Two-electron photo-oxidation of betanin on titanium dioxide and potential for improved dye-sensitized solar energy conversion

Fritz J. Knorr, Deborah J. Malamen, Jeanne L. McHale, Arianna Marchioro, and Jacques-E. Moser

Department of Chemistry, Washington State University Box 644630 Pullman, WA 99164-4630 USA

Photochemical Dynamics Group, Institute of Chemical Sciences and Engineering École Polytechnique Fédérale de Lausanne CH-1015 Lausanne, Switzerland

ABSTRACT

The plant pigment betanin is investigated as a dye-sensitizer on TiO$_2$ with regard to its potential to undergo two-electron oxidation following one-photon excitation. Electrochemical, spectroelectrochemical and transient absorption measurements provide evidence for two-electron proton-coupled photo-oxidation leading to a quinone methide intermediate which rearranges to 2-decarboxy-2,3-dehydrobetanin. Time-resolved spectroscopy measurements of betanin on nanocrystalline TiO$_2$ and ZrO$_2$ films were performed on femtosecond and nanosecond time-scales and provide evidence for transient species with absorption bands in the blue and the red. The results shed light on previous reports of high quantum efficiencies for electron injection and point the way to improved solar conversion efficiency of organic dye-sensitized solar cells.

Keywords: TiO$_2$, betanin, interfacial electron transfer, solar energy

1. INTRODUCTION

Dye-sensitized solar energy conversion based on earth abundant metal oxides, especially TiO$_2$, is potentially an economical and environmentally friendly means to convert sunlight into electricity. However, reliance on synthetic organic and metalorganic dyes as sensitizers offsets these advantages, especially in the case of widely used ruthenium containing dyes. The advantageous long term stability of Ru-based sensitizers derives from the facile one-electron oxidation and regeneration of the Ru$^{2+}$ center. All-organic molecules, on the other hand, are prone to proton-coupled two-electron oxidation which may lead to undesirable bond-breaking which hinders regeneration. In this work, we investigate a plant pigment, betanin, which evidences enhanced quantum efficiency for converting photons to electrons as a result of two-electron oxidation at the aromatic hydroxyl group.

We reported the first betanin-based dye-sensitized solar cell (DSSC) in 2008, and more recently we presented evidence for incident photon-to-current conversion efficiencies (IPCEs) which at their maxima exceed 100% after correction for reflection/absorption losses of the conductive glass. The betalain family of plant pigments, which includes betanin from red beet root and cactus pears, has attracted the interest of several groups, who in agreement with our work find fairly high IPCE, good photocurrents, but low overall energy conversion efficiency as a result of low photovoltages. For example, our best betanin-based DSSCs give photocurrents as high as 14 mA/cm$^2$, energy conversion efficiencies of 2.7%, and uncorrected IPCE maxima near 100%. However, the open-circuit voltage $V_{OC}$ is typically less than 0.4 V, much less than the thermodynamic maximum of about 1 V. Since the maximum voltage is limited by recombination, and IPCE maxima near 100% would preclude recombination if one photon leads to one injected electron,
we hypothesize that betanin undergoes two-electron oxidation from its excited electronic state when adsorbed on nanocrystalline TiO$_2$.

The food science community has had a long-standing interest in anti-oxidant activity of betalains, and it is known that this activity is correlated to the presence of phenolic hydroxyl groups. Wybraniec et al. have studied the oxidation of betanin (Bt) and its aglycone betanidin (Bd). They report that in the pH range 3 to 5, electrochemical oxidation of Bd is a reversible two-electron, one-proton process, while at somewhat higher pH this changes to a two-electron, two-proton process. Oxidation of Bt by the radical anion of ABTS, 2,2'-azinobis(3-ethylbenzthiazoline-6-sulfonic acid), was reported to result in the two-electron oxidation product shown in Fig. 1, a quinone methide (Qm), which rearranges to 2-decarboxy-2,3-dehydrobetanin (dhBt). The semiquinone radical of Bt, the one-electron oxidation product, was presumed to be an intermediate. The enzymatic oxidation of Bt, on the other hand, was observed to take place by a single-step, two-electron process that also resulted in the formation of dhBt, a stable yellow compound with wavelength of maximum absorption $\lambda_{\text{max}}$ at 445 nm. Thus there is evidence that Bt is capable of single-step, two-electron oxidation in aqueous media, but the mechanism of photo-oxidation of the adsorbed molecule on TiO$_2$ has not been investigated until now.

