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Conformal Single-layer Encapsulation of PEDOT at Low Substrate Temperature

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Abstract

In this work, we demonstrate a single-layer encapsulation method for poly(3,4-ethylenedioxythiophene) (PEDOT). This method is achieved by initiated chemical vapor deposition (iCVD) process, which is scalable and employs solvent-free and low-substrate temperature conditions. The encapsulant used, poly(divinylbenzene-co-maleic anhydride) (PDVB-MA), was first time synthesized via vapor phase process. This cross-linked iCVD polymer can be rapidly deposited (40 nm min$^{-1}$) with uniform and conformal morphology. In the test of PEDOT degradation, the encapsulation extended the half-life of PEDOT to 900 hours at 30°C in air, which is more than ten times of the counterpart without encapsulation.

Keywords

iCVD; PEDOT; degradation; cross-linked polymer; conformal encapsulation
1. Introduction

Conducting polymers are rapidly developing as electromaterials used in organic solar cells, organic light-emitting diodes (OLEDs), and biomedical applications.[1-6] They are attractive due to their easy processing, flexibility, low cost and high porosity, which is important for many commercial applications. Among all organic conducting materials, PEDOT has shown a great promise as a result of its high conductivities (>1000 S cm\(^{-1}\).[7, 8] However, like many organic materials in this field, PEDOT is chemically unstable. The instability of PEDOT, which limits its application in industry, registers as the deterioration of its electrical conductivity in air.[9-12] Thus, there is great need to improve the stability of PEDOT. PEDOT in air can undergo multiple degradation mechanisms. Previous studies indicated oxidization by permeated oxygen can decrease the conductivity of PEDOT,[10] and water vapor adsorption resulted insulating patches may also increase the resistivity of PEDOT.[13] At elevated temperature, thermal shrinkage of the conducting area can also cause conductivity loss of PEDOT.[9] Last but not least, scientists found sizes of dopant molecules in PEDOT films have a large impact on its conductivity loss, which suggests that a loss of dopants is also a mechanism of PEDOT degradation.[14, 15] Previous reports indicated that acid-rinsed PEDOT has enhanced stability as well as increased conductivity as the removal of the excess hygroscopic oxidant and improved chain packing, which provides a better barrier for oxygen and moisture.[15] However, the acid-rinse cannot prevent escape of dopants, although it can reduce the degradation caused by permeated oxygen or water vapor. Therefore, a cross-linked network preventing loss of dopants may be an efficient way to encapsulate and enhance the stability of acid-rinsed PEDOT.
Previous achievements on the encapsulation of PEDOT and its copolymers include parylene\cite{16, 17}, silicon nitride\cite{4}, and et al.. These materials have been proved to provide excellent barrier to water vapor and oxygen. However, there is still a need of a solvent-free and low-temperature encapsulation method, which is simple to apply and easy to scale up, since PEDOT applications are usually sensitive to heat, solvent, and plasma. iCVD is a solvent-free deposition method for conformal and functional polymer thin films. In this process, monomer and initiator flow into a vacuum chamber, passing through a hot filament, where initiators get thermally activated. On a cooled substrate beneath the filament, adsorbed monomers and radicals from activated initiator react and form a uniform layer of polymer thin film. In iCVD, thermal activation and polymerization happened on different sites, so the temperature of substrate can stay cool, and functional groups of the monomer can be retained. Films deposited via iCVD polymerization are highly conformal and uniform\cite{18-21}. Moreover, the iCVD process is scalable and easily integrated with other techniques, such as roll-to-roll processing\cite{22} and plasma enhanced CVD\cite{23}. These unique advantages make it an ideal way of encapsulation.

2. Material and Method

2.1 PEDOT synthesis

PEDOT was deposited directly on glass substrates via an oxidative chemical vapor deposition process (oCVD) described elsewhere\cite{15, 20}. Here, we only summarize key details. 3,4-ethylenedioxythiophene (EDOT, Aldrich 97\%) vapors was delivered to vacuum chamber at flow rate of \~5 sccm. FeCl$_3$ oxidant (Sigma Aldrich, 99.99\%) was evaporated from a crucible, which is heated to 170 °C with a constant heating rate of
1.5 °C min⁻¹. Glass substrates at a substrate temperature of 150 °C were exposed to monomer and oxidant under the chamber pressure of 10⁻⁴ Torr for 40 minutes. After synthesis, PEDOT samples were rinsed by hydrobromic acid (HBr, Aldrich 48%, diluted with deionized H₂O to 5 mol L⁻¹) for 5 minutes then followed by drying for 30 minutes before a final rinse with methanol. All rinsing steps were done in ambient conditions. Two 4mm-wide strips of silver (Alfa Aesar, 1~3 mm particles, 99.9999%) were thermally evaporated onto edges of PEDOT film in order to avoid damage to encapsulation layer during measurements of resistance. The resistance (R) of PEDOT samples was measured by a multimeter (Keithley, model 2000, 4-wire mode), and then converted to conductivity (C) using Eq. 1.

\[ C = \frac{L}{R \times W \times T} \quad (1) \]

where L, W, and T represent length, width and thickness of PEDOT film, respectively.

