Mechanical Recycling of Thermoplastics: A Review of Key Issues

Alae Lamtai 1,2, Said Elkoun 1,2,*, Mathieu Robert 1,2, Frej Mighri 2,3 and Carl Diez 4

Abstract: During the last decade, the consumption of plastics has increased highly in parallel with plastic waste. The transition towards a circular economy is the only way to prevent the environment from landflling and incineration. This review details the recycling techniques with a focus on mechanical recycling of polymers, which is the most known and developed technique in industries. The different steps of mechanical recycling have been highlighted, starting from sorting technologies to the different decontamination processes. This paper covers degradation mechanisms and ways to improve commodity polymers (Polyolefins), engineering polymers (PET, PA6), and bio-sourced polymers (PLA and PHB).

Keywords: mechanical recycling; sorting technologies; decontamination process; polyolefins; engineering (PET, PA6) and bio-sourced polymer (PLA and PHB)

1. Introduction

Plastic materials have gathered attention recently due to their omnipresence in the global economy. Since last century, plastics have become rapidly one of the most used materials in industry. In 2019, more than 400 million tonnes of plastics (Mt) were produced (Figure 1) [1].

Figure 1. Global plastics production: 1950 to 2019 [1].
If production continues to grow at a similar rate, plastics production will reach 1600 million tons (Mt) in 2050. The rapid growth of plastics production is due to the good properties and low cost of this material. Thanks to its versatility, this material is used in several fields, such as packaging, textile, transport, and construction. Polymers are widely used, depending on the final application (Figure 2).

The proliferation of plastic production contributes significantly to greenhouse gas emissions and generates pollution in the natural environment. Indeed, the production of virgin plastics requires the transformation of petroleum into monomers. This process is energy-intensive and generated 400 million tons (Mt) of greenhouse gas emissions in 2012 [2].

To protect the environment, some countries adopted a new economic model that aims to revalorize post-consumed plastic and avoid landfilling. The transition toward a circular economy is unavoidable to reduce the plastic footprint and promote recycling. To manage plastic waste, there are different gates that can be classified from the most to the least preferred (Figure 3) [3].

![Hierarchy of priority in plastics management](image)

**Figure 3.** Hierarchy of priority in plastics management [4].

<table>
<thead>
<tr>
<th>Collection</th>
<th>Sorting</th>
<th>Cleaning</th>
<th>Shredding</th>
<th>Processing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Most Preferred</td>
<td>Reduce</td>
<td>Recycle</td>
<td>Repurpose</td>
<td>Least Preferred</td>
</tr>
<tr>
<td>Avoid and reduce plastic waste</td>
<td>Direct reuse, repair, remanufacture, refurbish</td>
<td>Mechanical and chemical recycling, composting</td>
<td>Converting plastic wastes into energy</td>
<td>This should only be used as a last resort.</td>
</tr>
</tbody>
</table>
Waste management places reduction as the top priority. The idea is to prevent the unnecessary consumption of resources. Direct reuse of original products is the second-best practice in waste management. The third stage is recycling products to avoid landfilling. Repurpose is about energy recovery. If the material cannot be recycled and recovered to energy, it will be landfilled, but it is the least preferred stage in the waste management hierarchy.

To achieve the goal of 100% recovery of plastics, the waste management system should be extended to all fields using plastics. In the industry, there are four ways to recover plastics: primary, secondary, tertiary, and quaternary recycling. This review aims to highlight the different stages of mechanical recycling: collection, sorting, cleaning, shredding, and processing (Figure 4).

2. Sorting Technologies

Plastic separation faces a lot of challenges due to the huge quantity of plastics to collect and the complexity of identifying some types of plastics. The sorting is important to remove contaminants from plastics. This section will cover separation techniques that use density, surface charge transfer, and spectral analysis. The most used sorting methods are listed in Table 1.

<table>
<thead>
<tr>
<th>Sorting Method</th>
<th>Characteristics</th>
<th>Drawbacks</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual</td>
<td>Hand based sorting</td>
<td>Laborious, bad working environment</td>
<td>[5]</td>
</tr>
<tr>
<td>Tribo-electric</td>
<td>Based on electrostatic charge</td>
<td>Only for clean, dry, and non-surface-treated products.</td>
<td>[6]</td>
</tr>
<tr>
<td>Near-infrared Radiation (NIR)</td>
<td>Fundamental vibration</td>
<td>It is not adapted for dark plastics and is very expensive.</td>
<td>[7]</td>
</tr>
<tr>
<td>Flotation</td>
<td>It is related to a specific gravity of material.</td>
<td>It is not applied to High-density Polyethylene (HDPE) and Low-density Polyethylene (LDPE).</td>
<td>[8]</td>
</tr>
<tr>
<td>X-ray fluorescence (XRF)</td>
<td>Uses X-rays as a source</td>
<td>It is very expensive</td>
<td>[9]</td>
</tr>
</tbody>
</table>

2.1. Manual Sorting

Manual sorting is a technique that allows the identification by shape and color of the plastics visually. This technique is useful if the waste plastics are large and easily identified. Otherwise, it’s very laborious and inefficient due to human errors. Moreover, it is the cheapest sorting technique [10].
2.2. Near Infrared Radiation (NIR) and X-ray Technology

NIR (Near Infrared Radiation) involves irradiating the unsorted, unidentified plastic with near-infrared waves (600 to 2500 nm\(^{-1}\) in wavelength). When exposed to light waves, different polymer reflects an identification spectrum. NIR spectroscopy allows identification thanks to the plastic signature, but it’s not adapted for dark plastics since dark pigments mask the signature of plastic material [7].

