Label-free electrochemical sensor to investigate the effect of tocopherol on generation of superoxide ions following UV irradiation

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Abstract

Background: Generation of reactive oxygen species (ROS), triggered by ultraviolet radiation (UVR), is associated with carcinogenesis of the skin. UV irradiation induced superoxide anion (O2•−) is the key ROS involved in cellular damage. The cytoprotective efficacy of an unknown anti-oxidant compound can be evaluated by analyzing the production of O2•− from treated cells.

Methods: In this study, a glass carbon electrode functionalized with nanotube@DNA-Mn3(PO4)2 composite was applied to quantitative determination of generation of highly unstable O2•− from the melanoma A375 cell line following UVR(UV, UVA and UVB). In addition, the cytoprotective efficacy of anti-oxidant α-tocopherol was evaluated by quantifying the production of O2•−.

Results: The results showed that, UVR triggers generation of O2•− in melanoma A375 cells, and α-tocopherol is effective in diminishing the production of O2•− following UV irradiation. By comparing the conventional cell-survival assays results, we found that our simple and quick electrochemical sensing method can quantify O2•− generation through the biological activity of an anti-oxidant compound (α-tocopherol).

Conclusion: Our label-free electrochemical quantification method for ROS (O2•− major) in cells facing UVR stress demonstrates its potential application for high-throughput screening of anti-oxidation compounds.

Keywords: Skin, Electrochemical sensing, O2•−/ROS, UVR, Anti-oxidant screening

Highlighting points

- A carbon nanotube@DNA-Mn3(PO4)2 composite functionalized glass carbon electrode was applied for quantitative determination of O2•− generation from melanoma cell A375 following UV, UVA, UVB irradiation.
- Label-free Electrochemical Sensor was developed to quantify ROS produced in cells exposed to UVR.
- The anti-oxidation efficacy of tocopherol on melanoma cells towards UV, UVA and UVB were also investigated.

Background

Ultraviolet (UV) irradiation represents one of the most important environmental impacts for humans and recently became prominent because of the depletion of the atmospheric ozone layer, leads to increased UV irradiation exposure by the majority of population [1, 2]. It is well documented that UVR can stimulate the production of a series of ROS [3–5], which may cause cellular oxidative stress injury that is believed to be one of the key factors.
factors in carcinogenesis [6, 7]. For the most part, UV light from sunlight consists of three regions of wavelengths: UVC, UVB and UVA. UVC (100–290 nm) is absorbed by ozone ($O_3$) in the upper atmosphere but UVB and UVA reach to earth surface, are the major fractions linked to skin diseases. UVB (290–320 nm) is absorbed mostly by the epidermis and keratinocyte DNA, while UVA (320-400 nm) is primarily oxidative in nature and penetrates more deeply into the dermal layers of the skin [8, 9]. Since UV fractions reach different biological layers of skin and lead to skin pathology via different cellular pathways. The effects and mechanisms of action initiated by UVA and UVB have been studied extensively [10–12]. For instance, Petersen et al. reported that superoxide anion ($O_2^{-}$) production in HaCaT cells was probably linked to DNA damage induced by UVA [13]. UVB can also induce the formation of ROS, leading to cellular damage [14–16]. Among the many ROS species that have been studied, $O_2^{-}$ is one of the principal radical species [13, 17–19]. It is generated as a reduced intermediate of molecular oxygen in a variety of biological systems. It can easily form hydroxyl radical ($HO^-$) in the presence of transition metal ions such as $Fe^{2+}$ and $Cu^{2+}$ [20]. In addition, the reaction between $O_2^{-}$ and nitric oxide (‘NO) leads to the formation of highly reactive peroxynitrite (ONOO-) in the pathogenesis of atherosclerosis and neurodegenerative diseases [21]. As a consequence, efforts have been made to investigate generation of $O_2^{-}$ induced by UVR. Particularly, the cytoprotective efficacy of an unknown anti-oxidant compound can be evaluated by analyzing the production of $O_2^{-}$ from treated cells.

