LDPE Transformation by Exposure to Sequential Low-Pressure Plasma and TiO$_2$/UV Photocatalysis

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1. Introduction

On an everyday basis, one of the most employed polymers worldwide is low-density polyethylene (LDPE) with industrial, commercial, and domestic uses. Low-density polyethylene transformation and/or biotransformation is difficult and slow because it is a high molecular weight polymer with hydrophobic carbonated chains and low nitrogen content, thus generating a major environmental problem affecting the use of water, resources, soil, and air. During LDPE incineration and its subsequent disposal in sanitary landfills, gaseous emissions, such as CO$_x$ and volatile organic compounds (VOCs) are generated [1,2]. Therefore, to promote LDPE degradation different physical, chemical, and biological treatments can be implemented, such as direct-current low-pressure plasma (DC-LPP), photocatalytic oxidation, photo-oxidation, thermo-oxidation, and biological transformation [3,4]. Most of these treatments are performed in the amorphous region of the polymer to modify LDPE’s hydrophobic surface into a hydrophilic one and generate low molecular weight carbonated...
fractions (chain hydrolysis and formation of vinyl and carbonyl groups) [1,5]. In such a way its subsequent transformation is facilitated by implementing other technologies or its final disposal or recycling is carried out.

DC-LLP modifies LDPE surface (ablation) by randomly inserting onto the plastic’s surface carboxyl polar groups (-COOH) and hydroxyl groups (-OH) [6,7] that generate crosslinking with the polymer’s carbon chains [8,9], forming oxidized low molecular weight structures that become more susceptible for subsequent transformation [10–12]. To generate plasma through electric discharge, inert gases are employed, such as argon (Ar) [13,14] and reactive gases such as oxygen (O₂) [3,7,15] or a mix of both [8,11,16]. In the presence of O₂, plasma action results in double bond formation in LDPE’s polymeric chain, with groups such as vinyl and vinyldenes (R₁R₂C=CH₂) with hydrogen release (H₂) or with the participation of hydrogen in a second reaction to form water (H₂O) [16]. Vinyl group formation is also possible without H₂ release, where its formation occurs with the participation of methyl groups (CH₃⁻) present in LDPE [11]. At the same time, organic hydroperoxides are formed (CH₃-O-OH) on the material’s surface, due to the ease of breaking O-O bonds that later form free radicals, such as carboxy (C-O•), carboxyl (COO•) [7], hydroxyl (OH•), methyl (CH₃•), or alkoxy (CH₃-O•) [17] on the surface of the treated material [18]. Last, due to the elimination of H₂ atoms from the polymeric chain, various polar groups are formed on the LDPE surface, such as (hydroxyl (OH⁻), carbonyl (C=O), and carbonyl (C=OOH)) [19–21]. These groups contribute to LDPE’s increased surface hydrophilicity [17].

On the other hand, advanced oxidation processes, such as photocatalysis with TiO₂, are a physical/chemical alternative to modify or transform different types of plastic [22–25]. The photocatalytic process with TiO₂ begins when the oxide semiconductor is electromagnetically irradiated (λ < 400 nm) and a photoelectron excitation takes place, where photon energy (hv) is absorbed by the electron in the TiO₂ valence band (VB), which causes energy transitions between the VB and the conduction band (CB) to ensue. This dynamic generates vacancies or holes in the spaces previously occupied by VB electrons, producing an electron–hole pair (e⁻/h⁺) [6]. Electron–hole pairs (e⁻/h⁺) can produce a series of oxidation–reduction reactions, where photogenerated e⁻ in CB can form O₂•⁻, and the photogenerated h⁺ in VB can form OH•. Likewise, the photogenerated holes initiate oxidative stages where adsorbed organic molecules on the material’s surface can be oxidized to become CO₂ and water. Relating to plastic photocatalytic oxidation, various authors propose reactive oxygen species (ROS) randomly oxidize LDPE carbon chains, in the amorphous phase as well as the crystal phase [22,25,26].

Both DC-LLP and photocatalysis modify the LDPE’s chemical structure and mechanical properties, favoring its degradation, yet a combined action between both will notably increase the material transformation. Currently, this combination has not been studied. Therefore, the objective of this study was to select reaction conditions and/or operating conditions to perform plasma discharge and TiO₂/UV photocatalysis as technologies to transform LDPE sheets.

2. Results and Discussion
2.1. Initial LDPE Sheet Characterization

Clean 3 cm² LDPE sheets (pristine LDPE) results before being submitted to experimentation appear in Table 1.
### Table 1. Pristine low-density polyethylene (LDPE) sheet initial characteristics.

| Response Variables          | Value       | References |
|----------------------------|-------------|------------|
| Weight (mg)                | $4.3 \pm 0.5$ | Present work |
| Static contact angle ($^\circ$) | $87 \pm 1$ | Present work |
| Roughness (nm)             | $6 \pm 2$ | Present work |
| Young’s modulus (MPa)      | $34 \pm 1$ | Present work |
| Yield strength (MPa)       | $10 \pm 2$ | Present work |
| Spectra and vibrational band structure (FTIR) | $729 \text{ cm}^{-1}$ | $[27,28]$ |
| CH$_2$ rocking and deformation | $1460 \text{ cm}^{-1}$ | $[27,28]$ |
| CH$_2$ bending and deformation | $2921 \text{ cm}^{-1}$ | $[27,28]$ |
| CH$_2$ asymmetric stretching | $2843 \text{ cm}^{-1}$ | $[27,28]$ |
| CH$_2$ symmetric stretching | $2921 \text{ cm}^{-1}$ | $[27,28]$ |

#### 2.2. Plasma Discharge on LDPE Sheets–Static Contact Angle

In previous work by Gómez-Méndez et al. [29], O$_2$ plasma was used for LDPE pretreatment for consecutive biodeterioration with *P. ostreatus*. In this work, it was demonstrated that plasma discharge favored the material’s hydrophobicity and subsequent colonization and biodeterioration by the fungus [29]. In the present work, argon (Ar) in addition to O$_2$, was used for plasma discharge since various authors have described its use for LDPE surface modification [30–32].

