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## Local electrical characterization of PVDF textile filament

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### ABSTRACT

The piezoelectric behavior of Poly(vinylidene fluoride), PVDF, is known since several decades and is clearly related to its crystalline phases. Many works made on films or fibers have focused on the characterization of the phase transitions during various PVDF processing and on its electromechanical activity by combining several techniques. Piezo-force microscopy (PFM) is an interesting tool to underline the crystalline forms and piezoelectricity efficiency of PVDF at the local scale. However, this technique is little used on samples in the form of fibers and in this case, it is most often nanofibers. In this work, two conventional PVDF textile filaments, with different weak draw ratio, are produced and analyzed by FTIR, XRD and PFM. We demonstrate that the PFM analysis can be relevant for specimens presenting low signals during other characterizations. Therefore, the local piezo-/ferroelectricity into the fiber is highlighted underlining the existence of the polar phases of PVDF. Then, the effective piezoelectric coefficient  $d_{33}$  of PVDF fiber drawn with a ratio of 1.5 is estimated at 12 pm/V.

**KEYWORDS:** Poly(vinylidene fluoride) PVDF; melt-spun textile fiber; piezoelectric effect; piezo-force microscopy

## INTRODUCTION

Over the last decade, textile instrumentation has attracted the attention of several researchers [1]. Firstly confined to specific technical areas and dedicated to well-defined users (e.g. tracking for military [2, 3], sensors for firefighters [4, 5], etc.), this topic increasingly integrates mass-market applications due to the emergence of connected, e-textile or so-called “smart textiles” in various areas such home textiles [6], fashion [7], sport [8, 9], health [10, 11], etc. A better integration of “intelligent functions” in textile structures is particularly relevant in the latter sectors, where monitoring of body parameters is of particular interest. Due to the multiplication of sensors, process unit and communicative system, the need of energy and data of sector is substantial. One of the possible solutions is energy harvesting by textile structure itself [11, 12]. Some solutions based on the conversion of energy (thermal, mechanical, etc.) to electric voltage are nowadays investigated in laboratories.

Piezoelectric materials show reversible relationships between electric charge and mechanical deformation called converse and direct piezoelectric effect [13]. The latter leading to an electric charge generation during mechanical stress can be used in a wide variety of applications from neural stimulation [14, 15], sensors development [16, 17] or energy harvesters [13, 17]. As a consequence, such materials and more particularly piezoelectric polymers are perfect candidates to solve the main problem for e-textiles, i.e.: develop self-sustaining energy sensors. Among the piezoelectric polymers available, fluoropolymers such as poly(vinylidene fluoride) (PVDF) and its copolymers show the better piezoelectric behavior [18]. Although copolymers crystallize in polar form suitable for piezoelectric properties, PVDF is still preferred due to its relative low cost. Nevertheless, depending on the processing conditions it can crystallize in five different structures:  $\alpha$ ,  $\delta$ ,  $\epsilon$ ,  $\gamma$ ,  $\beta$  [19]. The  $\beta$  phase is the most polar phase [20].

Accordingly, this one is targeted to convert mechanical deformation into electric energy. The development of PVDF textile energy harvesters requires solving many scientific issues such as the generation of the  $\beta$  phase in the fibers but also its evaluation as well as its electromechanical activity.

