Expanding plastics recycling technologies: chemical aspects, technology status and challenges†

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Less than 10% of the plastics generated globally are recycled, while the rest are incinerated, accumulated in landfills, or leak into the environment. New technologies are emerging to chemically recycle waste plastics that are receiving tremendous interest from academia and industry. Chemists and chemical engineers need to understand the fundamentals of these technologies to design improved systems for chemical recycling and upcycling of waste plastics. In this paper, we review the entire life cycle of plastics and options for the management of plastic waste to address barriers to industrial chemical recycling and further provide perceptions on possible opportunities with such materials. Knowledge and insights to enhance plastic recycling beyond its current scale are provided. Outstanding research problems and where researchers in the field should focus their efforts in the future are also discussed.

1. Introduction

Plastics have revolutionized our society, allowing us to inexpensively make a variety of materials that have a myriad of uses throughout industry. The first three synthetic plastics were polystyrene (PS) in 1839, polyvinyl chloride (PVC) in 1835 and Bakelite in 1907.1 Industrial plastic production really began to expand in about 1950 when global plastic production reached 2 million tons per year.2 Current global plastic production is around 380 million tons per year and it is estimated that humankind will have produced more than 30 000 million tons of plastics by the end of 2050.3 The growth in plastic production is occurring for three likely reasons:2–4 (1) new plastic materials (made by both changing the structure and processing conditions of the plastics) allow for the use of plastics in new applications, (2) the developing world is using more plastics as their economy improves, and (3) the decreasing cost of plastic production due to cheap fossil feedstocks.

Fig. 1 shows a Sankey diagram for the production, utilization and recycling of packaging plastics adapted from the Ellen McArthur Foundation.5 This shows that 32 percent of all plastics is mismanaged (littered or inadequately disposed).6

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Between two to five percent of plastics are transported into the ocean through riverways. Any plastic waste on land has the potential to eventually reach the ocean as the ocean is the ultimate sink of the earth. In the environment, plastics will slowly degrade into microplastics, which are plastic fragments less than 5 mm in diameter. Recently, microplastics were even found in the troposphere. Microplastics can also enter the environment from the degradation of plastic clothing into wastewater. Degradation mechanisms include weathering, leaching, fragmentation and potentially assimilation and mineralization. Plastic leaks into the environment through improper disposal, transport of plastics from collection sites to a central facility, and leakage from the solid waste facilities. Ocean plastics have two well-studied impacts on marine and freshwater animals: waste ingestion–egestion of plastics and entanglement in plastic waste.

Fourteen percent of plastics is landfilled. Developed countries often have open landfills where plastic can easily leak into the environment. In the US, which has a modern solid waste disposal infrastructure, it has been estimated that 0.98–1.26 million metric tons of plastic (or 2.33–2.99% of plastic waste generated) leak into the environment. Total amounts of plastics wastes are also released into the environment during natural disasters. For example, in 2011 the Tohoku earthquake and tsunami in Japan deposited an estimated 5 million tons of debris into the ocean. By 2030, 58.4 million tons of plastics will be added to the oceans each year around the world.

Fourteen percent of plastics is burned for energy recovery. While this does not create a solid waste problem, it does generate greenhouse gases. Fourteen percent of plastics are “recycled” with 8% being “cascaded recycled” to lower quality materials and only 2% of plastics used in closed-loop recycling where virgin plastics can be used. The amount of plastics recycled varies by countries, with the US recycling 10% and the EU 31% of plastics consumed. As will be discussed in this

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review, the low amount of recycled plastics is due to several reasons including: (1) plastics have a wide variety of compositions, (2) the high cost of collecting and sorting different types of plastics, especially the removal of contamination, and (3) the high capital costs and technological uncertainties of more “advanced recycling options”. Recent changes in legislation along with the desire from consumers for more sustainable products has created a dramatic push by industry for more advanced plastic recycling technologies. IHS Markit has projected that $300 billion of capital spending that is earmarked for new plastic production capacity should be redirected to plastic recycling technologies to meet circular economy goals.¹²

Fig. 2 shows the current and future infrastructure for plastic recycling technologies. The first step in plastic recycling is the sorting of plastic wastes into different streams through the material recovery facility (MRF). The MRF sorts out the glass, metals, cardboard, and plastics, bails them, and then sells them to a downstream recycler. Plastic wastes can be divided up into four areas: post-industrial waste (PIW), post-consumer waste (PCW), plastics that are present in municipal solid wastes (MSW), and ocean plastics. PIW is typically of more uniform composition and lower in contaminants than the other types of plastic wastes. Oftentimes industry will recycle PIW in closed-loop recycling processes. However, PIW often contain multi-layer films which cannot be recycled by mechanical recycling due to material incompatibility. The current plastic recycling infrastructure is primarily focused on mechanical recycling of rigid #1 and #2 plastics. MRFs do not collect flexible films or textiles as this will clog the equipment. Ocean plastic waste is not currently collected at any large scale. MSW is primarily sent to landfills although some “dirty MRFs” can process these streams as well.

As shown in Fig. 2, several technologies currently exist to chemically recycle plastics. Plastics can undergo thermal degradation to produce a liquid known as pyrolysis oil. Pyrolysis oil can be converted into aromatics and olefins by either steam cracking or using a form of catalytic upgrading as discussed in sections 5 and 6. This produces aromatics and olefins which can then be re-used to make new recycled plastics that have the same properties as the virgin plastics. Multilayer-plastics can be processed via dissolution-based recycling approaches generating pure plastic flakes which can then be re-extruded into recycled plastic resins. Polyesters and polycarbonates can be chemically or enzymatically (section 9) converted into their monomers by methanolysis and other technologies. These monomers can then be re-used to remake the polymers. Plastics can also be gasified to synthesis gas which can be used to make methanol or transportation fuels as discussed in section 7. Methanol can then be converted into aromatics and olefins.¹³ Other approaches to recycle plastics, primarily being studied in academia at present, include hydrogenolysis to produce lubricants and oils, functionalization of plastics, and the creation of plastic alloys as discussed in section 10.

Sorting, recycling, disposal, and product redesign in the management of waste plastics have been reviewed.¹⁴⁻²² The objective of this review is to first describe current plastic recycling infrastructure followed by a discussion of the chemistry, engineering, and technology of advanced plastic recycling. We first review the manufacture of plastics (section 2), landfilling of waste plastics and MSW sortation (section 3), and mechanical recycling of waste plastics (section 4). We then summarize the technologies that are being used or developed including pyrolysis (section 5), liquefaction (section 6), gasification (section 7), dissolution-based approaches (section 8), PET conversion to monomers (section 9), followed by other catalytic processes for plastics conversion (section 10). Conclusions and outlook are provided in section 11.

Victor is a doctoral student and graduate research assistant in Food Science and Technology at Iowa State University (Ames, IA, USA) and is affiliated with the Polymer and Food Protection Consortium. His major professor is Dr Keith Vorst. He received his B.S. in Chemical Engineering from the Federal University of Sao Carlos (UFSCar, Brazil) in 2019, being a visiting scholar at the University of British Columbia (UBC, Canada) for one year. His current research is focused on the sorting, processing, and characterization of post-industrial and post-consumer plastic waste, including landfill-diverted waste streams from Materials Recovery Facilities (MRFs).

Julia Curley received her PhD in inorganic chemistry from Yale University under the supervision of Prof. Nilay Hazari in September 2021. Her graduate research included synthetic, catalytic, and mechanistic studies of iron and ruthenium pincer complexes used in reactions relevant to renewable energy storage. She began her postdoc at the National Renewable Energy Laboratory in October 2021 under the guidance of Dr Gregg Beckham and Dr Robert Allen. Her current work focuses on the development of efficient and scalable systems for polyester chemical recycling.
2 Manufacture of plastics and the environmental footprint of plastics disposal

2.1 Plastic industry supply chain

2.1.1 Supply chain of plastics. Fig. 3 depicts the current supply chain for production of virgin plastics. Natural gas and naphtha (a certain boiling fraction of crude oil) are the two main feedstocks. PET (#1) is formed from ethylene glycol (EG) and terephthalic acid (TPA), while both HDPE (#2) and low-density polyethylene (#4 LDPE) are produced from ethylene. Ethylene can also be used to produce vinyl chloride monomer, which in turn is the key precursor to PVC (#3). Polypropylene (#5 PP) and PS (#6) are made from propylene and benzene with ethylene, respectively. #1–#6 plastics meet the majority of demands, and the rest is fulfilled by other plastics (#7 plastic). Typically, these resins are inexpensive and easy to process.

The chemical process of polymerization is not the same for all plastics, since the reaction is dependent on monomer chemistry and controlled by several conditions such as temperature, pressure, monomer concentration, and desired end use as well as by structure-controlling additives including catalysts, activators, accelerators, initiators and inhibitors.23 There are two basic classifications that encompass most polymerization schemes: step growth (condensation) and chain growth (addition) polymerization. Linear and non-linear step growth polymerizations are processes in which the polymerization occurs through the reaction of more than one molecular species. On the other hand, chain polymerization processes occur with monomers that have an unsaturated group. For example, polyethylene (PE) is produced from the polymerization of ethylene (or ethene, olefin) monomers through polyaddition either from the radical polymerization of ethylene monomers or by using Ziegler–Natta or metallocene catalysts. Table 1 lists the various polymerization mechanisms and some examples of their polymer products.

PP, the most widely used thermoplastic globally, is exceptional due to its production cost-effectiveness and wide applications, such as flexible barrier film pouches, caps, closures, and containers. LDPE, the second most widely used resin, is typically used to make plastic films, and injection- or blow-molded products, such as wash bottles and computer hardware covers and packaging. HDPE is typically used as fibers for industrial plastics, pipes and tubing, and consumer products, such as containers. These polyolefins account for almost 48% of global plastic demand.24

Copolymerization is used to manipulate the properties of manufactured plastics to meet specific requirements and industrial needs, allowing for the improvement of mechanical and chemical properties.25 Through manipulation of monomer chemistry, more customized materials like copolymers have been formed and successfully introduced commercially, such as acrylonitrile–butadiene–stereyene (ABS), styrene–butadiene copolymer (SBR), nitrile rubber (NR), styrene–acrylonitrile (SAN), styrene–isoprene–stereyene (SIS), and ethylene–vinyl acetate (EVA), all of which are formed through chain growth polymerization. Linear low-density polyethylene (LLDPE) is produced by polymerizing ethylene with 1-butene and smaller amounts of 1-hexene and 1-octene, leading to a product that is structurally similar to LDPE but with short, uniform branches, leading to improvements to the polymer’s mechanical and chemical properties. Step growth polymerization has been used to produce most polyamides, e.g., nylon-12 and 66 polymers, as well as the PET family. These polymers consist of at least two types of constituents (monomers, e.g.,

![Harish Radhakrishnan](Image)

Harish Radhakrishnan, born in 1996, obtained his master’s degree in Mechanical Engineering from Iowa State University and has been working on his Ph.D. dissertation for the past three years. His dissertation concerns the thermochemical and plasma-based upcycling of post-consumer plastic wastes into fuels and chemicals for downstream processes. His research activities include pyrolyzing and liquefaction of polymers, modeling polymer pyrolysis and plasma-based deconstruction chemistries, and plasma reactor design. He also performs catalytic and non-catalytic experiments with gas-solids, liquid–solid, and gas–gas plasma interactions for various applications, including natural and synthetic polymer deconstruction, as well as CO₂ conversion and NO₂ production.

![Kevin L. Sánchez-Rivera](Image)

Kevin L. Sánchez-Rivera graduated from the University of Puerto Rico-Mayagüez in 2019 and joined Prof. George Huber’s group at the University of Wisconsin–Madison in the Chemical and Biological Engineering Department. His current work focuses on developing the Solvent-Targeted Recovery and Precipitation (STRAP) technology to recycle different types of multilayer plastics, as part of the efforts of the Chemical Upcycling of Waste Plastics (CUWP) center.
dicarboxylic acids and dialcohols) and can be classified depending on the repeat unit structure.

2.1.2 **Steam cracking to produce plastic feedstocks.** As shown in Fig. 3 the first step in the production of olefins is steam cracking of either ethane (for ethylene production) or naphtha. Fig. 4 is a simplified flowsheet diagram of a steam cracking unit when naphtha is the feed, which we will describe since it can be used in chemical recycling of plastic pyrolysis oils. The feed is mixed with steam and heated by flue gas from the process in the convection zone. Before leaving the convection zone, the hydrocarbon stream is heated to 500–680 °C, which is the incipient cracking temperature range. After entering the tubular reactor of the radiant zone, the stream is heated to 750–875 °C in 0.1–0.5 s. During this short residence time, the feedstock is cracked into smaller molecules, including ethylene and propylene, and other byproducts. In a steam cracker, the reaction follows a free radical chain mechanism. For this reaction scheme, there are hundreds of radical and molecular species and thousands of reactions between them. The chain reactions of ethane as the feedstock can be found in SI 1.0 in Electronic Supplementary Information (ESI†).

When the feedstock is naphtha, a similar free radical reaction mechanism occurs. Adding steam to the steam cracker facilitates the control over residence time, prevents coke formation, and reduces hydrocarbon partial pressure. Steam does not participate in the free radical reaction mechanism. Its main role is to react with coke to form CO and CO₂. Since the cracking process is highly endothermic, energy is input from burners in the radiant zone.

The effluent stream from the cracking furnaces is cooled by a transfer line exchanger (TLE) where feed water is vaporized, and high-pressure steam is generated. Then the cracked gas is further cooled by contact with cooled oil in a quench tower. The resulting stream is sent to a primary fractionation column where fuel oil is obtained at the bottom. The stream on the top is further cooled down using a quench water tower. On the

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**Fig. 2** Summary of current management system of waste plastics.
bottom of this quench tower, a three-phase separator splits water, liquid, and gas hydrocarbons. The liquid hydrocarbons are collected as gasoline-ranged products that are heavy in aromatics. These aromatics are also used to produce other types of plastics. On the top of the quench water tower, gas effluent is obtained and mixed with gas effluent of the three-phase separator. This stream contains light hydrocarbons, including ethylene and propylene.

After compressing the cracked gas with a multi-stage compressor, the gas is sent to an acid–gas removal unit where CO₂ and H₂S are removed. After drying the purified gas with a molecular sieve dryer, the cracked gas is sent to the cold box where it is cooled down to −156 °C. Under these conditions, hydrogen is recovered for use in hydrogenation of C₃H₄. The separation of hydrocarbons starts from a demethanizer where methane is removed. This methane stream is supplied to the cracker as fuel gas for providing the reaction heat. The bottom stream of the demethanizer is sent to a deethanizer where ethane or lighter components are obtained. To convert acetylene into ethylene, this C₂ stream is hydrogenated and fractionated cryogenically. The ethylene fractionator column splits ethylene and ethane. The ethane is mixed with propane and recycled back to the cracker. On the bottom of deethanizer, C₃ and heavier components are obtained and sent to a depropanizer where C₃ and C₄/heavy fractions are separated.

After hydrogenating methylacetylene and propadiene, the C₃ stream is split into propylene and propane. The propane is recycled back to the cracker. Meanwhile, the C₄ and heavier components stream is sent to a debutanizer where butylene/butane and gasoline-ranged products are obtained. When ethane/propane are fed, no primary fractionator is needed. The gas cleaning and separation processes are very similar to the above one when feeding the cracker with ethane only. Information related to steam cracking facilities in the US can be found in SI 1.0 in ESI†.

2.2 Plastics processing

The conversion of raw or virgin plastics into finished products involves a series of plastic manufacturing processes. These manufacturing processes will vary depending on the type of plastic (thermoplastic, thermosetting, or elastomeric) and the final product desired. Plastic manufacturing can be commonly divided into three stages. The first stage is the compounding of the raw plastic with additives to achieve the desired material properties. For example, PE and PP films often stick together, which makes them difficult to separate, as when opening a PE bag. Anti-blocking agents (such as limestone and talc) can be added to lower the blocking force between the film layers. Table 2 lists the most common additives and their classifi-

<p>| Table 1 | Polymerization classification and inhibitors²³ |</p>
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<th>Classification</th>
<th>Polymerization</th>
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The compounding process is highly complex and involves taking into consideration the effect of all the additives utilized on the final properties of the plastic product. More often, a plastic will contain several additives to aid the processing of the material, modify color, and achieve the required mechanical properties. These formulations tend to be proprietary and will vary depending on the manufacturer.

The second stage, called forming, involves imparting the desired shape on the material. An inherent requirement of this step is that the material must be able to maintain this shape after the step is completed. Synthetic fibers, typically polymerized from petroleum-based chemicals, are formed through extruding a plastic solution or melt through a spinneret, a process known as spinning. This liquid-to-fiber formation process can be compared to the spinning of cotton candy. Depending on the type of spinneret used, either solid or hollow fibers can be formed. Synthetic fibers account for about half of all fiber usage, with applications in every field of fiber and textile technology. Nylon, polyester, acrylic, and polyolefin-based synthetic plastics dominate the market with these four accounting for approximately 98% of synthetic fiber production. Polyester alone accounts for around 60%. The final stage involves finishing the piece, which includes machining, decoration, and assembly. Plastics processing and manufacturing has greater opportunities for cost-savings through mini-

mization of finishing processes as compared to the manufacture of more traditional materials. Table 3 lists common methods attributed to the three stages of plastics manufacturing.

For polymeric materials, it is typical for a particular product to require multiple forming processes for ease of transfer and shipping (such as extruding a tube, then pelletizing for shipping and further use) or to carry out a specific chemical process (such as crosslinking or vulcanization). These forming processes typically involve the softened stage of the polymer, which is achieved by heating the polymer material. Each forming process will impact a thermal history and, in the case of reactive polymeric systems, will impact a degree of cure. These repeated forming processes will influence the properties of the material, either through the degradation of the material or unwanted side reactions. Setting the shape of the product is typically achieved either by cooling (which can cause crystallization, further impacting material properties) or a chemical process (crosslinking) to achieve dimensional stability. Table 4 demonstrates the most common plastics processing techniques used to form and set material shapes.

### 2.3 Mechanical and thermal properties

Most plastics produced are specifically designed to serve very particular applications. Fig. 5 demonstrates material pro-
properties and pricing of thermoplastics, aluminum, and ceramics compared to steel. During the forming of the product, it is standard for manufacturers to use additives to manipulate the properties of the raw material to achieve specific characteristics or homogenize the characteristics of their material batches as shown in Table 2. These additives influence the mechanical and thermal properties of the plastics, either by manipulating the glass transition temperature, imparting higher chemical resistance or increasing impact strength and elasticity, among other things. Fig. 5 demonstrates average mechanical properties and pricing of common commercial plastics. Many commercial plastics will have different characteristics from those of the base plastic, even though they may have the same chemistry. In general, it is important to understand that many plastics are technically combinations of several materials. As a result, mechanical and thermal properties are influenced by each component. Often, manufacturers change additives or blend ratios due to environmental concerns, to enhance specific properties, or simply for more economical processing. These subtle changes and introduction of new materials and compounding may drastically change one or more engineering properties; because of this, it is of utmost importance to continually test and evaluate commercial plastics and their specifications. Table 5 contains thermal and mechanical properties for common commercial plastics. It is of note that these properties are averages and specific material formulations can manipulate these properties for specific applications.

Temperature can have varying effects on polymer products, from melting to degradation to morphological transitions. Temperature can also affect mechanical properties due to enhanced polymer chain movement and free volume. Two key temperatures describe these effects: the glass transition temperature and melting temperature. The glass transition temperature is a property of the amorphous region of the plastic, whereas the melting point is a property of the crystalline regions. At temperatures below the glass transition temperature, the molecules in the amorphous region exist in a frozen state where movement is minimized and the polymer chains can vibrate slightly. This is typically referred to as the plastic's glassy state. The rubbery state occurs when the plastic is heated above the glass transition temperature, allowing the polymer chains to move and shift around each other, which in turn allows the plastic to become soft and flexible. Glassy (amorphous) plastics exposed to heat eventually soften and flow into a viscous liquid.

Melting transitions in some plastics refer to the transition between a crystalline solid and a liquid. When a semi-crystalline plastic is heated above the melting temperature, it forms a viscous liquid and flows. Amorphous plastics do not experience melting since there are no crystal regions to melt. For semi-crystalline plastics, crystallinity is dependent on the type and structure of the plastic. Crystallinity is typically induced by cooling a melt, although strain-induced crystallization may occur due to the molecular chains disentangling, unfolding, and straightening above its yield point. Molecular weight plays a large role in crystallinity and most polymeric properties. A plastic’s tacticity will also influence its crystallinity, with isotactic plastics demonstrating higher crystallinity than syndiotactic plastics, and atactic plastics being considered uncry stallizable due to the lack of chain regularity. Strong intermolecular
forces and a stiff chain backbone favor the formation of crystals, while bulky side groups and branching have the opposite effect, due to the difficulty in folding and aligning the chains along the crystal growth direction. Crystallinity highly influences the thermal, mechanical, and physical properties since it impacts strength, toughness, and opaqueness depending on the degree of crystallinity. These properties are further varied by the plastic’s chemistry, composition, and compounding, which in turn influence the molecular weight of the plastic. Fig. 6 demonstrates the key characteristics of various amorphous and semi-crystalline plastics based on their performance and applications.

The mechanical properties of plastics are highly varied, although there are specific relations between plastic structure...
and their properties. A plastic’s strength (tensile, compressional, flexural, torsional, and so on) depends on the molecular weight, presence of crosslinking, crystallinity, and structure. In general, a linear plastic has less strength than a branched plastic, followed by cross-linked plastics having greater strength, and network plastics exhibiting the highest strength. Low molecular weight plastics exhibit low strength, regardless of crystallinity, due to weak van der Waals forces and the ease of movement between chains, while high molecular weight plastics exhibit a greater degree of entanglement, which greatly increases the plastic’s strength. Crosslinking increases the strength of the plastic by restricting chain motion, while crystallinity increases strength due to the significant intermolecular bonding. These properties are highly influenced by environmental and application factors such as loading, temperature, humidity, weather conditions, exposure, and time.

2.4 Environmental assessment on plastic production systems

Fig. 7 presents the supply chain greenhouse gas emission (GHG) arising in the production of major types of plastic. PET has the largest carbon footprint of these plastics. The equivalent GHG emissions for producing one kg of PET film is up to 4.5 kg, where 60% of the emissions result from the usage of processing fuel and 28% of emissions are from electricity usage. General-purpose (GP) HDPE has the lowest GHG emissions among the plastics listed. The production of injected molded (IM) LDPE tends to have a high emission associated with electricity usage (1.8 kg equivalent CO₂ per kg of plastic). In comparison with the GHG emissions of electricity and processing fuel, the emission associated with transportation is relatively low, and only corresponds to 9 to 18% of the emission of the plastic supply chain.

2.5 LCA on plastic recycling technologies

In the past 17 years, 13 major life cycle assessment (LCA)/environmental assessment studies on recycling technologies and plastic waste disposal options have been conducted. These recycling/disposal options could be classified into four types: mechanical recycling, chemical recycling, incineration, and landfilling. These studies explored the environmental impact of each technology from GHG emissions, water/air pollution, human health, and resource depletion perspectives. From these publications, some general conclusions can be made:

(a) Climate change: without considering the electricity credit, incineration of plastics has the highest GHG. The GHG emission of incineration ranges from 1.8–3 kg CO₂eq per kg plastic wastes. However, accounting for the electricity credit lowers the GHGs of incineration to around 1.4 kg CO₂eq per kg plastic wastes. Meanwhile, the chemical recycling technologies (e.g., pyrolysis, gasification) also have higher GHGs than the GHGs of mechanical recycling methods, due to the direct emission and electricity consumption.
(b) Acidification: chemical recycling technologies tend to have high acidification potential. For example, gasifying one ton of plastic waste results in 0.49 kg emission of NO\textsubscript{x} and 0.09 kg emission of SO\textsubscript{x}.\textsuperscript{48} However, after accounting for credit to avoid incineration of plastic waste, the acidification potential of chemical recycling options is lower than the acidification potential of producing virgin plastic from fossil fuel.\textsuperscript{49}

(c) Air pollution: incineration contributes to air pollution. The air emission of organic compounds is ten times higher than that of the air emission of pyrolysis (1.42 g emission of organic compounds per kg of plastic processed). Mechanical recycling has the lowest air emission among plastic disposal options.