There is another technique, like NIR, called X-ray fluorescence spectroscopy (XRF), which is used to identify flame-retardant materials (FR) and to determine the chemical composition of all kinds of materials (metal, cement, oil, and polymer).

This technology uses X-rays produced by a source to irradiate a sample. This latter produces fluorescent X-ray radiation with discrete energies that are characterized for these elements. Each element present in the sample produces a specific and unique set of characteristics of fluorescent X-rays [11,12].

2.3. Flotation Method

The flotation method aims to separate polymers depending on specific gravity [13]. The process is related to both the hydrophobicity and gravitational force of the material (Figure 5). This technique can be used to separate PET (Polyethylene Terephthalate) from other plastic packaging [14]. Furthermore, with this method, 95% of PVC (polyvinyl chloride) or PET (Polyethylene Terephthalate) can be separated [15].

![Figure 5. Process of froth flotation [14].](image)

In this method, calcium lignin sultanate and MIBC (methyl isobutyl carbinol) are used as wetting agents and frothing agents [16]. The materials are introduced in the first bin before being mixed with hot water. After that, an alkaline treatment is applied to the material in another tank, and pulp formation will take place and fed into a vibrating screen for rinsing with cold water [9]. Samples of different products may then be separated and extracted at predefined time intervals for analysis and approximation of the product weight. The inconvenience of this method is space; it requires a huge area, and the process is too long [17].

![Figure 6. Triboelectric separation technique [9].](image)
2.4. *Triboelectric Sorting*

This technique can be applied to complex mixtures. The separation happens due to the electrostatic charges of plastic mixture components (Figure 6). This technique can separate metallic parts, metallic parts from plastic parts, and different plastics based on material dielectric constant [18].

![Triboelectric Separation Technique](image)

*Figure 6. Tribo−electric separation technique [9].*

The materials are introduced on a rotating tube, and then the material presents two types of forces: the first is particle/particle forces, and the second one is particle/electrode force. When two materials stick against each other, then charge starts appearing on material particles. One gets a positive charge, and the second one gets a negative charge. Then, separation is initiated by forces acting in between them when a material particle passes through the intense electrostatic field [9].

3. Recycling Techniques: Overview

In the industry, there are several ways to recover plastics. They are classified into four categories: primary, secondary, tertiary, and quaternary recycling. Each type has its advantages and disadvantages. It depends on the application of recycled polymer. The four ways are listed in Table 2.

### Table 2. Recycling techniques.

<table>
<thead>
<tr>
<th>Type of Recycling</th>
<th>Process</th>
<th>Limits</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary recycling (Re-extrusion)</td>
<td>It is a process based on the extrusion of plastics. It does not require too much equipment and high investment.</td>
<td>It is adapted for plastic scrap with less contamination</td>
<td>[19,20]</td>
</tr>
<tr>
<td>Secondary recycling (Mechanical recycling)</td>
<td>The technique begins in the sorting center by separating, washing, and grinding plastics. After these steps, plastic materials are processed with extruders and pelletized.</td>
<td>Degradation of thermomechanical properties of plastics</td>
<td>[21–25]</td>
</tr>
<tr>
<td>Tertiary recycling (Chemical recycling)</td>
<td>Chemical recycling consists of converting polymer into molecules or monomers that can be used to manufacture new polymers.</td>
<td>It is an expensive process with a negative impact on the environment</td>
<td>[26]</td>
</tr>
<tr>
<td>Quaternary recycling (Energy recovery)</td>
<td>Quaternary recycling aims to generate energy heat or electricity from plastic scraps [20].</td>
<td>The emission of toxic gases.</td>
<td>[20]</td>
</tr>
</tbody>
</table>
3.1. Primary Recycling

Re-extrusion [19] is the process used for materials that are not too contaminated [27]. Currently, it is more suitable for post-consumption than post-industrial plastics. Therefore, this technique does not require expensive equipment, and it is easy to handle in case of recovering semi-clean plastic scrap.

3.2. Secondary Recycling

Secondary or mechanical recycling is dedicated to plastic scraps that are contaminated to reuse them to produce new products [28]. This type of recycling includes several steps, from the sorting to the pelletizing of post-consumer plastic such as Polystyrene (PS), High-Density Polyethylene (HDPE), Low-Density Polyethylene (LDPE), Poly Vinyl Chloride (PVC).

This technique faces a challenge related to the reduction of contaminants and impurities present in the resin, which affect the quality of recycled plastics [21]. The process begins with the separation, washing, and grinding of plastics in the sorting center. After these steps, plastic materials are processed and pelletized to form pellets [29]. Extrusion is one of the processes that is mostly used to manufacture polymers. In the industry, single or twin-screw extruders are both used to recycle used materials [3]. An extruder uses the rotation of the screw and heating elements to soften and mix materials. The high temperature and screw apply shear force on the polymer, which induces a scission in material crosslinking [22–24]. This chain degradation impacts the thermomechanical properties of the material. This last could be preserved by mastering the extrusion parameter and adding some additives such as carbon black and antioxidants. The machine melts the material and homogenizes it before entering the die (Figure 7) [9].