On one hand, various methods exist to determine the presence of  $\beta$  phase into PVDF. Usually this polar crystalline form is characterized by one or more thermal analysis techniques [21] (differential scanning calorimetry (DSC), modulated DSC [22], dynamic mechanical analysis (DMA) or thermal mechanical analysis (TMA)) as well as X-ray diffraction (XRD) and spectroscopy measurements (FTIR, Raman, NMR) [23]. Depending on the test used, it is easily to evaluate qualitatively and quantitatively the presence of  $\beta$  phase in PVDF fiber. DSC is a fast and controlled characterization method on fiber. However, as in thermal analysis techniques, melting and recrystallization phenomena during the test may disrupt the analysis of the results. Other limitations for easy characterization such as the size, the form and the morphology of produced fibers have to be taken into account in some other tests. Thus, it is often necessary to design special specimen holder for filaments and to consider the orientation of sample during the test [24]. To correlate mechanical stretching and  $\beta$  phase, characterizations can advantageously be made in situ by X-ray [25] or spectroscopy measurements [26]. Nevertheless, to our knowledge only few studies are made in line with real process conditions [27]. Even if FTIR experiment is easiest ex-situ spectroscopy technique available, the presence of relatively close characteristic peaks requires carefully conclusions and to correlate the results with other tests [28]. Furthermore, this technical analysis presents some limitations for specific multilayer materials.

On the other hand, characterization of piezoelectric behavior is essential to improve the design of textile energy harvester and to understand all important phenomena occurring during the processing of PVDF fiber. Numerous varied experimental methods based on direct or converse effect exist to determine piezoelectricity efficiency [29]. Among all these techniques Berlincourt type piezometer is a useful

solution for measuring piezoelectric coefficient on films. On textile fabric however, because of its small deformations, this test only allows soliciting the structure and do not give the piezoelectric response of the fiber. The evaluation of piezoelectric behavior of the textile can be made by output voltage measurements under large deformation using DMA for instance [30]. However, processing such structures is time and material consuming and gives only an average value at the fabric scale. Textile structures can be considered as a multi-scale material and a characterization of the piezoelectric behavior at each scale is relevant. Currently, there is no standardized test at the fiber scale for the measurement of piezoelectric coefficients and for the correlation with the  $\beta$  phase. Piezoelectric force microscopy (PFM) characterization is a relevant technical analysis to locally determine the presence of polar phase in the fiber and to evaluate its piezoelectric coefficient [31, 32]. PFM analyses have been carried out on materials at the dedicated measuring scale of the apparatus, i.e. on nanofibers [33-36]. To our knowledge, only one study has been published on the characterization of a textile filament [37]. However, this monofilament with a diameter around 150  $\mu\text{m}$  is not totally representative of the classical diameters used in textile applications. Indeed, in the work described by Soin *et al.* such large monofilament has been chosen to confer rigidity to the spacer fabric produced [37]. The textile fibers used in the yarns have conventionally diameters of less than 100  $\mu\text{m}$  and consequently a lack of knowledge on electromechanical behavior for such 30  $\mu\text{m}$  diameter fiber is underlined. Thus, the aim of this study is to leverage of PFM analysis for the development of micro-scale PVDF textile fiber by locally probing the physical properties. These first experimental trials on such fibers are particularly interesting. Indeed, they can have important implications for many researchers working on the development of multicomponent piezoelectric fiber systems for energy harvesting where conventional analyses could not be sufficient.

## EXPERIMENTAL

## Materials

Commercial grade of PVDF was obtained by Solvay (PVDF Solef 1006, Solvay, Belgium). The average melt flow index (MFI) is 40 g/10 min (load of 2.16 kg at 230 °C) and the density is 1780 kg.m<sup>-3</sup> in the solid state. Before melt spinning, PVDF is dried in oven for 24 h.

## Sample preparation

Multifilament yarns were produced using a spinning device Spinboy 1 from Busschaert Engineering. Four successive temperature of the spinning screw extruder are 80, 190, 200, and 210 °C. A volumetric pump ensured the injection of the molten polymer into two parallel dies each containing 40 holes (diameter of 400 μm) with a flow rate of 35 cm<sup>3</sup>.min<sup>-1</sup> in order to obtain a continuous multifilament yarn which was cooled down by air and coated with a surfactant. Finally, the multifilament yarn was hot drawn between two series of rolls. The ratio between rotation speed of the second roll (R2) and the first roll (R1) is called Draw Ratio (DR). Previous study has shown that different DR can be obtained up to a DR of 3 (R1: 150 m.min<sup>-1</sup> and R2: 450 m.min<sup>-1</sup>) leading to β-phase higher than 50 %. Due to future works and in particular that the development of multicomponent fibers having layers with different behavior under drawing (use of various polymers with or without fillers), a relatively low DR has been chosen for this study.[38] In addition, this low DR seems to be an appropriate strategy to highlight the powerfull of the PFM tool to detect piezo-activity in such textile fiber with relatively low polar phase content.

