Effect of Laser Induced Crystallinity Modification on Biodegradation Profile of Poly(L-Lactic Acid)

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Abstract

Poly(L-lactic acid) (PLLA) is of interest in drug delivery applications for its biodegradable and biocompatible properties. Polymer controlled drug delivery relies on the release of embedded drug molecules from the polymer matrix during its degradation. PLLA degradation exhibits an induction period, during which an insignificant amount of degraded products and embedded drug can be released. Due to this induction period, drug release is initially non linear, a complication in drug delivery applications. PLLA degradation is a function of crystallinity, such that control over its crystallinity tailors drug release over time. In this study, the effect of laser induced PLLA crystallinity reduction on degradation is investigated. Samples having lower surface crystallinity are shown to have higher rates of molecular weight reduction and earlier mass loss than non-laser treated samples, as observed from gel permeation chromatography and mass change. Wide-angle X-ray diffraction measurements show that crystallinity increases with degradation. A numerical model is implemented from hydrolysis and diffusion mechanisms to investigate the effect of laser irradiation on biodegradation. Controlled laser treatment of PLLA offers a method for constant drug release through the reduction of surface crystallinity.

Keywords: poly(L-lactic acid); laser treatment; biodegradation; crystallinity; drug delivery

1. Introduction

Poly(L-lactic acid) (PLLA) is attractive in drug delivery, food packaging, and tissue engineering applications because of its biocompatible and biodegradable properties. PLLA is especially of interest in drug delivery applications because it hydrolyzes in the human body into lactic acid, a product which is excreted by the body with no toxicity. In drug delivery applications, drugs are embedded in a polymer matrix and released as it degrades. Biodegradable polymers offer a means to control the drug delivery in time [1]. PLLA degradation exhibits an induction period, a duration of time required for water molecules to penetrate into the matrix before degradation can occur. During this time, embedded drugs are not released at a linear rate.

PLLA degradation in a physiological environment occurs via hydrolysis, in which water penetrates into the polymer matrix, attacking the ester bonds and causing chain scission. Water molecules readily penetrate into the amorphous region, but hardly into the crystalline region, because the polymer chains are highly packed and densely ordered in crystals [2]. Degradation of ester bonds occurs faster in the amorphous phase because of its more permeable structure. Hydrolysis in the crystalline phase begins at the fold surfaces and progresses inwards as controlled by chain scission [3]. Hydrolysis in the crystalline phase occurs less preferentially than in the amorphous phase. A sample composed of both crystalline and amorphous phases will more rapidly undergo hydrolysis in the amorphous phase than the crystalline phase.

PLLA hydrolysis is accelerated by autocatalysis [4]. The rate of hydrolysis increases as the concentration of reaction products increases. Hydrolysis of polyester produces shorter chains with acid and alcohol end groups. Acid end groups dissociate, leading to an acidic environment, which accelerates hydrolysis. Therefore, the diffusion of shorter chains out of the polymer

plays a key role in controlling the overall degradation rate. Another factor that complicates biodegradation is the increase of crystallinity during degradation [5,6]. Preferential degradation in the amorphous region leaves the crystalline phase behind, leading to a mostly crystalline material. Chain scission as a result of degradation also increases chain mobility which facilitates crystallization in the amorphous phase.

Because changing the surface crystallinity of PLLA has the potential to tailor the initial degradation rate, laser heat treatment to reduce surface crystallinity is of interest. PLLA crystallinity reduction by laser irradiation with infra-red (IR) and green wavelengths has been quantified through wide-angle X-ray diffraction (WAXD) measurements [7]. The same effects have been observed by excimer laser processes operating at ultra violet (UV) wavelengths with photon energies higher than bond energies [8,9]. It has been shown that surface crystallinity can be reduced with no measurable chemical modifications due to the low radical mobility in the crystal structure [9]. The effects of laser treatment on polymer degradation over time have yet to be studied.

The objective of this work is to investigate the effects of laser surface treatment on PLLA degradation, characterized by the change in mass, molecular weight (MW), and crystallinity. Since PLLA degradation behaves similarly at elevated temperatures [10], degradation is conducted at elevated temperatures to shorten the total test time. MW and crystallinity are determined through gel permeation chromatography (GPC) and WAXD, respectively. A numerical model is developed to capture the degradation process.

