THE EFFECT OF ATMOSPHERIC COLD PLASMA CLEANING OF FTO SUBSTRATES ON THE QUALITY OF TiO₂ ELECTRON TRANSPORT LAYERS FOR PRINTED CARBON-BASED PEROVSKITE SOLAR CELLS

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Abstract
In this work, TiO₂ electron transport layers (ETL) of printed perovskite solar cells with back carbon electrode (C-PSCs) were prepared on uncleaned, chemically cleaned, and plasma-treated FTO substrates. The effect of cleaning procedure type on the quality of TiO₂ ETL has been characterised by XPS and SEM. The C-PSCs with FTO substrates treated by low-temperature atmospheric plasma at ambient air for 10 s reached approximately the same efficiency (4.4%) as with the chemically cleaned FTO (4.5%). From optical microscope and SEM images, it was observed that the plasma cleaned the FTO substrate to some extent, thereby reducing the number of defects and pinholes occurring in the blocking layer. XPS measurements confirmed that chemical and plasma treatment significantly reduced organic species on the FTO surface.

Keywords: Atmospheric cold plasma, TiO₂, electron transport layer, carbon electrode, perovskite solar cells

1. INTRODUCTION
The role of the electron transport layer (ETL) of carbon-based perovskite solar cell (C-PSC) is to extract and transport electrons from the perovskite layer towards the transparent conductive oxide (TCO) electrode (glass or PET substrate with FTO or ITO layer). The quality of the ETL depends on two main parameters: 1) the preparation process and the composition of the used dispersion, which determine its thickness, roughness, and compactness; 2) the surface properties of the used substrate [1]. The most suitable ETL layer is very thin (<50 nm), which results in defects that cause internal short-circuits after the undesirable contact between the perovskite and TCO [2,3]. On the other hand, if the layer is too thick, the series resistance increases, the electrons recombine with the holes, resulting in a reduction of the current density and the fill factor of the solar cell [4,5]. The ETL with a thickness of several tens of nanometres is applied on the pre-treated substrates, and even the smallest impurity related to insufficient cleaning can significantly affect the quality of the ETL as well as the performance of the entire PSC. The most commonly used method of cleaning is chemical cleaning, which is very effective, but on the other hand, it is time-consuming (~1 hour) and requires the use of harmful organic solvents [6].

In this work, we showed that rapid (in order of seconds) atmospheric plasma treatment in ambient air has a potential to replace chemical cleaning of FTO substrates, and we examined the effect of plasma on FTO surface and the quality of TiO₂ ETL deposited on FTO. In addition to substrate cleaning, plasma also increases the surface energy and ensures better ETL homogeneity and charge transport at its interface with FTO.
2. EXPERIMENTAL SECTION

2.1. Materials
FTO/glass substrates (fluorine-doped tin oxide, 7 Ω/sq.), titanium diisopropoxide bis(acetylacetonate) (75 wt.% TiAcAc in isopropanol), isopropyl alcohol (≥ 99.7% (CH₃)₂CHOH), diethanolamine (≥ 98 wt.%, DEA) were purchased from Sigma-Aldrich. Titanium tetrachloride (≥ 97%, TiCl₄) was ordered from Merck KGaA. Ethanol (≥ 97%, CH₃CH₂OH) was from mikroCHEM. Mesoporous titania nanoparticle paste (Ti-Nanoxide T165/SP), mesoporous zirconium dioxide nanoparticle paste (Zr-Nanoxide ZT/SP), mesoporous carbon-black paste (Elcocarb B/SP) and perovskite solution ((5 – AVA)ₓ(MA)₁ₓPbI₃) were purchased from Solaronix.