![Figure 7. Schematic of a single screw extruder [9].](image)

To reduce the contamination rate of plastics, decontamination recycling lines are often equipped with decontamination systems such as degassing system and filtration system.

3.3. Tertiary Recycling

Tertiary recycling, also known as chemical recycling, is a process that aims to convert plastics into molecules (liquids or gas) that can be used to manufacture new polymers [26]. The products issued from chemical recycling are very profitable because they provide products with minimum waste. This type of recycling gathers: Pyrolysis, gasification, liquid-gasification, viscosity breaking, and catalytic cracking [25].

3.4. Quaternary Recycling

The last type of recycling is energy recovery or quaternary recycling, which aims to generate energy, heat, or electricity from plastic scraps [20]. Plastic materials have a very high calorific value after being burned compared to some oils. The burning of PSW also generates volatile organic compounds (VOCx) and smoke. Combustion is a harmful process that can be controlled by several techniques, such as acid neutralization flue gas cooling [30]. But still, energy recovery remains the last resort to recycling plastics.
4. Sources of Contaminations

In general, ballots obtained from sorting centers are composed of three components: the desired polymer, polymeric contaminants, and some residual wastes. Contaminants can be classified into two categories volatile organic contaminants (VOCs) and solid contaminants. This last can be a polymeric contaminant or another material (metal, wood…). In the next paragraph, structural inhomogeneities and residual impurities effects will be detailed.

4.1. Structural Inhomogeneities

Compared to virgin polymers, recycled resin presents heterogeneity due to the attacking environment during its lifecycle. Irreversible structural changes can happen at both molecular and morphological levels. This modification can be induced mechano-chemically or by irradiation. The oxidation of polymeric materials creates free radicals such as carbon-centered (alkyl) and oxygen-centered (alkoxyl). This transformation is enhanced by the formation of crosslinked structures caused by the radical recombination of low molecular fragments [31].

4.2. Impurities

Impurities are present in high concentrations in post-consumed plastics. Some of them are VOCs and depend on the polymer type. They come from additives added during the polymerization, such as phenolic antioxidants, consumed during the stabilization process. Furthermore, residues of titanium and aluminum polymerization generate colored salt. Resin absorbs contaminants, and the migration of some products to the matrix of packaging influences the quality of the material after being recycled [5].

5. Decontamination Techniques

In general, decontamination is performed by a degassing system or/and filtration system that is linked to an extruder. The material is melted at a high temperature, which generates VOCs (Volatile Organic Compounds) [32]. Numerous studies confirmed the effect of extruder profile heating on the extrusion performance. The ratio of VOCs was very high when heating reached 250 °C compared to 150 °C [33].

5.1. Degassing System

5.1.1. Without Chemical Agent

To remove VOCs contained in plastics, extruders are equipped with a devolatilization system. The vacuum present inside the extruder helps to remove volatile particles. Several studies highlighted the influence of using single and multi-degassing systems in a screw extruder [34]. The concentration of the odor was measured by dynamic olfactometry. The result shows that odor intensity decreases after one degassing step from 373 to 279 OU/m$^3$ and after the third degassing step to 235 OU/m$^3$ [35].

5.1.2. With Chemical Agent

The first type of chemical agents are adsorbent agents. They allow the control of polymer emission during extrusion. The addition of 0.30% of adsorbent based on silicate to HDPE virgin pellets reduces the amount of VOCs, and the intensities of odors also decreased [36]. Furthermore, some studies show that the introduction of these particles with post-consumer HPDE in the extruder decreases odor by 50% [34]. The second type of chemical agent is a stripping agent. Their addition to the melt improves the devolatilization of the VOCs contained in the polymer. Their role consists of creating a bubble inside the matrix so the free volume in the melt increases, which helps the diffusion of VOCs in the vapor so that they can easily quit the extruder through the degassing system [36]. The most used stripping agents are water, nitrogen, and air, which enhance the devolatilization of VOCs contained in polyolefins [37]. For example, the use of nitrogen with polyethylene during extrusion reduces VOCs by 50% [38]. In the same context, some researchers devel-
opped a mathematical model that showed that the uses of stripping agents such as methanol toluene decrease the time required for degassing, and this number can increase by rising solvent concentration. [39].

5.2. Filtration System

A screen changer is an important piece of equipment in a recycling line to remove solid particles from the melt. The most used screens in the extrusion process are manual and hydraulic. The common point between them is the interruption of the melt flow while changing the screen [40]. Large-area filtration systems are available in single vessel-discontinuous or dual vessel-continuous configurations, allowing to change screens without interrupting production [40]. The different types of screens are listed in Table 3.

Table 3. Types of filtration systems.