(d) Resource depletion: it is estimated that producing 1 kg of plastic requires 2 kg of crude oil, which is lost in cases where plastics are landfilled.\textsuperscript{48} Meanwhile, incineration also has high resource consumption, since the plastic is combusted for generating electricity.\textsuperscript{39} Among all the disposal options, mechanical recycling has the advantage in terms of natural resource depletion. After accounting for the credit of avoided landfill or incineration, the amount of resource consumption of mechanical recycling (crude oil/gas) is even negative.\textsuperscript{39}

Incineration causes air pollution, CO\textsubscript{2} emission, and acidification of the disposal of plastic wastes. Although landfills tend to have low air pollution, they can cause soil and groundwater pollution. The available space for landfills is limited, even if parts of these areas can be reused in some cases.\textsuperscript{49} Therefore, landfills are not a sustainable option. In comparison with chemical recycling, mechanical recycling has a lower energy consumption and a lower carbon emission. However, as will be discussed in section 4, only a portion of plastic wastes can be mechanically recycled (e.g., PET and HDPE bottles). For plastics that cannot be mechanically recycled (e.g., plastic films), chemical methods are the most environmentally friendly approach.

3. Landfills, disposal, and sorting of post-consumer waste plastics

3.1 Landfill and incineration

3.1.1 Municipal solid waste overview. MSW is the aggregate of materials discarded from households, businesses, and insti-
tutions and consists primarily of readily degradable organic materials (food wastes, green wastes such as landscaping debris) and less readily degradable solid materials with potential for recycling (e.g., paper/cardboard, plastic, glass, wood and metal). Landfills also accept a variety of other materials specific to the locale, which are referred to as “special wastes”. Wastewater treatment plant sludges, combustion wastes, and non-hazardous industrial wastes are examples of special wastes. Hazardous wastes and construction and demolition debris are not MSW and are generally managed differently. The composition of MSW varies geographically and by income level (Fig. 8), with the fraction of food and green waste decreasing as income level rises, replaced by a higher fraction of solid materials, particularly cardboard and plastics.

Plastics currently comprise 12% of all MSW generated globally. In the US, only 9% of plastics are recovered through recycling, 16% incinerated, and 76% landfilled. The EU recycles 33% of plastics, with 43% incinerated, and 25% landfilled. Estimates for the cumulative global fate of plastics indicate 79% of plastics produced have been landfilled or left in the open environment, whereas 12% have been incinerated, and 9% recovered for recycling.

The UN Environment Programme and US EPA have established waste hierarchies focused on shifting from a philosophy of waste management to a philosophy of resource management in the context of a circular economy. The recommended hierarchy (highest to lowest priority) consists of waste prevention, waste minimization, reuse, recycling, energy recovery, and disposal (Fig. 8). Most waste today, in the US and globally, is managed by disposal in landfills – the lowest priority in the hierarchy.

3.1.2 Landfills. Landfills and dumps are the largest disposition pathway for MSW (Fig. 8), primarily because this pathway is the least expensive and least cumbersome. Landfills and dumps have the common attribute of being a storage facility near the ground surface where waste is placed. However, a dump is an un-engineered facility with limited or no control on contaminated liquids or gases generated by the waste. Effect, waste is “dumped” and no longer managed. In contrast, landfills are engineered systems where liquids and gases are carefully managed so that impact on the environment remains within accepted norms. Dumps are more common in lower-income nations that lack the infrastructure and regulations for MSW management, although transitioning open dumps to engineered landfills has become a priority for some developing nations. The remainder of this section focuses on landfills.

A modern engineered landfill consists of an engineered containment system situated in an earthen depression (Fig. 9). The base is comprised of a multicomponent liner overlain by a leachate collection system. Waste is placed in layers approximately 3-4 m thick (called “lifts”) that are compacted with heavy machinery. A gas collection system is installed in the waste to collect landfill gas (LFG) for treatment or production of renewable energy. Interim cover (soil or synthetic material) is placed over the waste during filling to reduce ingress of precipitation, control emissions and odors, and limit vectors. Once final elevations are met, a final cover is placed to limit
the ingress of precipitation to *de minimis* amounts, facilitate gas collection, and provide separation from the surrounding environment for as long as the waste poses a threat. As with the liner, the final cover includes a multilayer hydraulic barrier overlain by a drainage layer. The surface is covered with vegetated cover soils for protection and aesthetics.\(^{10}\)

Leachate is generated when water percolating through the landfill contacts the waste.\(^{59}\) Leachate contains a variety of inorganic and organic contaminants, and therefore is treated before being discharged. Leachates are pumped from the leachate collection system, and generally are transported to a public wastewater treatment plant (a publicly owned treatment works, or POTW) where they are treated along with other wastewater (*e.g.*, sewage from a community, *etc*.). In some cases, leachates are treated on site or are recirculated in the waste to stimulate waste decomposition to promote more

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**Fig. 8** Municipal solid waste (MSW) generation and final disposition: (A) global generation in 2016 with fractions associated with national income, (B) US generation in 2018, and (C) waste management hierarchy. Inserted triangle illustrating how disposition pathways fit with preferred waste management hierarchy.

**Fig. 9** MSW landfill with liner, cover, leachate collection, leachate recirculation, and gas collection.
rapid waste stabilization and greater gas production for renewable energy.\textsuperscript{55,56}

Landfills operate under anaerobic conditions where the microbial community decomposes the degradable organic fraction and generates gas comprised of approximately equal parts methane (CH\textsubscript{4}) and carbon dioxide (CO\textsubscript{2}).\textsuperscript{56} MSW landfills are the third largest source of anthropogenic CH\textsubscript{4} in the US\textsuperscript{57} and contribute approximately 3\% of global GHG emissions.\textsuperscript{60} LFG also contains a variety of other contaminants in small quantities. Consequently, LFG must be treated before being released. Thermal treatment is most common, either through a flare or through combustion to create renewable energy (e.g., LFG to electricity). In recent years, landfills have also become a significant source of renewable natural gas (RNG), which is produced by treating and compressing LFG extracted from the landfill.\textsuperscript{56}

### 3.1.3 Incineration with energy recovery

Waste-to-energy (WtE) is the process of converting MSW into energy in the form of electricity, heat, or alternative fuels.\textsuperscript{61} Within a circular economy framework, WtE technologies have a crucial role in providing an alternative disposal option for MSW that cannot be reused or recycled.\textsuperscript{61} The technical, economic, and environmental aspects of common WtE have been evaluated extensively.\textsuperscript{62–66} Astrup et al.\textsuperscript{67} present a comprehensive review and comparison of WtE technologies. LCA methods were used to evaluate the environmental impacts of different technologies employed in WtE (e.g., plant type, energy recovery, type of energy output, flue gas cleaning, and residue types and management). Incineration is by far the most common technology for WtE, representing more than 88\% of the global WtE market\textsuperscript{68} with a valuation of $30 billion predicted by 2027.\textsuperscript{69}

The remainder of this section focuses on incineration.

Incineration is the combustion of MSW at temperatures at around 750–1100 °C, with the heat being used to produce steam for generating electricity and/or heating.\textsuperscript{70,71} Plants producing steam for heating and electricity can have efficiencies on the order of 80\%; whereas the efficiency is on the order of 20\% for plants producing only electricity.\textsuperscript{71} Most MSW incineration is conducted as mass burn (bulk MSW incinerated in a single chamber) or as refuse derived fuel (RDF) where screened and shredded MSW is combusted.\textsuperscript{72} RDF is common in the Portland cement industry and can represent up to 80\% of the thermal requirements in cement plants.\textsuperscript{61} Nearly 15\% of all MSW is incinerated globally. Incineration rates vary significantly, with incineration more common in developed land-constrained countries and islands,\textsuperscript{70} less common in developed countries without land constraints, and uncommon (<1\%) in lower-middle and low-income countries. For example, Japan has significant land constraints and incinerates 74\% of its MSW,\textsuperscript{73} whereas the US has virtually no land constraints and incinerates only 12\% of MSW. The effect of income is evident in the location of incinerators; 80\% of MSW incinerators globally are located in North America, Europe, and Asia Pacific.\textsuperscript{70} US incinerators manage 22\% of discarded food and 16\% of discarded plastics.\textsuperscript{74} Most WtE incinerators in the US (77\%) only generate electricity, 19\% produce electricity and heat, and 4\% export steam to local users.\textsuperscript{75}

Incineration reduces the mass and volume of MSW that is landfilled (70–85\% by mass and 75–90\% by volume)\textsuperscript{71} while generating energy for use or sale. Despite these advantages, incinerators are difficult to implement, especially in the US. Emissions from MSW incinerators can contain dioxins and furans, which are toxic persistent organic pollutants and human carcinogens.\textsuperscript{76,77} These pollutants are often of great concern to surrounding communities even when plants are in compliance with emissions criteria in the Clean Air Act. MSW incineration also generates ashes that need to be managed,\textsuperscript{73} generally through landfill disposal. Incinerator ashes tend to have higher levels of contaminants per unit mass than MSW, resulting in leachate with higher concentrations that requires more treatment. In some cases, ashes can be reactive, leading to heat generation that is problematic in landfills.\textsuperscript{78}

### 3.2 Material recovery facilities (MRFs)

MRFs (pronounced “murfds”) receive waste material and separate (or bail) these materials. Their origin can be explained when terms such as MSW and waste management began to be established in the 1950–60s, with the increased waste generation linked to economic growth and popularization of petrochemical derivatives. In the US, the first MRF was established in Groton, Connecticut in the early 1980s,\textsuperscript{79} with the number of facilities multiplying in the following years. There were 40 MRFs in operation or under planning in 1991, 166 in 1993, and 307 in 1995.\textsuperscript{80}

MRFs are classified based on the input waste material they receive: single-stream, dual-stream, mixed-waste, or pre-sorted, with the first being a typical design in the US.\textsuperscript{81} Single and mixed-waste MRFs (containing more organic contaminants from MSW) have similar inbound composition, receiving a single stream with paper, plastic, glass, and metals, usually from automatic or semi-automatic vehicles. Dual-stream MRFs have a two-category stream line: fibers, composed of paper and cardboard, and containers, composed of plastic, metal, and glass, whereas pre-sorted MRFs receive source-separated material streams.\textsuperscript{42,81} Fig. 10 is a block flow diagram for single-, mixed-, and dual-stream MRF frameworks (differences between operation modes are shown; otherwise, the process is the same for all) based on the work of Pressley \textit{et al.}\textsuperscript{42} who drew on common MRF designs around the US. The design of a MRF is unique to each location, with different specifications required and associated with inbound composition, investment capacity, and ownership and operation type.

Single and dual-stream frameworks have their advantages and disadvantages. Single-stream benefits from waste collection with larger volumes and lower costs but with increased difficulty and costs for processing at the MRF, in addition to an increased contamination rate.\textsuperscript{82} For example, during conversion from dual to single-stream, a MRF from Brookhaven (New York, US) observed a 25\% increase in volume of input recycling streams, which led to a diversion of recyclable material from landfills from 50 000 to 35 000 tons, along with an increase in non-recyclables in the single-stream.\textsuperscript{83} Another study that investigated GHG emissions observed benefits from
the conversion of three medium-size MRFs from dual-stream to single-stream systems, avoiding the emission of 711 kg-CO₂-equivalent per metric ton. The advantages of switching from a dual to single stream were not observed on the economic side, as observed in a study conducted using data from 223 municipalities in the province of Ontario, Canada. For example, it was found that single-stream recycling incurs 48.7% higher processing costs and 9.6% lower values of a recyclable saleable stream compared to multi-stream recycling systems, with only a 3% reduction in the collection costs.

Specifications were determined in the US in an effort to standardize the inbound recycling streams at MRFs, such as those stipulated by the Institute of Scrap Recycling Industries (ISRI). Table 6 describes the materials accepted, prohibited, and the possible contaminants in the majority of US MRFs. It should be noted that plastic bags and films are considered contamination due to the possibility of issues during sorting and are removed early in the process. A pilot study conducted by Resource Recycling Systems (RRF) for the Materials Recovery for the Future (MRFF) on an existing MRF in Birdsboro (Pennsylvania, US) looked at how to modify an MRF design to accept flexible packaging from residential single-stream inbound waste. It demanded additional equipment for separation, with a processing cost of $2.41 per ton and a break-even point compared with the sorted polypropylene market value.

3.2.1 Equipment overview. Over the years, several new technologies have been applied to the equipment used in a MRF to increase sorting efficiency, maximize the recovery of marketable materials, and deal with the increased complexity and diversity of inbound waste streams. As a connector for the whole processing facility, conveyor belts transfer material between different equipment, sort stations, and storage areas, including infeed conveyors that receive the material from the tipping floor up to the conveyors carrying the final bales of material for shipping.

Two types of conveyors are usually used: rubber and chain belt conveyors. Rubber belts are divided into two categories: slider beds, where the belt moves through a flat or shaped trough and is ideal for sorting with the material more spread-out, and idler conveyors, which have a belt over idler rollers perpendicular to the material movement, better suited for fine or gritty material. Chain belt conveyors are often employed when there is a need to move material through increased...
angles, such as feeding other equipment or weighty loads due to chain-link drive systems being more robust than rubber conveyors. The top speed for these conveyors is considerably slower.88

After entering the MRF environment, the waste stream passes through multiple separation equipment and operations. The simplest method is known as manual sorting, a labor-intensive step in the MRF operation and a source of potential health and occupational hazards,89 including physical, chemical, and biological concerns. The sorting can be divided into a negative sort when undesired materials are removed from the waste stream and a positive sort that looks for recyclables and placement into the correct bin.90 For the separation involving multiple material sizes into a specific range, different types of screens are commonly employed, such as vibratory, disc, trommel, ballistic, and, more recently, auger screens, as described in Table 7.

Air separation exploiting different densities of materials in mixed waste streams is another option at MRFs. A heavy fraction composed of metal, glass, and stones falls to the bottom of the equipment while a lighter fraction of paper, plastic, and dry organics is carried away by the air stream, requiring the use of a cyclone, for example, for the recycling of the air stream.90 Equipment in this category includes zig-zag classifiers, air-drum or rotary air, aspirator air or suction hood, cross-current air, and airbed classifiers.88,90

Another crucial piece of equipment typically used for material sorting is the optical sorter. The use of this type of equipment can substitute for manual labor and lead to high sortation efficiencies; however, a significant capital cost is required for its installation, which hinders small and locally owned MRFs from selling bales in a market with low tolerance to contamination. An estimate using data from the United Kingdom by Eule showed that a MRF with a single operation shift could take up to 3.14 more years to break even after installing an optical sorter compared to manual labor.92

Optical sorters use color-sensitive cameras and near-infrared (NIR) and ultraviolet (UV) spectroscopy sensors, or a combination of them, to identify different materials that pass through a conveyor belt. Once the material desired to be positively sorted is identified, air jets are activated to eject the piece to a different chute.90,92 Spectrometric techniques exploit the unique wavelength signature of different kinds of materials.88 NIR optical sorters are most commonly used for plastic separation as a non-destructive technique with high-speed measurement, penetration depth and signal-to-noise ratio.91 NIR optical sorter use has been growing for sorting fibers and cartons.93 Another advance in sorting technology is the use of hyperspectral imaging (HSI), which combines imaging from a digital camera with spectrometric analysis, to obtain a discrete spectrum for every pixel collected. This is then computationally analyzed using algorithms such as principal component analysis94,95 for the sorting of more complex plastic streams containing low and high density polyethylene, for example.96 Machine learning algorithms are being studied to overcome some of the challenges associated with NIR sorters,97 such as the difficulty of correctly sorting black plastics98 or avoiding the still high costs of HSI technology.99
Table 7 Common types of separation screens in MRFs. Reprinted from ref. 88 and 91, with permission from American Chemistry Council, CP Group Recycling Solutions, copyright 2016

<table>
<thead>
<tr>
<th>Types</th>
<th>Description</th>
<th>Illustration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vibratory (shaking)</td>
<td>The screen can be horizontal or inclined, with the waste stream cascading through a series of finger elements, perforated plates, or meshes, promoting material separation. Presents better performance when operating with dry and cleaner waste, as it can be easily clogged.</td>
<td><img src="image1" alt="Illustration" /></td>
</tr>
<tr>
<td>Disc</td>
<td>Rotating discs are positioned in shafts, moving the waste stream throughout the screen in horizontal or angled positions. The distance between discs (interface opening) is associated with the screened material size. Disc rolling might help by partially breaking the material on the screen. A specialized version of a disc screen that separates 2D (paper or film) from 3D materials (containers) can be called a polishing or planar screen.</td>
<td><img src="image2" alt="Illustration" /></td>
</tr>
<tr>
<td>Trommel (revolving)</td>
<td>Cylinder-mounted rollers in an inclined position contain holes in the sides and might have baffles. They are used for primary or final size screening. Factors that affect the input and separation efficiency include screen openings, trommel diameter, rotational speed, type and number of baffles, and cylinder inclination. Tumbling motion can be cascading, cataracting, or centrifuging.</td>
<td><img src="image3" alt="Illustration" /></td>
</tr>
<tr>
<td>Ballistic</td>
<td>Combines parallel paddles that vibrate (similar to a vibratory screen) with a 2D/3D screen framework. Three flows are collected: a fines stream with material that passed through the paddles, a 2D stream with paper, cardboard, and films, and a 3D stream mainly consisting of containers.</td>
<td><img src="image4" alt="Illustration" /></td>
</tr>
<tr>
<td>Auger</td>
<td>One of the newest types of equipment launched to the market consists of multiple cantilevered augers in corkscrew motion, which prevents the wrap of plastic films and hoses. Mainly employed as a pre-sort step for removing large flat fractions such as (old corrugated containers) OCC from smaller objects.</td>
<td><img src="image5" alt="Illustration" /></td>
</tr>
</tbody>
</table>

For the sortation of ferrous materials, magnetic separators are used to sort materials such as steel from the remaining waste stream. Multiple factors are considered to determine the system’s efficiency, including the distance between the conveyor carrying the waste stream and the magnet, the magnetic force employed, the conveyor speed, and the burden depth on the conveyor belt. Common types of setup include a drum holding magnet, a belt holding magnet, and a suspended type magnetic separator. For non-ferrous materials, such as aluminum and copper, eddy current separators are used, often after the magnetic separator. The sorting is based on the eddy currents formed when conductive objects are close to a magnetic field (created using rotating magnets), with another magnetic field of opposite direction. According to Lenz’s law, this creates a force that expels these objects from the initial magnetic field, giving them a different trajectory. Considering that less dense and conductive objects demand a lesser force, aluminum has the greatest potential for separation through this equipment since it has one of the highest conductivity/density ratios compared to other metals, such as lead, copper, zinc, and tin.

At the end of the sorting process, each material stream is compacted or densified to reduce its volume to minimize storage space requirements and increase payload for recyclable material and residue transportation, significantly reducing costs. Balers are commonly used to reduce the bulk density of the material and can be divided into two different types: vertical balers, an option with reduced output but low costs, or horizontal balers, which are the most utilized in MRFs. Horizontal balers are usually single-ram or dual-ram (two-ram), with the first mainly used for fiber, paper, and OCC output lines, while the latter is considered a multipurpose equipment utilized for plastic and aluminum outputs due to its capacity to produce more heavy and uniformly packed bales. Single-ram balers operate with the sorted material being pushed from a collection chamber into a narrow hopper, with the densified material moving continuously as more material is fed and tied with wires. On the other side, a dual-ram baler (Fig. 11) has the main ram that loads the material into a chamber until the desired amount is achieved, while a secondary ram is responsible for moving the bale to a tying process.

The Association of Plastic Recyclers (APR) has developed some specifications to standardize plastic bales and facilitate the sales between MRFs and recycled plastic reprocessors, brokers, or companies that can use them in their process. For rigid plastic containers bales, including natural and colored HDPE, PP, and #1-7 plastics, it is expected to have a bale density between 15 and 20 lb ft⁻³ (240 and 320 kg m⁻³), whereas, for a PET bottle bale, the targeted bale density is between 15 and 18 lb ft⁻³ (240 and 288 kg m⁻³), all using non-corrosive galvanized bale wires of 10–12 gauge (2.6–3.2 mm). The main products of a MRF are the bales of sorted material with the ultimate goal of producing saleable recycled material and the least amount of non-recyclable residues. In the optimal scenario, these residues can be used as...
fuel for waste-to-energy conversion, in anaerobic digestion or, in the worst case, be an expense for discard in a landfill. For plastic bales, the ISRI Scrap Specifications Circular defines standards for some of the most common types of bales from a MRF in terms of composition, contaminants, and prohibited items, as shown in Tables 8 and 9. The historical prices for some plastic bales are listed in Table 10.

### 3.2.2 MRFs in Europe vs. the US (typical size, numbers of MRFs in US)

According to the Environmental Protection Agency (EPA) there were 532 MRFs in the US in 2018 with an estimated daily throughput of 91 129 tons. The Recycling Partnership conducted another survey with data from Resource Recycling, initially finding 360 MRFs in the United States in 2019, a number that was updated in February 2020 to 367 after a verification conducted by The Last Beach Cleanup (a nonprofit 501(c)3 organization), with a spreadsheet available on their website and results published by Greenpeace. The distribution of MRFs across the U.S. is shown in Table 11. A list of the largest 75 MRFs in North America was published in 2019 by Recycling Today, with 65 located in the United States and 10 in Canada. The largest MRF was Sims Municipal Recycling’s Sunset Park, located in New York, which shipped 241 884 tons in 2018. In England, data from the last quarter of 2019 totaled 85 regulated material facilities (>1000 tonnes per year recyclables processed) with a total of 21 080 860 tonnes processed in 2017.

### Table 8 Outbound mixed plastic bale specifications (ISRI)

<table>
<thead>
<tr>
<th>Bale type</th>
<th>Material description</th>
<th>Main product</th>
<th>Allowed contamination</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1–7 bottles and small rigid plastic</td>
<td>Rigid plastics, consisting of at least 65% bottles, bulky rigid containers greater than 5 gallons should be avoided</td>
<td>Bottle and non-bottle containers</td>
<td>5 wt% total, with less than 2 wt% paper/cardboard and 1 wt% of metal/plastic bags/liquid/other residues</td>
</tr>
<tr>
<td>#3–7 bottles and small rigid plastic</td>
<td>Rigid plastics without PET (#1) and (#2), bulky rigid containers greater than 5 gallons should be avoided</td>
<td>Bottle and non-bottle containers</td>
<td>5 wt% total, with less than 2 wt% paper/cardboard/ metal and 1 wt% of liquid/other residues</td>
</tr>
<tr>
<td>Mixed bulky rigid (#2 and #3)</td>
<td>Large rigid plastics HDPE (#2) and PP (#5)</td>
<td>Bulky rigid plastic</td>
<td>15 wt% total, with less than 4 wt% for plastic/packaging items (#1, #3, #6, #7), 2 wt% of metals, glass, wood, liquid/other residues, paper/cardboard, plastic bags/sheets/films</td>
</tr>
<tr>
<td>Tubes and lids (#2, #4, and #5)</td>
<td>Any whole container of HDPE (#2), LDPE (#4), and PP (#5)</td>
<td>Tubes and lids</td>
<td>10 wt% total, with less than 2 wt% of metals, liquid/ other residues, paper/cardboard, injection-molded HDPE (#2), PET (#1) bottles and thermoforms, plastic/packaging items (#1, #3, #6, and #7)</td>
</tr>
</tbody>
</table>

### Table 9 Outbound PET and HDPE plastic bale specifications by grade (ISRI)

<table>
<thead>
<tr>
<th>Bale grade</th>
<th>Grade A</th>
<th>Grade B</th>
<th>Grade C</th>
<th>Grade F</th>
</tr>
</thead>
<tbody>
<tr>
<td>PET (#1)</td>
<td>&gt;94</td>
<td>93–83</td>
<td>82–73</td>
<td>&lt;72</td>
</tr>
<tr>
<td>Max. contamination (wt%)</td>
<td>6</td>
<td>7–17</td>
<td>18–27</td>
<td>&gt;28</td>
</tr>
<tr>
<td>HDPE (#2)</td>
<td>&gt;95</td>
<td>94–85</td>
<td>84–80</td>
<td>&lt;79</td>
</tr>
<tr>
<td>Max. contamination (wt%)</td>
<td>5</td>
<td>6–15</td>
<td>16–20</td>
<td>21</td>
</tr>
</tbody>
</table>

The distribution of MRFs across the U.S. is shown in Table 11. A list of the largest 75 MRFs in North America was published in 2019 by Recycling Today, with 65 located in the United States and 10 in Canada. The largest MRF was Sims Municipal Recycling’s Sunset Park, located in New York, which shipped 241 884 tons in 2018. In England, data from the last quarter of 2019 totaled 85 regulated material facilities (>1000 tonnes per year recyclables processed) with a total of 21 080 860 tonnes processed in 2017.
di selezione” - CSS) in 2020 associated with COREPLA (National Consortium for Collection and Recycling of Plastic Packaging) that receive a pre-sorted waste stream mainly composed of mixed plastics from 996 smaller and local sorting centers (“centri comprensoriali”).\(^{112,113}\) Germany’s federal statistics reported 987 waste sorting facilities in 2019, with a total input of 25 120 700 tonnes.\(^{114}\) In France, it was reported that 399 waste sorting facilities (“centres de tri de déchets”) were operating in 2016, receiving 11.1 million tons.\(^{115}\)

### 3.2.3 Waste pickers in Latin America

In less economically developed countries, recycled material is typically manually removed from MSW by human “waste pickers”. A key issue in MSW management and/or waste recycling in developing countries is the role of waste pickers, who collect, haul, sort, and sell recyclable commodities from MSW. There are around 20 million people worldwide informally dedicated to waste recycling, which represents approximately 50% of the workers involved in waste management. In developing countries, this informal recycling sector recovers the largest fractions of recycled waste and in some cases, it is the only organized form of recycling.\(^{116-118}\) In Latin American, waste pickers are recruited among vulnerable groups living under conditions of extreme poverty in communities that lack the educational, social, and psychological resources to prepare its citizens for the deficiency in MSW management systems in urban and rural areas, combined with a lack of recycling programs caused environmental and health problems, and the clogging of drainage systems in urban areas. Since the mid-1990s, the Mexican government has set voluntary PET recovery and recycling programs that were successful to some extent.\(^{121}\) PET recycling activities involve sorting, washing, re-granulating and pellet production by local enterprises. The pellets are then shipped to domestic Mexican plastic transformation industry (35%) or exported to other markets (65%). In Mexico, recycled PET facilities produce textile fibers, bottles for non-food uses and industrial strapping.

