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Improvement of electrical properties of ITO thin films by melt-free ultra-short laser crystallization

N Farid^{1,*}, A Sharif¹, R K Vijayaraghavan², M Wang³, H Chan⁴, A Brunton⁴, P J McNally², K L Choy³ and G M O'Connor¹

E-mail: nazar.farid@nuigalway.ie

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Abstract

We describe a novel solid state crystallisation method for optimising a thin film transparent conductive oxide when deposited on flexible polymer substrates. The method is based on ultra-short non-thermal laser sintering of indium tin oxide (ITO) thin films. In this study, we used commercial ITO thin films deposited on a flexible polyethylene terephthalate substrate with a relatively low melting temperature compared with ITO on glass. We demonstrate the use of laser scanning with high pulse overlapping at fluences seven times less than the threshold required for melting/damage of ITO. The results confirm greater than four times enhancement in the mobility of charge carriers of ITO thin films after laser scanning and sheet resistance can be reduced up to 25%. There is no reduction in optical transparency observed in laser treated samples. Surface morphology and x-ray diffraction analyses confirm the improvement in crystallite sizes by laser sintering, resulting in a greater than 37% increase in grain size due to enhanced crystallization. Comparison of experimental and simulation based on a delayed two temperature model confirms that ITO thin film crystallization occurred at about one-third of the melting temperature of ITO.

Keywords: ultra-short laser crystallization, melt-free, ITO, OLED, PFDs

(Some figures may appear in colour only in the online journal)

1. Introduction

Indium tin oxide (ITO) is an n-type degenerate semiconductor with a wide band gap. Its high electrical conductivity ($<10^{-5} \Omega$ cm), good optical transparency in the visible (<85%) and near infrared regions, and good adhesion to many types

of substrates make it crucial in a wide variety of applications [1]. It has been widely used as a transparent conducting layer in modern optoelectronic technology such as organic light emitting devices [2], transparent touch flat panel displays (FPDs) [3], touch sensors [4] and solar cells [5]. This conducting transparent oxide mainly serves as an electrode and often as an antistatic electromagnetic interference shield or an electric heater in FPDs [6]. It has been used as an ohmic contact material because it has high surface current [7]. The electrical properties of ITO depend on its atomic arrangement,

¹ National Centre for Laser Applications (NCLA), School of Physics, National University of Ireland Galway, Galway, Ireland

² Advanced Processing Technology Research Centre (APT), Nanomaterials Processing Laboratory, School of Electronic Engineering, Dublin City University Glasnevin, Dublin 9, Ireland

³ Institute for Materials Discovery, University College London, London, United Kingdom

⁴ M-Solv Ltd, Kidlington, Oxford, United Kingdom

^{*} Author to whom any correspondence should be addressed.

composition, surface structure and on the amorphous and crystalline nature of the film. ITO exhibits favourable properties such as low resistivity and high optical transparency in the crystalline state but not in the amorphous phase.

ITO thin films can be deposited by chemical vapour deposition [8], evaporation [9], spray pyrolysis [10], and commercially used magnetron sputtering [11] techniques. All these techniques require a deposition temperature of at least 300 °C–500 °C; higher temperatures (400 °C–700 °C) are required for further annealing to reduce the resistivity and to achieve improved transparency. The requirement for high temperature limits the use of heat sensitive polymers as substrate materials [12]. In most of the applications, the ITO materials are first sputtered/coated in the amorphous phase on the substrate and then thermally annealed to at least 300 °C to change to the crystalline state [13].

Lasers have become an emerging and alternative tool for material processing and industrial manufacturing; high power short and ultra-short pulse lasers with MHz repetition rates are set to transform micro- and nano-scale fabrication. Laser technology has replaced conventional and rapid high-temperature thermal annealing as in the case of Si crystallization [14]. Excimer [15] and nanosecond lasers [15] have also been investigated in ITO crystallization; the crystallized phase results from melting, the thermal budget for which is not suitable for application on heat sensitive flexible substrates. On the other hand, ultra-short lasers are promising in material processing because of their extremely short pulse durations which offer a smaller heat affected zone and thermal damage to the substrate compared to nanosecond laser pulses. Recently ultrashort lasers have also been used for selective ablation [16], patterning [17, 18] and direct laser crystallization [19]. A twostep method was also used for crystalline ITO patterns on amorphous ITO (a-ITO) thin films by femtosecond laser irradiation followed by chemical etching [20].