2.2. Device fabrication
Printed mesoporous perovskite solar cells with carbon back electrode (C-PSCs) were prepared on the commercial FTO/glass substrates cleaned by three different methods: chemically cleaned, plasma-treated, and uncleaned. The preparation process of solar cells with the structure (Figure 1): FTO glass substrate/electron transport – blocking layer TiO₂/mesoporous TiO₂ layer/scaffold ZrO₂ layer/carbon back electrode/perovskite ran under laboratory conditions. The chemical cleaning procedure was managed as follows: applied substrates were precisely and sequentially cleaned with a surfactant, acetone, and isopropyl alcohol in an ultrasonic bath for 20 min in every solvent. The drying of the substrates took place on the ambient air, and additionally before deposition of the functional layers were treated by ultraviolet radiation for 15 min to remove all organic residues. Plasma-treated substrates were exposed to 10 s low-temperature ambient-air surfaceplasma generated by RPS40+ (Roplass s.r.o., Czech Republic). The surface of the uncleaned substrates was not treated in any way. First, a compact TiO₂ sol-gel blocking layers prepared from the mixture of TiAcAc and TiCl₄ precursors were spin-coated on the different pre-treated FTO substrates and sintered at 500 °C for 30 min. Second, other functional layers of the C-PSC, such as mesoporous TiO₂, ZrO₂ and carbon back electrode were deposited by screen-printing. Third, the perovskite solution was directly dropped and infiltrated through the carbon layer and dried at 100 °C for 30 min. A detailed method of C-PSC and TiO₂ blocking layer preparation is described in the author’s previous work [7].

![Perovskite infiltration](image)

**Figure 1** The structure of the printed C-PSC

2.3. Characterisation
The conversion efficiency of solar to the electric energy of the C-PSC was determined by measuring photocurrent density-voltage load characteristics under 1000 W/m² of standard daylight exposition with a 1.5G air-mass (AM). Optical microscope (LEICA DM 2700 M) and scanning electron microscope (JEOL, JSM67500FA, Analytical field emission scanning electron microscope, Japan) were used for the analysis of the FTO/TiO₂ sol-gel surface. The X-ray photoelectron spectrometer Axis Supra from Kratos Analytical was used to study the surface chemistry of chemically cleaned, plasma-treated and uncleaned FTO samples.
3. RESULTS AND DISCUSSION

We analysed the reference, chemically cleaned and plasma-cleaned samples by X-ray photoelectron spectroscopy (XPS) to study the changes in the stoichiometry and bonding states on the surface of FTO glass. Table 1 presents the elemental composition of the surface of FTO for different cleaning methods used. The uncleaned sample was highly contaminated, containing as much as 74% of carbon on the surface. Chemical cleaning process led to a decrease in carbon concentration on the surface to 26%. The uncleaned surface also contained traces (below 1%) of potassium and calcium, which were also reduced and in case of potassium completely removed after chemical cleaning. The oxygen concentration on the surface also increased, which can be attributed to the relative change in carbon concentration. Plasma cleaning of FTO for 10 s led to a decrease in carbon concentration to around 14%. The increase of oxygen on the surface was more significant compared to the chemical cleaning process. Exposure of the surface to the ambient air plasma, which contains reactive oxygen species causes oxidation of the surface and is known to induce polar oxygen-containing functional groups to the treated surface [8-10]. The O:C ratio (Table 1) therefore increased significantly. Compared to chemical cleaning, plasma treatment was not so efficient for the total removal of the potassium and calcium contamination. Around 4% of nitrogen was introduced to the surface of plasma-treated samples, a known effect of treatment by plasma generated in ambient air. These results correspond well with the plasma-cleaning of ITO with the same plasma source reported by Homola et al. [11].

### Table 1 Atomic concentrations from XPS with standard deviation measured at two points for each sample

|            | C [%] ± | O [%] ± | Sn [%] | Ca [%] ± | K [%] ± | N [%] ± | O:C | O:Sn |
|------------|---------|---------|--------|---------|--------|--------|-----|------|
| No clean   | 74 ± 7  | 19 ± 4  | 5 ± 3  | 0.7 ± 0.1| 0.2 ± 0.0| 1.0 ± 0.3| 0.3 | 3.7  |
| Chem clean | 26 ± 1  | 49 ± 1  | 23 ± 1 | 0.3 ± 0.1| 0.0 ± 0.1| 2.1 ± 0.7| 1.8 | 2.1  |
| 10 s plasma| 14 ± 1  | 57 ± 0  | 23 ± 1 | 1.1 ± 0.3| 0.8 ± 0.2| 4.0 ± 1.0| 4.0 | 2.5  |

The high-resolution C 1s region in the XPS spectrum is shown in Figure 2a. Four components were used for deconvolution, specifically C—C—C—H, C—OH/C—O—C, C=O and O—C=O species at 285.0 eV, 286.5 eV, 288.0 eV and 289.0 eV of binding energy, respectively. The reference sample without any cleaning displayed a significant C—C—C—H component with around 82% relative intensity. Chemical cleaning led to the removal of C—C—C—H groups from the surface, while the relative intensity of the oxygen-containing functional groups increased. The oxidation effect of the plasma can be seen in the increase of the relative intensity of the oxygen-containing functional groups in the C 1s region of the plasma-treated sample.

