fig. 5 XPS high resolution spectra registered for the fTiNT, gTiNT, 25-Ag<sub>2</sub>S/fTiNT, 25-Ag<sub>2</sub>S/gTiNT, 45-Ag<sub>2</sub>S/gTiNT (a) Ti 2p, (b) O 1s, (c) C 1s, (d) Ag 3d, (e) S 2p.

such peaks can appear broader or less pronounced because of differences between the diameter and aggregations of Ag NPs.<sup>53</sup> The photoelectrochemical activity was tested by linear voltammetry under chopped visible light illumination for wavelengths above 420 nm in 0.5 M Na<sub>2</sub>SO<sub>4</sub>. It can be clearly seen than activity differs between the samples on glass and

foil substrates (Fig. 6c and d). The highest photocurrent of 21  $\mu$ A cm<sup>-2</sup> registered at 0.4 V  $\nu$ s. Ag/AgCl/0.1 M KCl was obtained for the 25-Ag<sub>2</sub>S/gTiNT electrode. This activity is correlated with the presence of silver oxides in various oxidation states as Ag<sub>2</sub>O, AgO, and Ag<sub>2</sub>O<sub>3</sub>. According to Sakar *et al.*<sup>54</sup> and Allen *et al.*<sup>55</sup> Ag<sub>2</sub>O, AgO, and Ag<sub>2</sub>O<sub>3</sub> is characterized by a

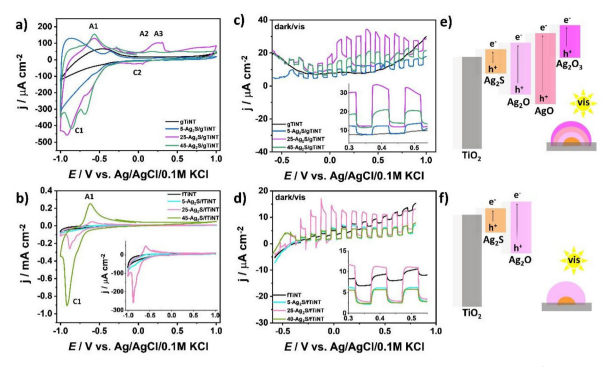


Fig. 6 Cyclic voltammetry curves for (a) 5-, 25-, 45-Ag<sub>2</sub>S/gTiNT, bare gTiNT, (b) 5-, 25-, 40-Ag<sub>2</sub>S/fTiNT, bare fTiNT (50 mV s<sup>-1</sup>) and linear voltammetry curves for (c) 5-, 25-, 45-Aq<sub>2</sub>S/qTiNT, bare qTiNT, (d) 5-, 25-, 40-Aq<sub>2</sub>S/fTiNT, bare fTiNT under visible light illumination registered in 0.5 M Na<sub>2</sub>SO<sub>4</sub> at a scan rate of 10 mV s<sup>-1</sup> and (e) energy band diagram for the 25-Aq<sub>2</sub>S/gTiNT electrode, (f) energy band diagram for the 25-Aq<sub>2</sub>S/fTiNT electrode.

bandgap energy of 1.5 eV, 2.25 eV, and 0.96 eV, respectively. Therefore, the combination of three oxides where silver(III) oxide has the smallest bandgap energy (Fig. 6e) is more favourable for photocurrent generation in contrast to presence of silver(1) oxide which has ca. 2 times higher bandgap (Fig. 6f). As can be seen, the most pronounced (A1), (A2), and (A3) peaks on CV were registered for electrode with the highest photoelectrochemical activity. In the case of 25-Ag<sub>2</sub>S/fTiNT electrode with the presence of the (A1) peak assigned to silver (1) oxide, the photocurrent density at 0.4 V is equal to 8 µA  $cm^{-2}$  which is ca. 3 times lower current than for samples coated by Ag<sub>2</sub>S in the same conditions but on gTiNT.