In most of the applications, ITO thin film is deposited on a suitable substrate (i.e. glass or polyethylene terephthalate (PET)) and then crystallized and patterned to form complex and multifunctional electronic circuits. PET being transparent, flexible, lightweight, and cost-effective is one of the candidates in flexible display technology [21]. However, the lower melting temperature of PET does not allow for the high-temperature thermal treatments required for ITO crystallization. Therefore, crystallization of ITO on PET must be carried out at temperatures below 200 °C which is insufficient to optimize ITO layers with good electrical conductivity and transparency [22]. Thus, a low temperature process is needed. In this work, we have developed a low temperature ultra-short laser annealing process to improve the electrical properties of ITO films on heat sensitive substrates.

2. Experimental details

As received commercial ITO thin films (Visiontek) deposited on 0.5 mm thick PET substrates were used as a target material. Two different ITO thicknesses of 50 nm and 100 nm deposited on PET were investigated to enhance the conductivity by

ultra-short laser treatment. A femtosecond laser of pulse duration 500 fs at a wavelength of 1030 nm which can operate from single pulse to 300 kHz was used for ITO crystallization. The sample position was controlled using a high precision 3D computer-controlled stage (Aerotech). The sample was scanned using industrial grade Galvo scanning system (Scanlabs). The scanning system controls the pulse to pulse overlap on the sample by adjusting the speed of steering mirrors of galvanometer-based beam scanning system. Optical and scanning electron microscopy, and atomic force microscopy (AFM) were utilized for analysing the surface morphology of the laser scanned area and single pulse crater depth profiles. The van der Pauw method was used to measure the sheet resistance and electrical conductivity of ITO before and after the laser scanning. The crystallinity of the ITO films was quantified using Cu-K α x-ray diffraction (XRD); $\lambda = 1.54$ Å. Optical transmission was measured using UV-vis spectrometer (SHIMADZU UV-2600). The laser spot diameter at the focused position and the damage threshold were calculated experimentally using the Liu method [23]:

$$D^2 = 2\omega_0^2 \ln\left(\frac{F_0}{F_{\rm th}}\right),\tag{1}$$

where, $F_0 = 2E_{\rm p}/\pi\omega_0^2$ and $F_{\rm th}$ are the applied and threshold fluence, ω_0 is beam waist radius of the Gaussian shaped beam at the focus, $E_{\rm p}$ is absorbed laser energy, and D is the measured diameter of crater.

3. Results

The spot diameter at the $1/e^2$ position was calculated using the above relation and found to be 49.5 μ m. The fluence at which the ITO thin film started to change as observed by optical microscopy is defined as the damage threshold, while the value of fluence at which ITO films start to ablate/delaminate is defined as the ablation threshold. A plot of D^2 verses log of fluence is provided for the 50 nm and 100 nm thick ITO films in figure 1. For 50 nm thick film, there was not a significant difference between the damage and ablation thresholds, which were found to be $0.26 \,\mathrm{J\,cm^{-2}}$. A fluence of $0.28 \,\mathrm{J\,cm^{-2}}$ and $0.51 \,\mathrm{J\,cm^{-2}}$ were measured as damage and ablation thresholds, respectively, for 100 nm thick ITO. There were no measurable changes induced in the film below $0.26 \,\mathrm{J\,cm^{-2}}$ for 50 nm thick film and $0.28 \,\mathrm{J\,cm^{-2}}$ for 100 nm thick film and fluence values higher than this resulted in physical damage.

To understand the single pulse ITO thin film ablation, different fluences were used to ablate the thin film. Figure 2 shows the surface topographic AFM images of laser irradiated 100 nm ITO thin film with different fluences higher than the damage threshold. At first, with a single laser pulse at low fluence, damage starts to appear on the surface and then at higher fluence a dome-like structure appeared. The dome-like feature results in the ITO film being ablated or delaminated at fluences of \sim 0.26 J cm⁻² and 0.51 J cm⁻² for 50 nm and 100 nm, respectively, as provided in figure 2 for 100 nm ITO thin film.