2.2 Encapsulation

The PDVB-MA thin film was deposited directly on top of the PEDOT film at a rate of 20~40 nm min⁻¹ using the iCVD process described elsewhere.[21] We only summarize key details in here. The deposition pressures (0.27 Torr and 0.67 Torr) were maintained using a throttling butterfly valve. The substrate temperature was controlled at 25 °C. Two monomers, divinylbenzene (DVB, Sigma Aldrich, 80%) and maleic anhydride (MA, Sigma Aldrich, 99%) were heated at 60 and 80 °C, respectively. Monomers were delivered into the chamber at ~1.0 sccm controlled by needle valves. The initiator, tert-butyl peroxide (TBPO, Sigma Aldrich 98%), was delivered by a mass flow controller (MKS Instruments, model 1479) at ~1.0 sccm under room temperature.
The filament array (Chromaloy O resistance alloy, Goodfellow) was heated to 270 °C in order to activate the initiator. An in-situ interferometry (JDS Uniphase, 633 nm HeNe laser) was used to monitor the film growth. Thicknesses of deposited PDVB-MA films were measured on bare silicon wafers (Wafer World, Inc., aside of PEDOT samples in the reactor) by a variable angle spectroscopic ellipsometer (VASE, J. A. Woollam M-2000) at incident angles of 65°, 70°, and 75°. Fourier transform infrared (FTIR) spectrum was measured by Thermo Nicolet Nexus 870 spectrometer. Morphology of films was investigated by scanning electron microscope (SEM, JEOL 6010L) and atomic force microscopy (AFM, Digital Instruments, model D3100). Root–mean–square (RMS) roughness of films was measured on a 5 μm × 5 μm surface area using AFM.

3. Results and Discussion

3.1 Characterization of PDVB-MA

We use FTIR spectrum to confirm the structure of the synthesized copolymer. In Fig. 1, strong absorptions at 1780 and 1870 cm\(^{-1}\) are due to stretching of carbonyl groups from the 5-member ring of maleic anhydride.[24] Absence of strong absorption at 903 cm\(^{-1}\) indicates vinyl groups from DVB were reacted during the polymerization.[25] Absorptions around 2900 cm\(^{-1}\) are from symmetric and asymmetric C-H stretching in the backbone of the newly formed polymer chain of PDVB-MA.[26, 27]

Cross-linked polymer networks are useful in many applications, including but not limited to sensors, micro-fluidic devices, shape memory films, and electronics protection.[28-30] PDVB and its copolymers have been used in many applications of cross-linked networks.[31-33] Previous report indicated copolymerization of styrene and MA has much faster deposition rate than that of polystyrene homopolymer.[34] Similar
results were observed in copolymerization of DVB and MA. Compared to the deposition rate of PDVB homopolymer in iCVD process (~4 nm min⁻¹), the PDVB-MA copolymer shows a rapid deposition rate (up to 40 nm min⁻¹). This higher deposition rate would correspond to lower overall cost. In order to clearly show the role of MA in accelerating the deposition, it is necessary to define a fundamental characteristic of the iCVD process, \( P_M / P_{sat} \), which is the ratio of the partial pressure of monomer to its saturation vapor pressure at certain substrate temperature. This ratio is directly related to the equilibrium surface concentration of adsorbed monomers based on Brunauer, Emmett, and Teller (BET) isotherm.[35] Previous report indicated 0.4~0.7 is an optimal \( P_M / P_{sat} \) range for rapid iCVD processes of vinyl monomers.[36] We summarize deposition rate of PDVB-MA at different \( P_M / P_{sat} \) of MA in Fig. 2. In the range of 0~0.7, \( P_M / P_{sat} \) of MA shows linear relationship with the deposition rate of the copolymer, which indicates participation of MA accelerated the reaction. In our experiments, the fastest deposition rates (40 nm min⁻¹) occurred at \( P_M / P_{sat} = 0.7 \). Based on following BET equations[37, 38] (Eq. 2 and Eq. 3), we can estimate an adsorption energy of 24.8 kJ mol⁻¹, which is in the typical range of 20 to 80 kJ mol⁻¹ for physisorption of small molecules.[39]

\[
V_{ad} = \frac{V_{ml}C(P_M/P_{SAT})}{(1-P_M/P_{SAT})(1-(1-C)(P_M/P_{SAT}))} \quad (2)
\]

\[
C = \exp\left(\frac{E_{ad}}{RT}\right) \quad (3)
\]

where \( V_{ad}, V_{ml}, C, \) and \( E_{ad} \) represent adsorbed volume, monolayer adsorbed volume, the BET constant, and adsorption energy, respectively. Besides the effect of \( P_M / P_{sat} \), temperatures of the filament (\( T_f \)) can also affect the deposition rate in the synthesis of PDVB-MA. Previous publication indicates the degree of decomposition of TBPO increases as \( T_f \) increases when \( T_f < 270 ^\circ C \), and higher \( T_f \) results methyl radicals[40],
which is more reactive than the tert-butoxy radicals and therefore accelerate the free-radical polymerization.[41]

