Chapter

Application of Atomic Force Microscopy in Organic and Perovskite Photovoltaics

Chandra Shakher Pathak

Abstract

Atomic force microscopy (AFM) has become widely used technique in air, liquids, or vacuum to generate high-resolution topographic images of a surface having a nanometer-scale resolution. AFM gives the information about the morphology, phase composition etc. Photovoltaic materials have been attracting intense interest due to their performance and the morphology as well as quality of these materials affects their performance. AFM is now a day widely used technique for morphology and other electronic properties measurements at nanoscale for photovoltaic materials to understand their relation with device performance. This chapter describe the brief introduction of Kelvin probe force microscopy (KPFM) and conducting atomic force microscopy (CAFM) and their application in electrical characterization at nanoscale of organic and perovskite photovoltaic materials.

Keywords: AFM, KPFM, CAFM, Photoconductive AFM, Nano scale, Electrical properties, Surface potential

1. Introduction

Scanning tunneling microscopy (STM) was first introduced in 1982 and it is based on tunneling of electron between the metallic tip and sample surface and it is limited to the study of conducting surface [1]. After four years in 1986, the solution of this restriction was provided by atomic force microscopy (AFM) and it is based on the detection of attractive or repulsive forces [2]. AFM was used to analyze the surface morphology and used to measure the force of interaction between the AFM tip and the sample. AFM consists of a sharp tip having nanometer dimension which is attached to a cantilever is used to scan the sample surface. A laser beam focused on the cantilever which detects the bending of cantilever. The reflection of the laser beam is focused on photodiode detector. Deflection of the cantilever is monitored during the scanning and converted into surface image. AFM is generally operated in contact mode and tapping mode. AFM is the widely used tool for the characterization of materials surface at the nanoscale. AFM was developed to a very versatile technique by combination with other measurements methods. This chapter will cover the application of electrical mode of AFM specifically Kelvin probe force microscopy (KPFM) and conducting atomic force microscopy (CAFM).
1.1 Kelvin probe force microscopy

Lord Kelvin proposed macroscopic Kelvin probe method in 1898 to determine the contact potential difference ($V_{CPD}$) between a metallic plate and sample [Kelvin L., Contact electricity of metals *Phil. Mag.* 1898, 46, 82–120]. Later in 1991, KPFM was first introduced by Nonnenmacher *et al.* [3]. The KPFM mode is basically two-scan process. In the first scan topography of the surface is acquired in tapping mode along a single line profile. Following this, the mechanical excitation of the cantilever is turned off and in the second scan this topography is retraced at a certain lift height (LH) above the sample surface, recording local variations in contact potential difference (CPD). During the second scan, the tip-sample distance is constant and it is equal to $d_{AFM} + LH$, where $d_{AFM}$ represent the tip-sample distance during the topographic scan. When the AFM tip is brought close to the sample surface, the electrostatic force is generated and it is proportional to the difference between their Fermi levels. If the work functions of the sample and the tip are different, electrons flow from the lower work function to the higher work function material to align their Fermi level and the system reach in charge equilibrium condition. The vacuum levels of the tip and sample are not the same and the surfaces of the sample and the tip are charged and an apparent $V_{CPD}$ will form. Due to the $V_{CPD}$ an electrical force acts on the contact area and this force can be nullified by applying a DC voltage and it is equal to the work function difference between tip and sample. The work function of the sample can be calculated when the tip work function is known from scanning a reference sample.

1.2 Conductive atomic force microscopy

Conductive atomic force microscopy (CAFM) is usually used to analyze the local variations in current of the sample. The AFM controller is used for applying dc bias through the substrate during measurements. The CAFM tip is connected through a low noise current amplifier to the AFM controllers to generate the current image. We can collect the current–voltage ($I$-$V$) characteristics by CAFM tip which can be contacted randomly at various positions of sample surface and $I$-$V$ characteristics are collected at each position.