2. Background

2.1 Laser melting of polymer

Polymer melting is an amorphization process in which crystalline polymer chains detach from crystals. Crystalline polymer chains are held by weak van der Waals forces and hydrogen bonds, which are broken during melting to form an amorphous structure. Melting initiates at the crystal fold surface, followed by the progressive unfolding of chains towards the center of the crystal. Polymer melting occurs within nanoseconds [11], and thus nanosecond laser treatment is used to induce polymer melting. Crystallinity is reduced after laser melting as a result of slow crystallization kinetics compared to the rapid melting and cooling during laser processing [7-9].

Polymer absorption of photons with energies exceeding the bond energy risk braking chemical bonds. In PLA, this reaction occurs between two CH₃CHCOO units. If no oxidation occurs during processing, the separated CH₃CHCOO units may recombine. According to the cage effect, high PLLA crystallinity has been shown to reduce the amount of bond breaking and lead to non-measurable chemical modifications under UV laser treatment [9].

2.2 Biodegradation of polyester

PLA is a biodegradable polyester which breaks down in the human body through hydrolysis. Water molecules attack the ester bonds via the following reaction

Hydrolysis causes chain scission, and produces shorter chains with carboxylic (-COOH) groups and alcohol (-OH). The rate of hydrolysis is proportional to the number of ester bonds present in each monomer constituting the polymer. The number of reactive bonds in the polymer decreases as hydrolysis goes on. Hydrolysis depends on the molar concentrations of the monomer still present in the polymer chain, C_e , and water, C_w [12], at the rate of

$$\frac{dC_e}{dt} = -k_1 C_e C_w \tag{2}$$

where the constant k_1 depends on temperature and does not vary with reaction. Carboxylic end groups generated from this reaction have a high degree of dissociation and can act as a catalyst to accelerate the hydrolysis. Hydrolysis of polyesters may become autocatalytic if carboxylic end groups remain in the bulk [13]. During autocatalyzed hydrolysis, the reaction rate depends on the concentration of the carboxylic end groups, C_{COOH} , as well. The rate of autocatalyzed hydrolysis is given by [14]

$$\frac{dC_e}{dt} = -k_2 C_e C_w (C_{COOH})^n \tag{3}$$

where k_2 is the rate constant for the autocatalysis reaction, and *n* accounts for the dissociation of the carboxylic groups and can be used as an empirical parameter to reflect the reactions [15].

Amorphous chains hydrolyze in three stages before generating monomers [16]. In stage 1, chain scission commences in the intact amorphous chains, resulting in the breaking of amorphous tie chains. In stage 2, chain scission occurs on already broken amorphous chains, creating unattached oligomers. In stage 3, hydrolytic reactions occur on short amorphous segments protruding from the crystals and the oligomers produced in stage 2. Hydrolysis in this stage generates highly mobile monomers which can diffuse out of the polymer matrix.

PLLA can be semicrystalline, and the amorphous region experiences a higher hydrolysis rate than the crystalline region [2,17]. Hydrolysis of the amorphous phase occurs randomly, but

occurs preferentially in the crystal phase at the fold surfaces. A schematic representation of semicrystalline polymer chains before and after extensive hydrolytic degradation is given in Fig. 1 [18]. Random hydrolysis generates chains with a wide length distribution; preferential hydrolysis on fold surfaces can lead to the chains representing the integral folds the crystalline residues [17,19]. Mobile segments reorganize themselves from a disordered to an ordered state due to intermolecular hydrogen bonds and van der Waals forces, leading to crystallization during hydrolysis [2].