<table>
<thead>
<tr>
<th>Filtration System</th>
<th>Advantages/Disadvantages</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual screen changers</td>
<td>It helps minimize labor and downtime for screen changes during line shutdown. The compact design eliminates the need for line disassembly and uses a hand lever to manually index the slide plate for screen change.</td>
<td>[40,41]</td>
</tr>
<tr>
<td>Hydraulic screen changer</td>
<td>It’s used in industry due to its wide range of sizes. It can reach 450 mm. Single screen operation limits filtration area, and the use of seals add to maintenance and the probability of leakage. The mesh size can reach 100 µm in some applications.</td>
<td>[40]</td>
</tr>
<tr>
<td>Rotary Disc type filtration system</td>
<td>This system offers high pressure with a lower residence time. System size can reach 250 mm and offer constant pressure operation but relatively high-pressure drops, with a lower residence time. It is a highly automated system with sophisticated controls and a backflush option. Leakage and disc lockup concerns are related to the clamping force of housing plates. It may require attention and to adapt system setup.</td>
<td>[40]</td>
</tr>
</tbody>
</table>

5.3. Supercritical Fluid

After being extruded, the polymer can be decontaminated by a purifying process such as extraction by supercritical fluid. Cristancho & Guzman [42] studied the supercritical extraction of VOCs using CO$_2$ and ethane. The process was performed with pressure in the range (7.6–20.7) MPa and two temperatures, 36 and 60 °C (Figure 8).

The result showed that using ethane was as effective as multiple extractions using CO$_2$. At a low pressure (7.6 Mpa) and medium temperature 60 °C, the extraction with CO$_2$ is effective. Higher pressure improves the extraction, but it will increase the operational cost. Both supercritical fluid help to decrease VOC concentration, but CO$_2$ remain safer and environment friendly than ethane. For this reason, CO$_2$ is the most used for the extraction of VOCs from polyethylene pellets [42].
6. Identification and Quantification of Contamination Rate

Contaminants can be classified into two categories: polymeric contaminants and volatile organic contaminants (VOCs). To identify and quantify polymeric contaminants present in the blends, Differential scanning calorimetry (DSC) and Fourier-transform infrared spectroscopy (FTIR) can be used. Concerning volatile organic compounds, chromatographic methods are the most adapted.

6.1. Differential Scanning Calorimetry (DSC)

This method is adapted to determine the fraction of polymeric contaminants based on the recording of heat exchange during heating and cooling. Nevertheless, this approach is not suitable to identify LDPE/HDPE or even LLDPE/LDPE because of the similarities between their microstructures and melting temperatures. For example, to identify the fraction of HDPE in isotactic polypropylene, we can model the blend with a known ratio of the virgin polymer. These blends are extruded and analyzed by DSC (Figure 9) [43].

Figure 8. Supercritical extraction apparatus. PS: pressure sensor, TC: temperature control, RTD: resistance temperature detector [42].

Figure 9. DSC thermograms of a 70/30 LDPE/HDPE blend (solid line), a 50/50 HDPE/i-PP blend (dashed line), and a 50/50 LDPE/i-PP (dotted line) (exo down) [43].
The presence of two melting peaks confirms the immiscibility of these polymers, which has been reported by many other authors [44].

The melting enthalpies were calculated using a linear peak integration and the results for each blend (Figure 10). This calibration curve can be used to determine the HDPE ratio in the PP/HPDE blend.

6.2. Fourier Transform Infrared Spectroscopy (FTIR)

This technique is used to quantify polymeric contaminants. Light is used to track molecular translations, rotations, and vibrations. The absorbed energy is specific for each chemical bond, and a spectrum is obtained that can be used as a fingerprint to identify polymers [43]. The spectra of the different compositions of blend based on PP and PE are shown in Figure 11.

![Figure 10. Peak area for HPDE/iPP model blends [43].](image1)

![Figure 11. FTIR-spectra of PP/HDPE blends [44].](image2)
The calibration curve was plotted based on the ratio of the absorbance (integrated area) of two peaks, 1168 cm\(^{-1}\) for methyl group in PP and a peak of 720 cm\(^{-1}\) for methylene in HDPE (Figure 12). \(A_{1168}/(A_{1168} + A_{720})\) was plotted as a function of PP content. The calibration curve can be used to determine the composition of the PP/HDPE blend.

Figure 12. Calibration curve for determination of PP content in PP/HDPE blends [44].

6.3. Chromatography Analysis

For the chemical analysis of recycled plastics, gas chromatography coupled with mass spectrometry (GC-MS) can be used to identify and quantify volatile organic compounds (VOCs). To detect organic contaminants, gas chromatography (GC) is equipped with a mass selective detector (MS). A capillary column with a film can be used for chromatographic separation. The GC oven can be programmed from 40 to 180 °C at 15 °C min\(^{-1}\) and then to 300 °C at 5 °C min\(^{-1}\), held for 12 min [45]. Organic contaminants can be identified by consulting the mass spectra libraries. The quantification of the contaminants can be performed by using external and internal calibration curves. The external standard method creates a calibration curve for a standard sample, and unknown samples are quantified using calibration curves. The internal standard method consists of adding a fixed amount of internal standard substance to an unknown sample when creating a calibration curve using a standard sample, and a calibration curve is created with the concentration ratio vs. peak area ratio for quantification [46,47].