Regarding preliminary assays, it was observed that only 24.8% (32 treatment combinations) met with conditions 1 and 2 previously described in the methodology (supplementary material 2). Out of the 32 combinations, 15% corresponded to 100% O$_2$ discharge, 41% to 100% Ar plasma discharge, and 44% to the (1:1) mix of gases discharge (Table S1).

For discharge of 100% Ar ($p = 0.0126$) (Table S2), the two best treatments were $2.4 \times 10^{-2}$ mbar, 900 V treatment and $2.4 \times 10^{-2}$ mbar, 800 V, presenting a static contact angle (SCA) of $(26 \pm 5)^\circ$ and $(30 \pm 5)^\circ$, respectively (Figure 2A). For 100% O$_2$ ($p = 0.4275$) (Table S2), the greatest decrease in SCA was achieved with $3.0 \times 10^{-2}$ mbar, 700 V with $(20 \pm 4)^\circ$ treatment followed by the $3.0 \times 10^{-2}$ treatment mbar, 600 V with $(21 \pm 3)^\circ$ (Figure 1B). The mixed discharge treatment Ar-O$_2$ (1:1) ($p = 0.0001$), (supplementary material 2) showed the greatest decrease in SCA for $3.0 \times 10^{-2}$ mbar, 600 V (SCA 28 ± 2°) followed by $2.6 \times 10^{-2}$ mbar, 700 V with an SCA of $(30 \pm 2)^\circ$ (Figure 1C). The comparison of means between treatments ($p = 0.0139$) (supplementary material 2) showed that the treatments with the highest hydrophilicity (lower SCA) were 100% O$_2$, $3.0 \times 10^{-2}$ mbar, 700 V $(20 \pm 4)^\circ$ followed by 100% O$_2$ treatment, $3.0 \times 10^{-2}$ mbar, 600 V $(21 \pm 3)^\circ$. However, no significant differences were observed between them ($p = 0.9981$). Therefore, the lowest voltage treatment was used (Figure 1D).

The main difference between Ar and O$_2$ discharge is that Ar results in chain cleavage and bond rupture on LDPE’s surface, generating Alkyl radicals (C•) by splitting of C-H and C-C bonds. These radicals prefer to interact with adjacent polymer chains rather than with polar groups, forming crosslinks [33,34], while O$_2$ discharge is very reactive and efficient in incorporating the surface polar groups, such as carbonyl (C=O), carboxyl (C=O-OH), and hydroxyls (-OH), which are responsible for decreasing SCA due to hydrophobicity changes [35,36]. Thus, in the present work, the highest hydrophilicity values are those for O$_2$ discharge. In addition, once the discharge ends, C• on the surface can react with atmospheric O$_2$ [30], and when removed from the reaction chamber, they oxidize the material and increase the presence of (C=O), (C=O-OH), esters (C=O-O-R) and (-OH) [14,37].
Abou Rich et al. [30] describe LDPE exposed to the environment after plasma discharge generates O• and hydroxyl radicals (OH•) that tend to take secondary hydrogens from the LDPE chains, resulting in the formation of alkyl radicals (CH₃•) [30]. These radicals can react with atomic oxygen (O₂) or ozone (O₃), allowing the formation of alkoxy (C-O•) and hydroxyl radicals (OH•), and form peroxide (C=O-O•) and hydroperoxide (H-O-O•) radicals [34].

Guruvenket et al. [38] observed when polyethylene (PE) was submitted to Ar and O₂ plasma discharge at different voltages and time of discharge, SCA decrease was greater in comparison with only O₂ plasma treatment. Likewise, they described how the time/voltage ratio required to decrease SCA was lower for O₂ in comparison with Ar discharge [38]. Ar–O₂ plasma forms oxidized structures crosslinked with LDPE’s surface, suggesting a more rapid transformation due to the incorporation of high-density functional groups [39]. In the present work, the highest SCA decrease was observed for treatments with O₂ plasma discharge at lower voltages (600 and 700 V) in comparison with Ar plasma treatment [38], as was described by Gómez-Méndez et al. [29].

Generally, LDPE suffers non-permanent physical and chemical modifications on its surface during and after plasma discharge [11]. Plasma-irradiated surface tends to reverse changes during discharge and return to its original chemical state [40,41] in a phenomenon known as “hydrophobic recovery” [12,33]. Hydrophobic recovery after the best O₂ plasma discharge treatment is illustrated in Figure 2A; at day 0, SCA was at zero degrees and seven days later it was at (65 ± 9)°, representing a 75.4% hydrophobic recovery (stored at 14 °C), while control without plasma discharge remained at (86 ± 3)° (Figure 2A). Pandiyaraj et al. [33] refer to LDPE hydrophobic recoveries of 82% after oxygen discharge of seven days later it was at (65 ± 9)°, representing a 75.4% hydrophobic recovery (stored at 14 °C), while control without plasma discharge remained at (86 ± 3)° (Figure 2A). Pandiyaraj et al. [33] refer to LDPE hydrophobic recoveries of 82% after oxygen discharge.
treatment and 15 days of storage at RT. Sanchis et al. [17] report 30° recovery after 170 h of storage with 30-min O2 plasma treatment. On the other hand, Abou Rich et al. [30] described SCA as close to 80°, even after 60-days post-treatment with oxygen plasma.

Hydrophobic recovery can occur through different mechanisms: (a) the reorganization of the polymer’s surface when low and high molecular weight molecules, not modified by the discharge, diffuse from the material’s core to the outer layers [34]; (b) diffusion of low molecular weight oxidized molecules, and (c) reorientation of chemical polar groups towards the core of the polymer [8,12,42]. According to Morent et al. [43], the degree of crosslinking affects hydrophobic recovery (ageing). The greater the crosslinking of LDPE, the more difficult chain movement will be, reducing the effect of post-plasma ageing [43].