In Mexico, recycling of the MSW is mostly in the hands of waste pickers, middlemen or brokers.\(^{122,123}\) Waste pickers are self-employed, sometimes resorting to collecting plastics from neighborhood garbage containers before they are collected by contracted waste management services. Waste pickers often organize themselves into associations, cooperatives, unions or microenterprises, through which they feel empowered to carry out their activities.\(^{116,121}\) Often local authorities promote and recognize the formation of associations or cooperatives, which are able to negotiate agreements for waste collection and recycling services. However, even with these associations the waste pickers income is not fixed but based on the quantity and quality of recycled materials and the selling price in local markets.

Currently, recycled PET price depends on factors such as impurities (i.e., amount of other plastics or residues), quality (i.e., clear vs. color PET), cleanliness, type of organization the waste pickers belong to, middleman or broker who buys the product, and even the geographical area (i.e., urban vs. rural) where PET is collected and sold. An average price for recycled PET ranges between $0.24–0.48 per kg,\(^{105}\) although it is likely that most waste pickers receive the lowest price. Waste pickers who recycle PET must work long hours to obtain sufficient income to support their families. The official minimum daily wage set by the Mexican federal government was $8.25 per day in January 2022. If an average recycled PET price of $0.36 per kg is considered, a waste picker must collect around 23 kg of PET every day to receive the equivalent of the minimum daily wage. A number of studies highlight the low monthly income due to the high consumption of soft drinks and bottled water.\(^{120,121}\) The high consumption of PET bottles, many of them discarded in public places, roads and transport systems and rural areas, combined with a lack of recycling programs caused environmental and health problems, and the clogging of drainage systems in urban areas. Since the mid-1990s, the Mexican government has set voluntary PET recovery and recycling programs that were successful to some extent.\(^{121}\)

### Table 10 Historical prices for some plastic bales\(^{105}\)

<table>
<thead>
<tr>
<th>Materials</th>
<th>National average price (cents lb(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2017</td>
</tr>
<tr>
<td>Plastics PET (baled)</td>
<td>11–16</td>
</tr>
<tr>
<td>Plastics film (grade A, sorted, 800 + lb bales)</td>
<td>8–11</td>
</tr>
<tr>
<td>Plastics colored HDPE (baled)</td>
<td>13–17</td>
</tr>
<tr>
<td>Plastics commingled (#1–7, baled)</td>
<td>2–3</td>
</tr>
<tr>
<td>Plastics PP post consumer (baled)</td>
<td>7–9</td>
</tr>
<tr>
<td>Plastics polystyrene EPS (baled)</td>
<td>2</td>
</tr>
</tbody>
</table>

### Table 11 Number of MRFs in the United States\(^{106,107}\)

<table>
<thead>
<tr>
<th>Region</th>
<th>Number of MRFs</th>
<th>EPA (2018)</th>
<th>The last beach cleanup/Greenpeace (2020)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Northeast</td>
<td>128</td>
<td>66</td>
<td></td>
</tr>
<tr>
<td>South</td>
<td>142</td>
<td>107</td>
<td></td>
</tr>
<tr>
<td>Midwest</td>
<td>139</td>
<td>76</td>
<td></td>
</tr>
<tr>
<td>West</td>
<td>123</td>
<td>118</td>
<td></td>
</tr>
<tr>
<td>Total (U. S.)</td>
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There is an urgent need to propose business schemes that improve the income of waste pickers, in addition to providing them health, safety, labor and social benefits through formal employment. The development of business models in Latin America that formalize the inclusion of waste pickers in the recycling value chain, and that limit the actions and profits of middlemen, will have social benefits for waste pickers and will bring favorable economic benefits for the Mexican recycled PET industry.

Some successful initiatives show that when waste pickers are organized by non-profit and for-profit organizations and they are formally included in the PET recycling value chain, a number of favorable social and economic benefits occurs.\textsuperscript{118} Social-oriented programs for waste pickers lead to their recognition and empowerment through the establishment of formal jobs, with fixed salaries and labor and health benefits with local government or private companies. Social-oriented programs may have a profound impact on the education and culture of waste picker’s communities. These social-based initiatives are emerging in the MSW management value chain of Mexico, Brazil, Argentina, Colombia and Chile, but the benefits are still limited to a few communities. In the current socio-economic situation in Latin America, the development of technologies and industries that facilitate the processing of plastic waste into products with higher added value in the market, is a critical factor to facilitate the inclusion of waste pickers in the value chain of plastic recycling.

MSW from urban and rural areas is typically processed by public services run by local governments to avoid negative impacts on health and the environment. In countries such as Mexico and Brazil, the management of MSW is regulated by federal and local laws, which establish guidelines for the collection, separation, recycling and final disposal of waste, using strategies that, in principle, contribute to the preservation of the environment and take care of the health and safety of the employees.\textsuperscript{118,119} Various technological, economic, administrative, social and cultural factors prevent risk and safety management from being carried out properly, especially in rural areas. As a result, it is estimated that around 40 million people (6\% of population) in Latin American communities lack trash service. Up to a third of MSW is disposed in open dumps.\textsuperscript{125–128} Consequently, the prevailing situation is that the MSW management in Latin America is a focus of concern in terms of public health, environmental risks, informal economy, security and social inequality, particularly for the communities where the MSW transfer units and final disposal sites are located.

A license is granted to private companies when MSW cannot be adequately collected by local governments. A growing number of companies in Latin America offer the following benefits in MSW management: a reduction in public spending, selective processing of domestic, commercial, and industrial waste, use of specialized technology and skilled workforce, generation of well-paying jobs, and management projects contributing to sustainable development.\textsuperscript{129,130} This recycling strategy is very distinctive from the circular economy and it has been identified as a vector contributing to the fulfillment of Sustainable Development Objectives set by the United Nations. For this reason, federal and local governments in Latin American communities have established legal frameworks to trigger waste recycling, as in the case of Brazil, which in 2010 established the National Solid Waste Policy, and Mexico, which in 2003 approved the Law for the Prevention and Integrated Management of Waste, and very recently the General Law of Circular Economy with the aim of promoting efficiency in the use of waste through reuse and recycling.\textsuperscript{119,131–133}

Poverty and the growing market value of recyclable products has induced many people in conditions of social and economic vulnerability to work as waste pickers. It is very common nowadays to find fixed human settlements around waste transfer and final disposal units. Entire families live in these settlements. It is common for waste pickers’ children to continue with the family tradition. The situation of waste pickers is similar in other countries that do not have a modern waste management system.\textsuperscript{134} The development of advanced waste management systems combined with new plastic recycling technologies will drastically improve the quality of life for waste pickers.

4. Mechanical recycling

Mechanical recycling is defined by the U.S. Food and Drug Administration (FDA) as secondary recycling and is the most common approach for recycling plastic solid waste (PSW). This process has an established infrastructure in most countries. In secondary recycling, the post-consumer plastic waste is reprocessed and converted into new materials.\textsuperscript{135} With the advancement and availability of mechanically recycled PSW, virgin plastics are substituted with post-consumer recycled materials or blends thereof in many industrial and consumer applications.\textsuperscript{136,137} Mechanical recycling involves collection/ segregation, cleaning and drying, chippering/sizing, coloring/ agglomeration, pelletization/extrusion, and manufacturing the end product. Both primary and secondary recycling involve similar process steps.\textsuperscript{138} Primary recycling as defined by the FDA recovers pre-consumer or post-industrial recycled (PIR) waste material and produces new products. Secondary recycling physically reprocesses post-consumer waste materials to produce new materials.\textsuperscript{139} Primary recycling maintains the same quality as virgin material, whereas downcycling is common in secondary recycling due to some degradation of the plastic’s properties.\textsuperscript{136} Post-consumer materials, in particular polyolefins, can have unpredictable physical properties due to the large range of polymer structures, molecular weights, and additive packages. Such variation and lack of grading can make using these materials more difficult. The variation in color further complicates usage in new products due to the inability to control the final colors. For example, plastic
4.1 Processing technologies in mechanical recycling

Mechanical recycling consists of a few general steps: shredding/granulation, contaminant removal, followed by flake segregation. The primary step involves the melting and re-extruding of the plastic into uniform pellets, which can then be utilized in cast/film extrusion, injection molding, blow molding, etc. The molecular weight of a plastic is highly influential to all the plastic’s innate properties. In general, an increase in molecular weight increases the glass transition temperature, the melting temperature, plastic strength, elasticity, stiffness, toughness, and viscoelastic properties. The processing residence time and reprocessing of the material also influence the molecular weight of the polymeric materials. Fig. 12 demonstrates the change in melt-flow index (MFI) of various plastics subjected to extrusion and injection molding. The repeated exposure to high heat, time, and shear rates can cause degradation of the polymeric material, and in some cases, unwanted side reactions that can either increase or decrease the molecular weight of the final product. These structural modifications lead to changes in both flow and mechanical properties of the material. Manufacturers may be inclined to utilize additives to recover material properties or homogenize the recycled material, although this typically adds cost to the recycled plastic and may further complicate future recycling attempts of the material waste. Different variations of mechanical and thermal processing technologies (Fig. 13) used in recycling are described below.

4.2 Extrusion molding

In extrusion molding, plastic pellets enter the hopper into the barrel of the extruder in a continuous process. The plastic is fed into the barrel in the screw feed zone, and the screw forces raw material through the hot barrel and causes melting. The constant moving of the plastic through the barrel restricts overheating and limits resonance time, which may degrade the plastic. Because of the heating during extrusion, some alteration to the polymer chain occurs. The plastic melt in the barrel passes through screen packs to remove gross contami-
nants during the extrusion process. The screen pack and breaker plate are found in commercial plastic extrusion between the extruder and the die. Melted plastic is re-shaped using a die. The breaker plate ensures the proper mixing in the screw of the plastic by creating backpressure and reorients the plastic creating a stable linear flow before entering the die. Uneven flow of the molten plastic into the die can create undesirable residual stress that can cause warping during the cooling stage. The extrudate is cooled with a water bath or cooled air stream. This process is complicated as plastics are good thermal insulators. The extrusion process for recycling is similar to a new product manufacturing technique. Details of extrusion molding, injection molding, blow molding, film molding, and fiber processing can be found in SI 2.0 in ESI†.

4.3 Mechanical reprocessing of PET

Bailed PET often contains a large number of contaminants including adhesives, sugars, and other plastics. It is important to clean the PET and separate it from other materials before it is melt-processed through extrusion to produce recycled PET pellets (rPET). Fig. 14 illustrates a typical PET bottle recycling process which is adapted from one of the PET bottle recycling companies in Europe. The delivered bottles are sorted depending on the type of material, color, processing method etc. The bottles are then shredded into small flakes followed by washing processes. The flakes go to the extruder after cleaning and drying.

Melt extrusion of clean washed PET flake or granules is used to manufacture pellets of uniform shape for subsequent manufacturing processes (injection molding, extrusion, injection stretch blow molding, etc.). When PET reprocessing uses an established facility, processing becomes more cost-effective, more straightforward, and possesses less environmental impact. Switching from virgin to recycled PET depends on material pricing, secure recycled material supply chains, and the ability to use existing facilities. Processing PET at high temperatures may cause chain degradation which influences crystallinity of the plastic. Plastic degradation from chain scission causes molar mass plastic decrease, which makes bottle-to-bottle recycling challenging.

rPET can be used for most applications instead of virgin materials provided that the properties meet the quality criteria. Clarity and yellowing are major property challenges for rPET. Like other plastics, PET’s quality is also affected by chain degradation because of thermo-oxidation. Some contaminants (such as PVC and polyvinyl alcohol) accelerate chain scission in PET plastic during melt processing due to acidolysis. Impurities in PET enhance chain scission reactions, and traces of PVC and polylactic acid (PLA) can accelerate acidolysis or hydrolysis. These reactions alter the plastic’s microstructure and the mobile amorphous region is altered causing thickening of the crystalline domains. Crosslinking can also occur, resulting in increased viscosity in PET, hindering processability for applications such as injection molding. The multipole extrusion cycles of mechanical recycling increase the rate of chain scission because of thermo-oxidation.

Compounding additives into rPET during reprocessing can protect the plastic from chain scission and improve mechanical properties. To minimize thermo-oxidation of the plastic, metal-based stabilizers are often used. The use of radical scavengers, such as organic phosphates, is another option to
hinder radical induced oxidation of the terephthalate ring, which is a source of yellowing.\textsuperscript{155,158} The most common problem in PET reprocessing is viscosity reduction (molecular weight reduction) due to hydrolytic degradation and thermal oxidation. Color change in the post-consumer PET is another vital problem and normally the industry solves the problem by “open-loop recycling” or “semi-closed-loop recycling”.\textsuperscript{154} The concepts of “open-loop recycling” or “semi-closed-loop recycling” are described in section 4.3.1.1.

If the molecular weight reduction is less severe, the plastic can be subjected to a process called solid state polymerization (SSP) to help rebuild the molecular weight.\textsuperscript{159} The plastic is heated to a temperature between the melting and glass transition temperatures to induce esterification and transesterification reactions.\textsuperscript{153} Industries are currently looking for a new stream of PET waste to reprocess, including colored PET.\textsuperscript{150} For example, in 2018 ‘Indorama Ventures’ partnered with ‘Unilever’ to recycle fifty billion bottles per year by 2025, including colored PET. According to them, traditionally non-recyclable colored PET is subjected to a chemical recycling process. These colored PET are reduced to monomeric species, and then the contaminants removed before being used as raw material for polymerization. This technology produces transparent food-grade PET plastic material from post-consumer PET.\textsuperscript{161,162} Recycling for food contact packaging requires a very demanding and clean plastic. No Objection Letters (NOLs) by the Food and Drug Administration are often sought by recyclers and manufacturers to demonstrate that their recycling process and materials are suitable for direct food contact applications.\textsuperscript{163}

Continuous improvements in plastic reprocessing is an ongoing effort with new technologies in degassers and filters to facilitate increased plastic melting quality.\textsuperscript{164} Degassing employs a vacuum vent on the extruder to enable volatile compound stripping from a plastic melt. This stage reduces hydrolysis by removing moisture and adds value by producing a high-quality plastic with maintained molecular weight and decreased contamination. The addition of a filter helps remove unwanted non-volatile contaminants from the plastic and increases the mechanical properties of the recycled plastic.\textsuperscript{165} When selecting recycling technologies in manufacturing processes, it is important to select the appropriate mechanism to minimize the degradation of the plastic for high-quality new parts.\textsuperscript{155}

4.3.1 Utilization of recycled PET waste

4.3.1.1 Closed-loop and open-loop recycling. The recycled raw material supply increases as plastic waste collection increases. These raw materials can be used to produce new material as ‘open loop’ or ‘semi-closed loop’ or ‘closed loop’ recycling.\textsuperscript{166} Closed-loop recycling means the recycled raw material is used for the same product and fully replaces the virgin material, e.g., PET bottle to bottle recycling. Open-loop recycling means the recycled material is used for a different product and fails to replace the virgin raw material in the original application, also referred to as downcycling. The inherent properties of raw materials for open-loop recycling are often too degraded by the service life and recycling process and the plastic cannot be used in the original application. Semi-closed recycling means the recycled raw materials are used to produce another type of product but cause no change in the inherent properties of the recycled materials.\textsuperscript{167}

4.3.1.2 Recycled PET products. rPET is incorporated to form a wide range of new products, for example, shoes, bags, fibers, boards, mechanical parts, etc., as open-loop recycling. Most rPET is used for fibers.\textsuperscript{164} Consumers can currently purchase rPET food packaging containers. According to a 2020 NAPCOR (National Association for PET Container Resources) Report post-consumer rPET used in the US and Canada has continuously increased since 2004. It has been predicted that by 2026 the market will reach up to $12.5 billion, representing an ~8% Compound Annual Growth Rate (CAGR) from 2019 to 2026.\textsuperscript{168,169} The NAPCOR Chairman mentioned that post-consumer rPET utilization is proceeding towards a circular economy. According to NAPCOR, fiber sectors made up more than 40% of the worldwide rPET market in 2019.\textsuperscript{170} The fiber market includes manufacturing mattresses, insulators, car equipment, etc. Asia Pacific plastic industries are increasing their dependency on rPET. Colored rPET is used for food and non-food applications for its stability and strength. These have high popularity as they are lightweight and provide strength for a long period.\textsuperscript{168}

4.4 Mechanical reprocessing of polyolefins

Mechanical processing of polyolefins also induces degradation at the molecular level and the formation of aldehydes, ketones, and short-chain hydrocarbons.\textsuperscript{171–173} The degradation mechanism is different in each type of polyolefin. The general degradation mechanisms begin with the generation of macro-radicals within the molecular chain. This radical creates many shorter, branched, or cross-linked polymer chains. When these radicals react with oxygen centered radicals, further degradation occurs.\textsuperscript{174} Previous studies have demonstrated that HDPE is more prone to crosslinking whereas PP shows a large number of chain scission events. These degradation reactions mainly occur in the amorphous phase below the plastic’s melting temperature because oxygen cannot diffuse into the crystalline phase.\textsuperscript{155}

As post-consumer plastic films are contaminated with various waste materials, an intensive pre-treatment is crucial before extrusion. The source of raw material in secondary recycling is MSW mixed with other plastics. This needs an initial separation step to separate the contaminants. The mechanical recycling process described here (Fig. 15) is an example of a European processing facility.\textsuperscript{175} The process starts with comminution of large plastic parts and shredding/sieving to remove as much of the contaminants (metals, glass, paper, etc.) as possible. The contaminant-free plastics are sent for pre-washing then vigorous washing to avoid micro-physical contamination. Afterward, the material enters the hydroclone to separate undesirable particles.\textsuperscript{176} A friction washer is used for washing to consistently remove contaminants from the plastic such as labels, glue, or organic dirt on the...
Fig. 15 Mechanical recycling process of polyethylene plastic film at an industrial plant.175

surface.176 After washing, the plastics are dried thermally to lower the moisture content before entering the extrusion and pelletizing stage.175 The function of extruder and blowing machines are discussed above. The quality of the recycled plastic film is degraded leading it to be used in applications such as products for constructions, public spaces, and traffic applications after injection molding or extrusion.177

4.4.1 Mechanical processing of HDPE. HDPE is a linear chain plastic with high molecular weight produced from carbon and hydrogen atoms. As this plastic has a straight chain, it has more strength than LDPE. It is an important component of MSW as this waste plastic has high recyclability potential.136 Commercially, two kinds of HDPE are often found in MSW depending on either the extrusion process or injection molding; each type differs in its viscosity.155 Like other plastics, chain scission and chain branching are also common in HDPE during the extrusion process. Generation of carbonyl groups in HDPE increases when the oxygen in the surrounding atmosphere is high. When oxygen is low, double bond formation is observed in the molecular backbone.178 Because of these reactions, the HDPE molecular weight is reduced. When the plastic chain length is lower, the plastic becomes vulnerable to degradation by radical attack. This scenario worsens with each extrusion cycle. Studies have also shown that the viscosity in the plastic reaches a very high level (five fold higher than the starting level) after sixty extrusion cycles due to crosslinking in chains.155 These degradation effects can partially be mitigated via controlling extruder parameters such as temperature, screw speed and screw design.179

Recycled HDPE and other recycled polyolefins have multiple applications in the food packaging sector and multilayer packaging applications. However, there is a concern with recycled raw material in direct food contact because potentially toxic small molecules (plasticizers, stabilizers, etc.) could diffuse and migrate into food products.180 As a result, recycled polyolefins are often utilized as the middle layer in multilayered packaging, and the surface layers are made of virgin plastics. Recycled HDPE has a large market as the processing cost can be relatively low. Post-consumer HDPE is a feasible option to produce plastic lumber which is a replacement for wooden lumber.

4.5 How legislation changes the scenario

The municipal waste management system and plastic recycling in most European countries has made remarkable advancement. EU countries with landﬁlling restrictions or bans have higher recycling and waste recovering percentages as shown in Fig. 16. Recycling of HDPE, LDPE, PET, PP, and PS plastics are performed together.181 Post-consumer plastic waste collection was more than 32% in several EU countries which was mostly comprised of mechanical recycling techniques. The highest recycling rate was found in Norway at 45% in comparison to ~9% in the United States.137,182

The number of recycling companies is increasing. There are almost 115 PET recycling and 40 PE recycling facilities in the US. For example, ‘77 recycling’ is a plastics (post-industrial plastic scrap) recycling company that accepts most materials (PET, PP, PS, PVC, HDPE, LDPE, etc.).183 China’s ban on imports of plastic waste increased the recycling facilities across the US.184 Good communication between product manufacturers and the recycling industries is needed to confirm the recycling of packaging materials. This communication can strengthen the use of recycled raw materials and create a new market for these recycled plastics.185

4.5.1 Utilization of mechanically recycled post-consumer materials as food contact materials (FCM). The most valuable type of recycled plastic is plastic that can be used for food packaging applications. In the US the FDA employs regulations regarding the safety of the use of recycled plastic materials as food contact materials (FCM).187 The FDA expresses their concerns and regulations maintained for using recycled plastic:

- When recycled plastic/material is used as FCM, the chance of contaminants in the final product may increase.
- Post-consumer recycled plastic/materials are permissible to use as FCM material.
- The adjuvants (for example petroleum waxes, synthetic petroleum wax, colorants, plastic modifiers) coming from the post-consumer recycled plastic/material may not be suggested for use in FCMs as the contaminant present in the recycled material can pass on to the new product and migrate into the food matrix.189
- The FDA regulates the plastic recycling processes to ensure that the recycled materials are not contaminated and that the recycling process is suitable for direct food contact.161 For example, in secondary recycling, the physical reprocessing steps, grinding, melting, and reforming, are documented and submitted for evaluation to obtain a favorable opinion letter.
through the No Objection Letter (NOL) program. For plastic depolymerization, it is necessary to confirm that the monomers are contamination-free and the produced plastic is the same as that from virgin monomers.190

- In the case of multi-layered packaging, sometimes, when proper contamination removal is complex, the recycled layer is used as the most outer layer. For example, for recycled PET, FCM is used as the outer layer in multi-layered containers, and the inner layer in contact with food is made of virgin material.155

Manufacturers of FCMs using post-consumer recycled materials often need special evaluation.163 The NOL determines the assessment of the plastic used for a FCM and whether the recycling process is physical or chemical. The earliest NOL191 was issued in 1990 for a physical recycling process of whole egg cartons.192 Many of the early NOLs that granted favorable opinions were for PET and PS plastics.193 However, in recent years, much of the focus has been on polyolefin materials; utilization of post-consumer polyolefin streams for direct food contact applications is challenging due to a large number of different molecular structures, additive packages, and uses for which consumer abuse can lead to incidental contamination.194 Such challenges are overcome through careful source control of the post-consumer material and challenge surrogate chemical testing to determine if the recycling process can reduce the concentration of different types of compounds, which is often a requirement for the NOL process.