To enable the improvement in the electrical properties using melt-free or all solid state crystallization, ITO thin films

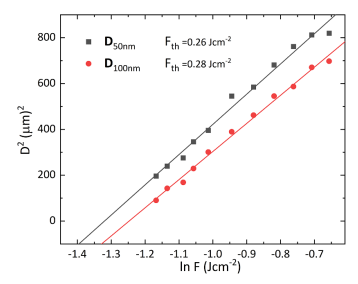


Figure 1. A linear relationship between squared diameter (D^2) and log of applied fluence for 50 nm and 100 nm thick ITO films. Data for 50 nm and 100 nm thick film is presented.

were scanned with high pulse overlapping and laser shots per area (SPA) while using a fluence approximately seven times smaller than the damage threshold for ITO. ITO thin films samples were scanned with 100 kHz laser repetition rate and 180 mm s⁻¹ laser scanning speed which gave 27 SPA and 96% pulse overlapping. The laser scanning parameters and fluence values are optimized to avoid any melting and visible damage on the ITO surface. The influence of laser scanning on the surface morphology of the ITO films was investigated by AFM. The topographic AFM images of the ITO films with thicknesses of 50 nm ((a), (b)) and 100 nm ((c), (d)) before and after laser scanning are shown in figure 3. The above provided AFM scanned area of 1 μ m \times 1 μ m is selected randomly from 10 mm × 10 mm ITO thin films samples before and after the laser crystallization. Before laser irradiation, the ITO films were composed of tiny particles of approximately 20 nm diameter and 45 nm diameter for the 50 nm and 100 nm films, respectively. The morphology of the surface was changed due to the laser scanning of the ITO films. The AFM images suggest that nanoparticle sintering has occurred during laser irradiation as shown in figures 3(b) and (d). In this socalled sintering process, the formation of a conductive neck is envisaged to be formed between the nanoparticles resulting in bigger particle sizes within the ITO films. The smaller particles agglomerate to form a larger particle with sizes roughly 60 nm for 50 nm thick film and 80 nm for 100 nm thick films. It is evident from the AFM images that the average particle sizes of the thin films have been increased after the laser scanning for both 50 nm and 100 nm thin films. For 50 nm thin ITO film, root mean square roughness is improved from 2.56 nm to 2.18 nm after the laser scanning while for 100 nm thin film the roughness increased from 1.87 nm to 2.13 nm because of the overall increase in the particle size. The changes in surface roughness might affect the surface reflection and transmittance of ITO films. It is evident from the AFM images that no melting of ITO thin film is observed in the laser scanning region and the only increase in the grain/particle sizes have occurred by the solid diffusion process.

Surface morphology and electrical properties are interconnected in the case of many thin films. To study the improvement in electrical properties of ITO after laser scanning, a commercial Four-Point Probe System (Ossila UK) was used to measure the sheet resistance of ITO thin films before and after the laser scanning. It was found that the sheet resistance of the 50 nm thin film decreased from 72 \pm 0.5 Ω sq $^{-1}$ to 54 \pm 0.2 Ω sq $^{-1}$ while that for the 100 nm film decreased from 52 \pm 0.2 to 40 \pm 0.15 Ω sq $^{-1}$.

In order to investigate if there is a phase change occurring in increasing the grain or particle size which leads to the reduction in the sheet resistance, Bragg-Brentano XRD analysis of the ITO thin films was performed before and after the laser scanning. Figure 4 shows XRD pattern of irradiated films in comparison to unirradiated films. The XRD analysis confirms the improvement in crystallinity as the peaks from the irradiated ITO film appear to have a reduced full width at half maximum (FWHM). The peak at 46.5° belongs to the PET substrate. The peak at 30.28° is the only ITO dominant peak that is observed; it is more prominent for the 100 nm thick film compared to 50 nm film. This peak corresponds to the coherent scattering from (2 2 2) reciprocal lattice vector in cubic In₂O₃. It is well known that peakshifting to higher angles indicate lattice contraction due to strain relaxation due to an increase in the substitutional Sn concentration because of the smaller ionic radius of Sn⁴⁺ compared to In³⁺ [24]. In our case, there was a slight peak shift from 30.28° to 30.29° after the laser crystallization. It predicts that the laser annealing could not induce changes of substitutional to interstitial Sn atoms in the ITO crystal structure upon laser annealing. The coherent scattering from ITO (2 2 2) is shown in more detail in inset of figure 4. This confirms that its FWHM decreases by 37% after laser scanning. The Scherrer relation can be used to estimate the grain size of the film [25]. It is found that the grain size calculated by this way correlates with AFM topographic results for ITO thin films before and after the laser scanning, indicating that the grain size increases from 19 nm to 30 nm. This increase in grain size is attributed to a corresponding increase in particle size by a laser sintering based process.