## Characterization

### ***FTIR***

A Thermo Scientific FTIR ATR Nicolet Spectrometer was used for recording the infrared spectra of as-spun (DR = 0) and drawn (DR = 1.5) fibers of PVDF. To perform the tests, fibers are lined up on a ZnSe substrate and analyzed in reflection mode. The samples were scanned eight times in the range of 600 to 1500  $\text{cm}^{-1}$ .

### ***XRD***

XRD measurements were carried out using a PANalytical diffractometer with Bragg-Brentano geometry. The radiation source was Cu K $\alpha$  with a wavelength of 1.5418 Å. The scan rate was set at 1 second per step of 0.028 (2 $\theta$ ).

### ***AFM/PFM***

Local characterizations were carried out under environmental conditions with a MFP-3D atomic force microscope from Asylum Research (USA). Surface morphology was characterized in AC mode, while PFM analyses were performed with Pt/Ir conductive tip coating (Nanosensors PPP-EFM probes,  $k \sim 2.8 \text{ N/m}$ ) and ground conductive substrates [39, 40]. Dual AC resonance tracking (DART) PFM was employed to enhance the piezoelectric signal [41]. The PFM probe was calibrated by using the GetReal procedure from Asylum Research, where both inverse optical lever sensitivity (InvOLS) and spring constant of the cantilever are calibrated in one step without touch the surface, then we probed our reference sample ( $\text{Pb}(\text{Mg}_{1/3}\text{Nb}_{2/3})\text{O}_3\text{--PbTiO}_3$  thin film) with known  $d_{33}$  piezoelectric coefficient.

## **RESULTS AND DISCUSSION**

### **Charaterization of polar phase by FTIR and XRD**

Absorbance spectra recorded during FTIR experiments for PVDF as-spun fiber (DR = 0) and fiber drawn with a ratio of 1.5 and XRD diffraction patterns of the same fibers are shown in figure 1. In the FTIR spectra, the characteristic peaks of the  $\alpha$  phase of PVDF at wavenumbers 760, 795, 855 and 976  $\text{cm}^{-1}$  can be observed for all both fibers. It can be seen from the spectra that drawn fiber (DR = 1.5) has distinct shoulder and peaks at wavenumbers 1234 and 1279 as well as 840  $\text{cm}^{-1}$ . The shoulder suggests the presence of  $\gamma$ -phase while the peaks represent  $\beta$ -phase of PVDF or more generally the polar phase ( $\beta/\gamma$ ) for this at 840  $\text{cm}^{-1}$  [42]. All these results are in agreement with previously reported studies on influence of drawing during the spinning process of PVDF. Stresses applied during the fiber production can transform the  $\alpha$ -phase of PVDF into  $\beta$  and/or  $\gamma$  polar phases [22, 38]. Usually XRD patterns are used to confirm the results obtained from FTIR tests. However, in the diffractogram of both fibers (drawn and undrawn) no characteristic diffraction peaks of  $\beta$  (peak at  $20.7^\circ$ ) and  $\gamma$  phases are present. The first peaks at  $2\theta = 17.9^\circ$  and  $2\theta = 18.4^\circ$  corresponds respectively to the (100) and (020) diffraction planes while the last peak at  $2\theta = 20.1^\circ$  is linked to the (110) plane. These two conventional analysis techniques show some of their limitations such as a lack of sensitivity and an average analysis of the considered systems. In these cases, the use of specific tools to perform local analyzes can be very relevant.