3. Numerical model

A 2D symmetric model is developed to investigate the effect of laser treatments on biodegradation profiles. Laser energy absorbed by semi-transparent bulk PLLA generates heat governed by the heat equation

$$\rho C_p(T) \frac{\partial T}{\partial t} = \nabla (k \nabla T) + q(z, t)$$
(4)

where ρ is mass density, $C_p(T)$ is specific heat as a function of temperature T, k is thermal conductivity, q(z,t) is the laser power density as a function of depth from the laser irradiated surface z and time t, expressed as

$$q(z,t) = Q_0 e^{-\alpha z + \beta \left(\frac{t}{t_p} - 2\right)^2}$$
(5)

where Q_0 is peak power density, α is absorption coefficient, t_p is pulse width, and β is -4ln2. Parameters used in Eqs. (4) and (5) and the change in $C_p(T)$ during the phase transition are further detailed in ref. [9]. Polymer crystals melt within a temperature range, from T_m to $T_m + \Delta T_m$. The crystal fraction melting in a temperature increment dT_m within T_m and $T_m + \Delta T_m$ can be expressed as $\phi(t, T_m) dT_m$. The total crystallinity is then $\Phi(t) = \int_{T_m}^{T_m + \Delta T_m} \phi(t, T_m) dT_m.$ The change in $\phi(t, T_m)$ during melting is given by [20]

$$\frac{d\Phi}{dt}(t,T_m) = -R_m(\Delta T)\Phi(t,T_m)$$
(6)

The melting rate coefficient R_m is a function of superheating, $\Delta T = T - T_m$, expressed as $R_m = R_m(\Delta T)$ if $\Delta T > 0$ and $R_m = 0$ if $\Delta T < 0$. Molecular dynamics simulations have suggested that crystallinity decreases within ns, and R_m is of the order of 10^9 s^{-1} [11]. Spatially resolved crystallinity after laser processing is imported into the degradation model as the initial condition.

Biodegradation is captured using a phenomenological model [15]. Semicrystalline PLLA is modeled to be consistent of 7 species during degradation: non-degraded amorphous chains, degraded amorphous chains in stages 1, 2, and 3 [16], crystalline chains, monomers, and water molecules. Non-degraded and degraded amorphous chains hydrolyze via the scission of the ester bonds contained in the species, but have zero diffuse-ability due to restricted mobility. Monomers have high mobility. Amorphous polymer chains crystallize during degradation. Crystalline polymer chains cannot diffuse and are assumed to hydrolyze ten times slower than amorphous chains [21], generating chain scission on the crystal fold surface. Water molecules are assumed abundant in time and space, and the size distribution of polymer chains is neglected. Acid end groups of the monomers, generated during chain scission, accelerate the hydrolysis through autocatalysis. Assuming water concentration is constant temporally and spatially, the molar concentration of the non-degraded amorphous chain is expressed as

$$\frac{dC_0}{dt} = -\gamma_0 C_0 \cdot \varepsilon_0 C_0 C_m^n \cdot \kappa_0 \frac{dC_c}{dt}$$
⁽⁷⁾

where C_0 and C_c are the molar concentrations of monomers in the non-degraded amorphous chains and crystalline chains, respectively. The dissociation of the acid end groups n is assumed to be unity. The first, second, and third terms on the right of Eq. (7) account for rates of non-autocatalysis, autocatalysis, and crystallization from amorphous chains, respectively. Values of γ_0 , ε_0 , and κ_0 are the corresponding phenomenological rate constants. Hydrolysis of non-degraded amorphous chains, Eq. (7), generates degraded amorphous chains in stage 1, which is then hydrolyzed into stage 2 and contributes to crystallinity. Similarly, hydrolysis of chains in stage 1 leads to stage 2, and hydrolysis of chains in stage 1 leads to stage 2. Part of the hydrolyzed chains also crystallize during hydrolysis. The molar concentration of monomers in each stage is expressed as

$$\frac{dC_i}{dt} = (\gamma_{i-1} + \varepsilon_{i-1}C_m^n)C_{i-1} - (\gamma_i + \varepsilon_iC_m^n)C_i - \kappa_i\frac{dC_c}{dt}$$
(8)