3.2 Conformality, uniformity, and surface roughness

Describing the ability to encapsulate the entire surface topography, conformality is an important standard to judge an encapsulation method. Conformal coating is highly desired since it can bring additional functionalities without changing the original morphology, which is important in many micro fabrication steps. In this work, we use cross-sectional SEM to demonstrate the overall profiles for PDVB-MA films rapidly growing (35 nm min\(^{-1}\)) inside a trench feature (aspect ratio = 4.2). Fig. 3 shows good thickness uniformity over the entire trench feature, which indicates the conformal nature of iCVD synthesized PDVB-MA films. Another important parameter of thin film coating is the uniformity of entire coating area. In order to demonstrate this uniformity, we deposited 350 nm PDVB-MA film on a bare silicon substrate and measured thicknesses at different positions in a 8 × 8 cm area by VASE. In Table 1, thicknesses at five different positions in the area are all within 5% of the averaged thickness, which indicates the iCVD deposition of PDVB-MA is uniform all over the entire 8 × 8 cm area. AFM is also used to characterize the surface morphology. Fig. 4 a) shows the AFM image of the bare acid-rinsed PEDOT, and Fig. 4 b) shows the same sample encapsulated with 200 nm PDVB-MA thin film. The RMS surface roughness calculated from a 5 × 5 μm area dropped from 2.35 nm before encapsulation to 0.63 nm after encapsulation. The reduced roughness suggests the PDVB-MA encapsulation smoothed the surface, which helps to reduce the absorption of water and oxygen.
3.3 PEDOT degradation

We tested the stability of PEDOT with and without encapsulation in air at 30°C. In order to characterize the performance of encapsulation, we define the time when PEDOT lose half of its original conductivity as its halflife. In Fig. 5, the PEDOT sample without encapsulation shows a halflife of 118 hours. With PDVB-MA encapsulation, the halflife of PEDOT can be extended to more than 850 hours, which is a 7-fold increase. We observed encapsulation with PDVB-MA films with different thicknesses, but we did not observe any obvious effect of thickness in the range of 200 nm to 800 nm, and a 4-fold increase in encapsulate thickness only brings less than 5% increase of the halflife. This suggests that the 200 nm film of PDVB-MA is sufficient to prevent loss of dopants, which is considered as one of major degradation mechanisms of acid-rinsed PEDOT. Also, the shape of curves in Fig. 5 indicates at least two mechanisms were involved in the degradation of PEDOT as the flat stage shown between 0 to 170 hours. However, it is still not enough to make a clear conclusion on the role of different mechanisms in PEDOT degradation. Further experiments are being carrying out to address this issue.

4. Conclusion

We have successfully synthesized PDVB-MA via iCVD process for the first time. This cross-linked polymer performs as a good single-layer encapsulant for PEDOT, and extends the halflife of PEDOT to 900 hours at 30°C in air. The rapid encapsulation method features with solvent-free and low-substrate-temperature conditions, which is ideal for many fragile PEDOT applications. The effect of monomer saturation ratio on the deposition rate is discussed, and a deposition rate of 40 nm min⁻¹ was achieved at P_M / P_{sat}
(MA) = 0.7. For morphology of the deposited films, SEM, VASE, and AFM investigations indicate the iCVD PDVB-MA film is conformal, uniform, and smooth, respectively. Overall, iCVD is an ideal method for conformal, solvent-free encapsulations at low substrate temperature.

Acknowledgement

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Reference


Figure 1. FTIR spectrum of 200nm PDVB-MA film on silicon substrate. The insert panel shows the structure diagram of PDVB-MA.
Figure 2. Effect of $P_M / P_{sat}$ (MA) on the rate of deposition at $T_{substrate} = 25^\circ C$ and $T_{filament} = 270^\circ C$. Error bars on data points are standard deviations observed in replicate measurements. The trendline is a linear fit of all data points.
Figure 3. Cross-sectional SEM image of iCVD deposited PDVB-MA on a trench feature.
Figure 4. AFM images of a) PEDOT on a silicon substrate; b) 200 nm PDVB-MA encapsulated PEDOT on silicon substrate.
Figure 5. Degradation curves of PEDOT samples at 30°C in air.