A survey of the literature shows that the main techniques for measuring $O_2^{-}$ are based on probe-labelling assays. Intracellular fluorescent histochemistry [22], flow cytometry [16] and spectrofluorometric analyses [23, 24] are the most used approaches to characterize ROS such as $O_2^{-}$ by using fluorescent dyes such as 2',7'-dichlorofluorescein diacetate (DCFH-DA), hydroethidine and dihydrorhodamine [1, 23]. The production of extracellular $O_2^{-}$ can also be measured by using the ability of $O_2^{-}$ to reduce ferricytochrome C that was added to the cell suspension [25]. Another advanced technique is electron spin resonance (ESR)-spin trapping, was applied for determination of $O_2^{-}$ generated by UV-irradiated skin cells [10, 24]. Apart from the expensive equipment and complicated assay procedures, these probe-labelling approaches are time-consuming, difficult to automate and highly prone to interference. The short lifetime of free radicals, such as $O_2^{-}$, particularly demands fast response of the analytical tool to the changes in concentration to obtain sufficient signal-to-noise ratios [20, 26].

Electrochemical biosensors have become promising candidates for real-time analysis of free radicals, since they provide the advantages of rather simpler equipment and operation protocols. Li et al. [20] found that an electrochemical biosensor can sense $O_2^{-}$ released from cancer cells, using potassium-doped multi-walled carbon nanotubes (KMWNTs)-1-butyl-3-methylimidazoliumhexa-fluorophosphate ionic liquid composite gels [20]. While hydrogen peroxide ($H_2O_2$) can directly be measured in cells growing in a 3D matrix environment [27]. These studies indicate that, with intimately coupled biological recognition elements and electrochemical transduction units, an electrochemical biosensor exhibits great potential for facilitating understanding of biological process. Also, the use of a small-volume sample allows expensive reagents, particular for rare clinical biopsy samples, to be conserved and makes the analysis more cost-effective.

In this study, we have designed and developed a label-free electrochemical sensor that can measure the generation of $O_2^{-}$ from UVR exposed melanoma A375 cells, though other ROS species e.g. $H_2O_2$ has not been totally excluded. To demonstrate the electrochemical analytical power of this sensor, the protective effects of model anti-oxidant ($\alpha$-tocopherol) was studied in melanoma A375 cells. The specific objectives to be addressed were [1] to measure generation of $O_2^{-}$ from melanoma A375 cells following exposure to UVR alone i.e., UV, UVA and UVB; [2] to quantify the production of $O_2^{-}$ from cells pre-treated with $\alpha$-tocopherol following UVR irradiation; [3] to compare the data from electrochemical measurement, the cell-survival assay and conventional ROS fluorescence staining therefore establish the potential use of the label-free electrochemical method for high-throughput screening of antioxidants.

**Methods**

**Materials**

The human melanoma cell A375 (purchased from ATCC) were maintained in RPMI 1640 medium (Gibco®) supplemented with 10% foetal calf serum (FCS, Gibco®) with 100 U mL-1 penicillin and 100 U mL-1 streptomycin at 37 °C in a humidified 5%CO2 incubator. $\alpha$-Tocopherol was purchased from Sigma-Aldrich. 3-[4,5-Dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide (MTT), 2-(4-amidino-phenyl)-6-indole carbamidine dihydrochloride (DAPI), 2',7'-dichlorodihydrofluorescein diacetate (DCFH-DA) and superoxide dismutase (SOD) were purchased from Beyotime Biotechnology (Beijing, China). All other chemicals were purchased from Sigma-Aldrich and were used without further purification, unless otherwise indicated. All solutions were prepared with de-ionized (DI) water produced by a PURELAB flex system (ELGA Corporation).
**Apparatus**

**UV irradiation**

Cells were irradiated with a broad-spectrum UV lamp (MUA-165, Japan). The proportion of UVA and UVB in delivered UV light from this lamp is around 20:1, a ratio close to solar light at mid-day [28]. The lamp exposure time was calculated using an ultraviolet radiometer (Photoelectric Instrument Factory, Beijing Normal University, China). Appropriate glass filters to block UVA, UVB were used respectively to provide fractions of UVA and UVB in the following experiments, according to previous studies [29–33].