7. Mechanical Recycling: Cases Studies

Polyolefins (i.e., PE and PP), Polyethylene Terephthalate (PET), Polyamide (PA), Poly(lactic) acid (PLA), and Polyhydroxybutyrate (PHB) are widely used in industry. These materials will be developed to highlight the different degradation mechanisms and ways to improve polymer properties.

7.1. Recycling of Commodity Polymers

Commodity polymers are used when higher properties are not required. This type of material is used for packaging, food contains, and films. The most known are Polyolefins, and their recovery is complex due to their similar densities, in particular High-density polyethylene (HDPE) and low-density polyethylene (LDPE). Furthermore, the recycling
process of this material decreases its thermomechanical properties. That’s why the incorporation of a stabilizer is needed to enhance the material’s properties.

7.1.1. Degradation of Polyolefins

The most used polyolefins are polyethylene (High-density polyethylene (HDPE), Low-density polyethylene (LDPE)) and polypropylene (PP) [48].

- **High-Density Polyethylene**: HDPE (0.952 g/cm³) is a polymer with a high degree of crystallinity, more rigid and less elastic than LDPE. During extrusion, two mechanisms can happen: either chain scission or chain branching (crosslinking) and sometimes both. Thermo-oxidation happens, and the dissolved oxygen in the melt reacts with chains, leading to chain scission with stable carbonyl. In regions with low oxygen, the chain scission produces two reactive chain ends. These macro radicals react to produce branching with higher molecular weight [49].

- **Low-density Polyethylene**: LDPE is more branched than HDPE, and its structure is susceptible to crosslinking and chain branching during extrusion. After several extrusion cycles, the complex viscosity increases (Figure 13) due to crosslinking [50].

![Figure 13. The Effect of repeated extrusion cycle on complex viscosity [51].](image)

- **Polypropylene**: PP has good optical and mechanical properties, which make it an essential material for packaging [50]. Several studies investigated the degradation of Polypropylene (PP). This material was introduced in a twin screw extruder. The screw speed was fixed at 50 rpm. The extruder had five temperature control zones. After five extrusion cycles at a high temperature (Die Zone: 270 °C), the molecular weight of PP decreases due to chain scission, and consequently, the degree of crystallinity increases. At lower temperatures (Die Zone: 240 °C), PP is stable in processing even after five extrusions. Beyond five cycles and at high temperatures, chain scission happens, and PP performance decreases [52].

7.1.2. Stabilizer Used in Polyolefins Recycling

- **Polyethylene**: To improve the thermomechanical properties of PEs and reduce thermo-oxidation, it is necessary to introduce stabilizers during extrusion. The choice of additives depends on their solubility and dispersion of the PE matrix [53,54]. Phenols and phosphate-based antioxidants are both effective in stabilizing hydrogen bonding. The use of both of them at the same time showed important results [55]. However, the combination of hindered amine light stabilizers (HALS) with phenolic antioxidants
negatively influences the stabilization of nitroxyl radicals produced from photooxidation reaction with the phenolic groups. To overcome this problem, carbon black is used as an inorganic UV stabilizer [56]. Moreover, sustainable antioxidants have been used, such as caffeic acid, curcumin, and vitamin E, to prevent polymer degradation. For instance, the addition of 12 wt % of coffee ground (CG) and 8 wt % of turmeric waste (TW) increases the elongation at the break of LDPE (Figure 14) [55]. Moreover, the addition of antioxidants combined with carbon black improves the thermomechanical properties of PEs and protects them against thermo-oxidation during extrusion [50].

![Figure 14. Elongation at break of LDPE extruded with antioxidants agro-waste [50].](image)

- **Polypropylene**: To avoid degradation of polymer chains during extrusion, PP also requires stabilization. Phenolic and hindered amine antioxidants can be used [57]. The spherulitic structure of PP allows a uniform distribution of stabilizers through the polymer matrix (Figure 15).

![Figure 15. Optical image of the spherulitic structure of PP [50].](image)
Lignin, which is a phenolic compound, can be used to stabilize PP at concentrations between 2–5 wt % (Figure 16). The lignin acts as an antioxidant and filler to increase the rigidity of the polymer matrix [58].

![Figure 16. Structure of flavonoid unit phenylpropanoid units of lignin (b) [58].](image)

7.2. Recycling of Engineering Polymer

Engineering plastics are polymers that have higher mechanical and thermal properties compared to other categories of plastics, which allow them to perform under mechanical stress at high temperatures and to resist a chemical environment. Polyethylene Terephthalate (PET) and polyamides (Pa) are the most used engineering polymers [59].

7.2.1. Polyethylene Terephthalate PET

PET is a thermoplastic that has good thermomechanical properties and processability [59]. Virgin PET has a high elongation at break values exceeding 80% and tensile strength exceeding 35 MPa. These properties decrease after several recycling cycles (Figure 17).

![Figure 17. Tensile Strength and Elongation at Break versus number of recycles for PET [50].](image)

The reduction of the properties is related to the degradation of chains due to thermo-oxidation [23]. The degradation of the chain induces the reduction of polymer molecular weights, and the presence of polymeric contaminants enhances this degradation. For instance, fewer traces of poly (vinyl alcohol) PVA or poly (lactic acid) lead to hydrolysis for
the polymer during extrusion [60,61]. A macroradical is formed and reacts with oxygen due to the screw and heat of the extruder. Radical hydrogen abstraction can produce hydroperoxide that decomposes to form two new radical species that can generate another macroradical chain that enhances thermo-oxidation [62].