The static contact angle for LDPE treated with O2 plasma for 6 min and control are in Figure 2A (ablation curve). An SCA decrease occurred for sheets submitted to discharge as a function of time until reaching zero (0°) after 4 min, while the SCA for the control was θ = (89 ± 2)°; demonstrating O2 plasma discharge under the conditions was effective in increasing the material’s hydrophilicity. Hydrophobic recovery after treatment appears in Figure 2B. One day after discharge, SCA was at zero degrees (0°) and day 7 after storing at 14 °C, SCA was at (65 ± 9)°, representing a 75.4% recovery. The control without discharge remained at (86 ± 3)°.

2.3. Other Response Variable Associated with Plasma Discharge

The following results were obtained from LDPE sheets submitted to the best treatment: (100% O2 3.0 × 10−2 mbar and 600 V) SCA (21 ± 3)°, final weight (4.2 ± 0.1) mg, roughness (10 ± 3) nm, Young’s modulus (41 ± 6) MPa, and yield strength (11 ± 2) MPa. In comparison, pristine LDPE-treated material presented a 76% SCA decrease (p = 0.0139) with a 100% increase in roughness, whereas Young’s modulus and yield strength increased by 21% and 12%, respectively, but not significantly in comparison with pristine LDPE sheets (p = 0.05234).

Complete Fourier-transform infrared spectroscopy (FTIR) spectra for pristine LDPE sheets (black line) and post-plasma treatment with 100% argon (red line), argon–oxygen gas mix (brown line), and 100% oxygen (blue line) appear in Figure 3. Complete spectra characteristic of LDPE sheets presented signals at 2920 cm−1 (CH2 asymmetric stretching), 1460 cm−1 (CH2 band bending), and 729 cm−1 (CH2 rocking deformation). It is important to highlight that a decrease in these signals appeared after treatments (Figure 3B–D). For the 100% O2 treatment, the decrease in the signal at 906 cm−1 was observed, corresponding to vinyl groups, and signals at 975 cm−1 and 1180 cm−1 appeared, indicating peroxide groups and alcohol bonds, respectively (Figure 3E). Figure 3B illustrates a decrease in the signal at 2920 cm−1, which can be attributed to differences in the material’s width, caused by the ablation process [27]. Rajandas et al. [28] describe how the 2920 cm−1 signal

![Figure 2](image-url)

**Figure 2.** LDPE static contact angle changes after plasma treatment and recovery. (A) LDPE sheet static contact angle (SCA) decrease after 6-min treatment with O2 plasma discharge (ablation curve). (B) Hydrophobic recovery in LDPE sheets after being submitted to O2 plasma discharge and stored for 7 days, represented as changes in SCA.
is directly proportional to LDPE’s concentration, which could indicate that the material exposed to O$_2$ plasma discharge when presenting a higher intensity would involve a loss of mass [28]. The same behavior was observed in Figure 3C,D. Shi et al. [44] specified that plasma active species, such as ions and electrons, bombard LDPE’s surface macromolecular chains, generating a cleavage in the C-H and C-C bonds producing oligomers and small organic molecules; hence, increasing loss of mass [44]. Shikova et al. [3] described O$_2$ plasma discharge on PE unsaturated vinyl groups (-CH=CH$_2$), vinylenes (R1R2C=CH$_2$), and trans vinylenes (-CH$_2$=CH$_2$), whose absorption bands appear between 890 and 960 cm$^{-1}$ (Figure 3E). In the present study, in the vicinity of 906 cm$^{-1}$, presence of vinyl was observed in the pristine LDPE, as well as Ar and Ar-O$_2$ mixed treatment, which could suggest that LDPE sheets employed in this study contained impurities of unsaturated nature. After, 100% O$_2$ plasma treatment bands in the 1180 cm$^{-1}$ and 975 cm$^{-1}$ range appeared, associated with alcohol and peroxide groups, respectively [45]. These results suggest LDPE’s sheet double bond loss and gain of carbonyl and peroxide groups after O$_2$ plasma treatment. Exposure of LDPE sheets to plasma produces mainly radicals, which are converted into peroxide radicals, subsequently transformed by auto-oxidation and/or crosslinking into hydroperoxides [18]. Peroxide radical formation takes place mainly in the amorphous region due to oxygen high diffusion capacity [8].

Figure 3. Cont.
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Figure 3. Fourier-transform infrared spectroscopy (FTIR) spectroscopy for LDPE sheets after plasma treatments with argon (Ar) (red line), oxygen (O2) (blue line), and Ar–O2 mix (brown line). The black line corresponds to LDPE pristine. (A) Complete spectra show LDPE characteristic signals. (B–E) Details characteristic of LDPE signals after plasma treatments.

LDPE’s morphological surface changes after plasma treatment were evidenced by SEM, whereas atomic force microscopy (AFM) revealed roughness results. Pristine LDPE surface without treatment is illustrated in Figure 4A. The pristine LDPE (Figure 4A) presented a homogeneous surface, while the sheets exposed to O2 (100%) exhibited much more change (Figure 4D) due to the presence of cracks and fissures on the surface (ablation). The AFM data revealed that the average roughness of the pristine LDPE was (5 ± 1) nm, while that of the 100% O2 treatment was (10 ± 3) nm (p = 0.0196). As determined by AFM after discharge, LDPE’s surface roughness changes doubled in comparison with pristine LDPE. Thus, this demonstrates the existence of processes leading to the material’s ablation, which removed the surface layers of the material [10]. These changes depend on the time of plasma exposure and voltage [32]. In this work, LDPE’s roughness increased by (5 ± 4) nm under 0.03 mbars of pressure with plasma generated at a cathode distance of 5.6 cm and six minutes of O2 gas exposure.