The high-resolution O 1s region is shown in Figure 2b. The SnO$_2$ in FTO typically displays a peak in the O 1s region at around 530.5 eV [12,13]. Furthermore, oxygen on metallic surfaces can be found in the form of metal-hydroxides (OH)$^-$, that are often found in around 531.5 eV and in adsorbed H$_2$O at around 533.0 eV [14,15]. Oxygen can also be bound to carbon in a multitude of environments of the organic species, positions of which can vary between 531.0 eV up to 534.0 eV. We can expect significant contributions from the organic oxygen to the (OH)$^-$ and H$_2$O components. Both chemical cleaning and plasma treatment led to a more pronounced SnO$_2$ component due to the removal of organic species. A shift of the SnO$_2$ component to higher binding energies can be seen after the plasma treatment. We believe that the surface of FTO prior to plasma treatment consisted of a mixture of Sn$^{4+}$ oxide and Sn$^{2+}$ oxide. Several author have shown that the signal in the XPS spectrum from the SnO is shifted from the signal from SnO$_2$ by around 0.4 eV towards lower binding energies [15-17]. The reactive oxygen species in plasma cause oxidation of the SnO into SnO$_2$ and therefore the shifted signal is observed.
The photocurrent-voltage load characteristics of C-PSCs with all types of treated and uncleaned substrates are shown in Figure 3. It is evident, that the purity of the substrate’s surface has a crucial effect on the quality of the deposited functional layers. Specifically on the blocking layers, as their thickness is several tens of nanometers and any impurity adversely affects all the photovoltaic parameters. The blocking layer with high quality without any cracks or failures prevents the recombination of electrons with holes, and also the internal short circuits what is necessary for high performance of C-PSC. Devices with substrates treated by the chemical way and by plasma for 10 s reached approximately the same efficiency 4.4 – 4.5% with relatively good shapes of photocurrent-voltage load characteristics, fill factor (FF) and short-circuit current density (J_sc) (Table 2). The open-circuit voltage (V_oc) has changed only partially from 0.65 to 0.70 V in all cases. Results achieved by the chemical cleaning can be attributed to a thin, homogeneous, and almost failure-free blocking layers, which was confirmed by optical and top-view SEM images (Figures 4, 5a). In the case of the substrates plasma-treated for 10 s, a small occurrence of impurities and defects in the blocking layers is seen in Figures 4, 5b. On the other hand, the effect of plasma improved the wettability of the FTO surface (contact angle decreased for plasma-treated FTO = 10° vs chemically cleaned FTO = 35°), and interfacial contact between the FTO and the blocking layer what led to better charge transport between the layers. The efficiency of C-PSCs with uncleaned substrates was at the level of 1% due to the presence of a large number of impurities, failures and pin-holes in the blocking layer (Figures 4, 5c). The presence of failures and pin-holes in the blocking layer created the trapping sites for charge recombination and current leakage what further led to a decrease in FF, J_sc, and to the linearisation of the load characteristics (Table 2, Figure 3). It can also be observed from optical images that the plasma, to some extent, cleans the surface of the substrate itself (Figures 4b vs 4c).

Table 2 Photovoltaic parameters of the C-PSCs with differently treated and uncleaned substrates

| Treatment    | J_sc (mA.cm²) | V_oc (V) | Efficiency (%) | FF  |
|--------------|---------------|----------|----------------|-----|
| Uncleaned    | 8.4           | 0.70     | 1.4            | 0.23|
| Cleaned      | 15.8          | 0.68     | 4.4            | 0.41|
| Plasma 10 s  | 17.0          | 0.65     | 4.5            | 0.41|

Figure 2 Comparison of narrow regions of a) C 1s and b) O 1s in the XPS spectrum of the uncleaned, chemically cleaned and plasma-cleaned surface of FTO
4. CONCLUSION

In this work, the mesoporous perovskite solar cells with carbon back electrode were prepared on chemically cleaned, plasma-treated and uncleaned substrates by screen-printing and coating technics. The obtained results showed that solar cells prepared on chemically cleaned substrates reached the approximately same efficiency and FF as on the substrates treated by 10 s low-temperature ambient air plasma. It means that the few second plasma treatments of FTO glass substrate can successfully replace protracted one hour chemically cleaning in different solvents and an ultrasonic bath.
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