The 25Ag<sub>2</sub>S/gTiNT electrode was exposed to light of different wavelengths to track its photoresponse in different light conditions. As a source of light, we used a LED revolver with light power in the range of 30 to 100 mW depending on the average wavelength. Fig. S4† presents photocurrent vs. time for different wavelengths recorded at +0.5 V and +1.0 V  $\nu s$ . Ag/AgCl/0.1 M KCl. The highest photocurrent was recorded for wavelength 373 nm, followed by 405 and 424 nm. As the wavelength increased, the photocurrent value decreased in accordance with the absorbance profile. It is worth mentioning that the higher the potential, the greater the contribution of the shortest investigated wavelength, namely 373 nm, to the overall photocurrent since the material absorbs light near the UV range.

Analysis of electrode composition. In order to characterize the silver sulfides and silver oxides, Raman spectroscopy was used

to study the electrodes before and after cyclic voltammetry measurements carried out in 0.5 M Na<sub>2</sub>SO<sub>4</sub> (Fig. 7). The combination of Raman and CV is an advantageous approach for comprehensive study of silver compounds allowing monitoring of structural changes during electrochemical processes. All changes observed in Raman spectra can be correlated with the electrochemical behaviour offering insight into the mechanism and kinetics of silver containing compounds that transform upon electrochemical tests. According to Fig. 1e the 145, 394, 516, and 634 cm<sup>-1</sup> bands correspond to anatase TiO<sub>2</sub> phase. Therefore, Tables S4 and S5† present only the location of Raman bands for compounds with silver species. First, the peak located at 145 cm<sup>-1</sup> is assigned to the presence of the anatase phase that overlaps with the Ag<sub>2</sub>SO<sub>3</sub> band.<sup>56</sup> The measurements were performed in electrolyte containing SO<sub>4</sub><sup>2-</sup> ions which may be reduced to  $SO_3^{2-}$  by some reducing agent during the CV scans. As shown by XPS analysis, the 25-Ag<sub>2</sub>S/ gTiNT and 45-Ag<sub>2</sub>S/gTiNT electrodes are composed of Ag<sub>2</sub>S. Therefore, it can be assumed Ag+ can act as reducing agent for  $SO_4^{2-}$  where  $Ag_2S$  can be oxidized to  $Ag_2O$  and  $AgO.^{58}$ Subsequently non-oxidized Ag<sup>+</sup> can bind with SO<sub>3</sub><sup>2-</sup>. The Ag-S bands can be found at 205, 225, 243, 250, and 276 cm<sup>-1</sup>.<sup>57,58</sup> In the case of the 25-Ag<sub>2</sub>S/gTiNT electrode after 3 and 10 CV cycles all remain well visible on Raman spectra. For the 45-Ag<sub>2</sub>S/gTiNT electrode bands at 205 and 243 cm<sup>-1</sup> disappear after 10 cycles and a small new band at 276 cm<sup>-1</sup> developed. This result indicates that some silver sulfides are preserved on the surface of the 25-Ag<sub>2</sub>S/gTiNT electrode after 10 cycles. This

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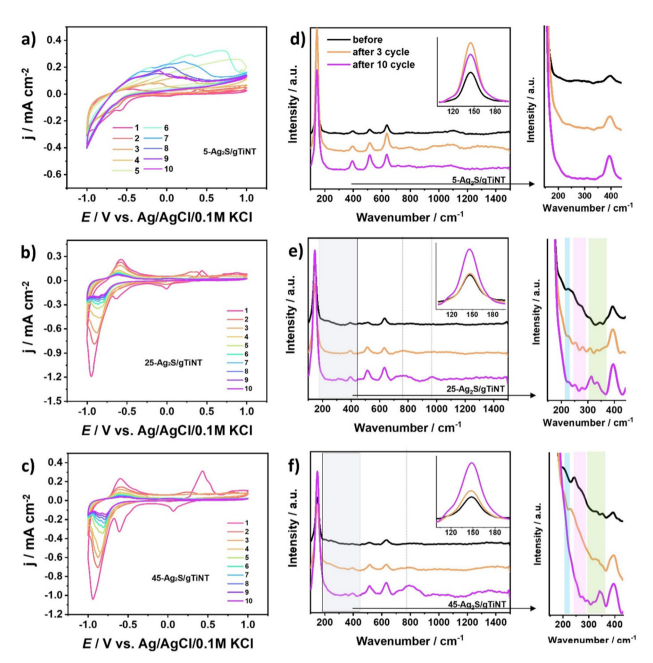