Figure 4. LDPE scanning electron microscopy (SEM) images submitted to Ar and O2 plasma treatments. Pristine LDPE (A). LDPE submitted to Ar plasma (B). Ar–O2 (1:1) mix (C). O2 plasma treatment (D).
As shown by scanning electron microscopy (SEM) images, the greatest change was observed for O₂ treatment (Figure 4D), followed by Ar–O₂ mix (Figure 4C), and Ar last (Figure 4B). Pandiyaraj et al. [33] achieved greater roughness with an Ar–O₂ plasma mix treatment for 10 min, 0.2 mbar pressure, and a 5 cm cathode distance. Švorčík et al. [10] reported an increase in PE roughness from 4 nm to 7 nm s after Ar plasma. Ataeefard et al. [11] reported roughness changes after exposing LDPE to Ar, O₂, N₂, and CO₂ plasma discharge. Moreover, Sanchis et al. [17] observed an increase in roughness from 21 nm to 30 nm after submitting LDPE sheets to 30 min of O₂ discharge. According to Ataeefard et al. [42], defined surface morphology can vary depending on the discharge voltage, time of exposure, and the type of gas employed [42]. According to Sanchis et al. [17], roughness augments LDPE’s wettability. In the present study, O₂ plasma discharge increased LDPE roughness five-fold and generated greater hydrophilicity (four-fold higher than pristine LDPE), demonstrating O₂ plasma discharge was more effective in generating ablation processes on LDPE’s surface in comparison with Ar discharge.

For the viscoelastic properties, Young’s modulus and yield strength values increased after O₂ plasma discharge. An augmentation in LDPE’s viscoelastic properties suggests it becomes more rigid after treatment due to two facts; crosslinking between chains generated by plasma discharge [9] and polar group incorporation [11], restricting their relative movement, enhancing the polymer’s resistance. Other reasons for the increase in material stiffness are that LPDE is a semi-crystalline polymer, and after being plasma exposed (under the conditions of this study), the crystalline zone remained unchanged, but the weak bonds of the amorphous region were lost.

2.4. TiO₂/UV Photocatalysis
2.4.1. Commercial TiO₂ Characteristics

Figure 5 depicts SEM images at different magnifications (5000× and 10,000×). At 5000×, a uniform material was observed with 1.23 ± 0.19 µm particle size. Observed spaces corresponded to cracks or sites where no TiO₂ was deposited (Figure 5A). At greater magnification (10,000×), formation of aggregates composed of different particle sizes (polydisperse material) and forms (polymorphic) was observed, while spherical shapes predominated in Aldrich brand TiO₂ (Figure 5B).

Figure 5. TiO₂ (Aldrich) SEM and XRD analysis. (A) TiO₂ (Aldrich) SEM images at 5000× TiO₂ (Aldrich). (B) SEM at 10,000×. (C) EDS analysis. (D) XRD results.
EDS analysis revealed that the Aldrich™ TiO$_2$ contained titanium (Ti) (8.90 wt%) and oxygen (O$_2$) (89.62 wt%) (Figure 5D). On the other hand, the XRD profile (Figure 5C), shows how the peak with the highest intensity is the orientation (A101), which corresponds to the anatase phase of TiO$_2$ [46,47].

2.4.2. 2$^2$ Factorial Design and Photocatalysis at 300 h

According to Table S3, the static contact angle, final weight, Young’s modulus, and yield strength were significant ($p < 0.0001$). Therefore, the main factors and interactions among factors were evaluated. For SCA, the $R^2$ was 0.8223, while for the final weight it was 0.8795. For the viscoelastic properties of Young’s modulus and the yield strength, the $R^2$ were 0.9800 and 0.8100, respectively. There was a high correlation among predicted and observed values with precision values greater than 4.0, confirming results were not the result of experimental noise (Table S3).

pH and TiO$_2$ concentration were factors that influenced SCA ($p < 0.0001$), with contributions of 45% and 40%, respectively. Additionally, standardized effects with their respective signs were +43.6, −6.5, −6.1, and +0.5; indicating the aforementioned factors could be employed at their low level. pH and TiO$_2$ concentration were also influential factors ($p < 0.0001$ and $p = 0.0002$, respectively); regarding LDPE sheet weight loss, the highest contribution percentage was obtained with pH (60%) (Table S3).

On the other hand, LDPE sheet viscoelastic properties were included in the statistical analysis (Young’s modulus and yield strength), which were significant ($p < 0.0001$). For Young’s modulus, pH and TiO$_2$ concentration and their interaction were significant ($p < 0.0001$, obtaining 33% contribution percentages (Table S3)). These results indicate a decrease in Young’s modulus could be favored by low levels of these factors. Last, for yield strength, the most influential factor was pH (factor A, $p < 0.0001$), followed by AB interaction ($p = 0.0007$), and TiO$_2$ concentration (factor B, $p = 0.026$), with contribution percentages of 70, 20, and 30%, for factor A, B, and AB interaction, respectively. Again, it was observed that a decrease in yield strength was favored by low levels of each factor and even their interaction was employed (Table S3).

Mean comparison among treatments demonstrated significant differences for SCA ($p = 0.0032$), final weight ($p = 0.0001$), Young’s modulus ($p < 0.0001$), and yield strength ($p = 0.0095$), were observed (Figure 6). Based on obtained results, treatment one was selected (1 g L$^{-1}$ TiO$_2$ and pH 4.5) to perform photocatalytic transformation curves because the lowest contact angle and lowest weight at 300 h of UV light exposure were observed.