where *i*=1, 2, and 3, representing degradation stages 1, 2, and 3. C_i is the molar concentration of the monomers in stage *i*. γ_i , ε_i , and κ_i are the phenomenological rate constants accounting for non-autocatalysis, autocatalysis, and crystallization due to hydrolysis in stage *i*. During degradation, the crystallization of linear polyesters are modeled using the Avrami equation [5], $\phi=1-exp(-k_c t^m)$ where ϕ is monomer fraction in the crystalline domain, k_c is the Avrami constant and *m* is the Avrami exponent [22]. Hydrolysis of the stage 3 species generates monomers which do not connect to other monomers and have high mobility to diffuse out of the polymer matrix. Assuming Fick's second law for monomer diffusion, which predicts the change of monomer concentration with space and time, the molar concentration C_m of the monomers with high mobility is modeled by

$$\frac{dC_m}{dt} = (\gamma_3 + \varepsilon_3 C_m^n) C_3 + \nabla (D \nabla C_m)$$
(9)

where D is the monomer diffusivity. Appropriate units are used for the phenomenological rate constants so that each term in Eqs. (7)-(9) is in mole per volume per time. The values of the rate constants are selected to capture experimental results [15].

The coupled partial differential Eqs. (4) and (6) are solved to determine crystallinity change due to laser treatments. Solutions of these equations are used as initial conditions to solve the coupled Eqs. (7)-(9) to capture degradation profiles. The two sets of coupled equations are solved through the finite element method in COMSOL Multiphysics 4.1. A 2D axial symmetric domain is used, in which the 1 mm by 5 mm film is immersed in a 10 mm by 10 mm aqueous medium. Laser treated area covers both sides of the film. The film is initially composed of non-degraded crystalline and amorphous chains with crystallinity determined experimentally. Monomers generated during degradation can diffuse out of the film into the surrounding aqueous medium. The time domain in the simulation is 14 days.

4. Materials and methods

PLLA granules were provided by PURAC and used as received. PLLA samples were prepared through thermal compression of PLLA granules under 5.7×10^4 Pa at 180°C for 4 hours, and cooled down in air. Crystals develop during the cooling process. The obtained sample is around 1 mm thick. Laser treatment was conducted on both sides of the sample by a KrF excimer laser with 248 nm wavelength 248 nm and 25 ns pulse width 25 ns. The homogenized excimer laser beam has a spatially uniform intensity favorable for a uniform surface treatment. The sample is radiated by a single pulse in argon atmosphere with a flow rate of 15 standard cubic feet per hour to prevent oxidation. To determine crystallinity, WAXD measurements are made using an Inel X-ray diffractometer to determine crystallinity. Monochromatic CuK α radiation with wavelength λ =0.15418 nm at 40 kV and 30 mA is used. The chemical compositions of the PLLA samples are measured using X-ray photoelectron spectroscopy (XPS, PHI 5500 ESCA). From XPS measurements, the O_{1s} and C_{1s} spectra are captured, and the take-off angle is 45°.

During degradation tests, each sample was placed in a vial and fully immersed in a 10 mL phosphate buffered saline (PBS) with a pH of 7.4 purchased from Life Technologies. Vials were placed in a water bath at 70°C, and the PBS was changed every two days. After degradation, samples were dried in vacuum for two days. The sample mass was recorded before and after vacuum drying. The weight-average MW (M_w) and number-average MW (M_n) were determined in tetrahydrofuran (THF) from gel permeation chromatography (GPC) at room temperature. Crystalline PLLA was first dissolved in methylene chloride and rapidly dried with a rotary evaporator to obtain the amorphous form with higher solubility in THF. PLLA/THF solutions were prepared with 1 mg/1 mL concentration for GPC measurements. Crystallinity is determined by the WAXD, and morphology is observed using stereomicroscopy and optical profilometry.

5. Results and discussion

5.1 Effect of laser irradiation on crystallinity and chemical modifications

Images of samples before and after laser treatment as well as after 14 days of degradation are presented in Fig. 2. The non-laser treated sample is semi-transparent because of the existence of crystalline phase. Laser treatment generates opaque spots, a result of strong light scattering from the roughened surface as confirmed by optical profilometry. Roughness is induced by laser surface melting and resolidification. WAXD profiles are given in Fig. 7. The 16.7° WAXD peak becomes less prominent for the samples treated with higher laser fluences, due to crystallinity reduction. Crystallinity is calculated based on ref. [23], as given in Fig. 3, which shows that the crystallinity decreases at 2.4 J/cm² as a result of laser melting.