Several studies show that PET can be recycled three times. Beyond this limit, the material properties decrease slowly due to the polymer size [63]. This result is related to the relationship between molecular weight and polymer degradation. Larger polymer chains enhance degradation [64]. The addition of additives during PET extrusion helps to improve the mechanical properties of recycle [65]. To reduce thermal oxidation, the use of solid-state polymerization (SSP) avoids the formation of hydroxyl and carboxyl end groups. Furthermore, chain extenders are efficient. They help repair the damage caused by chain degradation. The most common chain extenders are epoxides, carboxylic acids, and phosphates [65,66]. For instance, triphenyl phosphate (TPP) extends chains through its carboxyl and hydroxyl terminal groups [65]. The addition of pyromellitic dianhydride (PMDA) branching to PET promotes increased intrinsic viscosity ($\eta$) and a decrease in melting temperature [67]. The addition of an epoxide-based chain extender such as glycidyl methacrylate (GMA) increases the Elongation at break and Izod impact of recycled PET in reactive extrusion (Figure 18).

![Figure 18. The effect of GMA concentration on Elongation at break and Izod Impact [68].](image)

**7.2.2. Polyamides**

Polyamides (PAs), known as Nylon, are one of the most important engineering polymers. PA6s are widely used thanks to their versatility, competitive price, excellent strength and stiffness, low friction coefficient, and high dimensional stability [69].

To recycle this polymer, secondary recycling is the most widely used method, but during the extrusion process, PA6 is exposed to chemical change and degradation due to heat and shear forces. This change impacts thermomechanical properties and hence limits their use. To enhance the properties, chain extenders are used during the extrusion of recycled PA6 to increase the molecular weight of the material [70–72]. Chain extenders react faster with either amine or carboxyl end groups of polyamides and can link polymer chains to increase the molecular weight [70]. Selin Celebi O, Guralp O [73] studied the addition of several amounts of alternating copolymer of ethylene and maleic anhydride (EMA), multi-functional epoxy-based oligomeric chain extender (EPO), polyester wax with reactive caprolactam groups (CW) and dimeric 2,4-toluene diisocyanate (DTDI) in a twin screw extruder. To evaluate the effect of the chain extender on the mechanical and physical properties of rPA6, viscosity, and tensile tests were performed. The results showed that
Elongation at break increased at least by 4.4 times, even at a low rate of chain extender (0.3 wt %). Ethylene and maleic anhydride (EMA) showed higher elongation at break compared to the other investigated chain extenders. The highest elongation at break was reached with 1 wt % of EMA incorporation, which increased the elongation at break by 6.3 times (Table 4).

Table 4. Tensile properties of rPA6 before and after adding chain extended [73].

<table>
<thead>
<tr>
<th>Sample</th>
<th>Elastic Modulus (MPa)</th>
<th>Yield Strength (MPa)</th>
<th>Elongation at Break (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>rPA6</td>
<td>2788 ± 211</td>
<td>60 ± 0.7</td>
<td>7 ± 2</td>
</tr>
<tr>
<td>rPA6-EMA-0.3</td>
<td>2810 ± 65</td>
<td>63.9 ± 0.5</td>
<td>38 ± 3</td>
</tr>
<tr>
<td>rPA6-EMA-0.5</td>
<td>2908 ± 112</td>
<td>62.4 ± 0.6</td>
<td>41 ± 2</td>
</tr>
<tr>
<td>rPA6-EMA-0.7</td>
<td>2871 ± 156</td>
<td>62.2 ± 0.5</td>
<td>42 ± 1</td>
</tr>
<tr>
<td>rPA6-EMA-1</td>
<td>2850 ± 119</td>
<td>61.7 ± 0.8</td>
<td>44 ± 1</td>
</tr>
<tr>
<td>rPA6-CW-0.3</td>
<td>2718 ± 111</td>
<td>62.5 ± 0.4</td>
<td>33 ± 2</td>
</tr>
<tr>
<td>rPA6-CW-0.5</td>
<td>2727 ± 76</td>
<td>62.7 ± 0.5</td>
<td>34 ± 2</td>
</tr>
<tr>
<td>rPA6-CW-0.7</td>
<td>2563 ± 81</td>
<td>60.6 ± 0.4</td>
<td>35 ± 3</td>
</tr>
<tr>
<td>rPA6-CW-1</td>
<td>2650 ± 61</td>
<td>60.6 ± 1.3</td>
<td>36 ± 2</td>
</tr>
<tr>
<td>rPA6-EPO-0.3</td>
<td>2700 ± 89</td>
<td>63.1 ± 1.7</td>
<td>31 ± 1</td>
</tr>
<tr>
<td>rPA6-EPO-0.5</td>
<td>2784 ± 141</td>
<td>62.2 ± 0.8</td>
<td>34 ± 3</td>
</tr>
<tr>
<td>rPA6-EPO-0.7</td>
<td>2712 ± 157</td>
<td>61.4 ± 1.2</td>
<td>37 ± 2</td>
</tr>
<tr>
<td>rPA6-EPO-1</td>
<td>2648 ± 105</td>
<td>60.7 ± 0.5</td>
<td>37 ± 2</td>
</tr>
<tr>
<td>rPA6-DTDI-0.3</td>
<td>2878 ± 154</td>
<td>63.7 ± 0.4</td>
<td>33 ± 3</td>
</tr>
<tr>
<td>rPA6-DTDI0.5</td>
<td>2805 ± 106</td>
<td>64.1 ± 0.5</td>
<td>36 ± 2</td>
</tr>
<tr>
<td>rPA6-DTDI-0.7</td>
<td>2710 ± 111</td>
<td>64.2 ± 0.4</td>
<td>40 ± 3</td>
</tr>
<tr>
<td>rPA6-DTDI-1</td>
<td>2729 ± 79</td>
<td>63.4 ± 0.4</td>
<td>43 ± 1</td>
</tr>
</tbody>
</table>