ITO thin films microstructures and optical properties are very sensitive to the electrical and crystalline state. The microstructural changes as a result of laser annealing not only affect the electrical but optical properties as well. Higher electrical conductivity of ITO can be achieved but often at the expense of optical transmission. The variation in optical properties such as transmission and band gap energy after the laser scanning were measured by UV-vis spectrometry. There is an overall small improvement in the transmittance of 50 nm thin film after laser scanning as shown in figure 5. For 100 nm thin film, transmittance improved from 85% to 87% in the UV and decreased from 81% to 79% in the visible region. In the visible region, the transmittance of ITO thin films depends on the degree of crystallinity as the grain boundaries in the thin films scatter the visible light, resulting in lower optical transmittance. The increase in grain size after the laser scanning resulted in a decrease in the number of grain boundaries in ITO thin

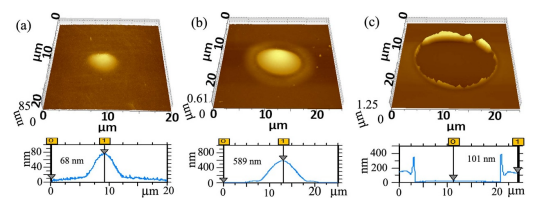


Figure 2. AFM topographic images of 100 nm thick ITO film with single fs laser pulse at (a) 0.36 J cm⁻², (b) 0.41 J cm⁻², and (c) 0.51 J cm⁻² and the images at the bottom are their respective surface profiles taken from the centre of crater.

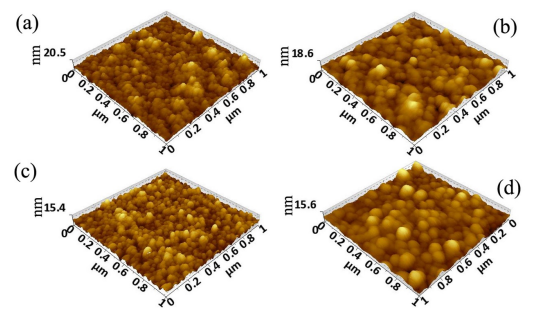


Figure 3. Topographic AFM images of (a) untreated 50 nm thick ITO film and (b) after laser scanning while (c) is untreated 100 nm thin film and (d) is after laser scanning. The samples were scanned with laser repetition rate of 100 kHz and at 0.04 J cm⁻² fluence with 180 mm s⁻¹ scan speed which gave 27 SPA and 96% overlapping. A hatching of 5 μ m was used.

films. Hence the increase in optical transmission after the laser scanning is due to the improvement in the crystallinity of the thin films as confirmed by AFM and XRD analyses. For the IR region, the interaction of incident radiation with the high density of free electrons in the material cause the decrease in transmission due to enhanced reflectivity. For the UV region, the absorption is due to the fundamental bandgap $(E_{\rm g})$ transition. The transparency of ITO thin films exhibits a sharp decrease in the UV region and a shift in absorption edge occurs after laser scanning.

3.1. Modelling

The dynamics of a free electron in ITO are metal-like and can be described by the Drude model. ITO is an interesting material as recent studies show it to have a nonlinear response [26]. It was found that the nonlinear response of ITO in the epsilon near zero regime is dominated by free

electrons. This nonlinear mechanism is well described by the so-called delayed two temperature model (TTM). The incident laser energy is absorbed, generates hot electrons with a non-thermal energy distribution and acts as a delayed heating source. Hence, the response of ultra-short laser interaction with ITO thin film is different from metals. This can be explained by three reasons. Firstly unlike metals, ITO has no interband transition resonance in the visible or infrared range of the optical spectrum; secondly the free-electron density in ITO is smaller than metals resulting in much smaller electron heat capacity, and thirdly the Fermi level is quite low in the conduction band (\sim 1 eV for ITO) due to the relatively small free-electron density [27]. After absorption of the laser pulse, the generated hot electrons acquire a nonthermalized energy distribution and act as a delayed source of heating while the lattice is considered cold. The overall dynamics of the conduction electrons and energy transfer to lattice when there exists a thermal non-equilibrium

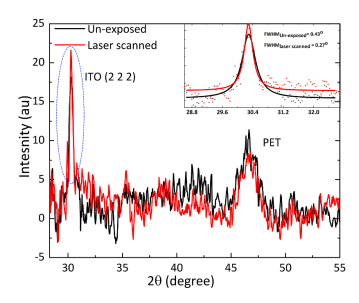


Figure 4. XRD patterns of unexposed and laser exposed ITO thin films of thickness 100 nm. The inset shows the peak fitting of ITO (2 2 2).