The treatment that presented the best results in terms of weight and SCA decrease was T1 pH 4.5 ± 0.2 and 1 gL$^{-1}$ TiO$_2$ (Figure 6). pH is a relevant photocatalytic factor as reported in the literature, since below the isoelectric point (pH of 6.5 ± 0.2) at acid conditions [6], TiO$_2$ acquires a positive charge with $h^+$ generated as the dominant species with the highest oxidative capability, which allows attraction of compounds of the opposite charge (negatively charges), enabling their oxidation [48]. Therefore, plasma pretreatment negatively charged LDPE sheets because of forming polar groups on the surface. This in turn allowed for transitory adsorption phenomena to take place, generating the LDPE/TiO$_2$ composite. These phenomena permitted more and better contact between the photo-exciter catalyzer and LDPE sheets, bringing about higher oxidation and photoinduced wear by the action of free radicals, and hence, photocatalytic degradation of the material. In the present work, the initial pH was 4.5 ± 0.2 and ended at 7.9 ± 0.2, an increase that could be attributed to (-OH) ion increase in the solution [6]. After exceeding the isoelectric point, TiO$_2$ was a negative charge, thus separating from the material due to charge similarity. However, as observed in SEM images, cavity generation could have facilitated TiO$_2$ particle capture, continuing with the photocatalytic degradation, which suggests that the photo-generated e$^-$ are transferred to adsorbed O$_2$, whose reduction allows the superoxide anion radical production (O$_2$•$^-$), which in turn favors hydroxyl radical formation (•OH) as dominant species [49], and this species was perhaps responsible for the degradation of the material.
LDPE photocatalytic degradation was evident from total organic carbon (TOC) and final weight data. Total carbon concentration increased from (4.07 ± 1.04) mg L$^{-1}$ (at 0 h), to (6.73 ± 0.77) mg L$^{-1}$ at 400 h, which could have been associated with fragmentation and hydrolysis of certain carbonated fractions of the polymer. Furthermore, the final weight decreased to 0.2 mg (Figure 7C).

Figure 6. 2$^2$ Factorial design response variables. (A) Static contact angle. (B) Weight. (C) Young’s modulus. (D) Yield strength. For SCA (35 ± 9)$^\circ$, T1 was significant whereas for weight, T1 (3.4 ± 0.5) mg and T3 (3.6 ± 0.6) mg were significant. For Young’s modulus, T4 was significant (21 ± 1) MPa and yield strength (2 ± 1) MPa. T1: pH 4.5, TiO$_2$ 1 g L$^{-1}$, T2: pH 9.0, TiO$_2$ 1 g L$^{-1}$, T3: pH 4.5, TiO$_2$ 10 g L$^{-1}$, and T4 pH: 9.0, TiO$_2$ 10 g L$^{-1}$. Letters a, b, c, d correspond to heterogeneous groups.

TiO$_2$ was a critical factor since high concentrations can minimize radiation reaching the material’s surface. Too high concentrations can decrease the effect UV can have on the system and photocatalytic degradation velocity, generating in the solution a “screen”-like effect [50].

Since the system operated with UV light it generated chemical photolysis, whose energy was sufficient to cleave the polymer’s covalent bonds, forming high-energy carbon-free radicals that reacted with O$_2$ to form aldehydes and carboxylic acids [51], in addition to chain crosslinks. The photocatalytic effect was favored by low TiO$_2$ concentrations in T1 treatment (Figure 6).
This work innovated in performing a photocatalysis with TiO$_2$ in solution and the contaminant (LDPE sheets) in a solid phase, and not the other way around, as it is described for most reports in the literature [46,47]. This condition could have favored photocatalytic processes, since according to Bouna et al. [52], TiO$_2$ adhered to a surface that can form films, which diminish the photocatalytic activity since the photocatalytic activity is less in a state of suspension [52]. For treatment of T1 during the first 100 h of the process, TiO$_2$ particles were in an acid solution. According to Wang et al. [53], acid pH increases the number of aggregated particles, which drastically decreases TiO$_2$ solubility [53]. Although this situation could have been present at the beginning of the process, it was minimized by the constant use of aeration within the photoreactor.

As in this work, others researchers have published that a mixture of the anatase–rutile phases, (Figure 5), present a greater yield or activity in comparison to using the anatase phase [54,55]. Results have been attributed to a higher redox potential due to electron transfer from the anatase conduction band to the rutile phase conduction band [56].

2.4.3. Photocatalytic Transformation Curve at 400 h

While the process of photocatalysis took place, a decrease in SCA was observed, with a final value of $(16 \pm 3)^\circ$, which after 400 h of treatment represented an 82% reduction in comparison with pristine LDPE. For Young’s modulus, an increase was observed after the first 100 h, which gradually decreased to end in $(47 \pm 5)$ Mpa, representing a 38% increase (Figure 7A). Yield strength and pH behavior are presented in Figure 7B. As the process developed in time, LDPE sheets were modified and yield strength decreased with a final value of $(9 \pm 2)$ Mpa. In comparison with pristine LDPE value, a 10% decrease was observed. Concerning pH, the initial value was $4.5 \pm 0.2$. After 400 h of treatment, it ended in $7.9 \pm 0.1$ (Figure 7B). Furthermore, LDPE sheets presented a slight decrease in weight, which indicated the process of photocatalytic degradation was taking place (Figure 7C). Last, for TOC as was observed for SCA treatment, a sharp increase was observed during the first 100 h, with a gradual decrease (Figure 7C). Overall, TOC increased from $4.07 \pm 1.04$ mg L$^{-1}$ (at 0 h), to $6.73 \pm 0.77$ mg L$^{-1}$ at 400 h.

For SCA photolysis control, values were $(61 \pm 9)^\circ$, a value greater than treatments, a final weight of $(4.2 \pm 0.3)$ mg, like T4, and Young’s modulus of $(48 \pm 4)$ MPa, and yield strength of $(4 \pm 1)$ MPa, comparable with T2. Collectively, these data indicate photolysis by itself is not sufficient to generate a hydrophilic event on LDPE sheets and facilitate their degradation, yet in combination with plasma discharge, they can significantly increase the transformation effect.