Elongation at break results is coherent with relative viscosity results. The addition of EMA increased relative viscosity by 41% (Figure 19) because molecular weight is linked to elongation at break and relative viscosity. Long branches cause an increase in chain entanglement and chain straightening prior to break.
7.3. Recycling of Biodegradable Polymer

Bio-sourced plastics are manufactured from renewable resources (microorganisms, plants...). They are classed into three categories. The first type is called partially bio-based (bio-based and nonbiodegradable), such as bio-based polyolefin. The second category is bio-based biodegradable plastics such as Polylactic acids (PLA) or Polybutylene succinate (PBS). The last one is conventional plastics that are biodegradable, such as Polycaprolactone (PCL) or Polybutyrate adipate terephthalate (PBAT) [74]. The most used bio-based and bio-degradable polymers are Poly(lactic) (PLA) and Polyhydroxybutyrate (PHB).

7.3.1. Acid Poly(lactic) (PLA)

Poly(lactic) acid is a bio-sourced material that presents an alternative to PEs. This plastic is widely used in food packaging and 3D impressions. PLA is an aliphatic polyester produced by the polymerization of lactic acid, which is obtained from the fermentation of sugar [75]. However, the management of PLA waste has not yet been mastered. There are several valorization techniques of PLA, such as composting chemical and mechanical recycling. The first technique has some drawbacks related to the slow degradation of some PLA, which will lead to the accumulation of Plastic waste [76]. Chemical recycling is more expensive than using virgin bio-based plastics [77]. The mechanical is the most suitable recovery technique for PLA due to its cost and lowest environmental footprint.

The recycling process of PLA has been cited in the literature in several studies [78,79], and all show that during the extrusion of PLA, some oligomers appear due to hydrolysis, such as hydroxyl and carboxyl [80]. In general, during recycling, the molecular weight of PLA decreases due to chain scission [81,82]. Moreover, after 10 successive extrusion cycles [79], the results show that the mechanical and rheological properties of PLA decrease (Figure 20). Tensile stress and impact resistance varies slowly. At the same time, the MFR (Melt flow rate) increases significantly after the third extrusion. This fact can be explained by the increase of broken chains, which raise PLA fluidity [80].

![Figure 20. Tensile stress impact resistance and MFR after 10 cycles of extrusion [80].](image-url)
There are several ways to improve the thermomechanical and physical properties of recycled PLA. The first technique is the annealing of recycled PLA, which is a thermal treatment that consists of exposing extruded PLA to high temperatures (120 °C) for 6 h. The goal of this treatment is to increase the material crystallinity, which has been decreasing during extrusion. This treatment improves young modulus, flexure modulus, and yield but will reduce the elongation at break [83]. The upgrade of PLA can be reached by the addition of chemical agents. For instance, the incorporation of plasticizers helps to improve the processability [81].

To prevent the problem of degradation during reprocessing, the incorporation of antioxidants can minimize degradation and improve properties such as impact resistance, flowability, or UV protection [84]. Furthermore, the use of chain extenders with PLA blend improves the mechanical and rheological properties. For example, glycylid methacrylate enhances mechanical, thermal, and rheological properties due to the stabilization of molar mass by phase dispersion [85]. Also, the use of a chain extender based on phosphites increases the molar mass and complex viscosity of recycled PLA. Meng Xin [86] studied the complex viscosity of raw PLA, virgin PLA, and PLA stabilized by different Phosphite chain extender PLA-TPP (Triphenylphosphite) which is PLA stabilized by TPP chain extender, PLA-168 which stabilized by Irgafos 168 chain extender and PLA-M46TBPP is PLA stabilized by M46TBPP chain extender (Figure 21: Complex viscosity of different PLA [86]).

![Figure 21. Complex viscosity of different PLA](image)

The curves of complex viscosity show that all complex viscosity values of PLA stabilized by phosphites are higher than those of virgin PLA at every angular frequency. However, the complex viscosity values of PLA-TPP at every angular frequency are the best.

7.3.2. Polyhydroxybutyrate (PHB)

Polyhydroxybutyrate (PHB) is a member of the Polyhydroxyalkanoates (PHAs) family. They are bioderived aliphatic polyesters obtained by the polymerization of hydroxy alkanoic acids, which are produced from the fermentation of sugar and lipids [87]. Although PHB is not as widely used as PLA, they are a very interesting class of biobased and biodegradable polymers, and their production capacity is expected to triple in the next five years.
years [88]. Several studies investigated the mechanical recycling of PHB. In one of these studies, the results show that tensile strength decreased after only two extrusions, and the degree of crystallinity increased due to chain scission (Figure 22).