**Electrochemical detection setup**

The experimental setup is shown in Fig. 1. A three-electrode system was employed, with a nano material-functionalised glass carbon electrode (GCE) as a working electrode (WE), an Hg/HgCl₂/KCl electrode as a reference electrode (RE) and a platinum wire as a counter-electrode (CE). To avoid interference from direct UV irradiation of the electrodes, the electrodes were assembled on the side-wall of the detection well. In addition, a piece of aluminium foil with a hole was placed on top of the detection well to confine the UV light to impacting only on the cells. Electrochemical detection was carried out with an electrochemical station (CHI 760e, Chen Hua Instruments Co. Ltd., China).

**Electrochemical detection of O₂⁻ released by the cells**

**Preparation of an O₂⁻ electrochemical sensor**

The construction of electrochemical sensor for O₂⁻ detection was detailed in our recent paper [34, 35]. In brief, single stranded DNA (2.1 mg) was added to MnSO₄ (1.0 mL, 0.10 M) under constant stirring at 60° Cstirring until the solution became transparent. The pellet of DNA-Mn₃(PO₄)₂ composite was recovered by centrifugation. To prepare the O₂⁻ sensing electrode, carbon nanotube (CNT, 0.5 mg mL⁻¹) and DNA- DNA-Mn₃(PO₄)₂ (0.20 M) were drop-casted on a polished glass-carbon electrode. To detect O₂⁻, electrodes modified with CNT@DNA-Mn₃(PO₄)₂ composite were exposed to aliquots of KO₂ and cyclic voltammetry curves were recorded. The peak current value at 0.70 V was recorded and was plotted against the concentration of KO₂ to draw a current-dose calibration curve.

**In situ detection of production of O₂⁻ from UV-irradiated cells**

Confluent A375 cells were seeded into 24 cell plates, at a density of 1 × 10⁵ per well, and were incubated for 6 h. Then α-tocopherol (25 μM was chosen due to optimize concentration [33]) was added into cells and incubated for another 12 h. The cell-culture disc with adherent cells was transferred to the electrochemical detection cell (Fig. 1), to which serum free medium (2.5 mL) was added. Then the cells were exposed to UVR and the electrochemical signal arising from O₂⁻ were recorded by CHI. Cells without anti-oxidant were measured with the same schedule as controls.

**Fluorescent staining of oxygen radicals**

**Intracellular ROS**

A375 cells were collected and seeded at a density of 1 × 10⁵ per well in 24-well plates and were incubated overnight with or without α-tocopherol. Intracellular
generation of ROS upon UV irradiation was determined with a DCFH-DA (10 mM) assay kit (Beyotime Inc. S-0033). In brief, following UVA irradiation, cells are washed twice in phosphate buffer saline (PBS) and allowed to incubate in 10 μM/L DCFH-DA (1:1000 dilution in PBS) at 37 °C for 20 min according to the manufacturer’s instructions (Beyotime Inc.). The excess DCFH-DA was rinsed off with phosphate buffer saline (PBS) (1.0 mM). The cells were then observed and imaged under a fluorescence microscope (IX-71, Olympus Corp., Tokyo, Japan).

**O2**\(^{-}\)-colorimetric assay kit**

Superoxide assay kit (S-0060 Beyotime Biotechnology Beijing Inc.) was conducted following the product instructions. In brief, A375 cells were seeded (1 × 10\(^4\)) in 96-well plates and 80–90% confluence when treated. Now the cells were washed twice in PBS, and allowed to incubate in 10 μmol/L DCFH-DA (1:1000 dilution) at 37 °C for 3 min (Beyotime Biotechnology, Beijing Inc.). The excess DCFH-DA was rinsed off with PBS (1.0 mM). Now the cells were irradiated with respective UVR (UV, UVA or UVB) stimulus. Prepare the three wells without irradiation should be used as a blank control. Select 2 wells to add 2% SOD (Beyotime Biotechnology, Beijing Inc.) in the wells of the stimulus to verify the entire assay system. The cells were then observed and imaged under a fluorescence microscope (IX-71, Olympus Corp., Tokyo, Japan). The absorbance values taken at 450 nm were compared with the reference absorbance values at 630 nm following to manufacturer’s instructions. These data were expressed as the percentage of increase of absorbance value compared to sham controls (sham-irradiated groups).