To investigate possible chemical modifications under laser irradiation, XPS measurements are

performed. Chemical reactions caused by laser irradiation with photon energies higher than the bonding energies produce small molecules such as CO and CO₂, and reduce the total ratio of oxygen to carbon [24]. The ratio of O_{1s} and C_{1s} , shown in Fig. 3, remains constant below 3.0 J/cm², and drops when fluence reaches 3.2 J/cm², indicative of chemical modifications due to laser treatment. PLLA samples remain chemically unmodified below 3.0 J/cm² due to the cage effect, which states that free radicals generated by the dissociation of molecules cannot move apart because of the confinement of surrounding molecules [9]. This results in the recombination of the dissociation products, which then return to the initial state. A fluence of 3.0 J/cm² induces the maximum amorphization with non-measurable chemical modifications, and is used as the energy level to treat samples in this study.

5.2 Effect of laser irradiation on degradation

5.2.1 Modification of morphology

Laser treated samples after having been degraded for 14 days is shown in Fig. 2. A uniformly opaque morphology is observed as a result of higher crystallinity developed during degradation. Detailed morphology investigations are given by stereomicroscopic images of laser treated samples as a function of degradation period in Fig. 4. A laser treated sample before degradation is given in Fig. 4(a). The 1 mm by 1 mm square spot is generated by laser irradiation. As degradation progresses, spot edges become less defined, Fig. 4(b), suggesting the degradation and erosion (mass loss) of laser melted layer. After 8 days, Fig. 4(c), spot edges become more indistinct than samples degraded for shorter periods. During prolonged testing, the laser melted layer is eroded and degradation starts to occur in the non-melted volume below. Eventually, the morphology of laser treated sample, Fig. 4(d), is similar to that of the non-laser treated sample.

It is observed that light scattering is not uniform over the surface. Surface roughness is caused by the selective degradation on the amorphous phase, leaving the crystalline phase with strong local light scattering.

The cross section of a laser treated sample after being degraded for 14 days is given in Fig. 5. It is observed that the bulk remains solid, which suggests that autocatalytic degradation is not dominant as a result of easy diffusion of acidic monomers out of the sample. Spatial degradation profiles of monomer molar concentration are given in Fig. 6. After 0.5 days, monomer concentration on the surface is higher than that in the bulk, suggesting accelerated degradation in the 80-µm thick laser melted layer. In the later stage of degradation, Fig. 6(b), monomer concentration in the non-melted bulk starts to increase. The simulation results capture that laser melted layer experiences a higher degradation rate.

5.2.2 Modification of crystallinity

To investigate the changes in crystallinity during degradation, WAXD measurements have been conducted on samples after regular degradation intervals, as depicted in Fig. 7. Laser treatments cause surface melting and a reduced crystalline peak, as discussed in Sec. 5.1. For both types of samples, crystalline peaks become more prominent with time, suggesting the occurrence of crystallization during degradation. Sample crystallinity, as calculated from WAXD measurements based on ref. [23], is given as a function of time in Fig. 8, which shows that crystallinity increases with degradation period, in agreement with Fig. 7. For both types of samples a significant increase of crystallinity is observed at day 0.5, when the sample MW begins to decrease, as is discussed in Sec. 5.2.3. The decrease of MW favors chain reorganization and crystallization, and increases crystallinity. Lower crystallinity and greater

amounts of crystallizable material in non-degraded samples also favors crystallization [25]. Crystallization slows between day 0.5 and day 5 for the non-laser treated sample, because part of the amorphous chains is less crystallizable, including the rigid amorphous phase (RAP) [26] and entangled chains. The RAP and entangled chains hinder chain movements and make it difficult for amorphous chains to relax and crystallize. Crystallinity also significantly increases at day 8 for the non-laser treated sample, and day 3 for the laser treated sample. Crystallinity increase at this stage is mainly caused by the loss of amorphous chains, Fig. 11, as discussed in Sec. 5.2.3.