2.4.4. Fourier-Transform Infrared Spectroscopy (FTIR)

FTIR spectra for pristine LDPE (black line), O$_2$ plasma-treated (red line), and sequential plasma followed by photocatalysis treatment (T1, P + P, blue line) appear in Figure 8A. LDPE distinctive signals (2920 cm$^{-1}$, CH$_2$ asymmetric stretching; 2851 cm$^{-1}$, CH$_2$ symmetric stretching; 1471 cm$^{-1}$, CH$_2$ bending and deformation; and 719 cm$^{-1}$, CH$_2$ rocking and deformation, were also present in spectra with plasma treatment (P + F), although with less intensity (Figure 8B–D)). Signals observed in pristine material and post-plasma material increased their signal after photocatalysis, at 610 cm$^{-1}$ as a signal associated with the C-OH bond (twisting deformation) (Figure 8E). Moreover, signals at 1186 cm$^{-1}$ and 1000 cm$^{-1}$ (Figure 8G) were related to quaternary carbon and C-OH stretching bond, respectively. After plasma and photocatalysis, sequential treatments observed LDPE changes appear in Figure 8. As aforementioned for Figure 5, characteristic LDPE signal decrease at 2920 cm$^{-1}$ (Figure 8B), could be attributed to ablation processes [27], which release carbon-containing material, decreasing LDPE weight [28]. During plasma discharge followed by photocatalysis (P + P), presence of two signals was observed (1200 cm$^{-1}$ and between 1600–1800 cm$^{-1}$) (Figure 8F), which corresponds to C-O vibrations and carbonyl signal (C=O), respectively. These later disappeared after 400 h of the photocatalytic process. In this latter one, an ample signal was observed in the vicinity of 610 cm$^{-1}$ (Figure 8E), corresponding to deformation
in the C-OH bond [45]. The presence of (C=O) groups during plasma discharge [23] and hydroxyl (OH-) during photocatalysis suggests the existence of LDPE oxidation processes for both treatments.

![FTIR spectra for untreated and treated LDPE sheets](image)

**Figure 8.** FTIR spectra for untreated and treated LDPE sheets. Pristine LDPE (black line), post-plasma (red line), and after sequential plasma and photocatalysis (blue line). (A–D) Regions characteristic of LDPE, where after the treatments, a decrease in signal appeared. (E–G) Regions where changes after the sequential process occurred.

LDPE SEM images obtained from the $2^2$ factorial design are depicted in Figure 9, without analysis (9A) and analyzed through image processing (9B). It can be observed that treatment T1 (pH 4.5 ± 0.2 and 1 gL$^{-1}$ TiO$_2$) (Figure 9B,E) resulted in the highest surface modification (cavity appearance) of 8.86% (Figure 9B’) and 11.94% (Figure 9E’) for the same treatment at 400 h. Table 2 shows the response variables during sequential plasma-photocatalysis.
Figure 9. LDPE SEM images after different photocatalytic degradation treatments. (A–E) LDPE SEM images after different photocatalytic degradation treatments ($2^5$ factorial design for 400 h) without analysis. (A’–E’) Same treatments with image analysis. In red, percentage quantification of generated cavities for different treatments. (A) Photolysis control. (B) T1. (C) T2. (D) T4 and (E) T4 at 400 h.

Table 2. Comparison of response variables during sequential plasma-photocatalysis treatment.

| Response Variables | Pristine | Plasma | Photocatalysis |
|--------------------|---------|--------|---------------|
| Weight (mg)        | 4.3 ± 0.5 | 4.2 ± 0.1 | 4.0 ± 0.1 |
| Static contact angle (°) | 87 ± 1   | 21 ± 3  | 16 ± 3       |
| Young’s modulus (MPa) | 34 ± 1  | 41 ± 6  | 46 ± 5       |
| Yield strength (MPa) | 10 ± 2  | 11 ± 2  | 9 ± 2        |

Scanning electron microscopy revealed LDPE’s superficial modifications after different photocatalytic treatments (Figure 9). Substantial topographical changes were observed when comparing pristine material (Figure 4A) with treated LDPE or even with LDPE submitted only to photolysis (Figure 8A). Out of the factorial design treatments, treatment T1 (pH 4.5 ± 0.2/1 gL$^{-1}$ TiO$_2$) (Figure 8B) presented the greatest surface changes, which were superior in the 400-h curve (Figure 8E), displaying cavities and a very rough surface. White particles on the surface corresponded to TiO$_2$. The presence of cavities resulted from volatile product release from the polymeric matrix [57], due to TiO$_2$/LDPE adsorption processes and later oxide reduction reactions during photocatalysis. These types of alterations have also been observed by other authors [57–59].

When comparing LDPE’s physical properties after the sequential plasma-photocatalysis processes with pristine material, it was observed that treatment generated a complementary effect that maintained hydrophilicity, attaining an 81.6% decrease for SCA. Another important change observed was a 38% increase in Young’s modulus, demonstrating these processes increased the material’s rigidity by restricting atom movement by the incorporation of polar groups, by crosslinking and bond cleavage in LDPE amorphous region, evidenced as cavities on the surface, which increase the crystalline zone proportion [29].

3. Materials and Methods

3.1. LDPE Sheets

LDPE sheets were purchased from a local market in Bogotá, D.C., Colombia. They were prepared for experiments as described [29].

3.2. Direct-Current Low-Pressure Plasma (DC-LLP) and Plasma Discharge

Within a high vacuum chamber (18 cm × 18 cm) with glass walls and stainless-steel lids and flanges, two flat electrodes, anode, and cathode, were placed with parallel cylinders separated by varying “s” distances to generate a low-temperature self-sustained DC plasma. Argon and oxygen reactive gas mix were rarefied at different concentrations (supplementary material 1). Plasma discharge condition selection that favored LDPE sheet modification included: gas effect (Ar, O$_2$, and a mix of Ar–O$_2$ at a 1:1 ratio), voltage (V),...
and plasma chamber pressure (mbar). Selection of operating conditions and their effect on LDPE are detailed in supplementary material 2.

Once treatments that met these conditions were identified, operation settings that would decrease static contact angle (SCA) in LDPE sheets were selected as follows: 5 treatments for O$_2$, 13 for Ar, and 14 for the Ar–O$_2$ mix (Table S1). Differences among conditions for SCA-obtained results were analyzed, as well as mean comparison among treatments for each gas and mix (independent blocks), where the most significant for each one was selected ($p < 0.05$). The analysis was performed by ANOVA employing SAS® software (SAS Institute 2017, version STAT 14.3, Cary, NC, USA: SAS Institute). Following, a new statistical analysis was performed for each block using the same tests previously selected (O$_2$, Ar, and Ar–O$_2$ mix). With the obtained results, subsequent experimental conditions were established [17,33,43,60].