**Cell viability assay**

The protective effects of α-tocopherol on UV-irradiated melanoma cell A375 were evaluated using the MTT assay. In brief, cells were seeded at a density of 1 × 10\(^4\) per well in 96-well plates and incubated with or without the α-tocopherol for 12 h. Following challenge with UVR, the cells were cultured in conditional culture medium for a further 24 h. Each well then received 10 μL of MTT solution (0.5 mg mL\(^{-1}\)) and were incubated for 3–4 h at 37 °C. Then 100μLlysisbuffer (mixture of 2-propanol, Triton-100 and hydrochloric acid) was added to each well to dissolve the purple-coloured formazan produced by viable cells, before colorimetric analysis. The absorbance at 570 nm was measured by a microplate reader and the results were expressed as the percentage of surviving cells compared to the sham-irradiated vehicle control group.

**Statistical analysis**

Results are expressed as means±standard error of the mean (SEM). The data were analysed by Student’s t-test using Origin Statistic software (Origin Lab Corporation, USA). P<0.05 was considered significant. All experiments were performed three independent times in triplicates.

**Results and discussion**

**Quantification of UV-induced O2**\(^{•−}\) in melanoma cells by label-free electrochemical sensor**

To evaluate the feasibility of using an electrochemical method to quantify production of O2**\(^{•−}\)** induced by UV irradiation from melanoma cells, an electrochemical sensor was fabricated and characterised. Initially, cyclic voltammetry (CV) was performed with PBS to characterize the CNT@DNA-Mn3(PO4)\(_2\) functionalised electrochemical sensor. KO2 was used as the source of O2**\(^{•−}\)** as reported in literature [19]. The peak current of the oxidative state/stress at 0.7 V increases when KO2 was added to PBS (blue line inset of Fig. 2a). The production of superoxide anions (O2**\(^{•−}\)**) is linked with various oxidative stress mediated pathologies. The electrochemical determination of superoxide ion is solely depending on SOD enzyme, hence check it’s reproducibility as well as not a cost effective, and advancement in nanosciences solved certain limitations with this procedure. Previously we demonstrated that Mn3(PO4)\(_2\) nanosheets which are a biomimetic enzyme were template-synthesized with DNA and further assembled on carbon nanotubes (CNTs) that form nanocomposite sheets (DNA-Mn3(PO4)\(_2\)-CNT) [34].

The dismutation of superoxide ion was catalyzed by Mn3(PO4)\(_2\) sheets, but the CNTs allows quick shifts of electrons to maximize its sensitivity, specificity and reproducibility for O2**\(^{•−}\)** detection [34].

The increased peak current gradually decreased to PBS level after the introduction of SOD to decompose/neutralize O2**\(^{•−}\)** (red line inset of Fig. 2a). Different concentrations of KO2 were added to vary the amount of O2**\(^{•−}\)** and the peak current at 0.7 V was recorded that was plotted against the concentration of O2**\(^{•−}\)**. The corresponding calibration curve of current response vs. O2**\(^{•−}\)** concentration showed a sensitivity of 70 nA nM\(^{-1}\) (Fig. 2b). The CV curves showed that, the change in current (μA) is associated with changing in the production of O2**\(^{•−}\)** at different UV treatments (UV, UVA, UVB) and in sham-irradiated control (Fig. 2c). Furthermore, to verify the electrochemical-sensing efficacy, the O2**\(^{•−}\)** assay kit was used to measure O2**\(^{•−}\)** generated from melanoma cells challenged with UVR; a similar increase was observed in both methods. Incubation of cells with SOD eliminates increased O2**\(^{•−}\)** in...
the culture medium, while the electrochemical sensing measures a similar pattern of generation of UVR-induced \( \text{O}_2^{-} \) as did conventional fluorescent staining for \( \text{O}_2^{-} \) (Fig. 2d), the former with less consumption of sample and shorter experimental time. These suggested that the electrochemical sensor provides a possibility of recording production of \( \text{O}_2^{-} \) in groups of A-375 cells facing different doses of UVR, without disturbing the cellular metabolism.