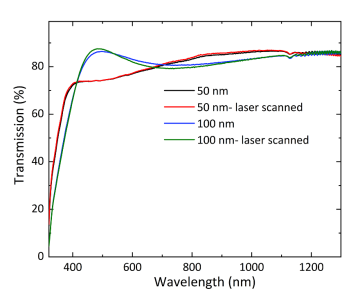


Figure 5. Optical transmittance of the 50 nm and 100 nm thick ITO thin films before and after laser scanning.

between electrons and phonons is described by a delayed TTM [26, 27]

$$C_{\rm e} \frac{\partial T_{\rm e}}{\partial t} = \nabla \cdot (k_{\rm e} \nabla T_{\rm e}) - G(T_{\rm e} - T_{\rm l}) + \frac{N}{2\tau_{\rm ee}}, \tag{4}$$

$$C_{1}\frac{\partial T_{1}}{\partial t} = \nabla \cdot (k_{1}\nabla T_{1}) + G(T_{e} - T_{1}) + \frac{N}{2\tau_{ep}}, \qquad (5)$$

$$\frac{\partial N}{\partial t} = -\frac{N}{2\tau_{\rm ee}} - \frac{N}{2\tau_{\rm en}} + S,\tag{6}$$

$$C_{\rm s} \frac{\partial T_{\rm s}}{\partial t} = \nabla \cdot (k_{\rm s} \nabla T_{\rm s}), \qquad (7)$$

where $C_{\rm e}$, $C_{\rm l}$ are electron and lattice heat capacity, $T_{\rm e}$ and $T_{\rm l}$ represents electron and lattice temperature, $k_{\rm e}$ and $k_{\rm l}$ are the electron and lattice conductivity, respectively, G ($T_{\rm e}$) is electron—phonon coupling factor, N is non-thermal electron energy density which is stored in excited electrons. Where $\tau_{\rm ee}$ and $\tau_{\rm ep}$ are the electron—electron and electron—phonon relaxation time and S is laser source term, respectively. $T_{\rm s}$, $C_{\rm s}$, and $k_{\rm s}$ are the lattice temperature, heat capacity and thermal conductivity respectively of ITO substrate.

Equation (7) represents the lattice heat diffusion from ITO thin film to the PET substrate. The energy transport at the ITO-PET interface is accounted by considering the interaction between electrons in ITO thin film with phonons in the PET substrate using the boundary condition as

$$(T_{\rm s} - T_{\rm e}) \Big|_{z=d} = R_{\rm es} \frac{\partial T_e}{\partial z} \Big|_{z=d},$$
 (8)

where $R_{\rm es}$ is the thermal boundary resistance.

The electron–phonon coupling factor $(G(T_e))$ describes the strength of energy transfer between the excited electrons and lattice vibrational modes of ITO and can be estimated as [28]

$$G(T_{\rm e}) = \frac{\pi^2 m_{\rm e}^* v_{\rm s} n_{\rm e}}{6\tau (T_{\rm e}) T_{\rm e}},$$
 (9)

where $m_{\rm e}^*$ is effective electron mass, $v_{\rm s}$ is speed of sound, $n_{\rm e}$ is free-electron density, and τ is the temperature dependent relaxation time. Electron–electron collisions lead to an energy relaxation within the electron gas called thermalization and the mean time required for electrons to change their states is termed as electron–electron relaxation time ($\tau_{\rm ee}$). Through this process of electron–electron collisions, the absorbed energy is redistributed among the free electrons resulting in electron gas thermal equilibrium and thus to a Fermi distribution [29]. In other words, τ_{ee} is measure of redistribution of energy to the electrons and is given by [30]

$$\frac{1}{\tau_{\rm ee}} = \frac{\pi (k_{\rm B} T_{\rm e})^2}{(8\hbar E_{\rm F})} + \frac{\sqrt{3}}{2\hbar \sqrt{E_{\rm F}}} \left(\frac{k_{\rm B} T_{\rm e}}{k_{\rm F} l}\right)^{3/2},\tag{10}$$

where, $k_{\rm B}$ is Boltzmann constant, $T_{\rm e}$ is electron temperature, \hbar is reduced Plank constant, $E_{\rm F}$ is Fermi energy, $k_{\rm F}$ is Fermi wavevector and l is electrons mean free path. Electron–phonon relaxation time ($\tau_{\rm ep}$) can be estimated by electron–phonon coupling factor and the temperature dependent specific heat capacity of free electrons as [26]

$$\tau_{\rm ep} = 2 \frac{C_{\rm e} \left(T_{\rm e} \right)}{G(T_{\rm e})}.\tag{11}$$