In this work, we pursue the electrochemistry of Bt in aqueous solution as a function of pH, spectroelectrochemistry of Bt on a nanocrystalline TiO$_2$ electrode in contact with nonaqueous electrolyte, and transient absorption spectra of Bt on TiO$_2$ on femtosecond and nanosecond time scales. The results point the way to improved efficiency of solar energy conversion using sustainable plant-based sensitizers.

2. EXPERIMENTAL

Betanin was extracted from red beet root and subjected to medium-pressure liquid chromatography (MPLC) as described in Ref. 4, except that a masticating juicer rather than extraction was used to obtain the beet juice. The purple fraction from MPLC was freeze-dried and the resulting powder subjected to recrystallization from 10% methanol, 0.1% HCl in water at −10 °C. HPLC confirmed that the resulting fuschia fraction was composed of the pigments betanin and its C-15 epimer isobetanin.

The voltammetric results were measured on a 0.1 mM solution of betanin in degassed 0.2 M NaClO$_4$ supporting electrolyte. The pH was adjusted with HCl. Digital pulsed voltammetry (DPV) was measured with a BAS 100A electrochemical analyzer. The working electrode was a 3 mm glassy carbon electrode. The working electrode was polished with 1 µm alumina between measurements. The reference electrode was Ag/AgCl and the counter electrode was Pt wire. The scan rate was 4 mV/s, pulse amplitude 50 mV, pulse width 50 ms, and sample width 17 ms.
Mesoporous TiO$_2$ on TEC 15 conductive glass was prepared with the doctor blade method from Solaronix T paste. Similar films have been measured to be 4 µm thick using the same technique. The film was sensitized from a 0.1 mM solution of betanin in water at pH 2.8. The film was dried and immersed in acetonitrile with 0.2 M NaClO$_2$ electrolyte. The reference electrode was silver wire. The potential of the silver wire was measured against ferricene/ferricinium after the spectrometric measurement, and the reported potentials adjusted to Ag/AgCl reference. The spectra were measured with a Shimadzu 2501 spectrophotometer.

Transmission-mode nanosecond transient absorption spectroscopy experiments were conducted using a frequency-tripled, Q-switched Nd:YAG laser (Continuum, 20 Hz repetition rate, 7 ns pulse duration). The second harmonic (532 nm) was attenuated through neutral density filters and used to excite the sample. The fluence was 35 µJ/cm$^2$. The cw probe light from a Xe arc lamp was passed through various optical elements, the sample, and a monochromator to select the desired probe wavelength, before being detected by a fast photomultiplier tube and recorded by a digital oscilloscope. Averaging over at least 1,000 laser shots was used. Dynamics were recorded over 10,000 points and a 2$^{{st} \text{ order Savitzky-Golay}}$ smoothing algorithm on 33 points was applied.