Fig. 7 Cyclic voltammetry curves of (a)  $5-Ag_2S/gTiNT$ , (b)  $25-Ag_2S/gTiNT$ , (c)  $45-Ag_2S/gTiNT$  and Raman spectra before CVs and after 3 or 10 CVs cycles for (d)  $5-Ag_2S/gTiNT$ , (e)  $25-Ag_2S/gTiNT$ , (f)  $45-Ag_2S/gTiNT$  in 0.5 M  $Na_2SO_4$  electrolyte (scan rate 50 mV s<sup>-1</sup>). All the graphs on the right are insets of the Raman graphs.

is consistent with the photoelectrochemical results where the best photoelectrochemical activity was obtained for samples with 25 SILAR deposition cycles. Furthermore, Raman bands located at 295, 310, 325, 340, and 350 cm<sup>-1</sup> are assigned to Ag–O, whereas 790 cm<sup>-1</sup> corresponds to O–Ag–O. <sup>58</sup> As CV cycles increases in number, bands on Raman spectra attributed to silver oxide becomes more and more intense.

Electrochemical activity in 0.5 M NaOH. The electrochemical activity of Ag<sub>2</sub>S-coated TiO<sub>2</sub> nanotubes fabricated on glass and foil substrates were tested by cyclic voltammetry not only in

0.5 M Na<sub>2</sub>SO<sub>4</sub> but also in 0.5 M NaOH. The results are presented in Fig. 8. The anodic peaks are associated with oxidation of Ag to Ag<sub>2</sub>O and Ag<sub>2</sub>O to AgO named (A1) and (A2), respectively. Whereas cathodic peaks correspond to reduction of AgO to Ag<sub>2</sub>O and Ag<sub>2</sub>O to Ag described as (C2) and (C1), respectively. It should be highlighted that electrodes fabricated on ITO exhibit higher currents than those obtained on Ti foil, especially towards OER. The tendency in both cases for Ag<sub>2</sub>S-coated gTiNT and fTiNT is similar, which means that as the number of SILAR cycles increases the current density value

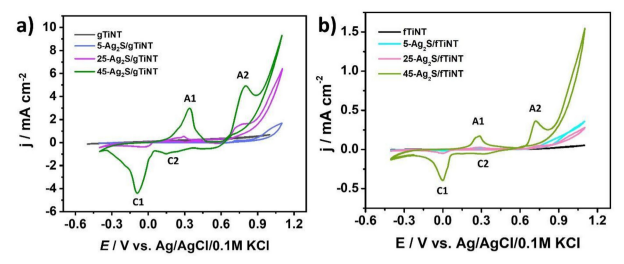


Fig. 8 Cyclic voltammetry curves for (a) Ag<sub>2</sub>S-coated and non-coated gTiNT and (b) Ag<sub>2</sub>S-coated and non-coated fTiNT in 0.5 M NaOH electrolyte (scan rate 50 mV s<sup>-1</sup>).

increases. The highest current density of 9.3 mA cm<sup>-2</sup> registered at +1.1 V vs. Ag/AgCl/0.1 M KCl was obtained for the 45-Ag<sub>2</sub>S/gTiNT electrode. It should be underlined that the 45-Ag<sub>2</sub>S/gTiNT electrode reached ca. 6 times higher current density associated with activity towards OER at +1.1 V than the 45-Ag<sub>2</sub>S/fTiNT electrode. As was already underlined by XPS analysis as well as Raman spectra, Ag<sub>2</sub>S species are present on the electrode before the immersion in the electrolyte and electrochemical measurements. However, after 10 CV cycles, new compounds as Ag<sub>2</sub>O and AgO arose on the surface. Therefore, it can be claimed that not only metal sulfides, but also metal oxides can still be responsible for the overall catalytic performance.<sup>59</sup> The first three cycles from the CV measurement for Ag<sub>2</sub>S-coated gTiNT and fTiNT are in Fig. S5.† Additionally, the results for the 45-Ag<sub>2</sub>S/gTiNT in 0.1, 0.5, and 1 M NaOH electrolytes are presented in Fig. S6.†