For the surrogate chemical challenge testing, the plastic is soaked in a chemical cocktail containing a volatile polar compound (e.g., chloroform, chlorobenzene, 1,1,1 trichloroethane, diethyl ketone), a non-volatile polar compound (toluene), a heavy metal compound (copper(II) 2-ethylhexanoate), non-volatile polar (benzophenone or methyl salicylate), and a non-volatile non-polar compound (tetracosane, lindane, methyl stearate, phenylcyclohexane, 1-phenyldecane, 2,4,6-trichloroanisole) using an isopropyl/hexane solvent delivery mixture to artificially contaminate the plastic.163 After the artificial contamination soaking of two weeks at 40 °C, the spiked plastic is subjected to recycling. The concentration of the artificial contaminants is quantified in the recovered, recycled plastic and the starting spiked plastic to demonstrate the ability of the recycling process to reduce contamination. These results, along with a description of the recycling procedure and post-consumer material feedstocks, are submitted to the FDA to evaluate...
5. Thermal decomposition of plastics

5.1 Non-catalytic thermal decomposition of plastics

Pyrolysis is the thermal decomposition of organic compounds typically in an oxygen-deficient atmosphere. The decomposed products are then condensed and collected as oils, waxes, gases, and char, with distributions dependent on feedstock, reaction conditions, and reactor type. Temperatures in the range of 350–700 °C are typical for plastics pyrolysis, with the general trend of higher yields of gas at higher temperatures. Pyrolysis is generally classified according to heating ability to convert plastic into lower molecular weight products to be used as fuels or feedstock for new chemicals or plastics. Pyrolysis also produces less emission than plastics incineration. Unlike mechanical recycling, pyrolysis can handle highly heterogeneous feedstocks with a higher degree of contamination.

5.1.1 Fundamental chemical processes during thermal pyrolysis of plastics

5.1.1.1 Polyolefins. PE follows a free radical thermal decomposition that takes place through sequential random bond-scissions. The degradation begins in weak links (mainly peroxide groups formed during preparation and storage of PE) and proceeds in the pristine segments of the plastic. Peterson et al. employed thermogravimetric analysis to demonstrate that the activation energy depends on the extent of the decomposition reaction and assumes values between 150 kJ mol$^{-1}$ and 240 kJ mol$^{-1}$. The latter number corresponds to the random scission process for pristine PE. Huber and coworkers reported a similar activation energy (234.8 kJ mol$^{-1}$), and argued that only models that include two random scission steps are capable of capturing the details of weight loss curves as obtained from thermogravimetric analysis. The complexity of the free radical thermal decomposition of PE is substantial, and this is reflected in the large variety of pyrolysis products. The main products are alkenes, alkynes, and alkanes, while Diels–Alder reactions may also lead to the formation of small amounts of cycloalkanes.

The initiation of PE pyrolysis involves a homolytic C–C cleavage reaction that happens at a random position along the PE chain, and gives two macroradicals (1) – (Scheme 1). Each of these radicals may undergo a β-scission to produce ethylene monomer and another primary radical. However, this unzipping process is not kinetically favorable at the early stage of PE decomposition (ca. 400 °C), and as a result no or little ethylene is produced in experiments (<1%). In contrast, radical (1) prefers to undergo a backbiting (often 1,5-backbiting through a six membered transition state) reaction, thereby generating a secondary internal radical (2) (Scheme 1). Subsequently, a β-scission at (I) gives rise to an alpha–olefin (3), while if the same process occurs at (II), a terminally unsaturated polymer residue (5) is produced, as well as a radical that transforms into an alkane (6) after abstracting a hydrogen atom. The formation of alkadienes (8) takes place either via a hydrogen abstraction reaction from (5) (Scheme 1) or via chain scission followed by a backbiting reaction. These processes give rise to (7) that, after a β-scission, is converted into an α,ω-alkadiene (8) (Scheme 1) and a new primary radical (1), which likely undergoes the processes of Scheme 1 anew.

Similar to PE, the thermal decomposition of PP (9), Scheme 2, proceeds via random scissions along the backbone of the polymer with an activation energy of about 250.0 kJ mol$^{-1}$. Each random scission produces a primary (10) and a secondary (11) radical (Scheme 2), which, in turn, are converted to alkanes, alkenes, and alkadienes. The major decomposition products are alkenes and alkadienes, and their formation takes place in a manner similar to what we described in Scheme 1.

The highest products from PP pyrolysis (i.e., 2-methyl-1-pentene, 2,4-dimethyl-1-heptene) are produced via the decomposition of the secondary radical (11), while only minor quantities of products are derived from the primary radical (10). On this basis, Scheme 2 focuses on the formation of alkenes and alkadienes via (11), which after a backbiting reaction and a β-scission at (IV) forms an alkene product (13) and another secondary radical (11). The new secondary radical is decomposed following the same sequence of steps. Alternatively, the β-scission may occur at (V), which results in a terminally unsaturated polymer (14). The latter species may undergo an atomic hydrogen abstraction reaction followed by a β-scission at (VI), thereby forming another secondary radical (11) and a series of alkadiene products (17).

5.1.1.2 Polyvinyl chloride. The thermal decomposition of PVC (18), Scheme 3, follows a completely different mechanism compared to polyolefins. According to thermogravimetric analysis studies, PVC (18) starts decomposing at much lower temperatures (<250 °C) than other common polymers, and its decomposition involves two distinct stages.

The dominant reaction in the first stage (250 °C–350 °C) is the dehydrochlorination of the backbone that leads to a 60% weight loss. The activation energy for this process has been determined in several studies, and values in the range of 112.0 kJ mol$^{-1}$–150.0 kJ mol$^{-1}$ have been reported. It has been conjectured that the initiation occurs in thermally labile defects like butyl branches associated with tertiary chloride. Once the reaction is initiated, its propagation proceeds via a quasi-ionic mechanism whereby HCl is removed in a concerted manner via a four-center transition state (19). Subsequently, the propagation reaction is relatively fast owing to the continuous formation of internal allylic chloride structures [see (20) in Scheme 3], which are more thermally unstable and susceptible to C–Cl heterolysis via HCl.
catalysis [i.e., autocatalysis – transition state (21) in Scheme 3] than the pristine PVC polymer segments. Remarkably, first-principles calculations by Yanborisov and Borisevich showed that the HCl-catalyzed dehydrochlorination of small chlorinated molecules (e.g., 2-chlorobutane) can be 60.0 kJ mol\(^{-1}\) more facile than the uncatalyzed process. The main dehydrochlorination products are HCl gas and \(\pi\)-conjugated polyene sequences (22). The latter products are highly unstable under pyrolytic conditions and therefore undergo intramolecular and intermolecular reactions as soon as they form. The intermolecular reactions require the interaction between two polyene chains (22), and give rise to crosslinked polyene chains (23), which are the precursor of char (24). Char is extensively formed in the second stage of the thermal pyrolysis. Alternatively, a polyene chain may undergo an intramolecular electrocyclic reaction, thereby giving rise to \(\pi\)-conjugated 1,3-cyclohexadiene rings (25). The latter are converted to benzene (26), and subsequently polyaromatics (27), after two consecutive homolytic C–C cleavages that occur at temperatures beyond 350 °C having activation energies larger than 220.0 kJ mol\(^{-1}\). C–C bond breaking also gives rise to small amounts of non-condensable gases (e.g., CH\(_4\) and H\(_2\)). Importantly, we note that the formation of aromatics can be promoted by applying a high-temperature pyrolysis step in the presence of a solid catalyst (e.g., ZSM-5 zeolite).

5.1.1.3 Polystyrene. The thermal degradation of PS (28) follows a radical chain mechanism (Scheme 4), with styrene (35) being the dominant product. By means of thermolysis experiments, Wall et al. demonstrated that the thermal stability of PS (28) depends on the polymerization method (i.e., anionic or radical polymerization) used in its synthesis. Preparation through a radical polymerization gives rise to polymer chains with randomly distributed “weak links”, which perhaps are peroxidic in nature and are the sites where the thermal decomposition of PS starts. Yet, anionic polymerization yields PS of high thermal stability, the general thermal decomposition mechanism for this case is described in Scheme 4 and the following text.

The experimentally measured activation energy for the thermal degradation of PS is 200.0 kJ mol\(^{-1}\) and remains constant regardless of the extent of the reaction. This is indicated by the Arrhenius plot (Fig. 2).
tive of a single rate-determine step during the reaction: the \( \beta \) scission that forms styrene (35) (see Scheme 4).\textsuperscript{211} According to Guyot and coworkers,\textsuperscript{234,237} a likely initiation pathway for “anionically” prepared PS is via an end chain scission reaction that forms a benzyl radical (30) and a primary radical (29). The two radicals go through a disproportionation reaction and give toluene (32) and a chain with a thermally unstable unsaturated end (31). In turn, (31) decomposes into \( \alpha \)-methylstyrene (33) and a secondary radical (34) that leads the degradation process thereafter. DFT work by Huang \textit{et al.} revealed that the latter reaction is exothermic by 80.8 kJ mol\(^{-1}\) and its activation energy is 288.2 kJ mol\(^{-1}\).\textsuperscript{238} The propagation occurs by a \( \beta \)-scission process at (IX), thereby generating styrene (35) and another secondary radical (34). According to DFT calculations on a PS trimer, this reaction has an activation energy of \textit{ca.} 117.0 kJ mol\(^{-1}\), but it is strongly endothermic (+105.4 kJ mol\(^{-1}\)) and therefore not favorable at temperatures below 250 °C.\textsuperscript{238} Interestingly, the computed activation energy for the unbuttoning reaction (\textit{i.e.}, 117.0 kJ mol\(^{-1}\)) is in stark contrast to the experimental value (200.0 kJ mol\(^{-1}\)) and this can, to some degree, be justified by the considerably smaller size of the simulated PS trimer as compared to actual PS chains. (34) can either undergo further unbuttoning to give extra monomer (35) or a backbiting reaction followed by another \( \beta \)-scission reaction at (X) and give a dimer (37) along with another secondary radical (34). Based on DFT, such backbiting and \( \beta \)-scission reactions require kinetic barriers of 174.0 kJ mol\(^{-1}\) and 134.5 kJ mol\(^{-1}\), respectively, and the whole process that forms (34) and (37) is endothermic by 94 kJ mol\(^{-1}\).\textsuperscript{238} We note that the formation of trimer or tetramer species takes place in an analogous manner to that for dimer production (37) (Scheme 4) with the only difference that...
5.1.1.4 Poly(ethylene terephthalate). PET (38) is a thermally stable polymer thanks to the combination of short aliphatic chains and aromatic rings. Its thermal decomposition follows a molecular mechanism and the initiation reaction involves a scission at the ester linkage (Scheme 5). This reaction takes place through a six-membered transition state (39), and recent DFT studies predict activation and reaction energies of 184.0 and +48.0 kJ mol$^{-1}$, respectively.\(^{(242)}\)

The initiation reaction leads to the formation of an oligomer with an olefin end group (40) and another oligomer with a carboxylic end group (41). These two species decompose further to produce a wide spectrum of products with acetaldehyde (44) and CO\(_2\) (43) being the most abundant at temperatures around 300 °C.\(^{(240)}\) The former can be produced directly from (40) and (41) or via the degradation of EG end groups as proposed by Holland and Hay.\(^{(243)}\) While CO\(_2\) is released by decarboxylation of (41). More than 45 wt% of the PET is degraded into CO\(_2\) with trace CO.\(^{(244)}\)

5.1.2 Variables affecting product distribution. Pyrolysis begins with free-radical initiation in the plastic melt, or condensed phase. Volatile products are released from the melt as depolymerization progresses, leaving behind ash, fillers and (sometimes) char. Secondary reactions occur in the vapor phase that cause further cracking. Gas, liquid, and wax are produced, with distributions dependent on reaction conditions. These products are usually categorized according to their carbon length and/or phase:\(^{(245)}\)

- C1–5 = gas
- C6–20 = naphtha or oil, liquid at ambient conditions
- C21+ = wax, solid or highly viscous liquid at ambient conditions

Al-Salem et al.\(^{(246)}\) published an in-depth review addressing plastic pyrolysis variables and their influences on reaction and product distribution.
Scheme 4  Fundamental reactions for the thermal decomposition of PS leading to different products. Arabic numerals indicate chemical species, while Roman numerals indicate the occurrence and position of the scission of a chemical bond.

Scheme 5  Fundamental reactions for the thermal decomposition of PET. The symbol † indicates a transition state. Arabic numerals indicate chemical species and intermediates formed during the pyrolysis of PET.
product distribution and asserted that the main decomposition behavior of the plastic is controlled by temperature. In PE pyrolysis, early studies showed that only liquid and gas – no wax – are produced above 600 °C. Increasing temperature leads to increased aromatic production, as gas is converted to aromatics through a Diels–Alder reaction and unimolecular cyclization reactions, although a study by Kulas et al. showed that the reaction pathway from gas to aromatics has low temperature dependence.

The composition of naphtha is vastly different than the composition of plastic oil (Fig. 17). The main products of naphtha are paraffin with around 10% aromatics, while the main products of plastic pyrolysis oil are olefins and cycloalkanes/alkadienes with a much wider range of carbon numbers.

Increasing vapor residence time increases the yields of oil and non-condensable gases. As primary products are exposed to heat for longer periods of time, secondary reactions and further cracking to smaller molecules are encouraged. Thus, the product distribution shifts from high molecular weight products to lower molecular weight hydrocarbons (MWHCs). For polyolefin feedstocks, longer reaction times decreased product olefin concentrations and increased polyaromatic hydrocarbon concentrations.

Polyolefins have received the most attention in the field of plastics pyrolysis. PE and PP decompose into a distribution of hydrocarbons, with decreasing product molecular weights as reaction temperature and time increase. Waxes and heavy oil olefins dominate product distributions at low temperatures, while gases and light oils are the main products at high temperatures. Polyolefins such as HDPE, LDPE, and PP decompose into a distribution of hydrocarbon products that have high concentrations of olefins and dienes (>70%). Mixed plastic feedstock with high PE content will lead to higher paraffin concentrations. Pure PP yields higher amounts of olefins, with lesser amounts of paraffins, naphthenes, and aromatics. Due to the tertiary carbon present on the backbone of PP, multiple types of secondary radicals are formed, shifting liquid product composition towards branched olefins.

PS decomposes primarily into the monomer styrene and its oligomers with the polymer undergoing chain scission followed by random scission. PS pyrolysis produces liquid products at yields greater than 90 wt% in both batch and continuous reactors. Liu et al. pyrolyzed PS in a fluidized bed reactor at 600 °C and achieved 98.7 wt% oil, of which 78.7 wt% was styrene. Secondary reactions and increased reaction temperature decrease the concentration of styrene in the liquid product and increases the formation of coke and gas. PS accelerates the pyrolysis of polyolefins due to higher concentrations of free radicals. Green Mantra Technologies is commercializing PS pyrolysis to recover styrene. A thorough review on the pyrolysis of PS can be found elsewhere.

The pyrolysis of PET yields oxygenated compounds from the presence of oxygen in the backbone of the polymer chain. Fast pyrolysis of PET produces mostly mixtures of aromatics and the organic acids benzoic and phthalic acid, char, and gas. Formation of TPA is particularly problematic due to its high boiling/sublimation temperatures, which clogs reactor and condenser internals. The presence of phthalic acids in product oils increases viscosity and acidity and lowers calorific value, resulting in low-quality oils. Pyrolysis temperature has little effect on calorific values of PET products. PET in mixed plastics increases the CO and CO₂ concentrations in product gases from pyrolysis reactions. While some investigators have concluded that PET is not suit-
able for plastic pyrolysis, others advocate for it based on the valorization of product char to activated carbon and product oil to fuel. US-based pyrolysis company Agilyx has undertaken pyrolysis of PET with some success, but PET is rarely used as a feedstock in commercial pyrolysis processes. Other possible chemical recycling processes of PET are detailed elsewhere and in section 9.1.

Many commercial pyrolysis processes have low tolerance to PVC, which produces significant concentrations of organic chlorine, HCl and a solid residue. The HCl causes corrosion issues with downstream processing of the products. Thermal decomposition of PVC is initiated by a dechlorination reaction, resulting in the formation of hydrogen chloride and benzene. The polymer chain then undergoes cyclization, yielding aromatic and alkylaromatic compounds that are often chlorinated. Pyrolysis oil containing chlorinated compounds has limited use. Most previously reported pyrolysis processes had a specification that the PVC content of the plastic feeds be less than 2 wt%.

Plastics can also be co-processed with waste tires, wastepaper, and agricultural residues, which has been shown to enhance products for fuels and value-added chemicals. The majority of waste plastics will contain high levels of contaminants. Additives, heavy metals, and pigments are often added to plastic to enhance its physical appearance or performance. Specific problematic elements include nitrogen, oxygen, sulfur, chlorine, iron, sodium, and calcium. In low-temperature pyrolysis, sulfur presents in the pyrolytic oil, which reduces its ability to be used as ultra-low sulfur diesel. Produced liquid may also contain amines or alcohols if the feedstock contains N, O, or S. Siloxanes can be present in the pyrolysis oil as well. Dechlorination is possible through the use of sorbents, absorption and neutralization, subcritical water treatment, catalysts, or stepwise pyrolysis, but the latter was shown to modify the composition of liquid products. Untreated plastic waste pyrolysis oils with high levels of contaminants are not feasible feedstocks for steam cracking unless sufficiently diluted. Contamination may lead to catalyst deactivation if catalytic upgrading of products is desired. For example, a municipal plastic waste pyrolysis plant in Idaho, U.S.A. produced an oil only suitable for low-value applications due to severe contamination.

Thermal oxo-degradation (TOD) involves adding small amounts of oxygen to the pyrolysis process to increase chain cleavage. Plastics subjected to TOD are decomposed to oxygenated products and hydrocarbons, with exothermic partial oxidation reactions releasing energy to provide heat back to the process. This process mirrors that of autothermal pyrolysis developed for the process intensification of biomass, with the possibility to achieve autothermal operation for plastics.

5.1.3 Reactor configurations. Various reactor configurations have been used for plastic pyrolysis, including fixed beds, fluidized beds, rotary kilns, auger reactors, conical spouted bed reactors, and melting vessels (Fig. 18). Detailed information on reactor types for plastic pyrolysis can be found in recent review papers. Fluidized bed reactors have excellent heat and mass transfer characteristics for pyrolysis of plastics and are the reactors that are most typically used.

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Fig. 18 Schematics of various plastic pyrolysis reactors including fixed bed, fluidized bed, conical spouted bed, stirred tank, auger, and rotary kiln reactors.
for continuous large-scale processing of solids. In a gas–solid system, fluidized bed reactors have vigorous solid contact with different gas flow rates. An increasing of flow rate in a bubbling fluidized bed (BFB) leads to larger instabilities and bubbling channels of gas with a more vigorous solid movement.\textsuperscript{311} In turbulent fluidized bed, with a sufficiently high gas flow rate, the terminal velocity of the solids is exceeded and solids undergo a turbulent motion so that the upper surfaces of the solid disappear.\textsuperscript{315} Fluidized bed reactors were used extensively in the groundbreaking work in plastics pyrolysis by Walter Kaminsky at the University of Hamburg.\textsuperscript{288,289,316–318} Both BFBs and circulating fluidized beds (CFB) have been commonly employed in both laboratory studies\textsuperscript{254,256,257,317,319–322} and industrial deployment.\textsuperscript{323} Fluidized beds usually employ smaller feedstock particles than other reactors, which, along with their inherently high convective coefficients, enhances heat transfer rates. Feedstock injected into a fluidized bed is rapidly dispersed and efficiently contacted with gas, resulting in high mass transfer rates.\textsuperscript{311,313} The combination of high heat and mass transfer rates are responsible for significant yields of light hydrocarbons and non-condensable gas from the pyrolysis of plastics in fluidized beds.\textsuperscript{305,324} Disadvantages include comminution of plastics to achieve small feedstock particles, and complicated reactor setups.\textsuperscript{375} Additionally, the high gas velocities required for fluidization can cause entrainment of char particles, requiring particulate separation downstream of the reactor.\textsuperscript{196} Fluidized beds may also see agglomeration of melted plastic that can lead to bed defluidization.\textsuperscript{325}

Conical spouted bed reactors are a type of fluidized bed with a cylindrical geometry and a conical base. Cyclic and vigorous movements by sand particles in the bed prevent agglomeration upon melting and subsequent bed defluidization.\textsuperscript{259,326} Conical spouted beds have been employed in laboratory-scale pyrolysis studies of PS\textsuperscript{327} and polyolefins,\textsuperscript{326,328,329} which report high yields of wax. Orozco \textit{et al.}\textsuperscript{255} identified stable operating conditions for conical spouted bed reactors for individual plastic feedstocks.

Fixed bed reactors are inexpensive, simple to operate, and have low maintenance requirements. However, these types of reactors would be difficult to operate for solid feedstocks on the industrial scale due to large amounts of solid coke/char formation. Vertical fixed beds use gravity to force products through the reactor. Tubular reactors are very similar to fixed beds.\textsuperscript{267,345,346} Stirred tank reactors.\textsuperscript{312} Stirred tank reactors are primarily used for batch reactions, which is not economical on a large continuous scale process.

Pyrolysis reactors can be heated by a heat carrier (Fig. 19a) or with an external heat exchanger (Fig. 19b). The heat balance when a heat carrier is used is shown in eqn (1), where $M_{\text{cat}}$ is the mass flow rate of catalyst (or heat carrier), $C_p$ is the heat capacity of the gas, $T$ is the temperature, $M_{\text{feed}}$ is the mass flow rate of reactant, and $\Delta H_{\text{rxn}}$ is the net heat of the heat of the pyrolysis reaction. When using a heat carrier, the reactor can be scaled to larger industrial sizes without limitations to the heat transfer area by use of an external fluid. In contrast if an external fluid is used for heating the energy balance is dependent on the heat transfer area as shown in eqn (2) and (3).

\begin{equation}
M_{\text{cat}}C_p(T_{\text{out}} - T_{\text{in}}) = M_{\text{feed}}(\Delta H_{\text{rxn}} + C_p(T_{\text{out}} - T_{\text{in}})) \tag{1}
\end{equation}

\begin{equation}
\dot{Q}_{\text{in}} = UA\Delta T = M_{\text{feed}}(\Delta H_{\text{rxn}} + C_p(T_{\text{out}} - T_{\text{in}})) \tag{2}
\end{equation}
PE pyrolysis requires around 260 kJ kg$^{-1}$ of heat. In a larger reactor, the heat transfer efficiency decreases due to the larger reactor diameter. Brown et al. reported that the maximum diameter of a biomass fast pyrolyzer is 9.4 mm, and an analogous calculation for a PE pyrolyzer obtained a maximum diameter of 6.4 cm. The reaction rate of PE used for the calculation is obtained from Zhao et al. In addition, the overall heat transfer will be affected by the bulky nature of the feed plastic, and the real diameter of the plastic pyrolyzer may be smaller than 6.4 cm. Autothermal pyrolysis can help to reduce the amount of heat required for pyrolysis by adding a certain amount of oxygen to partially oxidize the hydrocarbon feed, which makes the rate of energy release from the oxidation reaction equal the rate of energy consumed by the plastic pyrolysis. However, one main problem of the auto-thermal process is that it causes hot or cold spots due to the heat absorbed or released from the autothermal pyrolysis. The key to eliminating the hot and cold spots is again improving heat transfer within the reactor.