Femtosecond transient absorption spectra were recorded using femtosecond pulsed laser pump-probe spectroscopy. The pump beam (480 or 580 nm) was generated with a two-stage non-collinear optical parametric amplifier (NOPA) from the 778 nm output of a Ti:Sa laser system with a regenerative amplifier providing 150 fs pulses at a repetition rate of 1 kHz. The pump beam was compressed with a pair of SF-10 prisms to sub-50 fs pulses. The fluence was 20 µJ/cm$^2$. The probe consisted of a white light continuum (WLC) between 400 and 800 nm, generated by passing a portion of the 778 nm amplified Ti:Sa output through a 5 mm-thick CaF$_2$. The probe intensity was always less than the pump intensity and the spot size was much smaller. The probe pulses were time-delayed with respect to the pump pulses using a computerized translation stage. The probe beam was split before the sample into a signal beam (transmitted through the sample and crossed with the pump beam) and a reference beam. The signal and reference were detected with a pair of spectrographs (Princeton instruments, SpectraPro 2150) equipped with 512x58 pixels back-thinned cameras (Hamamatsu S07030-0906), assembled by Entwicklungsbüro Stresing, Berlin. The pump light was chopped at half the amplifier frequency, and the transmitted signal intensity was recorded shot by shot. It was corrected for intensity fluctuations using the reference beam. The transient spectra were averaged to the desired signal-to-noise ratio (typically 3000 acquisitions). The polarization of the probe pulse was at magic angle relative to that of the pump pulse. All spectra were corrected for the chirp of the white-light probe pulse.

3. STEADY STATE OPTICAL SPECTRA

![Absorption and Fluorescence Spectra](image)

Fig. 2a compares the normalized absorption spectrum of an aqueous solution of Bt to that of 2.5 µm thick TiO$_2$ and ZrO$_2$ films sensitized from this solution. The absorption maximum of Bt shifts from 534 nm in solution to 529 nm on ZrO$_2$ and 518 nm on TiO$_2$. Fig. 2b compares the weak fluorescence emission of Bt on TiO$_2$ and ZrO$_2$ at excitation wavelengths of 480 and 530 nm. Though a scattering background from the metal oxide film is apparent in all the data, clearly...
the Bt emission is quenched on TiO$_2$ but not on ZrO$_2$, a consequence of electron injection into the conduction band of TiO$_2$ but not ZrO$_2$. Fluorescence data were gathered at a wavelength near the maximum absorption of Bt (530 nm), as well as at 480 nm because the latter wavelength was expected to excite the strong fluorescence of any residual yellow betaxanthin pigments which are highly fluorescent.$^{13}$ The coincidence of the absorption spectrum of Bt with the excitation spectrum of the emission at 610 nm (not shown) confirms that the weak emission is from Bt and not from an impurity such as residual betaxanthin.

4. ELECTROCHEMISTRY AND SPECTROELECTROCHEMISTRY

![Digital pulsed voltammetry (DPV) of aqueous Bt as a function of pH. The DPV scans for the highest and lowest pH values used are labeled. b) Nernst plot of redox potential versus pH.](image)

Digital pulsed voltammetry (DPV) of aqueous Bt was performed in buffered solutions as a function of pH and is displayed in Fig. 3. The main oxidation occurring at 0.65 V vs. Ag/AgCl at the lowest pH (1.77) shifts to less positive potentials as the pH is raised, with a slope of $-0.034 \pm 0.003$ V/pH. Considering that the oxidation of betanin to give Bt$_{\text{ox}}$ involves the loss of $m$ protons and $n$ electrons, the standard redox potential $E_{\text{red}}^\circ$ applies to the half-reaction

$$\text{Bt}_{\text{ox}} + ne^- + mH^+ \rightarrow \text{Bt}.$$  

This leads to the Nernst equation:

$$E_{\text{red}} = E_{\text{red}}^\circ - \frac{59 \text{ mV}}{n} \log \left[ \frac{[\text{Bt}]}{[\text{Bt}_{\text{ox}}]} \right] - m \times \frac{59 \text{ mV}}{n} \text{pH}$$

Thus the observed slope of the redox potential versus pH suggests a one-proton, two-electron redox reaction. The scheme shown in Fig. 1, adapted from Ref. 10, suggests such a process for the oxidation of Bt to the quinone methide. The oxidation was found to be more facile at low pH. Though some smaller peaks at lower potential than the main peak were observed in the DPV data at lower pH, there was insufficient signal to track these as a function of pH. The above results can be compared to those obtained by Butera et al.$^{14}$ who found three oxidations of Bt at 0.404, 0.616, and 0.998 V, presumably at near neutral pH (PBS buffer). The first of these compares to the value of 0.44 V obtained from the best fit to our data assuming a pH of 7. While this is in rough agreement with the first peak reported in Ref. 14, in our hands the oxidation was not observed to take place at pH values near 7 and above.
Fig. 4 shows the spectro-electrochemical absorption spectrum of Bt on TiO\textsubscript{2} in contact with 0.2 M NaClO\textsubscript{4} in acetonitrile. See text for explanation.