**Measurement of UV-irradiation induced \( \text{O}_2^{-} \) and its diminution with \( \alpha \)-tocopherol by using label-free electrochemical sensor**

To demonstrate the feasibility of electrochemical sensor as an analytical tool to evaluate the anti-oxidative capability of compounds, we pre-treated UVR-challenged cells with an antioxidant compound (\( \alpha \)-tocopherol). The electrochemical sensor was used for direct measurement of \( \text{O}_2^{-} \) generated from A375 cells pre-treated with \( \alpha \)-tocopherol followed UV irradiation. In addition free-radical-scavenging effects of the antioxidant \( \alpha \)-tocopherol in A375 cells given UVA, UVB and bulk UV were also investigated with the same electrochemical sensor. For this purpose, the normalized current change at 0.7 V was calculated in sham-irradiated cells as a control by using following equation.

\[
\Delta \text{Current} = \left[ \frac{(C_i - C_0)}{C_0} \right] \times 100\%
\]

Where \( C_i \) is current value measured upon UV irradiation, \( C_0 \) is current value of sham control (sham-irradiated).

The \( \Delta \text{Current} \% \) was plotted against irradiation energy dose of UV fractions. A higher change in current indicated a higher generation of \( \text{O}_2^{-} \) when compared with sham-irradiated control. The changes of production of \( \text{O}_2^{-} \) in melanoma A375 cells irradiated with UVs (UV, UVA, UVB) was comparatively measured by the electrochemical sensor (Fig. 3a-c). Different UV doses (0–105 kJ m\(^{-2} \)) induced the production of \( \text{O}_2^{-} \), that altered the currents by 13.1% (50 kJ m\(^{-2} \)) to that of the sham. Tocopherol (25 \( \mu \)M) treatment to UV-challenged cells reduces the production of \( \text{O}_2^{-} \) to 8.2% (50.25 kJ m\(^{-2} \)) when compared to UVs-irradiation only (Fig. 3a). Similarly, pre-treatment of cells with
tocopherol efficiently decreased (8.5%) the production of $O_2^{-}$ induced by a UVA (100 kJ m$^{-2}$) dose to 1.6% compared to sham (Fig. 3b). In contrast, pre-treatment of cells with α-tocopherol followed by UVB irradiation, showed little effects on the production of $O_2^{-}$ (Fig. 3c).

The electrochemical sensor is not only able to quantify the generation of $O_2^{-}$ at acute phase following UV-irradiation, but also enabled quantification at post-UVR challenge period. After 60 min. Following UVR, in melanoma A375 cells $O_2^{-}$ generated current was decreased to half (8.5%) than at zero time point (17%), while at 120 min. The change in electric current faded to the background level (Fig. 3d).

The effect of UV on $O_2^{-}$ production are not the sum of the total effects of UVA and UVB, the production of $O_2^{-}$ from cells challenged by various UV doses (0-105 kJ m$^{-2}$, Fig. 3a) is lower than the sum of changes induced by UVA (Fig. 3b) plus UVB (Fig. 3c). Next, we found that α-tocopherol plays a distinct role and lessen the generation of $O_2^{-}$ in pre-treated A375 cells followed by UV irradiation, however, UVA and UVB showed different capacities to trigger the generation of $O_2^{-}$ in melanoma A-375 cells.

The pre-treatment of α-tocopherol reduced the UVA and UVB induced $O_2^{-}$ generation by almost 85% and 20%, respectively (Fig. 3b,c). This difference may due to the generation of $H_2O_2$, or interference of some other unknown reason yet, hence need further optimisation of the method. Furthermore, antioxidant tocopherol showed different behaviour against UVA and UVB induced $O_2^{-}$ is due to the difference in wavelength energy. In addition to variable UV wavelength effects, antioxidants are distinct chemical units with different modes of action for their effects and possess a unique biochemical profile hence making it different from other related compounds. Based on the results of electrochemical sensor, we speculated that α-tocopherol showed different protective properties in melanoma cells exposed to different UV wavelengths. In general, the electrochemical sensor is able to characterize the $O_2^{-}$ production, as cellular oxidative status is one of the indicators.