For T_1 much smaller than Debye temperature (T_D) , the number of interacting phonon modes and the lattice heat capacity (C_1) is a function of lattice temperature (T_1) and asymptotically approaches a constant value as T_1 approaches T_D , the temperature of highest mode of vibration of lattice. We used the temperature dependent values of C_e and C_1 in our model as provided in figures 6(a) and (b), reported in the supplementary information of [31].

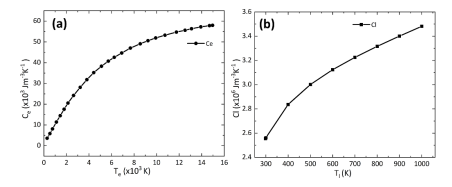


Figure 6. Temperature dependent (a) electrons heat capacity and (b) lattice heat capacity of ITO used in modelling.

The temporal and volumetric energy from the laser source of Gaussian profile can be defined as

$$S(\mathbf{r},t) = \frac{\alpha_{\rm eff} \times 0.94 \times F_0 \times (1-R)}{t_{\rm p}} \times \exp\left\{\frac{-2x^2}{\omega_0^2} - 4\operatorname{In}2\left(\frac{t-t_{\rm r}}{t_{\rm p}}\right)^2\right\} \exp\left(-z\alpha\right), \quad (12)$$

here F_0 is laser fluence, R is reflectivity of material, t_p is laser pulse duration, ω_0 is beam radius, t is time, and t_r is reference time. The wavelength dependent nonlinear effective absorption coefficient ($\alpha_{\rm eff}$) is estimated for 1030 nm wavelength using z-scan technique and is given as [32]

$$\alpha_{\text{eff}} = \alpha \left(\lambda\right) \frac{1}{1 + I_0/I_s} + \beta I,\tag{13}$$

where $\alpha(\lambda)$ is linear absorption coefficient, I_0 and I_s are laser intensity at the focus and saturation intensity, respectively, β is two-photon absorption coefficient which depends on the laser wavelength and laser intensity. The value of β for 1030 nm wavelength at the intensity of 80 GW cm⁻² is found to be $\sim 9.3 \times 10^{-6}$ cm W⁻¹. The values of linear and effective nonlinear absorption coefficient were found to be 1.2×10^6 m⁻¹ and 1.94×10^6 m⁻¹. Thermophysical parameters used in simulations are provided in table 1.

In ultra-short laser interactions, electrons re-establish the Fermi distribution after the absorption of the energy from the laser pulse while the lattice remains at the same temperature. Initially excited electrons are localized within the optical absorption depth and diffuse into deeper parts due to established temperature gradient. Electron–electron relaxation time characterizes whether the transport of electrons is diffusive or ballistic while electron–phonon coupling defines the diffusion length of electrons into sample. Thermal electrons transfer energy to lattice through electron–phonon coupling and establish electron–phonon equilibrium. The simulation results describing the evolution of electron and lattice temperatures are presented in figure 7; the fluence used corresponds to the values used in the XRD results.

The model predicts that the electron temperature rises fast because of their lower heat capacity compared to the lattice. The lattice temperature rises after the electron–phonon coupling starts. For $100 \, \text{nm}$ ITO thin film, at fluence of $0.04 \, \text{J} \, \text{cm}^{-2}$,

Table 1. Thermodynamical and optical properties parameters of ITO and PET used in modelling.