Fig. 4 shows the spectro-electrochemical absorption spectrum of Bt on TiO\textsubscript{2} in contact with 0.2 M NaClO\textsubscript{4} in acetonitrile. The data of Fig. 4 have not been normalized. Note that the initial spectrum at open circuit shows a shoulder at ~450 nm that we believe to be pre-oxidized dye that forms under room light. Two subsequent scans at open circuit gave the same spectrum, with a maximum absorbance of 1.65 at 508 nm. Following these two measurements, a potential of ~400 mV was applied giving the trace shown in green in Fig. 5. The reduction in the shoulder at 450 nm at this potential confirms that the oxidation is at least partly reversible. However, a decrease in the Bt absorbance peak at 508 nm was also observed. Following this, a potential of +2200 mV was applied and the spectra scanned 10 times at approximately 1 minute intervals. With each successive scan there is an increase in the blue shoulder and a decrease in the Bt peak, with an imperfect isosbestic point observed at about 475 nm. The potential was then returned to ~400 mV and a series of five spectra recorded. As seen in Fig. 4, this cathodic potential was capable of reducing the blue shoulder associated with the oxidation product, but the absorption at 530 nm does not recover fully. Finally, seven spectra were recorded while holding the film at a potential of +2200 V. Successive scans resulted in diminished absorbance at 530 nm, but the spectral changes to the blue of the main peak suggest that the 450 nm oxidation product can undergo further oxidation.

5. TRANSIENT ABSORPTION MEASUREMENTS

Fig. 5 shows the global fits of the femtosecond transient absorption data of Bt on TiO\textsubscript{2} and ZrO\textsubscript{2}. In each case the change in absorbance, \(\Delta A\), as a function of wavelength was fit to a model of bi-exponential decay plus a long-time (residual) component. The amplitudes of the two time-dependent components plus the time-independent residual component are shown as a function of probe wavelength. In Fig. 5a is seen the result for Bt on TiO\textsubscript{2} in contact with methoxypropionitrile (MPN). There are two decay components at 0.8 ps and 6 ps. The faster decay is associated with a strong positive absorption in the red and a negative absorption at longer wavelengths. The dip with a minimum at 530 nm is obviously the ground state bleach, while the negative absorbance at longer wavelengths is likely stimulated emission (SE) from Bt, in agreement with the fluorescence spectrum of the sensitized film shown in Fig. 2b. The amplitude of the slower (6 ps) decay in Fig. 5a shows a diminished positive absorption in the blue and reduced stimulated emission. The persistent bleach in the region 500 to 600 nm in the residual component in Fig. 5a (and in Fig. 5b) may correlate to the irreversibility observed in the spectrophotocatalytic measurements discussed above. The data in Fig. 5b are for Bt on TiO\textsubscript{2} in the presence of reducing agent LiI. Similar decay times, 0.9 ps and 8 ps, and similar transient features are seen for the data with and without LiI, but the relative amplitudes of the fast and slow components of the blue transient absorption (TA) are different from what was observed in the absence of LiI. Fig. 5c shows the global fit for Bt on ZrO\textsubscript{2} in contact with MPN. Again, there is a fast (0.8 ps) decay showing positive transients in the blue and negative \(\Delta A\) at longer wavelengths, and a slower (6 ps) decay with diminished SE at red wavelengths. The flat residual component...
shown in Fig. 5c reveals that there is no apparent persistent oxidation of Bt on ZrO₂, in accord with visual observation of the film after laser exposure and spectra recorded before and after laser exposure.