3.3. Ablation Curve and Hydrophobic Recovery

Employing the glow discharge treatment that generated the lowest SCA, a new assay was performed in the wet chamber [29] to determine the effect plasma treatment would have on LDPE’s hydrophobicity changes as a function of time (ablation curve) [2]. To this end, five LDPE sheets were submitted to O$_2$ plasma discharge for 30, 60, 120, and 240 s of exposure. As control, pristine LDPE sheets were used and results were expressed as final SCA in degrees ± standard deviation.

Additionally, following Mortazavi and Nosonovsky [12], LDPE sheet hydrophobic recovery [2] was evaluated for seven days. To this end, a new set of 10 pristine LDPE sheets was divided into two lots. The first five sheets were treated with plasma discharge and the remaining five were used as controls. To determine hydrophobic recovery percentage after ablation treatment, LDPE sheets were placed within a plastic box and left at room temperature (14° C), and for seven days SCA was measured [12].

3.4. TiO$_2$/UV Photocatalysis

For photocatalysis, titanium dioxide (TiO$_2$ Aldrich™) mixture of rutile and anatase (99.5% purity) was employed. To observe morphological characteristics and semi-quantitative composition, scanning electron microscopy images were captured, coupled with energy dispersion spectroscopy (SEM/EDS). A Jeol™ (Tescasn Brno, Czech Rep.) JSM 6490LV scanning electron microscope with 30kV, SEI signal, and 5000–1000 × magnification was used. Observations were performed by depositing 0.5 g TiO$_2$ on a glass substrate [61]. The sample was oven-dried at 50 °C for 24 h. To determine TiO$_2$ crystal structure, X-ray diffraction was performed (XRD) using an X-Ray diffractometer Siemens D-5000 (Munich, Germany) equipment operated with a Cu K$_{α}$ = 1.5418 Å anode [47,61,62].

For the photocatalysis experiments, a 2.5 L photoreactor (Sorvirel™-Bogotá, Colombia) with a work effective volume (WEV) of 1.2 L was employed. A 15 W UV lamp (Phillips™) with UV-C (280–100 nm) light emission capacity located within the photo-reactor and protected with a quartz glass jacket was employed. Within the reactor’s lower part, three ports for air injection were located and the air was injected at 10 ft$^3$/h (SCFH) (4.7 L min$^{-1}$). Positioned within the reactor and in parallel with the UV lamp, inert clamps were installed to hold previously O$_2$ plasma-irradiated LDPE sheets (14 sheets of 3 cm$^2$) [29] (Figure 10).

Condition selection for LDPE sheet photocatalytic transformation was performed through a $2^2$ factorial design where evaluated factors included Factor A: pH with a (−1) level of 4.5 ± 0.2 and a (+1) level of 9.0 ± 0.2. Factor B was TiO$_2$ in gL$^{-1}$, with a (−1) level of 1.0 and a (+1) level of 10 gL$^{-1}$ (TiO$_2$ was added within the reactor and homogenized by air injection with UV lamps off).
The design generated four codified treatments as follows: T1: $-1 - 1$ (pH: 4.5, TiO$_2$ 1 gL$^{-1}$), T2: $+1 - 1$ (pH: 9.0, TiO$_2$ 1 gL$^{-1}$), T3: $-1 + 1$ (pH: 4.5, TiO$_2$ 10 gL$^{-1}$), and T4: $+1 + 1$ (pH: 5, TiO$_2$ 1 gL$^{-1}$), performed in triplicate during 300 h of photocatalysis. After each treatment, LDPE sheets were removed to analyze response variables: SCA, final sheet weight, Young’s modulus, and yield strength [29]. Significant treatment selection was determined by an analysis of variance and a mean comparison using Design-Expert$^\text{®}$ (Stat-Ease Inc. 2017 software, version 11.0. Minneapolis, MN: Stat-Ease) and SAS$^\text{®}$ (SAS Institute 2017, version STAT 14.3. Cary, NC, USA: SAS Institute) [63].

The selected $2^2$ factorial design treatment was employed to perform photocatalytic degradation curves during 400 h; every 100 h, three LDPE sheets were removed to analyze changes associated with final weight (mg), SCA ($^\circ$), Young’s modulus (Mpa), and yield strength (Mpa). Additionally, SEM and Fourier-transform infrared spectroscopy (FTIR) studies were performed. As controls, LDPE sheets exposed only to photolysis and plasma discharge were employed. Determinations for controls were performed at the beginning and the end of the process. Moreover, to determine total organic carbon (TOC) [63], pH aqueous solutions containing TiO$_2$ were sampled, separating the chemical compound by sedimentation. Results obtained at 400 h of treatment were analyzed by ANOVA to determine differences among treatments. Additionally, multiple variable correlations with a confidence interval of 95% were performed using SAS$^\text{®}$ software (SAS Institute 2017, version STAT 14.3. Cary, NC, USA: SAS Institute) [63].

3.5. Variables and Analytical Techniques Associated with LDPE Sheet Changes

3.5.1. Static Contact Angle

To obtain LDPE’s surface microscopic information (atomic and/or molecular) in terms of its macroscopic properties, a correlation between static contact angle and tension or surface adhesion was established (hydrophilicity or wettability) [64,65] (supplementary material 3).

3.5.2. Viscoelastic Properties

LDPE sheet Young’s modulus [39] and yield strength [66] were determined from stress-strain curves employing a Cobra 4 dynamometer, a THORLAB$^\text{TM}$ (Newton, NJ, USA) MTS25-Z8 motorized translation stage and Phywe Measure 4 (version 1.4, 2010) and ATP User (version 1.0.28, 2008) software for supervision. Control operating conditions were as follows: 0.1 mm s$^{-1}$ testing speed, 250% maximum strain and 5 N maximum stress.