KPFM generates 3D mapping of surface electric potential and measure the local work function while CAFM generates the current map. CAFM is one of the simplest ways of characterizing electrical properties at high resolution is by applying voltage between the sample and a CAFM probe. Photoconductive AFM; current mapping under illumination is useful for correlating high resolution current mapping with the photovoltaic device performance. Now a day these two modes of AFM in electrical mode are widely used in all area of research; nanoelectronics field, solar cell, 2D materials and semiconductor industries, biology etc. This chapter consists the collection of some published work as well as some new results.

2. Organic photovoltaics

Conducting polymers have been attracting attention after their discovery by Shirakawa, MacDiarmid, and Heeger in 1977, who were awarded the Nobel Prize in Chemistry in 2000 for the discovery of conductive polymers [4, 5]. They used organic polymer polyacetylenes, which is a conjugate polymer and insulator and increased the conductivity of polyacetylene films by several orders of magnitude by chemical doping [4]. In recent years, there has been lot of research activity in the field of polymer electronics and attracted a lot of attention because of its high...
flexibility, light weight and solution process ability [6]. Applications of conducting polymers include organic light emitting diodes, organic thin film transistors, organic solar cells, actuators and sensors etc. [7–13]. AFM is usually used to measure the roughness, morphology and phase analyses of conductive polymer [13]. KPFM is used to measure the work function [14–20] and CAFM [18, 21, 22] used to measure the current of the organic solar cell materials. These two methods gives valuable insights in the structure and working mechanism of organic photovoltaic devices.

Poly(3,4-ethylenedioxythiophene)-poly (styrenesulfonate) (PEDOT:PSS) is a well-known conducting polymer because of its high conductivity, excellent thermal stability, transparency, structural stability and processability [23–25]. PEDOT:PSS polymer is a promising candidate as a transparent electrode for optoelectronic devices. PEDOT is made from ethylenedioxythiophene (EDOT) monomers. PEDOT is insoluble in many common solvents, and it is unstable in its neutral state. To improve its processability, water-soluble polystyrene sulfonate (PSS), can be added and the addition of PSS causes it to become soluble. During the polymerization, PSS acts as charge balancing dopant to yield PEDOT:PSS.

Figure 1 shows the height images and Figure 2 shows the surface potential images of PEDOT:PSS films with co-solvents (N-Methyl-2-pyrrolidone (NMP) and methanol (MeOH)). The average root-mean-square roughness (RMS) values were found in the range 3.2–5.5 nm. Pristine PEDOT:PSS film was quite smooth with a RMS roughness of 2.6 nm. RMS roughness values of PEDOT:PSS films increases after co-solvents addition. Variation in the RMS roughness after addition of co-solvents indicates the morphological change that arises from the conformation of the polymer chain [16]. The estimated average work function of co-solvents doped PEDOT:PSS thin films range from 4.63 to 4.82 eV as compared to 4.9 eV for the pristine PEDOT:PSS film [16]. Work function can be calculated with the following Equation [16, 27].

$$V_{CPD} = \frac{\phi_{tip} - \phi_{sample}}{-e}$$ (1)
where $V_{CPD}$ is the contact potential different measured by KPFM in volts, $\phi_{tip}$ and $\phi_{sample}$ are the work functions of tip and sample in eV, respectively, while $e$ is the electronic charge.

Figure 3 shows the current map of bare and NMP doped PEDOT:PSS films. Bare PEDOT:PSS has less current as compared to NMP doped PEDOT:PSS film as shown in Figure 3 [28]. This shows the reduction of more PSS from the surface for NMP doped PEDOT:PSS film. Hosseini et al. [22] also reported higher current for ethylene glycol (EG) doped PEDOT:PSS film.

Figure 4 shows topography, surface potential images and corresponding line profile of bare and NMP doped PEDOT:PSS films. Bare PEDOT:PSS has work function of 4.90 eV and 4.77 eV for NMP doped PEDOT:PSS film [16, 28]. Line profile shows the homogenous distribution of surface potential. We also showed that the work function was reduced with the addition of dimethyl sulfoxide (DMSO) solvent in PEDOT:PSS [15].