5.2.3 Modification of molecular weight and mass

Degradation is characterized by sample MW and sample mass. MW distribution profiles as a function of degradation period are given in Fig. 9, which plots the mass fraction of chains (w) per increment of MW in a logarithmic scale (logM) versus MW. The number average MW (M_n) and weight average MW (M_w) are determined from MW distributions. The polydispersity index (PDI), defined as M_w/M_n , is calculated as a measure of distribution of MW. M_n , M_w , and PDI for the non-laser treated and laser treated samples are given in Fig. 10. Sample mass as a function of degradation period is given in Fig. 11. Simulated sample MW and sample mass for both types of samples are given in Figs. 12 and 13.

For non-laser treated samples, the GPC profile before degradation is centered at around 120000 g/mol and the distribution is narrow, as shown in Fig. 9(a). As the degradation period increases, days 2, 3 and 5, the distribution gradually becomes wide and shifts left. A small hump is developed at around 60000 g/mol at day 5. Sample MW decreases while PDI increases as a result of random chain scission, as given in Fig. 10(a). Random chain scission before day 5 occurs in the amorphous region, causing chains to degrade into segments with various MWs [17].

There is no obvious mass decrease before day 5 for non-laser treated sample, as given in Fig. 11, because the degraded segments are not small enough to diffuse out of the sample. The degraded segments have higher mobility, and reorganize themselves from a disordered to an ordered state as a result of intermolecular hydrogen bonds and van der Waals forces during degradation, which increases the crystallinity for the first 5 days, as shown in Fig. 8.

At day 8, the GPC profile of the non-laser treated samples shifts left significantly, Fig. 9(a). A distinct peak, corresponding to a smaller MW of 10300 g/mol, is observed. This shift is due to the selective scission of the crystal fold surface. Preferential scission generates chains with MW representing the integral folds the crystalline residues [17,19]. The left peak is mainly composed of the MW of crystalline residues, and the right peak mainly comes from the amorphous region experiencing random chain scission. As degradation approaches 11 and 14 days, the right peak starts to diminish due to the continuous scission of amorphous chains, and the overall distribution is mainly composed of a peak centered at 10300 g/mol. The PDI thus decreases, as shown in Fig. 10(a).

The lowest MW, around 3000 g/mol, is detected after day 8 in Fig. 9(a), while mass decrease also occurs at day 8, Fig. 11. This finding suggests that chains with MW smaller than 3000 g/mol are diffused out of the sample. Figure 8 shows that crystallinity increases significantly at day 8 for the non-laser treated sample. Thus, the materials diffusing out of the sample are mainly amorphous. The reduction of amorphous material increases the crystallinity in the sample and leads to stronger local light scattering shown in Fig. 4(d).

GPC profiles of the laser treated sample are given in Fig. 9(b). Before degradation, the GPC profile of the laser treated sample has a peak at 120000 g/mol similar to the non-laser treated

sample, which suggests that laser treatment does not modify PLLA MW. When compared with the non-laser treated sample, MW decreases at a high rate at the early stage for the laser treated sample. On days 2 and 3, significant changes of GPC profiles are observed in Fig. 9(b). At day 2, the peak at 120000 g/mol decreases, while the distribution extends to the smaller MW region. The peak diminishes at day 3, and a hump at smaller MW region is developed. During this period, a random chain scission of the laser melted layer occurs. Non-homogeneous degradation at the laser melted layer and the bulk widens the MW distribution and thus increases the PDI, Fig. 10(b). The high degradation rate of the laser treated sample is caused by water molecules penetrating into the less ordered structure caused by laser melting. Faster water penetration into the bulk is also responsible for the diminished original peak at day 3.

At day 5, the hump is split into two peaks for the laser treated sample, as shown in Fig. 9(b). The distinct distribution of two peaks is suggestive of selective chain scission on crystal fold surfaces. Same phenomenon is observed in the non-laser treated sample at day 8. The earlier occurrence of this selective scission for the laser treated sample may be due to the less ordered structure of the partially melted crystals. For partially melted crystals, the crystal thickness is diminished, which increases the molecular loop length on the fold surface [27]. This longer loop length enlarges the free volume, the volume not occupied by polymer molecules in a polymer matrix. A larger free volume allows for easy water penetration, and facilitates hydrolysis on the fold surface. Toward the late stage of the degradation, the right peak diminishes and the left peak increases, as shown in Fig. 9(b). This left peak is composed of the chains with MW representing the integral folds the crystalline residues, as well as the smallest MW distribution of chains which can remain in the matrix. The narrowing MW distribution at

the late degradation stage decreases the PDI, as given in Fig. 10(b) after day 5.