Femtosecond transient absorption spectra were also collected at a pump wavelength of 480 nm and are shown in Fig. 6 for a Bt-sensitized TiO₂ film compared to Bt on ZrO₂. The results are similar to what was obtained at a pump wavelength of 580 nm. The data are fit by a bi-exponential decay with time constants of about 1 ps and 6 to 10 ps. There is again a blue transient absorption with nearly equal amplitudes of the fast and slow decays for Bt on ZrO₂, but a reduced amplitude of the slow TA compared to the fast for Bt on TiO₂. Stimulated emission in the red is observed to decay mostly on the faster time scale, and the bleach in the green is observed on both time scales. One notable difference compared to the data with the 580 nm pump is the blue shift in the bleach in the early time amplitude. The dip at about 510 nm in the early time amplitude shifts to 530 nm in the longer time trace.
Transient absorption data were also recorded on the nanosecond time scale as shown in Fig. 7. It is significant that on this time scale, no signal was observed for Bt on ZrO₂. On TiO₂, on the other hand, transient absorptions at 400-450 nm and 650-750 nm were observed and are assigned to absorption of oxidized Bt. The bleach in the green is also apparent at time scales of 200 ns. An exponential fit to the transients at 425, 600, and 670 nm gave decay times of 2.0, 2.5, and 2.2 μs, respectively, which are the same within the error of the measurement.

The films of Bt on ZrO₂ were found to be quite stable during the femtosecond measurements, with no significant change in the absorption spectrum of the film. The absorption spectrum of the Bt-sensitized TiO₂ film, on the other hand, was found to undergo a significant blue shift after laser exposure. Visual inspection of the exposed parts of Bt/TiO₂ films revealed yellow coloration consistent with formation of oxidized dye with an absorption band in the blue.

6. DISCUSSION

The femtosecond transient absorption data of Bt-sensitized TiO₂ and ZrO₂ consistently revealed bi-exponential decay with approximately 1 ps and 5-8 ps time constants, referred to here as the fast and slow decay components. Common features in the femtosecond pump-probe data are the blue TA, red SE, and a green bleach, all observed on both Bt/TiO₂ and Bt/ZrO₂ films. The green bleach is also observed in the long-time component for the Bt/TiO₂ film but not for Bt/ZrO₂. The relative amplitudes of the slow and fast components of the blue TA are significantly different for the two films. On ZrO₂, these amplitudes are roughly equal, while on TiO₂ in the presence of MPN, the slow component of this TA has an amplitude which is only 40% that of the fast component. This ratio is further reduced to about 20% for the Bt/TiO₂ film in contact with LiI in MPN. Since no transient absorption is observed in the nanosecond pump-probe exper-
ments for Bt/ZrO\(_2\), the blue (~450 nm) and red (~650-750 nm) TA features seen for Bt/TiO\(_2\) on this time scale (Fig. 7) are assigned to oxidation products. The former is certainly consistent with the oxidation product dhBt shown in Fig. 1. However, since the blue TA is also present in the fast and slow decays in the fit to the fs data for Bt/ZrO\(_2\), we must consider the positive absorption in the blue seen in the femtosecond experiments can also result from excited state absorption, i.e. the transition S\(_1\) \(\rightarrow\) S\(_2\). The red SE, observed for Bt on both TiO\(_2\) and ZrO\(_2\), is coincident with the fluorescence spectrum of Bt.

A possible scheme explaining these results is presented in Fig. 8. Consider first the transient events taking place on ZrO\(_2\), where we assume there is no photo-oxidation of Bt on any time scale. The bimodal decay of the blue TA could be a consequence of fast thermal relaxation of the vertical (Franck-Condon) excited state to the relaxed excited electronic state, accompanied by slower parallel relaxation of the vertical excited state directly to the ground state. This would explain the weak negative absorption (from SE) in the red at longer time scales, and the observation that the negative absorption in the green decays on both time scales as the ground state population recovers. Given that the fluorescence quantum yield of Bt in aqueous solution is on the order of \(\phi_f \approx 0.001\) (unpublished work), and with \(\tau_{rad}\) on the order of 5 ns (typical value for a strongly allowed S\(_1\) \(\rightarrow\) S\(_0\) transition with oscillator strength near unity), a nonradiative relaxation time \(\tau_{rad}\) of about 5 ps is expected in accord with our observations.