Inspection of electrode stability. In order to explore the activity of silver sulfides and silver oxides towards OER and their stability in 0.5 M NaOH, Raman spectroscopy was combined with cyclic voltammetry measurements (Fig. 9). The 25-Ag<sub>2</sub>S/gTiNT and 45-Ag<sub>2</sub>S/gTiNT were chosen for further analysis because of their catalytic activity. Focusing on CV, 10 curves were recorded for the 25-Ag<sub>2</sub>S/gTiNT electrode. It can be seen that anodic and cathodic peaks related to oxidation and reduction of silver oxides disappear. However, the electrode is still active towards OER. Furthermore, the current density value registered for the 25-Ag<sub>2</sub>S/gTiNT at +1.1 V vs. Ag/AgCl/0.1 M KCl for 5, 6, 7, 8, 9, and 10 cycle is the same, while between 1 and 5 cycles slight differences can be observed. The variations between current density at those potential conditions is stopped together with the absence of redox peaks originating from silver oxides. The Raman spectral peaks for the 25-Ag<sub>2</sub>S/ gTiNT located at 225, 240, 255, 270, and 280 cm<sup>-1</sup> are assigned to the Ag-S bond. 48,57 However, after 10 cycles those peaks disappear or become less pronounced but simultaneously the

peak located at 145 cm<sup>-1</sup> becomes more intense. This observation can be related with a silver lattice vibrational phase transition.<sup>60</sup> In the case of 45-Ag<sub>2</sub>S/gTiNT this peak become even more prominent which can be correlated with the higher number of SILAR cycles and thus the amount of Ag<sub>2</sub>S on the surface. As can be seen in Fig. 9b for 25-Ag<sub>2</sub>S/gTiNT, the peaks attributed to faradaic reaction occurring at the potential associated with silver oxide disappear, indicating instability of the electrode material, whereas, for 45-Ag<sub>2</sub>S/gTiNT such peaks do not change in position or intensity for numerous cycles. Therefore, it can be claimed that for 25-Ag<sub>2</sub>S/gTiNT, Ag<sub>2</sub>S is the main catalyst for OER and for 45-Ag<sub>2</sub>S/gTiNT the activity is additionally enhanced by the presence of silver oxides formed during the measurement.58 The most visible changes in the Raman spectra, namely peaks which are not observed before electrochemical measurements but appear after 10 cycles, are at 790, 980 and 1077 cm<sup>-1</sup> for O-Ag-O, S-O and Ag-O bonds. 58,61

The stability of electrochemical performance was verified by chronoamperometry measurements recorded at +1.0 V vs. Ag/ AgCl/0.1 M KCl and the multicyclic voltammetry where the potential was changed from -0.4 to +1.1 V vs. Ag/AgCl/0.1 M KCl (see Fig. S7†). As one can observe the electrode performance is stable throughout the whole CA measurement duration. Sharp peaks appearing cyclically are due to the gas bubbles that are formed at the applied potential on the electrode surface. The evolution of oxygen bubbles proceeds and in consequence appearing gas bubbles at the electrode/electrolyte interface change the contact surface area between the electrode and electrolyte. Taking into account CV measurements from cycle to cycle the redox peaks diminish and finally at the 50<sup>th</sup> cycle we do not observe any oxidation and reduction signals that were noted at the first CV run. This change occurs due to the release of sulfur ions, and oxidation of all silver as described in Table S5.†