Since PHB is expensive and does not have good mechanical properties, it is generally blended with other polymers or used as an additive [77]. X. Yang pointed out in his research [90] that degraded PHB could be used as a plasticizer of PLA. Initially, PHB was thermally degraded in an extruder at 220°C to generate oligomers with functional groups. This last will be grafted in the main chain of PLA through a second extrusion (Figure 23).

The results of PLA with 20% grafted PHB increased the elongation at break by 66. In parallel, WAXD measurements showed that grafting significantly increased the crystallization of PLA [90].

7.4. Summary of Degradation Mechanism of Common Polymer

The degradation mechanism and ways to upgrade the polymer covered in the previous section are listed in Table 5.
Table 5. Degradation mechanism of common polymer.

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Degradation Mechanism</th>
<th>Additives</th>
<th>Main Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Commodity (Polyolefins)</td>
<td>- Chain scission</td>
<td>- Phenolic and hindered amine antioxidants</td>
<td>- The addition of 12 wt % of coffee ground (CG) and 8 wt % of turmeric waste (TW) showed elongation at the break of LDPE [55].</td>
</tr>
<tr>
<td></td>
<td>- Thermo-oxidation</td>
<td>- Agro-waste antioxidants: Coffee ground, turmeric waste</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Carbon black (CB)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Carbon black (CB)</td>
<td>- Chain extender: Epoxide, Ethylene, and anhydride maleic (EMA)</td>
<td>- During reactive extrusion of PET, The addition of 20% of glycidyl methacrylate (GMA) increases by 150% Elongation at break and Izod [68].</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>- The addition of 1 wt % of EMA during PA extrusion increases the relative viscosity of the polymer by 41% and increases Elongation at the break by 6.3 times [73].</td>
</tr>
<tr>
<td>Engineering (PET, PA)</td>
<td>- Chain scission</td>
<td>- Chain extender: Epoxide, Ethylene, and anhydride maleic (EMA)</td>
<td></td>
</tr>
<tr>
<td>Biodegradable (PLA, PHB)</td>
<td>- Chain scission</td>
<td>- Chain extender: Phosphate chain extender such as Triphenyl phosphate (TPP)</td>
<td>- The addition of a chain extender based on phosphite such as Triphenyl phosphate (TPP) increases the molar mass complex viscosity of recycled PLA.</td>
</tr>
<tr>
<td></td>
<td>- Phosphite chain extender</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

8. Conclusions

Polymer recycling can be performed using either mechanical or chemical processes. Mechanical recycling involves processes such as shredding, melting, and reprocessing plastics into new products. Unlike the chemical recycling process, mechanical recycling requires nearly no chemical solvents or byproducts, and it is, by far, the simplest and most affordable way to recycle plastics. It requires widespread, simple, well-established, less energy-consuming, and greenhouse gas emission equipment.

Mechanical recycling faces two main challenges: material degradation during reprocessing and the purity, regularity, and homogeneity of polymers to be recycled. The degradation phenomenon can be overcome by using suitable additives such as chain extenders and antioxidants, while the continuous improvement of sorting and decontamination technologies leads to recycled polymers with comparable properties as virgin polymers and makes them suitable for a wide range of applications.

In this paper, the mechanical recycling of commodity, engineering, and bio-based polymers is reviewed in detail to give researchers the state of the art of what has been done in the field of polymer mechanical recycling. The ultimate objective of this review paper is to convince readers that mechanical recycling is an easy and effective way to quickly reduce the amount of plastic waste going to landfills and incineration, environmental pollution, and resource depletion associated with producing new polymers.

Author Contributions: Literature review A.L.; original draft preparation A.L.; writing—review and editing A.L. and S.E., Final reading A.L., S.E., F.M., C.D. and M.R. All authors have read and agreed to the published version of the manuscript.

Funding: Soleno Inc. and Mitacs IT23689.
Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: Not applicable.

Acknowledgments: The authors acknowledge the Soleno and Mitacs program for financial support.

Conflicts of Interest: The authors declare no conflict of interest.

References


13. Alter, H. Application of the critical surface tension concept to items in our daily life. J. Adhes. 1978, 9, 135–140. [CrossRef]


19. Al-Salem, S.M.; Lettieri, P.; Baeyens, J. The valorization of plastic solid waste (PSW) by primary to quaternary routes: From re-use to energy and chemicals. Prog. Energy Combust. Sci. 2010, 36, 103–129. [CrossRef]


27. Al-Salem, S. Establishing an integrated databank for plastic manufacturers and converters in Kuwait. Waste Manag. 2009, 29, 479–484. [CrossRef]


**Disclaimer/Publisher’s Note:** The statements, opinions and data contained in all publications are solely those of the individual author(s) and contributor(s) and not of MDPI and/or the editor(s). MDPI and/or the editor(s) disclaim responsibility for any injury to people or property resulting from any ideas, methods, instructions or products referred to in the content.