[Fig. 1 about here]

### **Charaterization of piezoresponse**

For PFM measurements, fibers were directly deposited on conductive disk with carbon paste in-between and the nano-probe tip was scanning individual fiber, as illustrated on figure 2a. Optical image of an individual fiber with 35  $\mu\text{m}$  in diameter is shown on figure 2b, while the characteristic morphology of the fiber is presented in figure 3a, as measured by AFM. No preferential direction is observed for such low draw ratio (DR = 1.5). The nanoscale tip was used as movable top electrode at the surface of the fiber in order to locally probe both the electromechanical activity and the ferroelectric polarization. The out-of-

plane PFM amplitude image shown in figure 3b displays contrasts corresponding to the local deformation of the fiber under electric field. Correspondingly, a strong contrasted out-of-plane PFM phase image is obtained suggesting several orientations for the polarization within the fiber, as seen on figure 3c. The obtained domains present sub-micrometer sizes, which could be correlated by the relatively low content of the  $\beta$ -phase resulting in small crystallites size. Black and white regions can be attributed to the out-of-plane component of the "upward" and "downward" polarization states, respectively, as related to the negative piezoelectric coefficient of the PVDF polymer. The various contrasts observed on both amplitude and phase PFM images reveal randomly oriented as-grown domains, which can be explained by the non-preferential orientation for polymer chains and crystals in the fiber at low draw ratio [43] correlated by the absence of preferential direction on AFM image. Besides, there is no correspondence between the topographic image (figure 3a) and the PFM signals, indicating that the nature of the contrasted regions is related to the intrinsic electromechanical activity of the fiber and not to the cross-talk with the topography (see figure 3d), to the electrostatic effect or to surface reactions. Therefore, these as-grown domain patterns clearly evidence the local piezo-/ferroelectricity into the fiber, demonstrating the existence of the polar crystalline phase. Besides, it is worth noting that the apparition of electro-active zones is for unpoled polymer fibers, as already observed in such PVDF-based fibers [33-35].

[Fig. 2 about here]

[Fig. 3 about here]

Switching behavior and vibration amplitude were then probed by recording phase and amplitude piezoresponse hysteresis loops, respectively. DC bias voltage applied to the fixed probing tip was swept between  $\pm 40$  V, i.e. high voltage PFM module was used to switch the polarization since the bulk coercive field of PVDF is known to be large (50 MV/m) [44]. Remnant signal (off-field mode) was measured in order

to eliminate electrostatic effect for the benefit of the electromechanical contribution [45, 46]. The figure 4 exhibits the piezoloops recorded over the free surface of the fiber. A square hysteretic behavior and two stable states of opposite polarization are observed on the phase loop pointing out the ferroelectric nature of the PVDF fiber. Indeed, a near  $180^\circ$  switching is observed for the domain phase under the DC bias voltage (the slight lower phase shift can be due to a small remaining electrostatic contribution). On the same figure is displayed the amplitude loop simultaneously measured showing a typical butterfly-shaped due to the inverse piezoelectric effect with clear saturation for higher applied voltages. These local PFM spectroscopic measurements reinforce the presence of the polar phase which presents ferro- and piezoelectric properties. As a remark, we note a slight asymmetry in switching process, as indicated by the shift of the piezoloops toward negative voltage values. This asymmetry is characteristic of an imprint behavior and point out the existence of an internal built-in electric field in the fiber [47, 48]. The main reason is probably our strong asymmetric configuration, for which the top and bottom electrodes are constituted of Pt-Ir nanometric probe and conductive disk covered with carbon paste respectively, leading to different boundary conditions. We can also note that the shape of the amplitude loop is slightly asymmetric. This asymmetric shape can be also explained by the difference between the work functions of the tip and the bottom electrode, resulting in an internal bias field [49]. From the phase PFM loop, we determine a coercive voltage of 15 V. Considering the fiber diameter of  $35\text{ }\mu\text{m}$ , we estimate a coercive field of 430 kV/m. This value is much lower than 50 MV/m in the case of bulk PVDF mentioned above. The difference can be explained by the micrometric size of our fiber which enables to decrease the voltage required to switch the polarization. Indeed, for such electroactive polymers with relatively high coercive field, a possible way to decrease the switching voltage is to synthesize these compounds as thin film form [50]. In addition, the nanometric size for ferroelectric domains seen on the figure 3c can disturb the cooperative coupling among these domains, leading to a decrease of the spontaneous polarization and thus to a reduction of the coercive voltage [51]. Also, such small ferroelectric domains are more efficiently switched