![Figure 2. Potential images of PEDOT:PSS films modified with (a) 0% NMP 1% MeOH, (b) 0.1% NMP 1% MeOH, (c) 0.3% NMP 1% MeOH, (d) 0.5% NMP 1% MeOH, (e) 0.8% NMP 1% MeOH, and (f) 0% MeOH 0.5% NMP. Reproduced with permission from [26].](image1)

![Figure 3. CAFM images of bare and NMP doped PEDOT:PSS films. Reproduced with permission from [28].](image2)
3. Perovskite photovoltaics

The last 10 years has seen a new photovoltaic (PV) technology being discovered and developed at a rate greater than any previous energy harvesting technology with research fuelled by the facile, low cost large area solution processing routes available for device fabrication. These devices, known as organic–inorganic metal halide perovskites have certified power conversion efficiencies (PCEs) > 25% [29], comparable values to the incumbent technologies but available at a fraction of the materials and processing costs. The perovskite thin films are typically polycrystalline ones, comprising microstructures such as grains and grain boundaries (GBs). Recently some of research groups have performed the microscopic investigation and suggested that the grain boundaries (GBs) in planar perovskite solar cells have beneficial [30, 31]. Hence engineering of the perovskite films and the microscopic investigation is essential for the further improvement in the properties of perovskite photovoltaics. CAFM and KPFM have been widely used to characterize the local properties of perovskite thin films. Such studies suggested that GBs have beneficial effects due to efficient photogenerated charge carrier separation and collection at GBs [30, 31]. Downward- as well as upward- band bending at GBs were reported from KPFM measurements, depending on the GB composition [41]. Zhao et al. [32] measured the photocurrent of FTO/compactTiO₂/mesoporousTiO₂ + CH₃NH₂PbI₃/CH₃NH₂PbI₃ film and found the photo current intensities were higher on the center of the grain than on the defect position, whereas the larger grain size leads to the higher photocurrent on the center. They also reported that the dark current intensities increased dramatically in the defect position, suggesting a high conductive character for defect position. Li et al. [31] fabricated CH₃NH₂PbI₃/mTiO2/c-TiO2/FTO typical device and performed c-AFM measurements under steady illumination of power 14 mW/cm² on CH₃NH₂PbI₃ surface from the top and showed that the photocurrent flows through the GBs are negligible at 0 V bias, while the major photocurrents form on the grains. However, with the low bias the photocurrents at the GBs become much higher than those of the grains. Xu et al. [35] also observed higher current near GBs for CH₃NH₂PbI₃- PCBM structure under high vacuum and dark conditions.
Kutes et al. [34] performed photoconductive AFM for methylammonium lead iodide (CH$_3$NH$_3$PbI$_3$)/c-TiO$_2$/FTO/glass structure and the schematic diagram of the photoconductive (CAFM under illumination) AFMs configuration shown in Figure 5. They observed higher current in grains than grain boundary (GB) as shown in Figure 6.

Yun et al. [30] also used CAFM and KPFM technique to investigate the role of GBs and the schematics shown in Figure 7. They observed the higher CPD at grain boundary with illumination and current near GBs for CH$_3$NH$_3$PbI$_3$/TiO$_2$/FTO/glass heterojunction structure than in the grains as shown in Figures 8 and 9.

They found that the higher current collection near GBs is consistent with KPFM results, which indicates that photogenerated carriers are more efficiently separated and transported along the GBs. Lower CPD at the GBs under the dark condition implies that downward band bending is present at GBs. KPFM is widely used for photovoltaics to analyses the work function of perovskite materials.