Parameter	Value
Debye temperature of ITO $(T_{\rm D})$ Melting temperature of ITO $(T_{\rm m})$	1000 K [27] 2100 K
Lattice thermal conductivity of ITO (k_1)	$3.95 \text{ W m}^{-1} \text{ K}^{-1} [33]$
Electron thermal conductivity ITO (k_e)	$1.4~{\rm W}~{\rm m}^{-1}~{\rm K}^{-1}~[34]$ at $300~{\rm K}$
Lattice heat capacity of ITO (C_1)	$2.6 \times 10^6 \text{ J m}^{-3} \text{ K}^{-1} \text{ at } 300 \text{ K}$ [27]
Electron heat capacity of ITO (C_e)	$4.53 \text{ J m}^{-3} \text{ K}^{-1}$
Fermi velocity of ITO (v_F)	$1.0 \times 10^6 \text{ m s}^{-1} [30]$
Fermi energy of ITO $(E_{\rm F})$	1.0 eV [30]
Fermi wavenumber of ITO $(k_{\rm F})$	$3.1 \times 10^9 \mathrm{m}^{-1}$
Electron mean free path of ITO (<i>l</i>)	8.3 nm
Effective mass of electron (m^*)	$0.35 \ m_{\rm e}$
Free electron number density of ITO (n)	$1 \times 10^{21} \text{cm}^{-3}$
Speed of sound in ITO film (C_s)	$4354 \text{ m s}^{-1} [35]$
Density of ITO (ρ)	7120 kg m^{-3}
Heat capacity of PET (C_s)	$1.42 \times 10^3 \text{ J m}^{-3} \text{ K}^{-1}$
Thermal conductivity (k_s)	$0.261 \text{ W m}^{-1} \text{ K}^{-1}$
Melting point of PET (T_{m-s})	533 K
Density of PET (ρ_s)	1.42 kg m^{-3}

a maximum lattice temperature of 608 K was observed while the electron temperature peaks at 16 800 K at 1.39 ps after the onset of the laser pulse. Thermal electron-lattice equilibrium was established at 8.4 ps. A maximum temperature of 420 K at the PET surface was observed which proves that ITO crystallization process was completed without damaging the lower PET surface. For 50 nm thin film, there was no significant difference in electron and lattice temperatures observed compared to those of the 100 nm film. The lattice temperature at the ITO–PET interface for 50 nm ITO thin film was 20 K higher than that of the 100 nm ITO film. While, at a fluence of ablation threshold (0.51 J cm⁻²) for 100 nm ITO thin film, model predicts maximum lattice temperature of 1380 K while PET surface temperature was 715 K. This suggests that the ablation

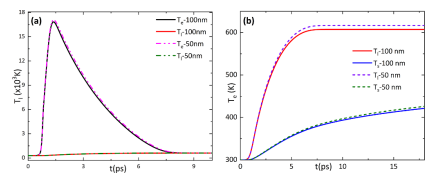


Figure 7. (a) The numerical simulation results for femtosecond laser induced evolution of electrons and lattice temperature for 50 nm and 100 nm ITO thin film on PET substrate as function of time and at a fluence of 0.04 J cm⁻². (b) Comparison of ITO lattice temperature and PET substrate temperature at the ITO-PET interface for the 50 nm and 100 nm ITO thin film.

of ITO was not because of melting or vaporization but due to different thermal expansion rates resulting in stress generation. The coefficient of thermal expansion of ITO $(7.2 \times 10^{-6} \text{ K}^{-1})$ is about ten times lower than of PET $(70 \times 10^{-6} \text{ K}^{-1})$.

4. Discussion

We have observed up to 23% improvement in conductivity for 50 nm and a 25% increase in 100 nm thick ITO thin films after the laser scanning. AFM characterization confirms an increase in average particle size after laser treatment, combined with a significant improvement in crystallinity as confirmed by XRD characterization. Transmission measurements are supportive of a more crystallized thin film. Simulation results predicts that crystallization occurred at low temperatures without melting of ITO and damage to the underlying PET substrate. It was identified that in ITO, the conductivity arises due to the charge carriers provided by Sn dopants and oxygen vacancies in ITO [36]. The conduction mechanism in a-ITO thin films, deposited at low temperature, is via the donation of electrons generated by the oxygen vacancies. It has been reported that when annealed at atmosphere, oxygen in the air reacts with the ITO films; the resulting decrease in density of oxygen vacancies reduces the free carrier concentration which in turn leads to an increase in the resistivity. However, in vacuum, there is less or no oxygen reaction with the ITO film surface, and even in vacuum oxygen atoms desorbed from the surface can escape to increase the carrier concentration [37]. Laser scanning in our work was done in atmosphere, we also consider decrease in density of oxygen vacancies along with the improvement in crystallization; together they result in an improvement in conductivity after the laser scanning. Obvious XRD peak changes due to crystallization can be seen from the XRD data for 100 nm ITO film but no obvious XRD changes were found for the 50 nm thick ITO film. It is likely that the as-deposited ITO film on PET is not well crystallized, so the 50 nm ITO is too thin and too amorphous to observe the XRD characteristic peaks of ITO. In order to clarify the mechanism for electrical property improvement, we performed Hall measurements on the 100 nm ITO thin films before and after laser treatment to measure the carrier mobility using the Ecopia Hall Effect Measurement system (HMS-3000). For 100 nm ITO thin film, the mobility increased from 4.36 cm² V s⁻¹ to 24.1 cm² V s⁻¹. Compared with the as deposited ITO thin film on PET, laser annealing led to decrease of the resistivity of ITO thin films and this decrease was mainly caused by an increase of carrier mobility in ITO films after laser annealing. Laser sintering leads to bigger sizes of ITO nanoparticles, which further leads to longer electron mean free paths and higher carrier mobilities. Hence, the enhancement in charge carriers' mobility is the key parameter along with the crystallization in conductivity improvement of ITO without losing its transparency in the infrared region [38, 39].