by external electrical bias, leading to narrow hysteresis loops. On the other hand, we have to keep in mind that the physical processes considered for measuring the coercive field are different when using a movable tip at the nanoscale level and planar electrodes at the macroscopic scale. From the amplitude loop, the higher deformation at the maximum bias voltage is about 600 pm, but it is enhanced by the resonance (DART PFM method). In addition, for such polymer material, the loss is considered as pretty high, which leads to a low quality factor  $Q$  of around 10 [52-54]. Considering the amplitude  $A$  detected by PFM is proportional to  $A = d_{33}V_{ac}Q$ , [52, 53] and the driving voltage used for the piezoloops measurements was  $V_{ac} = 5$  V, we can estimate the effective piezoelectric coefficient as  $d_{33} = 12$  pm/V (absolute value). This value is lower than others reported for PVDF-based fibers and determined by similar PFM analyses. Indeed,  $d_{33}$  coefficient of 30 pm/V was measured for  $\beta$ -phase PVDF fiber, which reaches 35 and 54 pm/V when added by carboxyl functionalized multiwall carbon nanotubes (CNTs) and silver decorated CNTs respectively [55]. M. Kanik *et al.* obtained an effective piezoelectric constant of 58.5 pm/V for  $\gamma$ -PVDF nanoribbons [36] while  $d_{33}$  of 45-50 pm/V was determined for BaTiO<sub>3</sub>/ $\beta$ -PVDF composite fibers [34]. However, for all these studies, the PVDF-based fibers are in the nanometric size range, which can reasonably explain the higher  $d_{33}$ , as compared to the microscopic size of our fibers. It is known that nanoscale confinement induces very high piezoelectric response, mainly due to the larger contribution of the domain wall motion in such nanofibers [36, 56-60]. In addition, the  $\beta$ -phase content in our fibers fabricated at DR = 1.5 is not optimized (the  $\beta$ -phase is estimated around 15%) compared to higher DR [38] limiting the piezo-active volume probed by the AFM tip and leading to lower electromechanical activity. However, beyond this value of 12 pm/V, the PFM technique was shown to be suitable to demonstrate the electromechanical response in such individual micrometer-scale PVDF textile filament, evidencing the existence of  $\beta$  and  $\gamma$  polar phases, which is essential in view of fabrication of complex multicomponent fibers.

[Fig. 4 about here]

## CONCLUSIONS

As-spun (DR = 0) and drawn (DR = 1.5) fibers of PVDF have been produced by melt spinning process and characterized during FTIR, XRD and PFM experiments. FTIR spectra revealed small intensity peaks than can be attributed to the polar phase of PVDF for samples having a draw ratio at 1.5. Nevertheless, the presence of such crystalline form could not be confirmed by XRD. It seems that the transformation of  $\alpha$  into  $\beta$  phase during the spinning process is too low to present a sensitive signal. The feasibility and the interest of PFM analysis on such fiber was investigated as a first step of the development of more complex piezoelectric fiber, *i.e.*: multicomponent fiber with concentric layers of different polymers around the PVDF. Different problems such as adhesion between the components will limit the drawability of final fiber and the possible conversion of crystalline phases of PVDF. Thus, it is supposed to obtain weak responses of the piezoelectric behavior of PVDF with conventional analyses. In this context, PFM can be relevant for future development of piezoelectric PVDF textile filament. In this work, PFM analyses have revealed phase and amplitude PFM contrasts for pristine regions over the surface of the fiber without correlation with the associated AFM morphology, which are related to the polar  $\beta$ -phase of the PVDF. From piezoresponse loops, local effective piezoelectric constant of about 12 pm/V was determined, evidencing the electromechanical behavior of the individual fiber at the local scale. These performances are required for considering such micro-scale functional fibers in textile applications.