Interpretation of the femtosecond pump-probe data for Bt on TiO\(_2\) is complicated by the apparent coincidence of the absorption of oxidized Bt (B\(_{ox}\)) with the S\(_1\) \(\rightarrow\) S\(_2\) TA, which both occur at about 450 nm. The fast and slow time constants of the bi-exponential relaxation are not significantly different for Bt on TiO\(_2\) versus ZrO\(_2\), however, the relative amplitudes vary greatly. For Bt/TiO\(_2\) in the presence of MPN, the amplitude of the slow component of the blue TA is about 40\% of that of the fast component. Excited state electron injection into the conduction band of TiO\(_2\) can be reasonably assumed to take place from the vertical excited state, leading to greater amplitude of the fast decay of the excited electronic state. However, if the result is offset by the appearance of positive blue TA from Bt\(_{ox}\), we would see increased amplitude of the fast decaying blue TA only if the oscillator strength of Bt\(_{ox}\) is less than that of the S\(_1\) \(\rightarrow\) S\(_2\) transition. Only the relative amplitudes are meaningful, and the additional fast relaxation channel resulting from electron injection from the vertical state reduces the relative amount of relaxed excited state dye and thus the relative amplitude of the slowly relaxing blue TA.

In the presence of LiI (Fig. 5b), we expect to see changes in the dynamics that reflect regeneration of ground state Bt from Bt\(_{ox}\), e.g., faster ground state recovery. This was not apparent, but we do observe that the amplitude of the slow decay of the blue TA is reduced to about 20\% of that of the fast decay. This effect could result from the well-known effect of Li\(^+\) on the TiO\(_2\) conduction band. A lower energy conduction band edge in the presence of Li\(^+\) could account for increased amplitude of the fast decay resulting from electron injection, the driving force for which is greater than in the absence of Li\(^+\). Thus, compared to Bt on ZrO\(_2\) and on TiO\(_2\) in the absence of LiI, the relative amplitude of the slowly relaxing blue TA is further reduced as electron injection from the vertical excited state is enhanced.

The nanosecond transient absorption data reveal a red absorption band from an oxidized form of Bt. This does not appear to be an additional transition of the same species that gives rise to the 450 nm absorption band, since it is not seen in the steady-state spectra of oxidized dye. A positive red TA was not seen in the femtosecond data, but might have been offset by the negative \(\Delta\)A resulting from SE. DFT calculations are presently underway to investigate the possibility that the red TA observed in Fig. 7 is an intermediate such as the quinone methide shown in Fig. 1, a degradation product, or a one-electron semiquinone oxidation product. The results of these calculations will be reported in a separate publication.

7. CONCLUSIONS

Transient positive absorption at 450 nm in femtosecond pump-probe measurements of Bt on TiO\(_2\) results from excited state absorption which coincides with the two-electron oxidation product 2-decarboxy-2,3-dehydrobatiman. Relative amplitudes of the fast and slow decay components of the blue TA are consistent with increased efficiency of electron injection in the presence of LiI. Nanosecond data of Bt on TiO\(_2\) is entirely due to oxidation products, and reveals the two-electron oxidation product and an as-yet unknown intermediate that absorbs in the red. The results of analysis of the nanosecond data will be presented in a future publication along with DFT calculations of potential intermediates in the oxidation of Bt.
ACKNOWLEDGMENTS

The support of the National Science Foundation through grant DMR 1305592 is gratefully acknowledged. FJK and JLM thank Jacques Moser and his group members for their gracious hospitality during their stay at EPFL.

REFERENCES