In ultra-short laser pulse semiconductor interaction, laser energy is transferred first to the electrons and then to the lattice. After laser irradiation, the semiconductor undergoes several stages before returning to equilibrium; these stages include carrier excitation, thermalization of carriers with themselves, thermalization of carriers with the lattice, carrier removal and thermal/structural changes [40]. If the photon energy is greater than the bandgap, then single photon absorption is the dominant mechanism for excitation of valence electrons to the conduction band. However, if the direct band gap (3.1 eV for ITO) is greater than the infrared laser photon energy (1.2 eV for IR photons), then multiphoton absorption is the most likely main excitation mechanism. If there are free carriers in the semiconductor, then free carrier absorption increases the energy of carriers and the electron (and hole) plasma dominates as it does in metals. The number density of free carriers does not change with absorption, but the energy of the free carrier gas is increased. If the laser energy is high enough to cause the excitation of free carriers above the bandgap (for a semiconductor) or above the Fermi level (for a metal) then impact ionization can generate additional excited carriers [41].

Covalent bonding is dominant for ITO and it is stable only if the electrons are in the ground state. However, this covalent bonding breaks down on excitation of electrons from stable valence band states to antibonding conduction states. If enough electrons populate the antibonding states in the conduction band during ultra-short pulse excitation, the lattice structure will become unstable; the cohesive energy of the lattice will be reduced by bond breaking. It has been reported that if 10% of the valence electrons are excited, the lattice begins

to deform due to the softening of certain phonon modes [41]. Hence it is plausible that an ultra-short laser pulse can induce such a lattice instability by dense photo-excited plasmas which weaken the lattice and thereby promote the ion movement over a significant fraction of the bond length without significantly increasing their thermal energy [42]. Therefore, we propose the weakening of the lattice by high density electron and hole plasmas to be the main mechanism for the non-thermal phase change in femtosecond laser annealing of ITO.

5. Conclusion

This study reports on improvements in the electrical properties and crystallization of very thin films on a heat sensitive PET substrate by ultra-short laser scanning. The experimental results have demonstrated that non-melting crystallization of ITO was performed at a fluence seven times smaller than the damage threshold of ITO films. An analysis of surface morphology revealed enhancement in the films grain size after laser scanning while XRD confirms the requisite improved crystallinity. We proposed ultra-short laser non-melt crystallization based on solid diffusion at low fluence. A decrease up to 25% in sheet resistance was measured after laser scanning. This improvement is primarily attributed to greater than four times enhancement in the mobility of charge carriers after laser scanning. The hypothesis for low temperature solid state crystallisation is supported by the predications obtained in the delayed two temperature numerical model; simulations predict that the lattice temperatures only reach \sim 33% of the reported melt temperature for the ITO lattice. Thus, low fluence ultrashort laser crystallization could be useful in the crystallization of thin films on a heat sensitive substrate. This process offers a selective crystallization in remote regions without affecting the nearby components.

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ORCID iDs

N Farid https://orcid.org/0000-0003-0556-6794
A Sharif https://orcid.org/0000-0001-5604-1475
R K Vijayaraghavan https://orcid.org/0000-0003-1096-

G M O'Connor https://orcid.org/0000-0002-4577-1023

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