5.1.4 Economic analysis. Hundreds of species of hydrocarbons may be present in the pyrolysis oil, including olefins, paraffins, aromatics, diolefins, iso-paraffins, and naphthenes. PE-derived oils have high concentrations of linear hydrocarbons, while PP-derived oils contain more branched components such as trimers, tetramers, and pentamers of propene. Many petroleum companies like Shell, BP, TotalEnergies, SABIC, and ExxonMobil, are planning to upgrade pyrolysis oil through steam cracking. Kusenberg et al. has reported that the uncertainty and contaminants in the pyrolysis oil hampered industrialization.

Various studies have shown that the profitability of plastic pyrolysis refineries depends on several factors including feedstock cost, yield rate, product type, and facility scale. Feedstock composition, pyrolysis yield, and product selectivity are critical drivers for process efficiency. Feedstock cost, capital investment, and market prices play a significant role in the profitability of a plastic pyrolysis facility. Table 12 summarizes the key assumptions and findings from several studies on plastic pyrolysis. Fivga & Dimitriou investigated the pyrolysis of plastic waste to produce wax/oil products that can be used as a heavy fuel oil substitute or as raw materials by the petrochemical industry. The study evaluated a base case scenario of a 0.7 kton per year pyrolysis pilot plant built by a recycling company in the United Kingdom. The base case analysis resulted in a capital cost of $1.34 million. The product cost from the base case scenario was $0.98 per kg of fuel produced, which was more than the market price of the heavy fuel oil product of ca. $0.8 per kg. Increasing the pyrolysis scale from 70.1 kton per year to 701 kton per year decreased the product price from 29.4 cents to 3.39 cents per kg. The pyrolysis plants can be designed as either closed-loop or open-loop. Larrain et al. also emphasized that to make an open-loop and

\[ A = 2\pi nrL \] (3)

Table 12 Summary of waste plastic techno-economic analysis pyrolysis articles by feedstock, products, region, capacity, capital cost, and return on investment (ROI)

<table>
<thead>
<tr>
<th>Technology</th>
<th>Feedstock</th>
<th>Major products</th>
<th>Region</th>
<th>Capacity (kton per year)</th>
<th>Capital Cost ($ million)</th>
<th>NPV ($)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyrolysis</td>
<td>PS, PP, PE</td>
<td>Heavy oil</td>
<td>United Kingdom</td>
<td>0.7–701</td>
<td>1.36–77.2</td>
<td>–0.44 per kg–0.71 per kg</td>
<td>356</td>
</tr>
<tr>
<td>Pyrolysis + upgrading</td>
<td>PS, PP, PE, PET</td>
<td>Petrochemical feedstock</td>
<td>Korea</td>
<td>260</td>
<td>118</td>
<td>0.062 per gal</td>
<td>278</td>
</tr>
<tr>
<td>Pyrolysis &amp; heat integration</td>
<td>PE, PP, PET</td>
<td>Hydrocarbon fuel</td>
<td>Australia</td>
<td>14.6</td>
<td>3.76</td>
<td>2.03 million</td>
<td>360</td>
</tr>
<tr>
<td>Fast pyrolysis (open-loop &amp; closed-loop)</td>
<td>HDPE</td>
<td>Diesel power char</td>
<td>United States</td>
<td>193</td>
<td>118.5–120.5</td>
<td>367.8 million–383million</td>
<td>359</td>
</tr>
<tr>
<td>Fast pyrolysis</td>
<td>Mixed polyolefins mainly LDPE and residual PP</td>
<td>Ethylene propylene</td>
<td>Belgium</td>
<td>120</td>
<td>Not disclosed</td>
<td>open loop: 32.5 per ton closed loop: 2.72 per ton</td>
<td>357</td>
</tr>
<tr>
<td>Pyrolysis</td>
<td>Plastic waste (PP, PE, PS)</td>
<td>Light oil heavy oil</td>
<td>Malaysia</td>
<td>120</td>
<td>58.6</td>
<td>20.9 million</td>
<td>358</td>
</tr>
</tbody>
</table>
closed-loop pyrolysis of mixed olefins attractive to investors, closed-loop and open-loop plastic pyrolysis plants should be scaled up to at least 70 kton per year and 115 kton per year, respectively.\textsuperscript{157}

Larrain et al. investigated the economic feasibility of open-loop and closed-loop pyrolysis of mixed polyolefins (mainly LDPE and residual PP) in a 120 kton per year facility in Belgium. In the open-loop system, the process only produces naphtha, while the closed-loop system produces naphtha and wax. The study recorded a base case net present value (NPV) per tonne of plastic waste processed of $32.5 per ton and $2.72 per ton for the open-loop and closed-loop systems, respectively. The market value of the products resulted in a significant difference of $24.75 per ton, thereby emphasizing that the type of products plays a vital role in the facility's profitability. The study went further to consider the uncertainties in oil prices that significantly affect the prices of wax and naphtha in the market and concluded that the open-loop system has about 98% probability of achieving positive returns compared to 58% recorded for the closed-loop system.\textsuperscript{157}

Gracida-Alvarez et al. conducted the economic analysis of a multi-product 193 kton per year pyrolysis-based refinery for the conversion of HDPE to value-added chemicals (ethylene, propylene, and aromatic mixtures) and hydrocarbon mixtures via pyrolysis. In this study, the authors estimated the capital cost, operating cost, and NPVs for a base case where HDPE is converted to the products and a heat integrated case. The authors reported a capital cost of $118.5 million for the base case and $120.5 million for the heat integrated case. The NPV can range from $320 million to $450 million depending on the selling price of low MWHCs and the internal rate of return (IRR).\textsuperscript{159} The study showed that plastic pyrolysis targeting multiple products could be very profitable.

5.1.5 Commercial activities. Table 13 lists the publicly made announcements about major plastic pyrolysis technologies that are in operation, were previously planned or are currently planned. Western Europe began to study plastic pyrolysis in the late 1970s.\textsuperscript{287} The first of these plants was designed by BP in 1994. Using mixed waste packaging plastics, research on a laboratory scale was followed by demonstration on a continuous pilot plant scale (nominal 50 kg h\textsuperscript{-1}) at BP’s Grangemouth site. BP Chemicals, VALPAK, and Shanks & McEwan formed a collaborative initiative (POLSCO) in 1998 to study the viability of a 25 kton per year facility with a fluidized bed reactor, which would include logistical infrastructure for delivering mixed plastics from Scotland.\textsuperscript{361} This plant was never built because BP claimed it would require a tipping fee of €250 per ton ($225 per ton) to fulfill a €15–20 million ($24.75–33 million) total investment cost of this plant.\textsuperscript{287}

In Europe, there are 3 operating pyrolysis plants owned by Quantafuels and Plastic Energy. Quantafuels’s first commercial production facility is in Skive, Denmark. The plant processes 20 kton per year. The liquid product is sent to BASF, who uses the liquid product to produce virgin plastic and other chemicals.\textsuperscript{362} Plastic Energy currently has two plants in Almeria (started in 2014) and Seville (started in 2017) in Spain. The plant in Seville has the ability to treat 5 kton per year.\textsuperscript{363}

Several companies have announced plans in the past few years to build plastic pyrolysis plants and convert the plastic oils into aromatics/olefins by steam cracking. In Germany, SABIC and BP announced in 2021 plans to build a plastic recycling plant in Gelsenkirchen.\textsuperscript{355} BP will build plants to steam crack the plastic oil obtained from SABIC and forward the cracking materials to SABIC for making new plastics.\textsuperscript{355} One goal of BP is to achieve 30% of propylene and ethylene produced from recycled material by 2030.\textsuperscript{364} Shell plans to work with Pryme to build a plastic pyrolysis plant and produce circular chemicals in the Netherlands and Germany with the capacity of 60 kton per year in 2022.\textsuperscript{365} Plastic Energy announced a collaboration with ExxonMobil in early 2021. Starting in 2023, ExxonMobil will use the plastic oil from Plastic Energy and upgrade the plastic to a naphtha-quality petrochemical feedstock.\textsuperscript{354} The company has started to construct a recycling plant in France next to ExxonMobil’s Notre Dame de Gravenchon petrochemical complex, which will have a capacity of 25 kton per year.\textsuperscript{366} QuantaFuel plans to build a 100 kton per year plant in Sunderland UK and to work with VITOL and VITTO to build another 100 kton per year plant in Amsterdam.\textsuperscript{367,368} Fuenix Ecogy raised €4 million ($4.53 million) to scale up its technology to recycle mixed plastic waste, and in a collaboration with Dow to build a plant to pyrolyse plastic in Weert, Netherlands in 2019.\textsuperscript{369,370} The pyrolysis plastic oil will be sent to Dow Terneuzen, Netherlands, for steam cracking and producing virgin quality plastics.\textsuperscript{371} The goal for both Dow and Fuenix is to generate 100 kton of the recycled plastics by 2025.\textsuperscript{372}

In the US, Brightmark is planning to build one of the largest plastic pyrolysis plants in the world, with a scale of 100 kton per year, in Ashley, Indiana.\textsuperscript{373} Freepoint Eco-Systems plan to invest $60 million to construct a plastic pyrolysis plant in Obitz, Ohio, with a capacity of 90 kton per year. The company claims that the plant will be operating in 2023 and will be able to convert 300 tons of waste plastic into 1500 barrels of oil.\textsuperscript{374,375} TotalEnergies is also working with the Freepoint Eco-Systems to build a 33 kton per year plant in Houston, Texas in mid-2024. The products will be sent to TotalEnergies’ steam cracker to produce monomer and manufacture high-quality plastics.\textsuperscript{352,376} Agilix and ExxonMobil created a joint venture together called Cyclyx International, which will be focused on creating novel ways to collect and pre-process massive amounts of plastic trash into feedstocks for higher-value products.\textsuperscript{374}

There are also some plastic pyrolysis projects underway in Asia. Shell decided to start producing plastic oil with a capacity of 50 kton per year at Pulau Bukom, Singapore in 2023.\textsuperscript{377} Plastic Energy collaborates with WWF-Indonesia and aims to help Indonesia to reduce 100 kton of plastic waste by 2025.\textsuperscript{378} Insights provided by McKinsey showed that pyrolysis could be the most profitable process in the coming years.\textsuperscript{379} Due to the large amounts of oil required to run the hydrocarbon
<table>
<thead>
<tr>
<th>Name</th>
<th>Plant location</th>
<th>Scale (kton per year)</th>
<th>Status</th>
<th>Reactor type</th>
<th>Feedstocks</th>
<th>Product</th>
<th>Collaborators</th>
</tr>
</thead>
<tbody>
<tr>
<td>BASF361</td>
<td>Ludwig-shafen (Polit plant)</td>
<td>15</td>
<td>1994–1996</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Duales System Deutschland</td>
</tr>
<tr>
<td>Agilyx376</td>
<td>Tigard (Polit plant)</td>
<td>3</td>
<td>Operational since 2018</td>
<td>Stirred tank reactor</td>
<td>PS</td>
<td>Styrene monomer</td>
<td>Toyo, Exxon Mobil, Braskem, AmSty, Lucite, and NexiChem</td>
</tr>
<tr>
<td>Quantafuel367,381</td>
<td>Skive (Polit plant)</td>
<td>3</td>
<td>Operational since 2017</td>
<td>Fluidized bed reactor</td>
<td>HDPE, LDPE, PP, PS, and PET</td>
<td>Liquid oil, non-condensable gas, and carbon rich ash. The liquid oil will be used to produce virgin-quality plastic</td>
<td>BASF, VITOL, and VITTI</td>
</tr>
<tr>
<td>Pryme353,380</td>
<td>Amsterdam</td>
<td>100</td>
<td>Under construction and plan to operate in 2023–2024</td>
<td>Fluidized bed reactor</td>
<td>PS, PE, and PP</td>
<td>Kerosine, naphtha, and wax followed with steam cracking to obtain petroleum products</td>
<td>Shell</td>
</tr>
<tr>
<td>Brightmark339,373,381</td>
<td>Ashley</td>
<td>100</td>
<td>Partially operational since 2021</td>
<td>Auger reactor</td>
<td>PET, HDPE, PVC, LDPE, PP, and PS</td>
<td>Ultra-low sulfur diesel, naphtha, and wax</td>
<td>BP, Chevron, clean fuel Partners, and Northeast Indiana Solid Waste Management District</td>
</tr>
<tr>
<td>Plastic energy354,366</td>
<td>Seville (Pilot plant)</td>
<td>5</td>
<td>Operational since 2017</td>
<td>Stirred tank reactor</td>
<td>LDPE, HDPE, PP, PS</td>
<td>Diesel and naphtha followed with steam cracking to obtain raw materials to make new plastic</td>
<td>SABIC, ExxonMobil, and freepoint Eco-Systems</td>
</tr>
<tr>
<td></td>
<td>Le Havre</td>
<td>25</td>
<td>Design phase planned to start up in 2023</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Almeria</td>
<td></td>
<td>Operational since 2014</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Freepoint Eco-Systems374</td>
<td>Houston</td>
<td>33</td>
<td>Plan to operate in Mid-2024</td>
<td>Fluidized bed reactor</td>
<td>LDPE, HDPE, PP, PS</td>
<td>Synthetic oil</td>
<td>Plastic energy, and TotalEnergies</td>
</tr>
<tr>
<td></td>
<td>Obetz</td>
<td>90</td>
<td>Plan to operate in 2023</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shell377</td>
<td>Pulau Bukom</td>
<td>50</td>
<td>Under construction and plan to operate in 2022</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Pryme</td>
</tr>
<tr>
<td>ExxonMobil374</td>
<td>Baytown</td>
<td>30</td>
<td>Under construction and plan to operate in end-2022</td>
<td>Fluidized bed reactor</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Cyclyx, agilyx, and plastic energy</td>
</tr>
</tbody>
</table>
5.2 Catalytic thermal decomposition of plastics

5.2.1 Catalysts. Numerous catalysts have been explored for catalytic pyrolysis with entire review papers written on this topic. A large number of catalytic pyrolysis studies focus on the use of aluminosilicate materials such as amorphous SiO$_2$-Al$_2$O$_3$, mesoporous SiO$_2$-Al$_2$O$_3$, and zeolites. Aluminosilicates offer the unique advantage of having tunable acidity which can be used to direct the product selectivity of catalytic pyrolysis transformations. By increasing the alumina content – and thus acidity – within these materials, improved selectivities towards gas fractions can be achieved, while lower acidity can yield greater liquid fractions. Aluminosilicates can increase product quality, and reduce residue production as compared to thermal degradation. Amorphous SiO$_2$-Al$_2$O$_3$ lacks a pore network that could induce higher control over product selectivity. Mesoporous catalysts have been explored for catalytic pyrolysis to increase mass-transfer. The presence of mesopores allows for improved diffusion of the plastic molecules, which accelerate the initial rate of degradation and thus enhance plastic cracking. Both pure silica (kanemite-derived folded silica, FSM, MCM-41) and aluminosilicate (Al-MCM-41 and SBA-15) mesoporous structures have been explored for the degradation of PSW with varying success. Pure silica mesoporous materials were found to yield higher quantities of liquid products – with higher selectivity towards kerosene and diesel fractions – during the catalytic pyrolysis of PE and PP as compared to thermal degradation. It was found that the siloxane oxygens lining the channels of the mesopores could interact with the hydrogen atoms of the polymers to induce cracking. The interaction allows for liquid oil product selectivity to be increased at the expense of heavy oil and wax fractions. Additionally, the mild acidic character of the pure silica materials was found to enhance the cracking rate of polymers as compared to thermal degradation while also inhibiting extensive secondary cracking. As such, the all-silica mesoporous catalysts were found to be coke-resistant throughout multiple runs.

For mesoporous aluminosilicates, the plastic decomposition product distribution was dependent on the acidity of the material. Low-acidity materials (Al-MCM-41, Al-SBA-15) have high resistance to coking and increased cracking activity compared to all-silica materials. The additional acidity induced by the presence of alumina sites facilitates plastic cracking and can yield higher selectivity towards lighter hydrocarbons. For example, the product distribution for a PP/PE mixture was shifted towards C$_6$–C$_{12}$ products for Al-MCM-41, while Al-SBA-15 yielded high-quality diesel oil (high selectivity towards C$_{13}$–C$_{20}$ fraction, Fig. 20). However, both Al-MCM-41 and Al-SBA-15 were found to degrade a mixed plastics feed (HDPE, LDPE, PP, PE, and PS) to a similar product distribution of near equal amounts of C$_1$–C$_5$, C$_6$–C$_{12}$, and C$_{13}$–C$_{40}$ hydrocarbons. The large pore size of mesoporous aluminosilicates allows for improved intercalation of polymer molecules to Brønsted acid sites but prevents the moderately cracked products from undergoing re-adsorption and secondary reactions that result in extensive cracking. However, it was found that the promising cracking ability of mesoporous materials was only effective with polyolefin and polyaromatic PSW – polymers with other heteroatoms form coke which causes catalyst deactivation.

While mesoporous aluminosilicates were found to be capable of producing favorable product distributions with limited amount of coke, zeolites have still been extensively explored for the catalytic pyrolysis of plastic waste. Zeolite Y was found to have high cracking ability due to its high accessibility to active sites, but the larger micropores and high acidity were found to facilitate condensation and secondary reactions leading to fast deactivation. Similarly, ZSM-5 has high cracking activity for polyolefins due to its strong acidity. The microporous network within the ZSM-5 structure is ideal for the aromatization of cracked intermediates, however this precludes coke formation and fast deactivation. β-Zeolite yields higher gas fractions at the expense of liquid products because it is highly active for secondary reactions. Additionally, β-zeolites were found to produce high quantities of residue and wax. Natural zeolites were found to have moderate cracking ability with higher selectivity towards the liquid fraction, but its efficiency was strongly correlated to the plastic/catalyst ratio. Given the uncontrolled structure of natural zeolites, the materials contain impurities and there is little consistency amongst the materials. As such, there is high variability in the effectiveness of natural zeolites. Lastly, doped zeolites have been explored to
enhance the acidity of the material.436,443–452 Doped zeolites were found to yield high selectivity towards olefinic products and were active for the removal of heteroatoms such as Br. One great challenge of the catalytic pyrolysis of plastic waste with zeolites is catalyst deactivation due to secondary coking reactions and limited diffusion of polymeric molecules into the microporous network.

Clays are another form of aluminosilicate that are of interest due to their weaker acidity, wide availability, and ability to produce medium distillate products such as diesel fuel.409,453–456 Clays are composed of silica and alumina or magnesium (or both) and have a macroporous structure which induces high coke resistance and moderate polymer cracking to heavier products. Structured clays, such as pillared clays (PILC), can include heteroatoms like metal oxides (Al, Fe, Ti, and Zr) to improve the textural properties and increase catalyst acidity.453–456 As such, PILCs have been found to have improved catalytic performance for HDPE and PE, as well as improved regeneration abilities. Clays, however, have lower activity due to the macroporous nature of the material, resulting in reduced contact of the feedstock with the weakly acidic sites within the structure.

Spray dried catalysts (typically Geldart Particle size A) are used in fluidized bed reactors because they are easier to flow in the fluidized bed. Spray dried catalysts (Fig. 21) are a mixture of clays or binders, zeolites, and silica-alumina matrices.457–463 Spray dried catalysts can have a bimodal pore structure with micropores in the zeolite framework and mesopores in the silica-alumina matrix. It was recently determined by Weckhuysen and coworkers that, in the conversion of PP, the fluidized catalytic cracking (FCC) matrix was responsible for aromatization of cracked products and the zeolite domain was responsible for coking.457 Zeolite and silica alumina catalysts can undergo structural changes during the reaction due to water vapor (dealumination) and impurities from the feed building up on the catalyst surface. Thus, in a circulating fluidized bed reactor, such as in fluid catalytic cracking, it is important to study the “equilibrium catalyst”. It is the equilibrium catalyst which is the actual catalyst in the reactor. Studies of the effect of the equilibrium FCC catalyst on PP conversion found the same aromatic content as fresh FCC catalysts with reduced coking. The zeolite domains in an equilibrium catalyst were found to be inaccessible – resulting in reduced coking – and the FCC matrix was found to have enhanced cracking activity due to leached metal deposits. This work by Weckhuysen and co-workers suggests that the strong acidity and microporous structure of the zeolite domain within an FCC catalyst is not necessary for the aromatization of polyolefin feedstocks.457 Further, the presence of zeolite domains may even be detrimental to the effectiveness of FCC catalysts for the catalytic pyrolysis of plastic waste.

Like FCC catalysts, multi-catalyst systems are zeolites mixed with MCM-41, NiO, muds, clays, salts, or metal silicates.456,464–468 The systems are typically formed by either the mechanical mixing of the two separate catalysts via mortar and pestle or by layering the catalysts within the bed. As with other catalytic pyrolysis systems, the presence of the catalysts reduces the required temperature for polymeric degradation as compared to thermal pyrolysis.464,467 The use of muds and clays with zeolites – such as a combination system of zeolite Y, metakaolin, aluminum hydroxide, and sodium silicate – has been found to improve the selectivity toward liquid products for a mixed plastic waste feed stream as compared to thermal pyrolysis.415,469,470 Additionally, red mud and calcium hydroxide was found to reduce chlorine content in the products. The combination of zeolites and MCM-41 has been found to induce synergistic effects on plastic degradation, where the MCM-41 provides enhanced surface area, and the zeolite provides strong acidity.464 For zeolite/MCM-41 systems, diffusional limitations were observed for the direct catalytic pyrolysis of HDPE. In two-stage pyrolytic-catalytic cracking with HZSM-5/MCM-41, a high yield of aromatic gasoline range products was formed.464 Similarly, HY-zeolite/NiO was explored in a two-stage reactor.466 The addition of NiO was found to increase gas selectivity at the expense of liquid products and reduce coke formation as compared to HY-zeolite alone. NiO was speculated to promote primary degradation of larger polymer fragments into radicals preventing coke deposition.466 In total, composite systems can induce favorable synergistic effects that can result in a narrowed carbon number distribution of products as compared to thermal pyrolysis, however the favorable effects are highly dependent on the catalyst mixture ratio. Further, diffusional limitations are increased in composite systems given the physical mixture of varying pore networks and may require two-stage pyrolysis-catalysis.

Like composite systems, hierarchical and core–shell catalysts combine the advantageous properties of various materials into one system. Unlike composite systems, hierarchical and core–shell catalysts are contained within one structure rather than in a heterogeneous mixture or across multiple reactors. Many of the studies to date of hierarchical or core–shell cata-

![Fig. 21. Structure of a spray dried FCC catalyst. Reproduced from ref. 457 with permission from Wiley-VCH, copyright 2021.](Image)
lysts for plastic waste deconstruction combine the micropores of zeolites with secondary mesoporosity.\textsuperscript{445,446,449,450,471–481} The goal is to allow for enhanced accessibility where faster intra-crystalline diffusion and lower steric hindrances occur. Catalysts such as these have been found to have higher resistance to deactivation and deactivation compared to traditional microporous zeolites.\textsuperscript{479} Further, the secondary porosity of such materials renders improved gas and aromatic hydrocarbon production than their microporous counterpart. Hierarchical and core–shell materials are capable of supporting metals to enhance acidity and therefore activity.\textsuperscript{446,449,450,471–476,478,479} However, these materials require refined and detailed synthesis – making scale up difficult – and the dual porosity must be well defined otherwise deactivation \textit{via} coking will occur.