**Prediction of protective effect of α-tocopherol on cell survival through electrochemical sensing**

To analyse the effect of α-tocopherol on cell survival following UVR, A375 cells were exposed to different doses of UV, UVA and UVB, and the anti-oxidant effect of α-tocopherol was assessed by calculating the cell survival rate by using following equation.

$$\text{Cell survival} = \left[ \frac{A_T - A_c}{A_c} \right] \times 100\%$$  \hspace{1cm} (2)

Where $A_T$ is the absorbance value of UV-irradiated cells and $A_c$ is absorbance value of sham-irradiated cells. This calculation, gives survival of cells under UV irradiation is normalized to the sham or control cells. The α-tocopherol had significant cytoprotective effects on melanoma A375 cells, and it enhanced cell survival (18%) following UV light exposure (50.25 kJ m$^{-2} = $ UVA 50 kJ m$^{-2} + $ UVB 0.25 kJ m$^{-2}$).
25 kJ m$^{-2}$) (Fig. 4a). During studies of UVA-induced cell damage, α-tocopherol appeared to be more cytoprotective as indicated by the survival rates of pre-treated (92.3%) compared with control melanoma A375 cells (80%), with 10% marked and significant difference in reduction of cell survival (Fig. 4b). On the other hand, UVB induced cell viability loss (72%) was significantly repaired by α-tocopherol pre-treatment (84%) compared with control, with 12% of protection revival (Fig. 4c).

Collectively, MTT cell-survival assay showed that α-tocopherol is effective in diminishing UV, UVA and UVB-induced damage in melanoma A375 cells, this is similar to results given by electrochemical sensing and colorimetric assay for O$_2^•$− levels (Fig. 3a-e). Interpreting the survival rate (Fig. 4a-c) of cells pre-treated with α-tocopherol followed by UV irradiation caused the generation of O$_2^•$− (Fig. 3a-c) may efficiently harbour the oxidative stress and lessen it promptly. The different cytoprotective efficacy of α-tocopherol against UVA- or UVB-induced cell death was decreased, thus could be specified as the distinct capability of α-tocopherol to eliminate UV fraction induced oxidative stress, since UVA is the major oxidative stress causing wavelength in UV.

### Comparison of methodologies towards UV-induced cellular oxidative stress

The MTT assay was conducted to evaluate the loss of viability induced by UV irradiation. It monitors the overall cell growth over a period of time, thus showing its strength in evaluating the long-term effect of a treatment on cells [34, 35]. A large body of work, spanning more than four decades, demonstrates fluorescence probe labelling for characterisation of changes of intracellular ROS [36, 37]. However, it is also clear that this probe-labelling method measures transient fluxes of oxidants only with great difficulty [38, 39]. Most importantly, the labelling of cells challenges the possibility to characterize the changes in situ of cells during a time period in responding to a treatment. The merits of label-free