Figure 10. Photoreactor’s assembly for the photocatalysis process.
3.5.3. Microscopy

Scanning electron microscopy (SEM) [67,68] (Jeol™ JSM 6490LV) with a 10 kV to 20 kV potency SEI signal and 500 and 6500× magnification was used to characterize LDPE’s surface. Samples were coated with gold in the Denton Vacuum Desk IV preparation system. Service was contracted with Universidad de Los Andes (UNIANDES), Bogotá, D.C., Colombia.

Atomic force microscopy (AFM) [30,64] [(Nanosurf™ easy scan 2) Liestal, Switzerland] was used to study the surface’s changes in roughness. Parameters were: Mode: contact, Size: 61.8 μm, Setpoint: 20 nN; P-Gain: 1000; I-Gain: 100; D-Gain: 0. For roughness calculation, three measurements at different locations of the sample were carried out and mean ± SD was determined according to (Equation (1)) [69].

\[ R = \frac{1}{N} \sum_{i=1}^{N} |Z_i - Z| \]  

where: \( N \) is surface height data number and \( Z \) mean height distance

3.5.4. Computational Image Processing

Computational SEM image processing techniques were used to measure surface cavities produced by sequential treatment (plasma + TiO\(_2\)/UV photocatalysis). All images were formatted to 1280 × 860 pixels and their noise was reduced by a Gaussian filter aided by a Euclidean curve shortening flow process to enhance edges [70]. The brightest sections were selected using the morphological Top-Hat transformation [71]. Lastly, a binary process based on morphology was used to select pixels, whose intensity was above a base threshold. The cavity area was calculated as the number of pixels selected divided by the total number of pixels in the image.

3.5.5. Fourier-Transform Infrared Spectroscopy (FTIR)

Fourier-transform infrared spectroscopy was used to evaluate LDPE chemical bond modifications and chemical group composition [42,64,67]. A Shimadzu™ (Kyoto, Japan) IR Prestige-21 spectrophotometer was used by following the parameters reported by Gómez-Méndez et al. [29].

3.5.6. Total Organic Carbon (TOC) and pH for the TiO\(_2\) Solution

For each sampling interval, 15 mL TiO\(_2\) solution was collected, and pH was measured in duplicate [63] using an Okaton™ (Skokie, IL, USA) pH meter. The sample could sediment for 24 h, 10 mL of the supernatant was collected and analyzed in triplicate with a TOC analyzer [63] (Shimadzu™ TOC-L).

4. Conclusions

LDPE sheets treated with 100% O\(_2\) plasma at 3.0 × 10\(^{-2}\) mbar, 600 V, six minutes of exposition showed in comparison with pristine LDPE a 75.58% reduction in SCA because of polar group incorporation (peroxides and alcohols) on the material’s surface, making the material hydrophilic, and a 94.89% increase in roughness, due to ablation and an increase in rigidity because of carbonated chain relative movement decrease from crosslinking and oxide reduction events. These oxidations facilitated the photocatalytic activity of a TiO\(_2\) solution (1 gL\(^{-1}\), pH 4.5 ± 0.2, 400 h), generating cavities on LDPE’s surface or expanding the ablation initiated by the plasma discharge, yet retaining high hydrophilicity and material’s rigidity, hence demonstrating a complementary effect of both processes. To the best of our knowledge, this is a pioneering work employing LDPE sheets in a sequential O\(_2\) plasma and TiO\(_2\)/UV photocatalysis in solution, maintaining low SCA. Our data suggest sequential treatment with physicochemical processes in a complementary manner should be evaluated on plastic transformation.
**Supplementary Materials:** The following are available online, Figure S1: Schematic diagram of the plasma reactor employed in this study. Figure S2: Profile of the spherical cap description for a sessile droplet. Table S1: Treatments employed for oxygen, argon, and argon-oxygen (50:50) mix. Table S2: ANOVA results for Ar, O₂ and Ar-O₂ (50:50) mix. Table S3: SCA; roughness and LDPE’s viscoelastic properties obtained during the 2²-factorial design. Table S4: Response variables obtained during the 2²-factorial design (300 h). Material S1: Direct Current low-pressure plasma (DC-LLP) - Assembly and operating conditions. Material S2: Plasma discharge conditions selection, Material S3: Static contact angle.

**Author Contributions:** Conceptualization, A.M.P.-R., J.C.S.-R. and R.A.P.-P.; methodology, L.D.G.-M., L.C.J.-B., A.P.-F., A.V., and J.M.B.; software, A.V. and J.M.B.; investigation, L.D.G.-M., A.V. and J.M.B.; data curation, L.D.G.-M., J.C.S.-R., L.C.J.-B. and A.M.P.-R.; writing—original draft preparation, L.D.G.-M.; writing—review and editing, A.M.P.-R., J.C.S.-R., A.P.-F. and R.A.P.-P.; project administration, A.M.P.-R.; funding acquisition, A.M.P.-R., J.C.S.-R. and R.A.P.-P. All authors have read and agreed to the published version of the manuscript.

**Funding:** This research was funded by Pontificia Universidad Javeriana Bogotá, D.C. Colombia, grant number 00006736, titled “Transformación de polietileno de baja densidad (LDPE) por medio de ablación con plasma y actividad enzimática de Pleurotus ostreatus”.

**Acknowledgments:** This work was supported by “Pontificia Universidad Javeriana Bogotá, D.C. Colombia” Grant ID: 00006736, titled “Transformación de polietileno de baja densidad (LDPE) por medio de ablación con plasma y actividad enzimática de Pleurotus ostreatus”. The authors thank Maria Lucia Gutierrez Ph.D., for English editing.

**Conflicts of Interest:** The authors declare that no competing interest exist. “Pontificia Universidad Javeriana” did not play any role in the study design, data collection, analysis, decision to publish, or manuscript preparation.

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