Silica and aluminosilicates have dominated catalytic pyrolysis because of their porous networks and acidity, however other metal oxides such as ZrO\textsubscript{2}, MgO, and CaO have shown interesting results.\textsuperscript{482–494} In particular, sulfated zirconia – despite suffering from coke deactivation – has shown high cracking activity and isomerization, esterification, and hydrocracking ability.\textsuperscript{485,487,492,494} By sulfating the metal oxide, the acidity of the material is enhanced, resulting in improved reactivity. Further, studies have explored promoting sulfated ZrO\textsubscript{2} with platinum and found that the material – in a two-step pyrolysis-catalytic hydrocracking process – can enhance the hydrocracking of LDPE into liquid products as compared to the non-doped material.\textsuperscript{483} Other metal oxides such as MgO and CaO have been shown to reduce activation barriers and enhance selectivity toward phenolic compounds for PC catalytic pyrolysis, as well as reduce production of light gases (H\textsubscript{2}, CO\textsubscript{2}, CO, and methane) for PC and LDPE degradation.\textsuperscript{482,484,488,489,491} It was determined that the basicity of CaO allowed for the reaction of CO\textsubscript{2} in the pyrolysis vapors, resulting in the production of CaCO\textsubscript{3}, while MgO was found to interact with H\textsubscript{2} and utilize the molecule for hydrogenation reactions.\textsuperscript{488,493} However, CaO and MgO produce significant amounts of solid residue, where the carbonaceous char inhibited the movement of free radicals during the cracking reaction, preventing propagation and degradation of the plastic molecules.\textsuperscript{484}

Outside of metal oxide catalysts, activated carbon has been tested for plastic catalytic pyrolysis but has been found to only minimally change product distribution.\textsuperscript{396,481,495–497} Further, activated carbon has variable acidity – which can be modulated by the addition of different acid groups (–OH, –OOH, and –P=O) – that dictates the product distribution.\textsuperscript{396,497} Weakly acidic activated carbons have been found to have high selectivity towards jet-fuel range alkanes within the liquid fraction and strongly acidic carbons (with P functionality) have improved selectivity towards jet-fuel range aromatics.\textsuperscript{481} Additionally, metal-doped carbons have been utilized for PSW decomposition due to the high thermal stability of the material and the cracking ability of transition metals. For example, Pt/C and Pd/C were found to decompose PET into monocyclic compounds within the liquid fraction by improving decyclization activity and promoting thermal free radical cracking mechanisms.\textsuperscript{498} Pt/C was also found to crack the carbonaceous compounds that formed during catalytic pyrolysis of PET, further inhibiting the production of undesirable products.\textsuperscript{498}

Like many of the catalysts described above, carbonates have been found to increase the rate, reduce operational temperature, and reduce residence time of plastic decomposition. However, carbonates are prone to decomposition under thermal treatment, resulting in the formation of oxide compounds and CO\textsubscript{2}. MgCO\textsubscript{3}, CaCO\textsubscript{3}, and CuCO\textsubscript{3} were all found to yield largely diesel fuel components in the liquid fraction for the catalytic pyrolysis of HDPE.\textsuperscript{499–502} At the same time, gas yields were increased by 10–15\% due to the decomposition of the material under reaction conditions.\textsuperscript{499} The basic carbonates were hypothesized to initiate the plastic cracking mechanism \textit{via} a different step than the acidic catalysts previously described, where the carbonates will accept a proton from a plastic defect site to create a carboxonium chain rather than the catalyst donating the proton to form a carbonium molecule. The carboxonium chain will undergo β-scission to form lower carbon number alkanes and alkenes, then further reaction will occur through carbonium ion attacks or interactions with the catalyst basic sites.

Ashes from plastic incineration plants have also proven to be effective in reforming plastic pyrolysis vapor using a pyrolysis-reforming reactor.\textsuperscript{503} Incineration ashes contain ppm levels of metals due to metal evaporation and adsorption to the surface of ash particles during incineration. When used for in-line upgrading of plastic vapor, flying ash increased the BTX content of liquid product compared to non-catalytic upgrading. The Fe content of ash promoted the formation of monoaromatics while inhibiting the formation of polynaromatics.\textsuperscript{455,504} Although the use of ash as a catalyst is attractive as it is a waste, the catalytic performance of this material is still inferior compared ZSM-5. Furthermore, pre-treatment of ashes is required to remove impurities that prevent access to active sites.

\textbf{5.2.2 Reactor design.} Two main approaches for catalytic pyrolysis have been proposed for plastic conversion: a single-stage catalytic-pyrolysis (\textit{in situ}) process and a two-stage pyrolysis-catalysis (\textit{ex situ}) process. In the single-stage process, the plastic waste and catalyst are directly mixed in a reactor (usually a fluidized bed reactor, Fig. 22) and heated to the degradation temperature. In the two-stage pyrolysis-catalysis process, the plastic waste is first thermally pyrolyzed and then the pyrolysis vapors are passed over a catalyst bed (which could potentially be a fixed bed reactor).\textsuperscript{505,506} Fluidized beds are the most commonly used industrial reactors for processing solid materials. They have high rates of heat transfer and therefore can be scaled to larger industrial reactors.\textsuperscript{507} Circulating fluidized bed reactors are commonly used in industry as they circulate the catalyst between a reaction and regeneration zone, where the coke on the catalyst can be continuously removed by oxidation. In contrast, fixed bed reactors must be shut down when the catalyst is regenerated leading to a decrease in capital efficiency. Fixed-bed catalytic reactors are preferred.
Liquefaction typically operates at temperatures between 200–450 °C under high pressure to convert the plastic feedstock into a liquid oil in the liquid phase. Heterogenous or homogenous catalysts are sometimes added.\textsuperscript{513,514} When liquefaction occurs in a solvent, it is called solvent liquefaction. Fuel liquids or monomers can be obtained from the liquefaction of plastics.\textsuperscript{513,515–523} During liquefaction, pressurized nitrogen or other gases keep the pressure high, helping maintain the products in the liquid phase. Pressurized hydrogen enhances polymer chain cleavages, caps the chain ends with hydrogen, and produces liquids with higher H/C ratios and/or low boiling point products.\textsuperscript{521} Various organic solvents, including hydrocarbons, alcohols, and water, are used to enhance the reaction rate and lower the activation energy of plastic decomposition, in addition to reducing unfavorable secondary reactions for repolymerization.\textsuperscript{524–527} Chemically-compatible solvents can facilitate product removal and dissolution, suppressing intermolecular coupling reactions.\textsuperscript{520} Hydrogen-donating solvents can enhance product dissolution and stabilize intermediate products.\textsuperscript{518}

When performed under supercritical states of solvents, solvent liquefaction also takes advantage of drastically increased reaction rates and enhanced solubilization capability of the polar and non-polar products.\textsuperscript{518,528} In the case of hydrothermal liquefaction (HTL), water is a low-cost green solvent. Table 14 summarizes some of the previous work on plastic liquefaction processes, which are grouped into liquefaction using pressurized gases, HTL, and liquefaction by organic solvents with the majority of this work in stirred-tank batch reactors.

### 6. Liquefaction of plastics

Liquefaction typically operates at temperatures between 200–450 °C under high pressure to convert the plastic feedstock into a liquid oil in the liquid phase. Heterogenous or homogenous catalysts are sometimes added.\textsuperscript{513,514} When liquefaction occurs in a solvent, it is called solvent liquefaction. Fuel liquids or monomers can be obtained from the liquefaction of plastics.\textsuperscript{513,515–523} During liquefaction, pressurized nitrogen or other gases keep the pressure high, helping maintain the products in the liquid phase. Pressurized hydrogen enhances polymer chain cleavages, caps the chain ends with hydrogen, and produces liquids with higher H/C ratios and/or low boiling point products.\textsuperscript{521} Various organic solvents, including hydrocarbons, alcohols, and water, are used to enhance the reaction rate and lower the activation energy of plastic decomposition, in addition to reducing unfavorable secondary reactions for repolymerization.\textsuperscript{524–527} Chemically-compatible solvents can facilitate product removal and dissolution, suppressing intermolecular coupling reactions.\textsuperscript{520} Hydrogen-donating solvents can enhance product dissolution and stabilize intermediate products.\textsuperscript{518}

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#### 6.1 Products

6.1.1 Liquefaction using pressurized gases. Direct liquefaction of plastics using pressurized gases has been investigated since the 1990s, mainly applied to producing synthetic fuel oils from individual non-polar polymers or co-mingled waste plastics. Increasing the reaction pressure during liquefaction enhances liquid production and reduces gas formation. Hydrogen can be added to the liquefaction process to increase product yields by hydrogenating the products in a single step. Williams et al.\textsuperscript{523} reported an increase in the oil yields from 71% to 77% for PS and 15% to 27% for PET by switching from a nitrogen to a hydrogen atmosphere, which almost entirely contributed to the decreased solid yields. In comparison, there were no significant increases in the oil yields from PE and PP by switching from a nitrogen atmosphere to hydrogen, as shown in Table 14. Liquefaction under pressurized gases also promotes the formation of aromatics from PE and PP. While PE pyrolysis at atmospheric pressure nitrogen almost exclusively produced aliphatic hydrocarbons, the oil produced using pressurized nitrogen contained 16.4% aromatics.\textsuperscript{523} The increased gas pressure promoted secondary reactions of PE decomposition, such as saturation, cyclization, and isomerization.\textsuperscript{529} Cyclization reactions were also promoted during hydrogen liquefaction, albeit less so than in liquefaction under nitrogen gas.\textsuperscript{523} The work of Williams et al. also showed that hydrogen liquefaction increased the selectivity of single-ring aromatics for PS,\textsuperscript{523} because the hydrogen atmosphere...
<table>
<thead>
<tr>
<th>Type of plastics</th>
<th>Temp. (°C)</th>
<th>Pressure (MPa)</th>
<th>Time (h)</th>
<th>Concentration (%)</th>
<th>Solvent/gas/catalyst</th>
<th>Products</th>
<th>Oil yield (wt%)</th>
<th>Year</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>PE, PP, PS (continuous reactor)</td>
<td>400</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>w/o w/o ZSM-5</td>
<td>Majority aromatics, some aliphatic liquids and gases</td>
<td>85% (PE, PP); 90% (PS)</td>
<td>1992</td>
</tr>
<tr>
<td>MDPE, HDPE, PP, PET or mix</td>
<td>420–450</td>
<td>5</td>
<td>1</td>
<td>66% plastics in tetralin/waste oil</td>
<td>H2; w/o w/o 1 wt% HZSM-5/ferrihydride catalyst</td>
<td>Hydrocarbon oil and gases</td>
<td>w/o to with catalyst; 11 to 96% (HDPE); 83 to 96% (PP); 33 to 93% (MDPE)</td>
<td>1994</td>
<td>516</td>
</tr>
<tr>
<td>Mix of HDPE, LDPE, PET and PS</td>
<td>400–440</td>
<td>5.6</td>
<td>0.5–2</td>
<td>50% plastics in tetralin, decalin, dodecane, C12–C20 alkanes</td>
<td>H2; 10–20 wt% HZSM-5/FCC catalyst</td>
<td>Hydrocarbon oil and gases</td>
<td>56.2–7.5% conversion (mixture); 90–100% conversion (individual plastics)</td>
<td>1996</td>
<td>513</td>
</tr>
<tr>
<td>PS and SBR</td>
<td>350–450</td>
<td>3.45–17.23</td>
<td>0.25–2</td>
<td>1–5 wt% Fe3O4/Sm2O3 and ZrO2/SO43–</td>
<td>—</td>
<td>Aromatics (PS); aromatics and C1-C5 paraaffins/cycloparaffins, C5–C10 hydrocarbons and gases</td>
<td>80.3% (PS); 72% (SBR)</td>
<td>1996</td>
<td>517</td>
</tr>
<tr>
<td>LDPE, PET, PVC</td>
<td>420–440</td>
<td>5.5</td>
<td>0.25–1</td>
<td>70% in tetralin</td>
<td>Hydrogen</td>
<td>Light, medium and heavy oils</td>
<td>&gt;90% (all plastics)</td>
<td>1996</td>
<td>518</td>
</tr>
<tr>
<td>MDPE, HDPE, PP</td>
<td>&gt;420</td>
<td>0.68–5.5</td>
<td>0.3–1</td>
<td>30–50% in tetralin; H2 or N2; HZSM-5 or Al2O3–SiO2/ferrihydrite</td>
<td>n-Octadecane; H2; 1–2 wt% Fe3O4/Sm2O3 and ZrO2/SO43–</td>
<td>Gasoline range paraaffins as major products</td>
<td>&gt;90%</td>
<td>1996</td>
<td>519</td>
</tr>
<tr>
<td>HDPE, PP, PB</td>
<td>350–450</td>
<td>3.5–13.8</td>
<td>0.5–3</td>
<td>—</td>
<td>Hydrogen</td>
<td>Light and heavy oils</td>
<td>up to 60%</td>
<td>1997</td>
<td>520</td>
</tr>
<tr>
<td>PE and PP</td>
<td>415–455</td>
<td>1.4</td>
<td>0.5–1</td>
<td>—</td>
<td>H2; 1–5 wt% of HZSM-5 and others</td>
<td>Higher gasoline range oil with catalysts</td>
<td>up to 85%</td>
<td>1998</td>
<td>521</td>
</tr>
<tr>
<td>Post-consumer plastic (PCW) mixture</td>
<td>500</td>
<td>0.79</td>
<td>0.5</td>
<td>Hydrogen</td>
<td>Hydrocarbon oil and gases with high concentrations of alkanes and single-ring aromatics</td>
<td>Calculated mix vs. PCW</td>
<td>2007</td>
<td>523</td>
<td></td>
</tr>
<tr>
<td>PE, PP, PS, PVC, and PET (standalone and mixed)</td>
<td>500</td>
<td>1</td>
<td>1</td>
<td>—</td>
<td>Nitrogen and hydrogen</td>
<td>None</td>
<td>—</td>
<td>1999</td>
<td>524</td>
</tr>
</tbody>
</table>

**Hydrothermal liquefaction**

- **HDPE**
  - 425 Corresp. to T
  - 0.16–3
  - None
  - Light and medium oil
  - 90.2% (120 min)
  - 1999 | 524 |
- **SBR**
  - 450–686
  - <1000
  - 0.022–0.5
  - 5/10% H2O2
  - Light and medium HCs
  - 25.9–66%
  - 2002 | 530 |
- **Hydrothermal liquefaction**
  - **HDPE**
    - 450–480
    - Corresp. to T
    - 0.016–0.5
    - 25–50
    - None
    - Light and medium oil
    - 91.4% (460 °C, 1 min)
    - 2004 | 531 |
  - **PVC**
    - 200–600
    - 1.6–55.7
    - 1
    - 0.1–2
    - None
    - Low-molecular weight aromatic and aliphatic compounds
    - C7–C11, C12–C18, C19–C24, >C24
    - 179 ppm (300 °C), 396 ppm
    - (400 °C)
    - 79%
    - 2004 | 532 |
- **HDPE (continuous reactor)**
  - 530
  - 25
  - 0.02–0.07
  - —
  - None
  - Chlorine content after NaOH-based dechlorination
  - Chlorine content after NaOH-based dechlorination
  - O1% solid% nil/50.8 (PBT), 99.8/nil (PC), /68.5 (PET), 48-nil
  - PMMA), 13.7/8.1 (POM), 78.9/8.8 (PPO), 35.4/2.9 (PVA), 80.8/1.2
  - (SB)
  - 40–120 ppm in oil (negl.)
  - 2011 | 534 |
- **Model mix of PE, PP, PS and PVC**
  - 200–400
  - 1–5
  - 100–200
  - Glass powder additive
  - None
  - Ethyl benzene (51.3 wt%), toluene (14 wt%) and other polycyclics (490 °C per 1 h)
  - 80% naphtha
  - Maximum carbon liquefaction rate of 77 wt% (490 °C, 1 h)
  - 91% (2 h per 425 °C; 1 h per 450 °C)
  - 2019 | 535 |
- **High impact PS (HIPS)**
  - 350–550
  - 30
  - 0.12–1
  - 1–9
  - None
  - Ethyl benzene (51.3 wt%), toluene (14 wt%) and other polycyclics (490 °C per 1 h)
  - 80% naphtha
  - Maximum carbon liquefaction rate of 77 wt% (490 °C, 1 h)
  - 91% (2 h per 425 °C; 1 h per 450 °C)
  - 2019 | 536 |
- **PP**
  - 425–450
  - 23
  - 0.5–4
  - —
  - None
  - Ethyl benzene (51.3 wt%), toluene (14 wt%) and other polycyclics (490 °C per 1 h)
  - 80% naphtha
  - Maximum carbon liquefaction rate of 77 wt% (490 °C, 1 h)
  - 91% (2 h per 425 °C; 1 h per 450 °C)
  - 2019 | 537 |
<table>
<thead>
<tr>
<th>Type of plastics</th>
<th>Temp. (°C)</th>
<th>Pressure (MPa)</th>
<th>Time (h)</th>
<th>Concentration (%)</th>
<th>Solvent/gas/catalyst</th>
<th>Products</th>
<th>Oil yield (wt%)</th>
<th>Year</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>ABS, PA6, PA66, PET, Epoxy, PC, PUR, HDPE, PVC, LDPE, PP, PS, PP, PS, PC and PET</td>
<td>350–450</td>
<td>25+</td>
<td>0.5–1</td>
<td>0.06–0.35</td>
<td>None</td>
<td>None</td>
<td>32% (PP, 425 °C, 30 min), 16% (PET, 450 °C, 30 min), 86% (PS, 350 °C, 30 min), and 60% (PC, 425 °C, 30 min).</td>
<td>2020</td>
<td>528</td>
</tr>
<tr>
<td>HDPE</td>
<td>350–450</td>
<td>Corresp. to T</td>
<td>0.03–1</td>
<td>5</td>
<td>None</td>
<td>None</td>
<td>57.7</td>
<td>2020</td>
<td>539</td>
</tr>
<tr>
<td>LDPE, HDPE</td>
<td>380–450</td>
<td>Corresp. to T</td>
<td>0.25–4</td>
<td>20</td>
<td>1% acetic acid</td>
<td>None</td>
<td>Alkanes, alkenes, cycloalkanes, aromatics, and negligible alcohols</td>
<td>2020</td>
<td>540</td>
</tr>
<tr>
<td><strong>Hydrothermal liquefaction</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mix of LLDPE and PP</td>
<td>400</td>
<td>25</td>
<td>1</td>
<td>16.67</td>
<td>None</td>
<td>None</td>
<td>Paraffins, olefins, cyclics and aromatics</td>
<td>2021</td>
<td>541</td>
</tr>
<tr>
<td>A waste mixture containing polyolefins, PET, nylon, PVAc, and cellulose</td>
<td>340</td>
<td>Corresp. to T</td>
<td>5</td>
<td>10</td>
<td>2% NaOH</td>
<td>None</td>
<td>Aromatics, oxygenated compounds, benzoic acid, phenols, caprolactones, fatty acids, and no polyolefin-derived products</td>
<td>2021</td>
<td>542</td>
</tr>
<tr>
<td><strong>Liquefaction with other supercritical solvents</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HDPE, PP and HDPE/PP mix</td>
<td>450–470</td>
<td>Corresp. to T</td>
<td>1</td>
<td>10%</td>
<td>Acetone</td>
<td>Aliphatic hydrocarbons</td>
<td>Oil yield: 88% (HDPE); 96% (PP); 90% (mix)</td>
<td>2021</td>
<td>543</td>
</tr>
<tr>
<td>PS</td>
<td>370</td>
<td>Corresp. to T</td>
<td>1.5</td>
<td>7–10%</td>
<td>n-Hexane</td>
<td>Styrene (primary); toluene and ethyl benzene (secondary)</td>
<td>90% conversion (380 °C/15 min)</td>
<td>2001</td>
<td>526</td>
</tr>
<tr>
<td>PS</td>
<td>340–420</td>
<td>Corresp. to T</td>
<td>0.25–1</td>
<td>—</td>
<td>Methanol</td>
<td>Styrene, toluene, ethylbenzene, dimer, and other 2-ringed aromatics</td>
<td>92% conversion (380 °C/15 min)</td>
<td>2008</td>
<td>527</td>
</tr>
<tr>
<td>HDPE; HDPE + spirulina microalgae</td>
<td>340</td>
<td>Corresp. to T</td>
<td>—</td>
<td>6–14%</td>
<td>Ethanol</td>
<td>Aliphatic hydrocarbons</td>
<td>7.55% (PE oil); ~50% (HDPE + spirulina)</td>
<td>2012</td>
<td>544</td>
</tr>
<tr>
<td>PS</td>
<td>350 (SC ethanol); 370 (HTL)</td>
<td>Corresp. to T</td>
<td>0.25–1.25</td>
<td>25–400%</td>
<td>Ethanol; water</td>
<td>Allyls, alkenes, and aromatics (SC ethanol)</td>
<td>Oil yield: 84.74% (SC ethanol); 78.3% (HTL)</td>
<td>2020</td>
<td>544</td>
</tr>
<tr>
<td>HDPE + sugarcane bagasse mix</td>
<td>240–280</td>
<td>Corresp. to T</td>
<td>1–1.5</td>
<td>5–10%</td>
<td>Ethanol</td>
<td>Aliphatic hydrocarbons</td>
<td>32.3% (bio-oil yield)</td>
<td>2020</td>
<td>545 and 546</td>
</tr>
</tbody>
</table>
enhanced both chain cleavage and secondary saturation reactions compared to nitrogen.

Solid acid catalysts are often used during the hydrogen liquefaction of plastics. In general, the addition of catalyst lowers reaction temperature, increases conversion, and produces oils with higher light fractions. Shabtai et al. reported that conversion of HDPE during hydrogen liquefaction at 350 °C increased from nearly zero without a catalyst to 30% with a 17 wt% of SO4/ZrO2 catalyst, and 64.7% using a 33% Pt/SO4-Al2O3 catalyst.520 Much lower metal loadings were used in other related studies. For example, Zmierczak et al. found catalytic hydrogen liquefaction of PS at 375 °C increased the conversion from 59.4% without a catalyst to 70.5% with a 5 wt% Fe2O3/SO42− catalyst and to 80.3% with a 5 wt% ZrO2/SO4 catalyst. Other catalysts were also evaluated for hydrogen liquefaction, as shown in Table 14. Shah et al. converted post-consumer plastic waste using 1 wt% HZSM-5 zeolite, a ZrO2-WO3 catalyst, nanoscale ferrihydrite treated with citric acid, ferrihydrite containing 5% Mo, a SiO2-Al2O3 binary oxide, and two TiO2-SiO2 binary oxides.522 They reported that while there were no significant differences in the oil yields, the presence of ZSM-5 reduced the boiling point distribution of the oil products resulting in the most gasoline-like products. Similar results were also observed by Feng et al.519

Chemically compatible or hydrogen donor solvents have been used to aid the hydrogen liquefaction of plastics. For example, adding n-octadecane during Pt/SO42−/ZrO2 catalyzed hydrogen liquefaction of HDPE at 350 °C increased the conversion from 64.7% without solvent to 92.5% with the solvent.520 The plastic conversion in the solvent without catalyst was only 2%.520 Tetralin has been studied as a hydrogen donor for coal liquefaction. As shown in Table 14, Taghiei et al. reported that tetralin-aided hydrogen liquefaction of PE and PP increased the oil yield from 11% without a catalyst to 83% with a catalyst.5 In contrast, the oil yield only increased from 62% to 64% for PET. Since PE and PP-derived products are non-polar hydrocarbons whereas PET-derived products are oxygenated polar products, chemical compatibility between the solvent and plastic decomposition products seemed to play a critical role in their study. Murty et al.518 reported that PVC absorbs tetralin and that this promoted PVC decomposition. Feng et al.519 reported that liquefaction of co-mingled waste plastics in the absence of solvent resulted in higher oil yields than the liquefaction in tetralin or waste motor oil, regardless of whether catalyst was added or not.

The liquefaction of plastics is highly sensitive to reaction temperature. Zmierczak et al.517 reported an increase in the distillable product yield from 71% at 350 °C to 93% at 450 °C during catalytic hydrogen liquefaction of PS. Increasing temperature also increased the formation of single-ring aromatics (i.e., benzene and alkylbenzenes) by further decomposing multi-ring products (e.g., diphenyl or triphenyl alkenes).517 However, increasing the temperature during catalytic liquefaction can also promote the cracking of liquid products to gases. Compared to catalytic liquefaction, reaction temperature seemed to have a greater effect on non-catalytic liquefaction. Feng et al.519 showed that the oil yield from mixed post-consumer plastics at 430 °C was only about 25% with no catalyst and 60–63 wt% with 1 wt% HZSM-5 or Al/Si as the catalyst. However, oil yields of over 90% were obtained at 460 °C both with and without catalysts.519 For all cases, gasoline and kerosene fractions increased with increasing temperature.