### Table 1 Comparison of methods used to study effect of UV irradiation induced oxidative stress

|                  | MTT assay | Fluorescent assay | Electrochemical detection |
|------------------|-----------|-------------------|---------------------------|
| Labeling         | Yes       | Yes               | No                        |
| Real time        | NO        | NO                | Yes                       |
| Sample consumption | Large    | Large             | Small                     |
| Assay time       | 24–72 h   | 1–2 h             | Within minutes            |
| Overall comparison | Alive cell catalyse enzyme to indirectly characterize cell growth. Evaluate the overall effect of treatment on cell survival, growth. | Fluorophore probe is used to label target molecules. Not feasibly for real-time characterizing a reaction. Staining process will affect the health of cells. | In site characterize a reaction. No damage to cells. Fast reaction time allows for high throughput screening. No expensive equipment is involved. |
electrochemical sensing have been discussed in the literature [27, 34, 40] (Table 1). To confirm the feasibility of electrochemical sensing as an analytical tool to determine the anti-oxidative capability of unknown compounds, we performed conventional $O_2\cdot^−$ fluorescent staining with a standard DCFH-DA fluorescent staining protocol that measures intracellular ROS from cells after exposure to UV light. It is clear from the results that UV irradiation can induce an increase of the intensity of the signal for green fluorescence. Figure 5a presents the quantitative changes of ROS ($O_2\cdot^−$) measured by fluorescent staining of three independent methods. The fluorescence signal is gradually increased in response to increasing UV dose and the maximum signal is given by cells irradiated with $10^5$ kJ m$^{-2}$ UV light. Increases of intracellular ROS following UV irradiation visualized in this study are well in line with previously reported studies [17, 23] and our present electrochemical sensing results. Further increase of UV energy ($21$ J cm$^{-2} = 210$ kJ m$^{-2}$) does not give further increased ROS fluorescence signal, perhaps due to significant cell death (Data no shown).

The protective efficacy of anti-oxidant α-tocopherol was assessed and much attention was paid to the UV dose of $10^5$ kJ m$^{-2}$, i.e. [UVA ($10$ J cm$^{-2}$) + UVB ($0.5$ J cm$^{-2}$)]. The $O_2\cdot^−$ assay kit was used to measure the colorimetric changes induced by $O_2\cdot^−$ in a 96-well plate experimental setting. The results showed that the anti-oxidant can reduce the generation of $O_2\cdot^−$ induced by UV in A375 cells. Different anti-oxidation efficacies of the model anti-oxidant are observed in melanoma A375 cells against UVA and UVB exposure (Fig. 5b). It is encouraging to see that standard labelling techniques and electrochemical quantification depict a response pattern of UV irradiation and the anti-oxidant effects of α-tocopherol which are well correlated. In addition, electrochemical sensing revealed the oxidation status, which can be used to predict UV-induced ROS efflux and ultimately cell damage. The fast and easy experimental procedures permitted us to test the efficacy of anti-oxidant α-tocopherol against UVA and UVB in melanoma cells [33, 40]. We demonstrate the potential of electrochemical methods for high throughput screening of potential anti-oxidant compounds, although further improvements in the signal-to-noise ratio are needed. We believe that electrochemical sensing would be a potent and economical candidate to provide meaningful complementary information for standard fluorescent staining experiments.

Conclusions
The results showed, in melanoma A375 cells electrochemical sensing both UVA and UVB-induced production of ROS e.g. $O_2\cdot^−$. The potential use of in UV (UVA, UVB) involved biological research is enhanced, since the electrochemical sensor quantified $O_2\cdot^−$ levels in medium could be correlated with the cell survival ability under the protection of anti-oxidant α-tocopherol as well. The electrochemical changes occurred with $O_2\cdot^−$ production, can be seen with this label-free method correlated with intracellular fluorescent staining as well. It highlighted the potential of the electrochemical method for high-throughput screening of anti-oxidant under UV irradiation, and for investigation of their related mechanisms.

Abbreviations
DCFH-DA: 2′,7′-dichlorodihydrofluorescein diacetate; H$_2$O$_2$: Hydrogen peroxide; kJ m$^{-2}$: Kilo Jouls per square meter; MTT: 3-[4,5-Dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide; $O_2\cdot^−$: Superoxide anion; ROS: Reactive oxygen species; SOD: Superoxide dismutase; UVA: Ultraviolet A; UVB: Ultraviolet B; UVR: Ultraviolet radiation.
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Availability of data and materials
No additional data is linked with these findings. Moreover, in any query please contact author for data requests.

Authors’ contributions
LG and CB performed the experiments and most of the lab work. MFN, SC, YL, CML, and JLZ designed the study and experimental procedures and statistical analyses. PJ, EH, MFN and JLZ drafted this study to a final form. All authors read and approved the final manuscript.

Ethics approval and consent to participate
Not applicable.

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All the authors mentioned in the author section, agreed to submit this paper for publication in Journal of Biological Engineering, and have no dispute for this submission and publication. None of the authors declare any conflict of interest.

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