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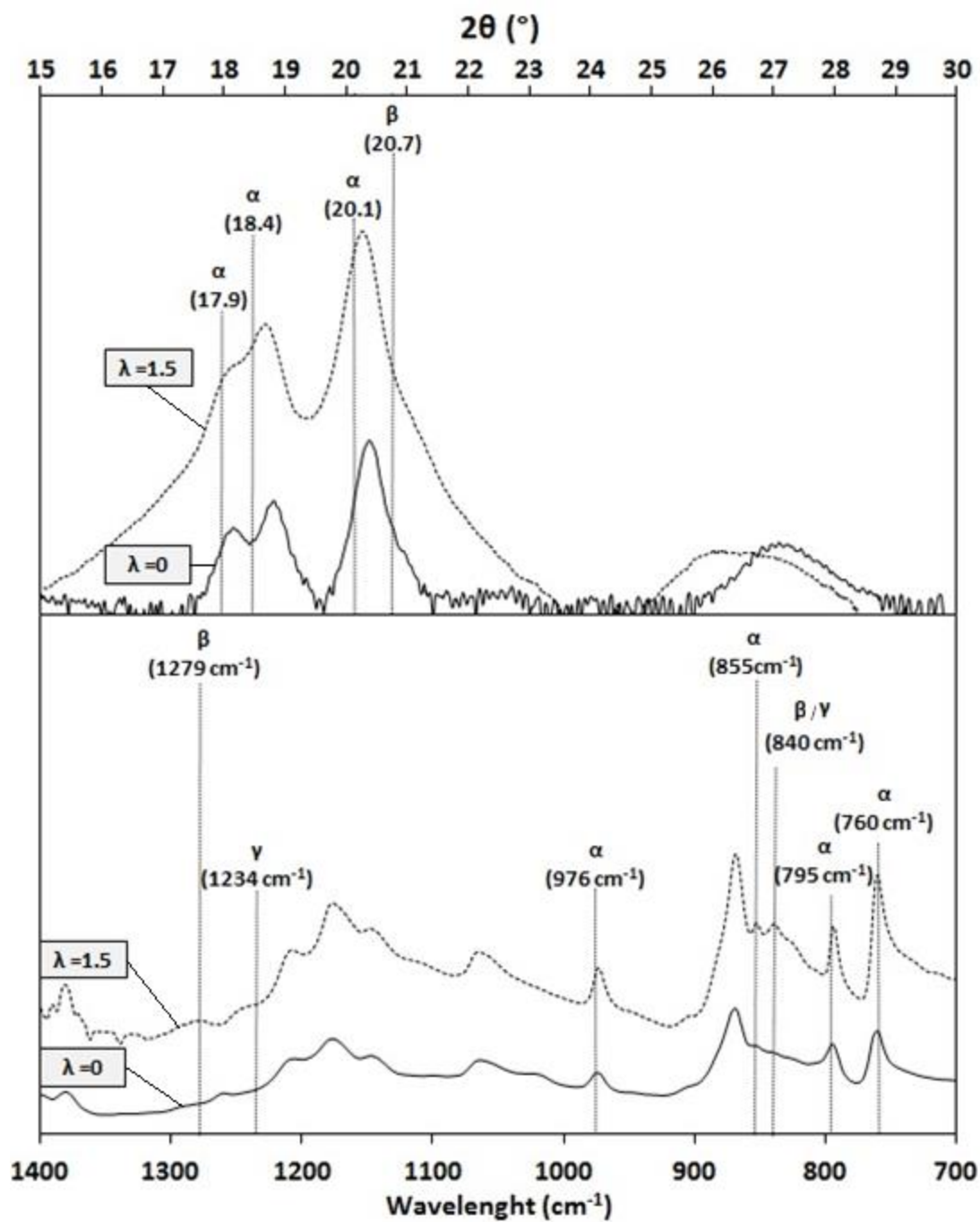
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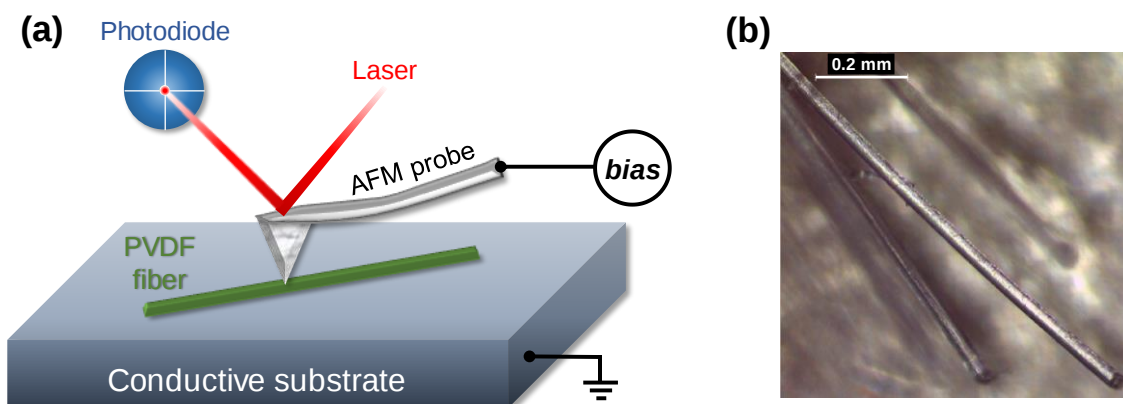
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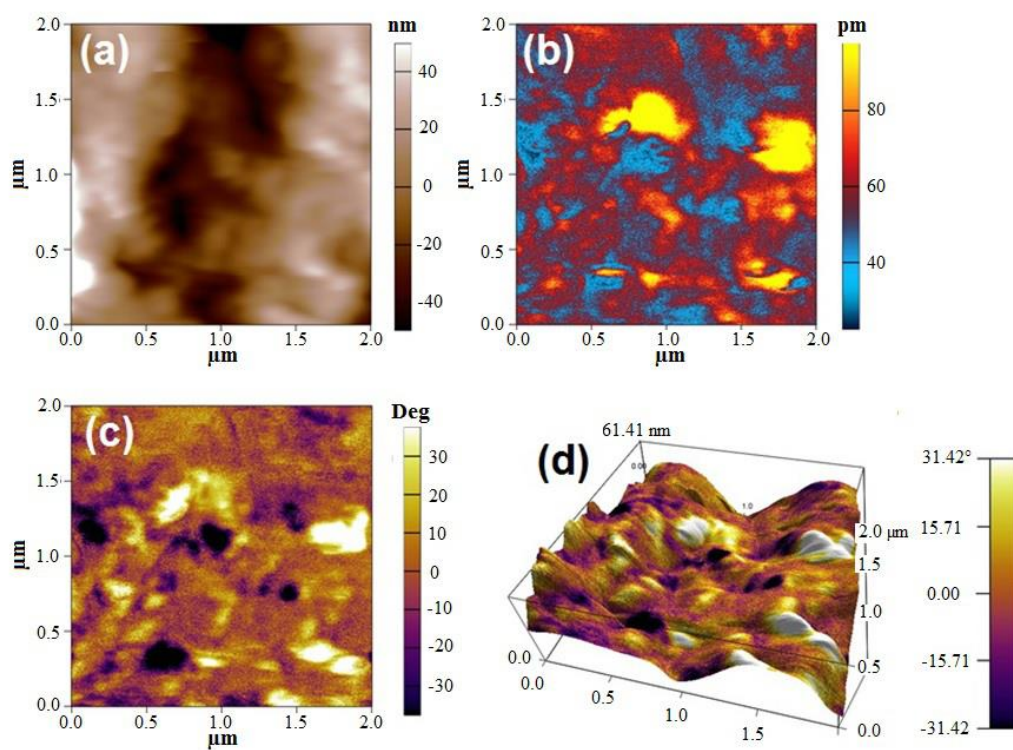
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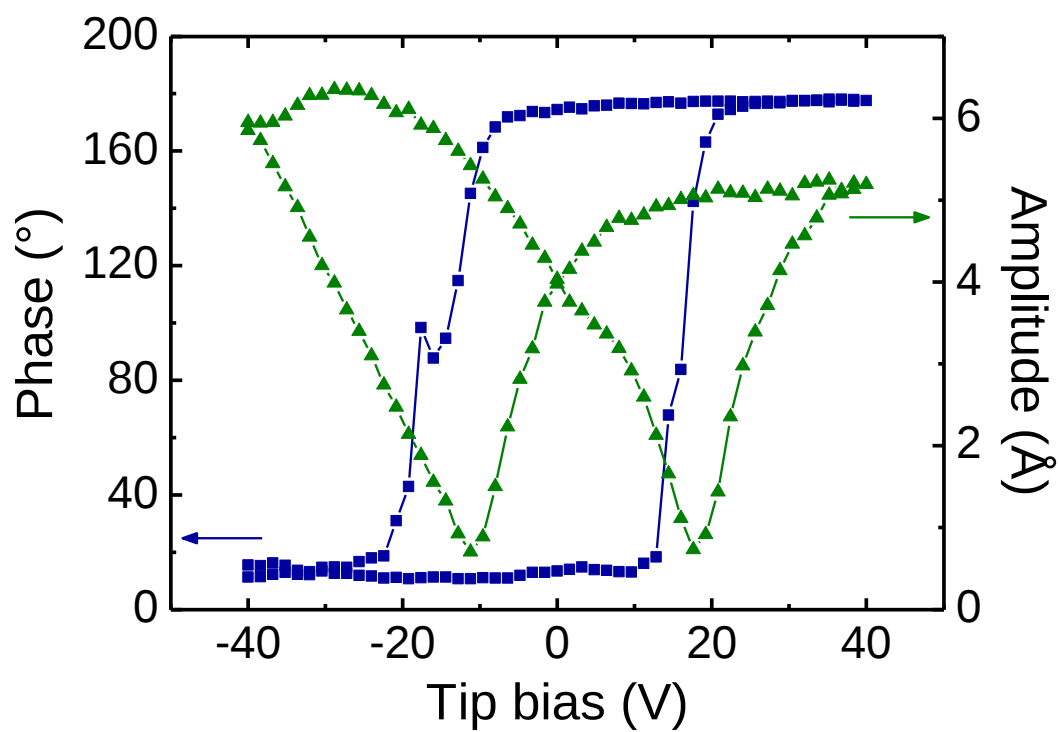
**Figure 1.** FTIR (bottom) and XRD patterns (up) of as-spun fiber (DR = 0) and fiber drawn with ratio of 1.5.



**Figure 2.** (a) Experimental setup for probing the local electromechanical response of single PVDF fiber (DR = 1.5); (b) optical microscope image of the PVDF fiber of 35  $\mu\text{m}$  in diameter.



**Figure 3.** (a) AFM morphology, (b) PFM amplitude and (c) PFM phase images simultaneously recorded on the PVDF fiber. (d) Phase piezoresponse overlaid on top of the 3D topography showing no correlation between the two signals.



**Figure 4.** Piezoresponse phase and amplitude loops measured on the PVDF fiber (DR = 1.5).