In general, increasing reaction time increased both conversion and secondary reactions. Shabtai et al.520 reported that increasing reaction time during catalytic hydrogen liquefaction of HDPE at 450 °C improved the conversion from 82.5% to 100% but also reduced the oil yield from 92.8% to 60.2% due to increased cracking at prolonged reaction times. A similar trend was also observed with PP.520 Increasing the reaction time also enabled the complete depolymerization of PS to single ring aromatics and promoted secondary cyclization of the primary monomers to form more naphthalene and indane as reported by Zmierczak et al.517 While the effect of hydrogen pressure is usually less significant than reaction temperature or time, it can improve both conversion and oil quality.519,520 For example, Murty et al.518 reported that the oil produced from LDPE at 440 °C using 800 psi hydrogen pressure was three times less viscous than the oil obtained using 100 psi hydrogen pressure because of the conversion from heavy oil to light oil at higher pressure. Increasing pressure also promoted single-ring aromatics and suppressed secondary cyclization during catalytic liquefaction of PS.517

Feedstock heterogeneity is the most challenging problem in converting real-world plastic wastes. Williams and Slaney523 evaluated the feedstock compositional variability under comparable liquefaction in this context. They conducted non-catalytic liquefaction of a mixture containing HDPE, PP, PET, and PVC. The product yields obtained from the mixture were then compared with the theoretical product yields calculated using those obtained when their constituent polymers were independently converted under the same condition. The oil yields produced from the mixed plastics were significantly lower than the calculated yields for nitrogen and hydrogen liquefaction. The effect of feedstock preparation methods showed that the PCW prepared in Belgium by Fost Plus via air blowing separation (obtaining a low density fraction of PCW) produced much higher oil yields and lower solid residues than the non-plastic removed PCW prepared by water washing (Waste DSD).521 Plastic wastes from Waste Fost Plus were expected to contain a high fraction of low-density plastics (LDPE, PP, and PS) and a lower fraction of high-density plastics (HDPE, PVC, and PET).523 Unfortunately, the authors investigated neither the product compositions nor the possible interactions among the compositions to elaborate on the crucial results. Contaminant content in mixed plastics also had a significant detrimental impact during hydrogen liquefaction. The catalytic effect was severely hindered during Shah et al.’s study when the dirtier PCW prepared by dry separation was converted.522 The authors found no noticeable changes in either yields or boiling point distributions of the products before and after using 1 wt% of HZSM-5 or other catalysts. In contrast, adding identical amounts of catalysts to a cleaner PCW prepared by
produced. Passosa et al. showed that adding KOH catalyst to the HTL of PA could significantly increase oil yield and promote complete depolymerization in reduced residence times. The presence of the base catalyst also accelerated the production of oxygenated products soluble in water and increased the selectivity of polyaromatics and oxygenated compounds derived from plastics like PC and PURs.

HTL of non-polar plastics (polyolefins and PS) usually requires supercritical or near supercritical conditions to be effective. According to Ciuffi et al., compounds from PE, PP, and PS depolymerization were absent from the liquid products obtained for subcritical HTL of mixed plastic wastes. The lack of heteroatoms (i.e., polar reactive sites) in the polyolefins and PS prevents the plastics from depolymerizing under mild conditions. As shown in Table 14, HTL of PS has been widely studied. Kwak et al. reported the complete and nearly instantaneous conversion of PS at 400 °C. Higher temperatures or prolonged reaction times at supercritical conditions will further decompose liquefied products by promoting secondary reactions. Seshasayee et al. reported a decrease in PS-derived oil yield from 86.2% at 350 °C to 38.3% at 450 °C after 30 minutes of treatment. HTP of PS usually produces styrene monomer, ethylbenzene, α-methylstyrene, styrene dimer, styrene trimer, and other dimer/trimer associated products. Toluene and ethylbenzene were the major liquid products under supercritical reaction conditions. Bai et al. reported yields for toluene and ethylbenzene at 490 °C to reach 14% and 51.3%, respectively.

HTL of polyolefins was conducted under supercritical conditions to produce an oil similar to naphtha. Several articles have reported 85–90% oil yields from HDPE and LDPE in supercritical HTL. As given in Table 14, oil yields higher than 90% from HDPE were reported by Murty et al. and Su et al. where HTL was performed in batch reactors. In comparison, the highest yield obtained using a continuous reactor system was 79% at 530 °C (Table 14). Supercritical HTL of polyolefins yields oils with a wide range of molecular weights. A narrower distribution can be obtained by increasing temperature and reaction time. HTL of PE produces n-alkanes as the major products, followed by 1-alkenes and alkadienes.

Intramolecular β-scission and intermolecular hydrogen abstraction (H abstraction) are generally considered the primary mechanisms for polyolefins to degrade during HTL. When the PE backbone undergoes homolytic cleavage, the random scission produces free radicals with a random number of carbon atoms. These radical ends can be terminated by hydrogen abstraction from within the molecule (by β-scission) or from other molecules (by H abstraction). The ratio between 1-alkenes and n-alkanes in the product can indicate which reaction is prominent. During HTL, polymer fragments with radical ends immediately dissolve in SCW, where large amounts of excited H radicals are available. Thus, H abstraction becomes more prominent than β-scission. This is also why previous studies frequently report the 1-alkene-to-n-alkane ratio to be lower in HTL of PE than in thermal pyrolysis of PE. At extended reaction times, n-alkanes can undergo isomerization, and alkenes can undergo secondary reactions like cyclization. Gas production is also increased under these conditions due to increased cracking or recombination of short-chain free radicals. Aromatization is suppressed under HTL, thus producing less char from plastics. When HTL was conducted in a continuous or a semi-batch
reactor system, the radicals were quickly removed from the reaction region. This led to a decreased opportunity for H abstraction, producing more alkenes than n-alkanes.\textsuperscript{533,551} In addition to promoting chain cleavage, HTL can also oxidize the hydrocarbon products to form alcohols and ketones. Products such as 2-propanol, 2-butanol, 2-propanone, and 2-butane were found in the aqueous phase products obtained during HTP of polyolefins by Moriya et al.\textsuperscript{524}

As shown in Table 14, oil yields up to 91% from supercritical HTL of PP were reported using reaction temperatures up to 450 °C.\textsuperscript{537} The oil fraction consisted of olefins, paraffins, and cyclic and aromatic compounds, with 80–90% of these compounds in the naphtha range. The authors also reported that the PP-derived oil is comparable to gasoline in terms of their H/C and O/C ratios, boiling point range, carbon numbers, density, viscosity, and surface tension.\textsuperscript{537} Similar product composition from HTL of PP was also reported by others,\textsuperscript{541} presenting an opportunity for PP-derived oil to be directly used as fuel feedstocks and blendstocks without further upgrading. Zhao et al. studied co-conversion of LLDPE and PP by HTL at 400 °C and reported the oil yield from the plastic mixture to be higher than the yield produced by converting LLDPE or PP alone, as shown in Table 14.\textsuperscript{541} In this study, PE produced more heavy oil than medium or light oils whereas PP produced equal amounts of naphtha (light oil), medium, and heavy oils. In comparison, the mixtures of PE and PP produced more heavy oil than naphtha or light oil. HTL co-conversion also increased the formation of cyclic compounds while reducing paraffin production.\textsuperscript{541} Thus, relatively lower quality oil was obtained by co-conversion, although the oil yield synergistically increased.

Subcritical HTL of PVC was also investigated because hydrochloric acid, produced from the PVC, is water soluble. PVC was dechlorinated in subcritical water at 200 °C in Takeshita et al.’s work to yield polyenes as solid residue without producing any hazardous organochlorine compounds.\textsuperscript{532} When the reaction temperature increased to 250–350 °C, a mixture of low-molecular-weight aromatic and aliphatic compounds was produced. As the temperature further increased to 400 °C, the increased production of light alkanes and alkenes was also accompanied by the formation of aliphatic ketones and oxygenated benzene compounds at higher yields.\textsuperscript{532} Inside the highly acidic solution created by the PVC-derived HCl, the PVC-derived polyene can undergo cyclization, hydrolysis, oxidation, and thermal degradation.\textsuperscript{532,538} As shown in Table 14, base catalysts, such as NaOH or KOH, were added during HTL of PVC to study their effect on dechlorination. Passosa et al.\textsuperscript{538} showed that although adding KOH did not affect the oil and solid yields produced from PVC at 350 °C, it caused more chlorine to leave as Cl\textsubscript{2} gas rather than remaining in the solution as HCl.

Increasing temperature and treatment time during HTL often facilitated conversion to increase liquid and gas yields.\textsuperscript{514,524,531,539,540} However, it has been noted that high reaction temperatures and a prolonged residence time can reduce the desired product yield and increase coke formation due to over-cracking of the liquefied products.\textsuperscript{530,532,536} The feedstock-to-solvent mass ratio also affected the HTL process. Increasing the feedstock mass loading caused the liquefaction reaction rate to first increase and then decrease. Increasing the solid mass loading to beyond the corresponding optimum led to increased heat transfer limitations and decreased mobility of the macromolecular free-radical fragments in the solution.\textsuperscript{539} Such limitations favored undesired secondary reactions like repolymerization. Jin et al. also reported that increasing temperature and mass ratio would eventually result in decreased liquefaction efficiency for the same reaction time. When high mass ratios were employed at higher reaction temperatures, increased reaction rate and higher H\textsubscript{2}/H\textsubscript{2} concentration in the solution caused the monomers and oligomers of the liquefied products to undergo further cracking to form gaseous products.\textsuperscript{539} The effect of pressure during HTL of plastics is generally considered less significant than reaction temperature and time, especially for continuous and semibatch reactor systems.\textsuperscript{533} Although the pressure effect was only observed in some studies, an increase in pressure increased the ion products of the water due to increased density, subsequently bolstering the cracking and hydrolysis process.\textsuperscript{536} An increase in pressure also enables enhanced interaction among different reaction species, promoting cyclization, hydrolysis, oxidation, and thermal degradation all at the same time.\textsuperscript{532} Depending on plastic types and the availability of reactive sites, any of these reactions may take precedence.

### 6.1.3 Liquefaction with other organic solvents.

Liquefaction of plastics in supercritical hydrocarbons and alcohol solvents has also been studied, as shown in Table 14. In general, the solvents were used to enhance product dissolution, increase reaction rates, and stabilize products. However, in these cases, the solvent is a co-reactant and the conversion of the solvent must be considered. Hwang et al.\textsuperscript{526} reported that at least 90% of PS was converted in n-hexane at 370 °C, while there was almost no conversion when PS was pyrolyzed at the same temperature.

Ethanol achieves its critical states at relatively milder operating conditions, and its low dielectric constant increases the solubility of both polar and non-polar products. Furthermore, ethanol can also act as a hydrogen-donor solvent to stabilize the intermediate products and reduce repolymerization reactions. Breaking alcohol solvent at the critical state produces excessive hydrogen and hydroxyl ions, aiding the hydrogenation reactions. Ahmad et al. reported that the oil yield produced from PS conversion in ethanol at 350 °C was 84.74%, higher than 71% from pyrolysis at 500 °C, and 78.3% obtained using HTL at 370 °C.\textsuperscript{544} Rather than producing styrene as the primary monomer, the liquid produced using ethanol solvent consisted of alkyls, alkenes, and aromatics.\textsuperscript{544} Compared to PS, less liquid was produced when PE was converted in ethanol. According to Pei et al., the oil yield of HDPE in 340 °C ethanol was only 7.55%.\textsuperscript{544} However, co-liquefying PE with microalgae or lignocellulosic biomass in ethanol was found to synergistically increase the liquid yield because the oxygenated feedstock enhanced the thermal decomposition of PE in the
solvent probably by hydrogen abstraction from PE by the oxygenated biopolymers.545,546

Methanol is also a hydrogen donor solvent with low critical conditions. Shin and Bae reported 92% conversion for PS by treating with 380 °C methanol for 15 min.527 Their comparative study showed that the activation energy for degrading PS in supercritical methanol was 117.2 kJ mol$^{-1}$, lower than 132 kJ mol$^{-1}$ in supercritical $n$-hexane, 157 kJ mol$^{-1}$ with supercritical water, and 224 kJ mol$^{-1}$ in pyrolysis. The authors reported the hydrogenation of unsaturated products to be the predominant reaction in methanol. Besides donating hydrogen, methanol also directly reacted with styrene monomer and $\alpha$-methylstyrene to produce 3-phenyl propanol and 3-phenyl-1-butanol.527

6.2 Reactor designs

Lab-scale liquefaction studies have predominantly been carried out in batch reactors with few accounts of continuous reactor systems. Industrial-scale processes have incorporated both batch and continuous reactor designs. Stirred or agitated tank reactors are the most employed batch reactor designs at both lab and industry-scale. The different continuous reactor designs adopted for industrial-scale processes are continuous flow stirred tank, rotary kiln, fixed bed, and tubular reactors, with gravity or auger/extruder-based material feeding mechanisms. Several other multi-stage upgrading and direct hydrogenation liquefaction projects of waste plastics were also established by German companies like RWE-Entsorgungs AG, Hiedrierwerke Zeitz GmbH and Bergakademie Freiberg, but later terminated.552

6.3 Commercial activities in plastics liquefaction

6.3.1 Hydrogenation plants in Germany. The Vega process was initially developed to convert coal into naphtha and gas oil. This process was pioneered by Vega Oel GmbH of Germany and then later adopted to depolymerize waste plastics into syncrude, hydrogenated solid residue, and HCl if a dechlorination step was involved.552 Later, in 1999, the technology was discontinued due to the higher cost involved than in more economical feedstock recycling processes and mechanical recycling. Several other multi-stage upgrading and direct hydrogenation liquefaction projects of waste plastics were also established by German companies like RWE-Entsorgungs AG, Hiedrierwerke Zeitz GmbH and Bergakademie Freiberg, but later terminated.552

6.3.2 Liquefaction plants in Japan. The growth of feedstock recycling in Japan was spurred by the Plastic Containers and Packaging Recycling Law passed in 2000. This law promoted sorting at collection and recycling of plastic waste containers. Less of the waste plastic bale materials were made available to liquefaction plants and at a higher tender price when compared to other feedstock recycling. Due to this, liquefaction had a lower combined plant capacity than other feedstock recycling approaches. The Niigata liquefaction plant was established with a capacity of 6 kton per year with support from the Plastic Waste Management Institute (PWMI) and the city of Niigata. The plant operated by Rekisei Koyu started commercial operation in April 1998 and demonstrated

Fig. 23 Example of a continuous tubular reactor design for hydrothermal liquefaction of plastics (extracted from the US patent US 8980143B2).554
that unsegregated and contaminated household waste plastics could be converted to oil.\textsuperscript{55,56} Mixed plastic bales comprising PE, PP, PS, PET, and PVC were allocated from the city of Niigata. The PWMI designed and TOSHIBA-built plant has a pretreatment stage to remove PET and other contaminants followed by a dechlorination stage. The liquefaction process occurs at 420 °C in a stirred tank reactor where 65% of the oil produced was utilized as process fuel within the plant.\textsuperscript{55,56} Details for this process can be found in the 2019 PWMI annual report.\textsuperscript{557}

The Sapporo Plastic Recycle Co. Ltd (SPR) was established in 2000 with the support of PWMI and Sapporo city with an initial capacity of 13.5 kton/year which later grew into 14.5 kton per year.\textsuperscript{553} The SPR plant treated mixed plastics at three stages. The first stage involved pretreatment, sorting and pellitizing, followed by dechlorination, which fed calcium hydroxide into the system. The third and final stage involved thermally decomposing the waste plastics in a rotary kiln reactor at 400 °C and 5 kPa overpressure to produce light, medium, and heavy fractions of oils.\textsuperscript{553,556} Approximately 65% of the total oil produced was utilized as a process fuel within the plant.\textsuperscript{553} Due to the tender system governed by the Japan Container and Package Recycling Association (JCPRA), only half of the baled plastics were used in the SPR liquefaction plant. Additionally, the small operation scale of the SPR plant also raised the cost of waste plastics recycling.\textsuperscript{553} In Mikasa city, another large scale (6 kton per year capacity) liquefaction plant started operation in 2000 but was shut down in early 2004 due to a lack of waste plastic feedstock to process.\textsuperscript{556} By 2010, all major liquefaction plants in Japan ceased, with the advanced SPR plant being the last of them. A 2019 PWMI report suggested that the need for the crude oil product to be further cracked and refined meant that the process was not commercially viable at that time.\textsuperscript{557} Furthermore, these facilities are prone to fire accidents due to the risk of an explosion associated with the liquefaction reactors. Since the 2000s, liquefaction research and development have improved direct liquefaction methods to convert waste feedstocks to naphtha and diesel and thus reduced the burden for further refining. Nevertheless, the issues related to the scale of the plants and businesses remain, and any new companies adopting this technology will face difficulties.\textsuperscript{557}

6.3.3 Plants with hydrothermal liquefaction. Licella Holdings, Australia, has carried out HTL research over the past two decades and has established the Cat-HTR technology.\textsuperscript{558} The first-ever continuous-flow Cat-HTR liquefaction facility, with a capacity of 20 kton per year, was established by Licella Holdings in Australia in 2007, now operated by iQRenew.\textsuperscript{558} Cat-HTR technology is being commercialized internationally via MURA Technology Ltd, Licella’s global joint-venture with Armstrong Energy based in the UK.\textsuperscript{558} MURA technology has proposed a process called Hydro-PRS adopting the Cat-HTR technology.\textsuperscript{559} In 2021, Renew-ELP, based in the UK, started construction of a similar plant in England after obtaining a license for the Hydro-PRS process from Mura Technology.\textsuperscript{559} The companies claim that mixed plastic waste with a limited PVC content can be treated using this process. The Cat-HTR/Hydro-PRS process involved treating waste plastics or other carbonaceous feedstock at sub-critical or supercritical water to produce naphtha, distillate gas oil, heavy gas oil, and heavy wax residue.\textsuperscript{559,560} Decontaminated waste plastic mixture is added to the hopper with an extruder (1), and the melted plastics exiting the hopper were pressurized (2). These molten plastics and SCW were mixed in a chamber (3) before the slurry was fed into the Cat-HTR reactors (4). Note that the Cat-HTR reactors are heated by both a heat carrier (hot water in the chamber) and an external heat exchanger. Here, the waste plastics are treated for 20–25 min at 350–420 °C and above 220 bar to produce over 85% liquid. The post-conversion contents are depressurized (5) before feeding into the flash distillation chamber, where the products are separated (6). The condensable products are then stored (7), and the gases are used as process fuel.\textsuperscript{558,560} The used water after the product separation was recycled in the same system. This process operated at a waste concentration of about 70%.\textsuperscript{558,561}

Another company called Aduro Clean Technologies from Canada has also developed an HTL-based commercial process. They call their process Hydrochemical Plastic Upcycling where the reactor operates at only 240–390 °C and takes about 25 min to completely convert the plastic mixtures.\textsuperscript{562,563} According to their claim, their water-based process operates under less severe conditions and allows the manipulation of chemical reactions associated with different types of plastics to form a comprehensive approach for chemical conversion of plastics.\textsuperscript{563}

7. Gasification of waste plastics

7.1 Fundamentals of gasification reactions

Gasification (or partial oxidation) of MSW converts solid waste into synthesis gas or producer gas (CO, H₂, CH₄) by the reaction shown in eqn (4), where CHₓOᵧ represents the feedstock; q and r are the hydrogen and oxygen to carbon ratios, respectively; m and b are the input air and steam to MSW ratios; xi is the mole number of molecular species.\textsuperscript{564} Producer gas (CO, H₂, N₂) is produced when air is used for the gasification while synthesis gas (CO, H₂) is produced when oxygen is used for gasification. Synthesis gas can be further processed to produce a range of chemicals and fuels. MSW and mixed plastic waste can be used as the feed for gasification.\textsuperscript{565} MSW gasification could be more profitable and environmentally sustainable than incineration.\textsuperscript{566,567} A comparative study estimates that the global warming impacts of waste gasification with a combined cycle powerplant are over 50% lower than incineration.\textsuperscript{568}

\[
\text{CH}_x\text{O}_y + m(\text{O}_2 + 3.76\text{N}_2) + b\text{H}_2\text{O} \rightarrow x\text{H}_2 + x\text{CO} + x\text{CO}_2 + x\text{CH}_4 + x\text{N}_2 + x\text{H}_2\text{O} \tag{4}
\]

Gasification of MSW occurs above 550 °C.\textsuperscript{569} The main MSW gasification products are gases including CO, H₂, CO₂, and CH₄. Solid residues (i.e., ash) are also produced. The
amount of ash depends on the equivalence ratio (i.e., the ratio of oxygen in the employed oxidant and the oxygen required for complete combustion). Ash can be minimized by applying equivalence ratios between 0.25 and 0.35. Accordingly, equivalence ratios within this range are usually used in commercial reactors. Chemical thermodynamics can be used to predict the gas phase products during gasification.\(^5\)\(^7\)\(^0\)\(^5\)\(^7\)\(^1\)

As shown in Table 15, at lower gasification temperatures (i.e., around 600 °C) there is a significant amount of CO\(_2\)\(^5\)\(^7\)\(^2\) formed from the highly exothermic CO oxidation (eqn (5)), the oxidation of carbon (eqn (6)) and the water–gas shift reaction (eqn (7)). These reactions are thermodynamically unfavorable at higher gasification temperatures (>700 °C) leading to a decrease in the amount of CO\(_2\) produced.\(^5\)\(^7\)\(^1\)\(^5\)\(^7\)\(^3\) Conversely, more CO is formed at these higher temperatures by the steam reforming of CH\(_4\) (eqn (8)), steam reforming of hydrocarbons, the steam gasification of char (eqn (9)) and the dry reforming of hydrocarbons (eqn (11)) all of which are highly endothermic reactions. The Boudouard reaction (eqn (10)) is yet another highly endothermic reaction that yields CO.\(^5\)\(^7\)\(^4\) Material and operational constraints such as tar condensation and accumulation\(^5\)\(^7\)\(^5\) often require maintaining gas temperatures above 700 °C.

### 7.2 Waste plastic gasification process flow diagram

Fig. 24 shows a general flow diagram for waste plastic gasification processes. The feedstock is fed into a gasifier where gasification takes place. The volatiles enter a combination of physical, thermal, or chemical gas cleaning technologies where char, slags, and ash are removed. The clean syngas is then cooled in a quench unit to the temperature required for a secondary gas cleaning step. The secondary gas cleaning step removes contaminants that can impact catalyst performance, such as sulfur, halides, HCN, NH\(_3\), and tar, to levels of less than 0.1 parts per million by volume.\(^5\)\(^7\)\(^6\) The cleaned syngas can either go to a turbine for power generation or for catalytic upgrading to other products. Synthetic gas consisting mostly of CO and H\(_2\) is the feedstock for many catalytic synthesis systems.\(^5\)\(^7\)\(^7\)\(^5\)\(^7\)\(^8\) The synthetic gas mixture can be catalytically converted to methanol and other alcohols. Platform chemicals such as ethylene, propylene, and other alkenes can be produced via the methanol-to-olefins process.\(^5\)\(^7\)\(^9\)

The design and configuration of waste plastic gasification facilities depend on the facility scale and targeted products. Some gasification facilities combust the synthetic gas and solid by-product streams to produce combinations of heat and

<table>
<thead>
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<th>Reaction</th>
<th>(\Delta H) (kJ mol(^{-1}))</th>
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<th>Eqn</th>
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<tbody>
<tr>
<td>CO + (\frac{1}{2})O(_2) → CO(_2)</td>
<td>−283.7 kJ mol(^{-1})</td>
<td>CO oxidation</td>
<td>(5)</td>
</tr>
<tr>
<td>C + (\frac{1}{2})O(_2) → CO</td>
<td>−394.1 kJ mol(^{-1})</td>
<td>Carbon partial oxidation</td>
<td>(6)</td>
</tr>
<tr>
<td>CO + H(_2)O ↔ CO(_2) + H(_2)</td>
<td>−41.0 kJ mol(^{-1})</td>
<td>Water–gas shift</td>
<td>(7)</td>
</tr>
<tr>
<td>(\text{CH}_4) + H(_2)O ↔ CO + 3H(_2)</td>
<td>+206.0 kJ mol(^{-1})</td>
<td>Methane steam reforming</td>
<td>(8)</td>
</tr>
<tr>
<td>C + H(_2)O ↔ CO + H(_2)</td>
<td>+131.0 kJ mol(^{-1})</td>
<td>Steam gasification of char</td>
<td>(9)</td>
</tr>
<tr>
<td>C + CO(_2) → 2CO</td>
<td>+172.0 kJ mol(^{-1})</td>
<td>Boudouard reaction</td>
<td>(10)</td>
</tr>
<tr>
<td>(\text{C}_x\text{H}_y + x\text{CO}_2 \leftrightarrow 2x\text{CO} + \frac{y}{2}\text{H}_2)</td>
<td></td>
<td>Dry reforming</td>
<td>(11)</td>
</tr>
</tbody>
</table>

![Fig. 24 Gasification of waste plastics and upgrading to heat, power, olefins, alcohols, and other fuels.](image)
power. Other gasification facilities include steam generation, catalytic upgrading, and other product recovery units to improve energy efficiency and target higher-value markets.