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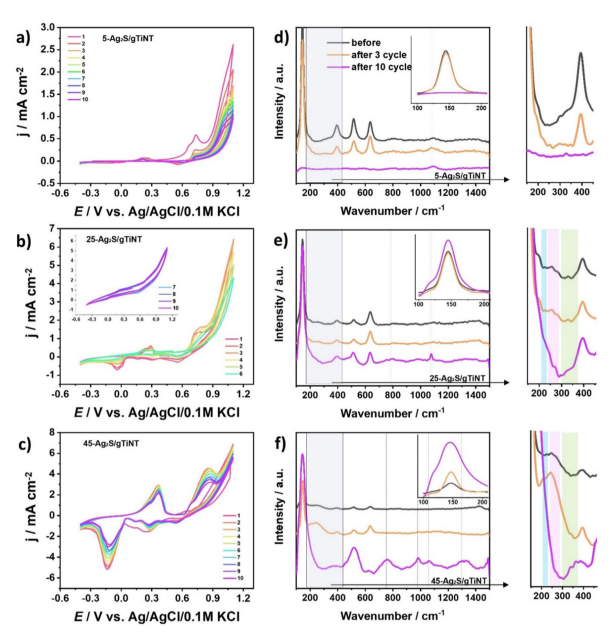


Fig. 9 Cyclic voltammetry curves of (a)  $5-Ag_2S/gTiNT$ , (b)  $25-Ag_2S/gTiNT$ , (c)  $45-Ag_2S/gTiNT$  and Raman spectra before CVs and after 3 or 10 CVs cycles for (d)  $5-Ag_2S/gTiNT$ , (e)  $25-Ag_2S/gTiNT$ , (f)  $45-Ag_2S/gTiNT$  in 0.5 M NaOH electrolyte (scan rate 50 mV s<sup>-1</sup>). All the graphs on the right are insets of the Raman graphs.

### 4. Conclusions

We showcase how adding silver sulfide to titanium dioxide nanotubes affects their shape, structure, and optical properties, as well as photoelectrochemical and electrochemical activity. This study explores these effects when the NT are grown on ITO substrates with a sputtered Ti layer or on Ti foil. The successive ionic layer adsorption and reaction method with the use of silver and sulfur precursors enables the formation of silver sulfide species inside the nanotubes. It has been shown that  $Ag_2S$ -coated transparent  $TiO_2$  nanotubes exhibit better photoelectrochemical activity under visible light

illumination and activity towards the oxygen evolution reaction than if the non-transparent material was used as a substrate. In both cases, the highest photocurrent was obtained by electrodes coated with 25 SILAR cycles while nanostructures fabricated on ITO substrates obtained *ca.* 3 times higher currents. In the case of Ag<sub>2</sub>S transparent samples, a combination of Ag<sub>2</sub>S, Ag<sub>2</sub>O, AgO, and Ag<sub>2</sub>O<sub>3</sub> compounds took part in the photoelectrochemical process. Whereas for Ag<sub>2</sub>S deposited on TiO<sub>2</sub> nanotubes and Ti foil only Ag<sub>2</sub>S and Ag<sub>2</sub>O were involved in photocurrent generation. The highest activity towards OER was obtained by transparent nanotubes coated by 45 SILAR cycles and reached of 9.3 mA cm<sup>-2</sup> at 1.1 V vs. Ag/AgCl/0.1 M

KCl which is six times higher than the non-transparent material also obtained *via* anodization of Ti foil. The significant catalytic activity was enhanced by a synergetic effect between silver sulfide and silver oxides. Considering both structural and electrochemical data, transparent TiO<sub>2</sub> nanotubes are characterised by more uniformly distributed nanotubes with smaller diameter and higher purity with lower charge transfer resistance than TiO<sub>2</sub> nanotubes on Ti foil. Ag<sub>2</sub>S deposition improved the overall electrochemical performance of transparent electrodes and warrants further exploration.

# Data availability

The data supporting this article have been included as part of the ESI.†

# Conflicts of interest

There are no conflicts to delare.

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