Laser surface melting accelerates MW reduction, which in turn shortens the induction periods of mass loss. Mass loss occurs at day 3 for the laser treated samples, compared to day 8 for the non-laser treated sample, Fig. 11. The decrease of sample mass is accompanied with the increase of crystallinity, Fig. 8, a phenomenon observed for both types of samples. The laser melted layer is composed of amorphous chains, part of which degrades and diffuses out at a higher rate, while the remainder crystallizes. At day 3, the degraded products in the laser melted layer have small enough MW such that diffusion out of the matrix occurs, which leads to the mass loss as shown in Fig. 11. After the erosion of the laser melted layer, the sample MW and mass decrease at similar rates as the non-laser treated sample, as observed in Figs. 10 and 11.

Simulation results of MW change as a function of degradation period are given in Fig. 12. Numerical results of the laser treated sample experience a larger MW decrease, in agreement with experiments. Experimentally, the MW of the non-laser treated sample does not decrease as significantly as predicted in simulation before day 1. This difference is because it takes time for water molecules to diffuse into the bulk in experiments. Water diffusion time is not considered in simulation, which causes the overestimation of the MW decrease.

The simulated mass changes with degradation, Fig. 13, capture the phenomenon that appreciable mass decrease occurs after an induction period. During the induction period in which mass loss does not occur, the original amorphous polymer chains degrade into fragments not small enough for diffusion out of the sample. Fragments experience further degradation into monomers, which can diffuse and lead to mass loss. Experimentally, the mass change is more significant when compared with simulation. This is because the erosion of crystalline material also occurs

on the surface during degradation [28], which is not considered in simulation, leading to a conservative prediction.

5.3 Concentration of species during degradation

The molar concentration of species considered in the simulation is plotted with the degradation period, given in Fig. 14(a). Amorphous chains start to degrade in the early stage of the degradation period, generating the species in stage 1. Stage 1 is sequentially degraded into stages 2 and 3. Chains in stage 3, upon further hydrolysis, are decomposed into monomers, which diffuse out of the sample. Only a small amount of monomers remain on the sample surface. Degradation also enhances chain mobility, which leads to crystallization. The simulated concentration of crystalline chains increases as observed in experiments.

Simulated crystallinity increases via two stages, as agreed with the experiments. The first stage is predicted by the Avrami crystallization theory [25]. Experimentally, crystallinity increases to a significant extent before day 0.5, not captured in the simulation. The discrepancy results from the fact that according to the Avrami theory, crystallization begins with an induction period due to the formation of nuclei. Pre-existing nuclei in the experiment samples favor crystal growth without experiencing the induction period. The second stage of crystallinity increase is due to the loss of amorphous material, which increases crystallinity as experimentally observed and numerically simulated. Hydrolysis of crystals occurs on the fold surfaces. Concentration of non-degraded crystalline chains decreases, which leads to the increased concentration of the hydrolyzed crystals and the reduced MW.

Simulation results of the species concentration in the laser melted layer are shown in Fig. 14(b). Laser irradiation melts the crystal structure, generating an amorphous layer near the surface. Amorphous chains are degraded to stages 1, 2, 3, and monomers in sequence. Because the species are near the surface, the major part of monomers readily diffuses out of the sample and thus the concentration is lower as compared to that in the sample bulk. Although most amorphous chains degrade, crystallization takes place in the laser melted layer, mainly at the early stage.