We prepared the MAPI films as described by Liu et al. [42] and they dipped PbI$_2$ film on CH$_3$NH$_3$I solution. Here we spin casted CH$_3$NH$_3$I in 1 ml isopropanol on mesoporous PbI$_2$ film and annealed the prepared films at 70° to 110°C for 10 minutes. Figure 10 shows the height image and current–voltage characteristics of CH$_3$NH$_3$PbI$_3$ (MAPI) films on FTO annealed at 70°, 90° and 110° C for 10 min respectively. Which shows the higher grain size for 70° C annealed perovskite film (Figure 10a) and the current was in pA for all three samples.

After the temperature variation CAFM studies, We prepared mesoporous PbI$_2$ scaffolds MAPI films on FTO and annealed at 100°C were analyzed in details. CAFM and KPFM with and without illumination measurements were done and the schematics of these measurements are shown in Figure 11 and results are shown in Figures 12 and 13. At dark and with the illumination of 532 nm laser having intensity of 0.6 W/cm$^2$ shows homogeneous higher current near GBs. KPFM also shows...
lower CPD value at GBs and mostly homogeneous mapping which are consistent with the results by Yun et al. [30]. Here we illuminated the sample from glass side not from the top (perovskite) side.

**Figure 13** shows the topography and surface potential images and line profile of the perovskite films under illumination. The change in the CPD value reflects the change in the work function of the perovskite surface. Charge generation occurs significantly in GBs and the higher photocurrent near GBs suggests that GBs acts as channels for current flow than strong recombination centers. As we know that the higher number of defects are present at the grain boundaries which will increase
the non-radiative recombination of electrons and holes. The charge accumulation or depletion between grains and GBs may cause band bending, which induces the charge carrier separation. Hence the investigation of electronic properties at grain boundaries is crucial. KPFM has been used to determine the surface potential at
Figure 9.
CAFM measurements performed on a CH$_3$NH$_3$PbI$_3$/TiO$_2$/FTO/glass structure over an area of 5 $\mu$m$^2$.
(a) Topographic image and (b) current image are taken in the dark at 0 V. Current images (c) under illumination at 0 V and (d) under illumination at 0.3 V. Insets in panels b – d are overlap of corresponding CAFM maps and a topography map of the region with a white outline in panel a. Wavelength and intensity of the illumination were 500 nm and 1.1 kW/cm$^2$. Reproduced with permission from [30].

Figure 10.
Topography, corresponding current-bias characteristics of MAPI/film annealed at (a) 70°, (b) 90° and (c) 110° C.
Figure 11. Schematic representation of CAFM and KPFM measurements.

Figure 12. Topography, current map and line profile of MAPI/FTO (a) dark and (b) under illumination of 532 nm laser having intensity of 0.6 W/cm².

Figure 13. Topography, CPD map and line profile of MAPI/FTO (a) dark and (b) under illumination of 532 nm laser having intensity of 0.6 W/cm².
grains and GBs. From the topography and CPD map we can see the individual grains and the CPD is higher in grains compared to GBs. The line profile plots are useful in quantitative analysis of CPD variations across topographical feature in perovskite films. CPD is higher at higher region (grains) and lower in lower regions (GBs) and it might be due to the presence of built in potential around the GBs.

4. Conclusion

AFM provides lots of required and interesting results with advanced modes and widely accepted technique to characterize all type of materials. CAFM and KPFM provides information about grains and GBs of photovoltaic materials which help us to understand the current transport and band bending to improve the performance and life time of photovoltaic materials. Combination of such characterization at nanoscale with macroscopic analysis can link the photovoltaic materials properties and optimization of device performance.

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Author details

Chandra Shakher Pathak\textsuperscript{1,2}

1 Ben-Gurion National Solar Energy Center, Department of Solar Energy and Environmental Physics, Jacob Blaustein Institutes for Desert Research, Ben-Gurion University of the Negev, Israel

2 Department of Chemical Engineering, Ben-Gurion University of the Negev, Beer Sheva, Israel

*Address all correspondence to: cspathak12@gmail.com
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