6. Conclusions

The effect of laser irradiation on PLLA biodegradation has been studied experimentally and numerically. Excimer laser irradiation has been shown to melt the PLLA surface. The melted layer showed lower crystallinity and no observable chemical changes according to WAXD and XPS measurements. Optical micrographs taken during degradation testing show that degradation initiates on the sample surface, and the melted layer allows for the accelerated initial degradation as is also captured through numerical simulations. Laser treated samples experience a faster initial MW reduction, which leads to a shorter induction period of mass loss as observed from GPC. Heterogeneous degradation of PLLA samples caused by inhomogeneous crystallinity is observed in both laser treated and non-laser treated samples from bimodal distribution of MW from GPC measurements. Laser treated samples exhibited higher crystallinity after degradation as a result of preferred degradation and higher chain mobility in the amorphous phase, as confirmed from the WAXD measurements. Changes in MW, sample mass, and species evolution during degradation are captured in simulation results of both laser treated and non-laser treated samples. Laser modification to the crystallinity of PLLA has been shown to reduce the degradation induction period, a desired property in drug delivery applications.

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(b) Crystalline residue of polymer

Fig. 1 Schematic representation of (a) semicrystalline structure and (b) crystalline residue of polymer after hydrolysis, which preferentially occurs in amorphous region and on crystal fold surface [18].



Fig. 2 Non-degraded PLLA sample (a) before and (b) after laser treatment with a fluence of 3 J/cm². Laser irradiated spots show less transparency due to increased surface roughness. The laser-treated sample degraded for 14 days is given in (c) and its high crystallinity reduces the transparency.



Fig. 3 PLLA crystallinity and the ratio of O_{1s} to C_{1s} as a function of laser fluence. The error bar represents the standard deviation of 3 data points.



Fig. 4 Surface morphology of the laser treated sample (a) before and after degradation for (b) 3, (c) 8, and (d) 14 days under the stereomicroscope. The squares in (a) are laser spots. The

edges of laser spots become less defined with degradation period, suggesting the erosion of laser melted layer. Degradation in the non-melted bulk volume occurs in the later stage.



Fig. 5 Cross section of the laser-treated sample degraded for 14 days. The bulk remains solid, suggesting that autocatalysis is not dominant.





Fig. 6 Simulated spatial distribution of monomer concentration in the laser treated sample degraded for (a) 0.5 days and (b) 6 days.







Fig. 7 WAXD profiles of the (a) non-laser treated and (b) laser treated samples degraded for regular periods. Intensity of crystalline peaks increases with degradation period, suggesting a higher crystallinity. Profiles are shifted in y direction for viewing clarity.



Fig. 8 Crystallinity as a function of degradation period determined from WAXD. Crystallinity increases with degradation period. A significant increase occurs on day 0.5 for both types of samples. Crystallinity also increases on day 8 and day 5 for the non-laser treated and laser treated samples, respectively. The error bar represents the standard deviation of 3 data points.



(b) Laser treated sample

Fig. 9 GPC profiles of the (a) non-laser treated and (b) laser treated samples after regular degradation periods. For (a), the distribution becomes wider and shifts left as the degradation period increases to day 5, representing the random chain scission in the amorphous region.

After day 8, a distinct new peak is developed due to selective chain scission of the fold surface of crystals. For (b), the MW distribution extends to the left before day 3, signifying the random chain scission of the laser melt layer. At day 5, two distinct peaks are developed due to the selective chain scission of the partially melted crystal fold surfaces. Profiles are shifted in y direction for viewing clarity.



(b) Laser treated sample

Fig. 10 M_w , M_n , and PDI of the (a) non-laser treated and (b) laser treated samples after regular degradation periods. M_w and M_n decrease at a higher rate for the laser treated samples, as a result of fast degradation in the laser melted layer. The non-homogeneous degradation of the melted layer and bulk increases PDI. The error bar represents the standard deviation of 3 data points.



Fig. 11 Experimental results of sample mass with and without laser treatments. Mass decrease is observed after day 8 for non-laser treated sample and after day 3 for laser treated sample. The error bar represents the standard deviation of 3 data points.



Fig. 12 Simulated MW of the non-laser treated and laser treated samples.



Fig. 13 Simulated mass change of the non-laser treated and laser treated samples.



(a) In the center of the non-laser treated sample



(b) In the laser melted layer of the laser treated sample **Fig. 14** Molar concentration of the simulated species as a function of degradation period.