### 7.3 Commercial activities scale

According to the Energy Information Administration (EIA), global waste gasification capacity is estimated to be less than 10 000 Megawatts thermal (MWth) compared to more than 200 000 MWth of coal gasification capacity.\(^{580}\) Despite the low total capacity, the number of global waste gasifiers in operation is estimated to be about 100 facilities which is similar to the number of biomass and natural gas gasifiers.\(^{580}\) These gasifiers primarily produce electricity and heat.

One of the leaders in gasification of MSW is Enerkem who is currently planning to build four commercial MSW recovery facilities. Enerkem’s facility, in Québec, Canada, cost $680 million and will convert 200 kton per year of MSW and forest residues to 125 million liters of biofuels and chemicals starting in approximately 2023. In Alberta, Canada Enerkem recently commissioned a $50 million waste-to-biofuel facility with a capacity of 38 million liters per year.\(^{581}\) In partnership with Shell and the Port of Rotterdam, Enerkem plans to convert about 360 kton per year of MSW to 80 kton of renewable products including aviation fuels. Finally, Enerkem, Agbar, and Repsol plan to build a facility processing 400 kton per year to 220 million liters per year of fuels in Tarragona, Spain pending final investment decisions.\(^{582}\)

### 7.4 Techno-economic analysis

The profitability of waste plastic gasification facilities depends on several process factors and market drivers. Feedstock composition, process yields, and energy efficiency are key drivers for process performance. Feedstock tipping fees, capital costs, and market prices have a significant impact on profitability. Several studies show that waste plastic gasification is more profitable than conventional waste management practices such as incineration and landfiling.\(^{58,583,584}\)

There are numerous techno-economic analysis studies on MSW gasification, but only a small subset focuses on waste plastic upgrading. For this review, we identified three comparative analyses that investigate the costs of waste plastic gasification as a primary strategy. Table 16 shows a summary of the key assumptions and findings from these studies.

### Table 16 Summary of techno-economic analysis estimates for waste plastic gasification

<table>
<thead>
<tr>
<th>Technology</th>
<th>Feedstock</th>
<th>Major products</th>
<th>Region</th>
<th>Feed capacity (kton per year)</th>
<th>Capital cost ($ million)</th>
<th>IRR (%)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical recycling + gasification + catalytic upgrading</td>
<td>MSW</td>
<td>Propylene, ethylene, LPG, others</td>
<td>Germany</td>
<td>76</td>
<td>158.2–271.2</td>
<td>Not reported</td>
<td>583</td>
</tr>
<tr>
<td>Bubbling fluidized bed gasifier</td>
<td>Package derived fuel</td>
<td>Syngas, electricity</td>
<td>Italy</td>
<td>4</td>
<td>0.00549–0.00841/kWe</td>
<td>0.5–29.8</td>
<td>584</td>
</tr>
<tr>
<td>Gasification to hydrocarbons</td>
<td>Waste polypropylene</td>
<td>Propylene, ethylene, aromatics</td>
<td>United States</td>
<td>100</td>
<td>49.09 (annualized)</td>
<td>16.91</td>
<td>58</td>
</tr>
</tbody>
</table>

Voss et al. compared the economic performance of chemical recycling and gasification to incineration as municipal waste treatment options in Germany.\(^{583}\) In their study, they assumed that their feedstock consists of “not differentiable mixed municipal waste” and “household waste, household-type commercial waste” as defined by the European Waste Classification Syste.\(^{585}\) Their MSW stream contains 6.9 wt% plastics. Most of Germany’s waste streams are incinerated directly via combustion to produce heat and power or indirectly via anaerobic digestion of the organic fraction and the refuse-derived fuel (RDF) combustion. The chemical recycling scenario assumes that 1 tonne of MSW is initially mechanically separated into the organic fraction (750 kg) and RDF (250 kg). The organic fraction undergoes anaerobic digestion followed by a biogas-powered combined heat and power system. The RDF fraction is gasified to produce syngas, and the syngas is catalytically upgraded to methanol and subsequently to olefins using sodium hydroxide as the catalyst. Hydrocarbons including propylene (32 kg), ethylene (30 kg), butadiene (4.2 kg), LPG (4.0 kg), and butane (1.9 kg) are recovered in the recovery stage. The chemical recycling unit has a capital cost of $176.6 million for a 76 kton per year facility. Under the basic scenario, this technology is not profitable. It yields a net present value of $180.8 million versus $8927 million for direct incineration. A combination of increasing the plant scale, environmental regulations, and favorable market conditions increases the net present value to $497.2 million.

Gregorio and Zaccariello investigated the economic performance of gasifying packaging-derived fuel (PDF) to produce power, heat and power, and district heating in Italy. PDF is a mixture of primarily plastic-derived materials. Their samples contained 53.9 wt% carbon, 7.7 wt% hydrogen, and 26.0 wt% oxygen. The material contains 0.1 wt% sulfur and 0.3 wt% chlorine, among other contaminants. The nominal capacity (4 kton per year) for the plant was 500 kilowatt-electric (kWe). Total plant costs ranged from $5500 for the power system to $8400 for the district heating scenario. The revenue generated from the three scenarios analyzed includes a tariff of $0.07 per kWe delivered to the grid and $0.09 per kWe for the sale of saturated or superheated steam production in the combined heat and power plant. Based on the revenues, the internal rate of return was estimated at 0.5% for power generation, 18.9% for district heating, and 29.8% for combined heat and power generation.\(^{584}\)
8. Dissolution-based approaches for plastics recycling

Dissolution-based approaches are methods to use solvents to separate and recycle plastics without chemically modifying their structures. The basic workflow of a dissolution-based polymer recycling process can be described as follows: first, the target polymer is selectively dissolved in a suitable solvent or solvent mixture at a defined temperature; second, the mixture is filtered to separate the solid plastic and the liquid phase that contains the solvated polymer; last, the polymer precipitates from the solution by adding an antisolvent (i.e., a solvent in which the polymer is insoluble) and/or decreasing the temperature. Dissolution-based polymer recycling has several advantages over chemical or mechanical recycling methods: it produces high-quality polymers by removing impurities and additives, it can deal with multicomponent plastic mixtures by selectively dissolving a single target polymer; it does not require a high-purity input stream, and it is typically more environmentally friendly than chemical recycling.586–588

8.1 Thermodynamics of selective dissolution and solvent screening

The key to a dissolution-based polymer recycling process is the selection of appropriate solvents. In a selective dissolution process, the solvent should dissolve the target polymer with a sufficient solubility without dissolving other unwanted components of the input stream. Since each polymer presents unique interactions that dictate solubility, and the composition of plastic waste is often complicated, theoretical approaches are valuable for systematically identifying suitable solvents. The thermodynamics of polymer-solvent mixing is governed by the free energy of mixing (eqn (12)),589 where \( \Delta G_{mix} \) is the molar Gibbs free energy of mixing, \( \Delta H_{mix} \) is the molar enthalpy of mixing, \( T \) is temperature, and \( \Delta S_{mix} \) is the molar entropy of mixing. A polymer will favorably mix with a solvent if \( \Delta G_{mix} \) is negative.

\[
\Delta G_{mix} = \Delta H_{mix} - T \Delta S_{mix} \quad (12)
\]

Key contributions to the Gibbs free energy of mixing can be understood using Flory–Huggins theory, a mean-field model for polymer solutions (eqn (13)),590 where \( k_B \) is the Boltzmann constant, \( \phi \) is the volume fraction of the polymer, \( N \) is the degree of polymerization, and \( \chi \) is the Flory interaction parameter that characterizes interactions in the mixture.

\[
\Delta G_{mix} = k_B T \frac{\phi}{N} \ln \phi + (1 - \phi) \ln (1 - \phi) + \chi \phi (1 - \phi) \quad (13)
\]

The \( \chi \) parameter is the critical parameter that governs how favorable a solvent system is for a selected polymer; smaller values promote mixing. The \( \chi \) parameter can be estimated by eqn (14), where \( v_0 \) is the molar volume of a monomer and \( \delta \) is the Hildebrand solubility parameter of compound \( i \), which is defined as the square root of the cohesive energy density. A polymer is expected to interact more favorably – and exhibit a higher solubility – in a solvent with a similar solubility parameter, reflecting the general axiom of “like dissolves like”. This conceptual framework thus highlights polymer- and solvent-specific solubility parameters as critical to the design of dissolution-based processes.

\[
\chi = \frac{v_0}{k_B T} \left( \delta_{\text{polymer}} - \delta_{\text{solvent}} \right)^2. \quad (14)
\]

8.2 Origin of polymer dissolution and precipitation processes

The simplicity and relevancy of dissolution/precipitation for the separation and purification of polymeric materials has been demonstrated experimentally and disclosed in patents since the 1930s. For example, the dissolution and purification of condensation products, mainly polyesters, in ethyl alcohol was disclosed in a 1936 patent by Sändig.591 In 1949, Young and Sparks of the Standard Oil Development Company patented a method to purify polyesters via the solvent-mediated separation of high-molecular weight compounds using paraffinic mineral oils.592 These approaches showed the feasibility of using solvents to dissolve and purify polymerization products, giving way to the idea that this could be used to separate dissimilar polymers. In the 1970s, Seymour and Stahl demonstrated the separation of PE, PVC, PS, polyvinyl acetate (PVAc), and polymethyl methacrylate (PMMA) using a series of solvents that included toluene, methanol, and petroleum ether.593 Around the same time, companies like DuPont filed patents for the recovery of solid scrap thermoplastic polymers like PET, pol(hexamethylene adipamide), poly(acrylonitrile), and poly(oxyethylene) by dissolving them in hexafluoroisopropanol.594 In 1977, Monsanto patented a process for recovering solid PET, poly(cyclohexanedimethanol terephthalate), and poly(butylene terephthalate) from fibers using naphthalene as a selective solvent and dimethylformamide (DMF), water, and acetone as quenching mediums (antisolvent).595

8.3 Applications of dissolution-based plastic recycling

Building upon these early materials recovery and purification processes, applications of dissolution and precipitation to plastic recycling has gained interest along the years. Nauman et al. proposed an approach to separate plastics in physically comingled solid mixtures by selective dissolution. Single solvents were used to dissolve plastics like PS, PVC, LDPE, HDPE, and PP, at different temperatures and pressures.596,597
process consisted of shredding the comingled solids, dissolution of the plastic, filtration, flash devolatization, and pelletization of the separated plastic. Model experiments of plastic recycling with the dissolution/precipitation method were performed by Kampouris, Papaspyrides, and collaborators to recover PVC, LDPE, PP and PS.\textsuperscript{598-602} In one study, Pappa et al.\textsuperscript{603} designed a pilot unit for the separation of LDPE/PP mixtures using the dissolution/precipitation method with xylene and i-propanol as solvent and antisolvent, respectively. They claimed that the costs of the plastics produced from the process were comparable to those of the virgin plastics.\textsuperscript{603}

Regarding the impact of solvent treatment to the plastic properties, early recycling experiments with LDPE, HDPE, PET and PP showed that applying these processes once or twice did not affect the rheological, thermal, or mechanical properties of the final plastic.\textsuperscript{587,598,602,604} Many of these reports established solvent/antisolvent systems for common plastics, including solvent combinations, as summarized in Table 17.

Recent and ongoing research efforts have developed dissolution/precipitation methods for increasingly complex sources of plastic waste. Researchers at the Illinois Sustainable Technology Center have developed a nontoxic energy-efficient chemical solvent process to recover plastics from blends of electronic waste that are converted to fuel oil by pyrolysis. The process has been demonstrated at the lab scale and can produce plastics with properties similar to virgin resins and will begin a pilot-scale project depending on the performance of the recovered plastics.\textsuperscript{605-607} Recently, Georgiopoulou et al.\textsuperscript{608} demonstrated a process to recycle Tetra Pak\textsuperscript{®} packaging materials composed of LDPE, paperboard, and aluminum. Their approach included a hydropulping process to recycle the paper component and a dissolution/precipitation process to recover LDPE, using xylene and i-propanol as the solvent/antisolvent pair. Researchers at the Center for the Chemical Upcycling of Waste Plastics at the University of Wisconsin–Madison have developed a solvent-based method called Solvent-Targeted Recovery and Precipitation (STRAP) that has been demonstrated for the recycling of different post-industrial multilayer films composed of PE, EVOH, PET and EVA.\textsuperscript{609,610} STRAP combines experiments, computational modeling, and process design tools to develop solvent systems to recycle multilayer plastics via selective dissolution. A technoeconomic analysis showed that STRAP can produce plastics at costs similar to virgin resins and can be more economically

<table>
<thead>
<tr>
<th>Polymer</th>
<th>Solvent</th>
<th>Antisolvent</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVC</td>
<td>Cyclohexanone</td>
<td>n-Hexane</td>
<td>601</td>
</tr>
<tr>
<td>PVC</td>
<td>Dichloromethane</td>
<td>Methanol</td>
<td>611</td>
</tr>
<tr>
<td>PVC</td>
<td>Toluene</td>
<td>Methanol</td>
<td>611</td>
</tr>
<tr>
<td>PVC</td>
<td>Tetrahydrofuran (THF), xylene</td>
<td>—</td>
<td>596</td>
</tr>
<tr>
<td>PS</td>
<td>Methyl ethyl ketone (MEK)</td>
<td>Methanol, n-hexane</td>
<td>599</td>
</tr>
<tr>
<td>PS</td>
<td>p-Xylene</td>
<td>n-Pentane</td>
<td>599</td>
</tr>
<tr>
<td>PS</td>
<td>Benzene, toluene</td>
<td>Water</td>
<td>600</td>
</tr>
<tr>
<td>PS</td>
<td>Toluene</td>
<td>n-Hexane</td>
<td>611</td>
</tr>
<tr>
<td>PS</td>
<td>Xylene</td>
<td>Methanol</td>
<td>611</td>
</tr>
<tr>
<td>PS</td>
<td>THF, toluene, xylene</td>
<td>—</td>
<td>596</td>
</tr>
<tr>
<td>PS</td>
<td>p-Cymene</td>
<td>—</td>
<td>612</td>
</tr>
<tr>
<td>EPS</td>
<td>n-Limonene</td>
<td>—</td>
<td>611</td>
</tr>
<tr>
<td>HDPE</td>
<td>Xylene</td>
<td>n-Hexane, methanol</td>
<td>611</td>
</tr>
<tr>
<td>HDPE</td>
<td>THF, toluene, xylene</td>
<td>—</td>
<td>596</td>
</tr>
<tr>
<td>DDPE, LDPE, PP</td>
<td>Turpentine</td>
<td>n-Hexane, petroleum ether</td>
<td>613 and (PetE) 614</td>
</tr>
<tr>
<td>PP</td>
<td>Turpentine/PetE</td>
<td>n-Hexane, PetE</td>
<td>613 and 614</td>
</tr>
<tr>
<td>PP</td>
<td>Turpentine/benzene</td>
<td>n-Hexane, PetE</td>
<td>613 and 614</td>
</tr>
<tr>
<td>LDPE</td>
<td>Xylene</td>
<td>n-Hexane</td>
<td>611</td>
</tr>
<tr>
<td>LDPE</td>
<td>THF, toluene, xylene</td>
<td>—</td>
<td>596</td>
</tr>
<tr>
<td>PE, PP, PS</td>
<td>n-Butane, propane</td>
<td>—</td>
<td>615</td>
</tr>
<tr>
<td>PP</td>
<td>Xylene</td>
<td>n-Hexane</td>
<td>611</td>
</tr>
<tr>
<td>PP</td>
<td>Tetrachloroethylene</td>
<td>—</td>
<td>616</td>
</tr>
<tr>
<td>PP</td>
<td>THF, toluene, xylene</td>
<td>—</td>
<td>596</td>
</tr>
<tr>
<td>PET</td>
<td>THF</td>
<td>—</td>
<td>596</td>
</tr>
<tr>
<td>PET</td>
<td>γ-Valerolactone (GVL)</td>
<td>—</td>
<td>617</td>
</tr>
<tr>
<td>PET</td>
<td>N-Methyl-2-pyrrolidone (NMP)</td>
<td>n-Octane, n-hexane</td>
<td>604</td>
</tr>
<tr>
<td>PET</td>
<td>Diphenyl, diphenyl ether, naphthalene, methynaphthalene, benzophenone, diphenylmethane</td>
<td>—</td>
<td>618</td>
</tr>
<tr>
<td>PET</td>
<td>Benzyl alcohol</td>
<td>Methanol</td>
<td>611</td>
</tr>
<tr>
<td>PCL</td>
<td>Dichloromethane</td>
<td>Acetone</td>
<td>619</td>
</tr>
<tr>
<td>PA 6</td>
<td>Dimethyl sulfoxide (DMSO)</td>
<td>MEK</td>
<td>620</td>
</tr>
<tr>
<td>PA 6,6</td>
<td>Formic acid</td>
<td>MEK</td>
<td>620</td>
</tr>
</tbody>
</table>
feasible when the use of antisolvents is reduced by instead inducing precipitation via temperature.609

Other than targeting specific plastic components in plastic waste, solvents can also be used to extract additives from plastics or to selectively dissolve or decompose adhesives in plastic structures to trigger delamination. While these topics are out of the scope of this discussion, they have been discussed at length in recent publications.621–624

8.4 Industrial demonstrations of dissolution/precipitation processes

A handful of technologies are being commercialized for the recycling of different plastic waste feeds via dissolution methods. Here, we focus primarily on technologies that are currently in the process of commercialization. We omit discussion of past efforts; for example, the Solvay VinyLoop process was developed to separate PVC from a polymer coating using a selective dissolution step in a proprietary solvent,625 but the process ceased operations in 2018.

APK AG’s NewCycling process is being demonstrated in a 8000 ton per year plant in Germany to produce polyamides (PA) and PE from PIW multilayer plastics.626–628 Their technology is based on dissolving a plastic using a solvent mixture from a group of alkanes, isoctane or cycloalkanes.629 The plastics are recovered from solution and then pelletized by extrusion.626,627 Centrifuges separate the solids from the plastic solution and solvent.630 APK AG claims that PP, PET, PS, PLA and aluminum could also be recovered with this process in the future.627

The Fraunhofer Institute is developing a solvent-based process called CreaSolv that produces plastics with comparable properties to virgin materials, effectively removing contaminants and additives.631 The process is likely based on the dissolution of a target plastic, mainly polyolefins, using a solvent with a Hansen parameter $\delta_H$ between 0.0 to 3.0 MPa$^{1/2}$, selected from a group of aliphatic hydrocarbons. An antisolvent made out of mono/polyalkyldioxy hydrocarbons, like 1-propanol or 1,3-propanediol, with a $\delta_H$ between 4.0 and 38.0 MPa$^{1/2}$ is then used to precipitate the polyolefin from the mixture.632 The institute has also studied the separation of PS633 and is researching the recycling of multilayer food packaging that can consist of PET, PE, PP, EVOH, PA and metalized layers.634 Currently, Unilever is implementing the CreaSolv process in a 1100 ton per year demonstration plant in Indonesia to recover PE from multilayer sachets.635,636

CreaSolv is also reported to have a demonstration plant in the Netherlands to separate EPS from POF, a pilot plant for recycling packaging and automotive compounds, and a demonstration plant in Germany for multilayer films.637

PureCycle Technologies is building a 54 000 ton per year facility that uses solvents to produce PP from PIW and PCW in collaboration with P&G.638,639 The method consists of contacting the plastic waste with an alkane at elevated temperatures and pressures to obtain the purified PP.640 The process removes impurities, undesired colors and odors and produces PP with comparable properties to the virgin resin.639 P&G has recently filed patents with methods to separate and purify adhesives, PP, PE, PET, cellulose, and polyacrylic acid (PAA).641 The company also has inventions in the purification of PE using a pressurized solvent consisting of hydrocarbons, primarily n-butane or pentane.642 Additionally, they have a process for the separation of PS, PP, PE and poly(dimethylsiloxane) under similar conditions.643

Polystyrene developed a dissolution process to recycle different types of PS, including expanded PS, extruded PS and high-impact PS. Polystyrene operates at 125 kg per hour with an output of 600 metric tons per year.605 In their technology $p$-cymene is used as a selective solvent for PS and a non-polar solvent is used as an antisolvent to precipitate the plastic and remove $p$-cymene.612 Through this approach they can separate and recover PS from other materials, as only PS dissolves in the solvent and other materials float or sink in the system.643

Dissolution-based recycling processes have also been applied to textile waste. For example, Worn Again Technologies has patented a technology to recycle PET into pellets. In their process, the polymers are dissolved in a solvent system with either benzyl acetate, benzyl benzoate, benzaldehyde or similar solvents, precipitated by cooling of the solvent system, and then separated via filtration. After the plastic is separated, it is washed and dried and then molded into pellets and/or converted into fibers.644 This process can be applied to cotton textiles, post-consumer PET bottles, and plastic packaging containing PET. Worn Again has also partnered with companies such as H&M and Kering to promote the reduction of textile waste.645 Another start-up company, Amberecycle, has developed a technology to purify and separate polymers from different garments and is planning a demonstration plant to produce a metric ton of recycled material per day.646,647

Other approaches are being used to recycle multilayer plastics using a combination of mechanical and solvent-mediated steps. For example, Saperatec has introduced a low-energy mechanical recycling process that involves shredding the multilayer film followed by use of surfactant-containing microemulsions to break up and separate the layers.648 The technology separates PE and aluminum film, PP from aluminum, and PE from PP. The technology was originally developed for rigid packaging and is being extended to flexible packaging and will be demonstrated at a 17 000 ton per year facility in Germany.649

8.5 Advanced solvent selection for complicated feedstocks

As presented above, several solvents and antisolvents have already been identified for many common plastics, enabling the development of dissolution/precipitation processes for single plastics by selecting solvents from past literature.588,650,651 However, an ongoing challenge is solvent selection for plastic mixtures which are ubiquitous in realistic plastic waste. Multicomponent polymeric materials are often made from two or more plastics, each of which is selected to contribute its own useful properties.652 For example, food packaging boxes are typically multilayer plastic films, which are extremely challenging to recycle because of their complex
compositions and the incompatibility of different polymer layers.\textsuperscript{610} Dissolution-based methods are promising for recycling such materials because selective dissolution enables the separation of different plastics and is tolerant to the additives and impurities present in waste. However, due to the variety of possible plastic combinations, as well as the necessity for choosing appropriate operating temperatures for dissolution and precipitation, solvent selection for plastic mixtures is a difficult task.

Based on the thermodynamic framework described above, one method for solvent selection is to compute solubility parameters for target plastics and solvents to predict dissolution. Hildebrand solubility parameters can be applied with reasonable accuracy to predict solubility in systems dominated by non-polar interactions, but are inaccurate for mixtures with strong polar or specific interactions such as hydrogen bonds.\textsuperscript{590,633} Similar to the Hildebrand solubility parameter, other parameter sets have been applied to identify solvents and antisolvents for plastics; examples include Hansen,\textsuperscript{654} Kamlet-Taft,\textsuperscript{655} Gutmann,\textsuperscript{656} and Swain\textsuperscript{657} parameters. These parameters assess the chemical similarity between species to predict dissolution behaviors.\textsuperscript{651} Among them, Hansen solubility parameters (HSP) are the most widely used to select solvents for plastic dissolution.\textsuperscript{588,651,658} The HSP system assigns a sphere in the HSP space. Solvents within this sphere defines a sphere in the HSP space. Solvents within this sphere have demonstrated that plastics recovered via this technology can be sold at prices that are comparable to virgin resins.\textsuperscript{610} Due to STRAP being a new technology, the size of the process required to achieve a competitive sell price is important, as large capital costs can make scaling up to an industrial level difficult. At a feed rate of 3000 tonnes per year, plastic recycled by the STRAP process can be sold at a minimum selling price (MSP) of $2.18 per kg, while delivering a discounted return on investment (DROI) of 10% to investors over a 20 year lifetime. The MSP can be further reduced through optimization of STRAP design parameters, as shown in Fig. 25.\textsuperscript{609}

Building upon the success of these solubility parameter methods, newer computational approaches for predicting polymer solubilities and screening solvents with minimal experimental effort have emerged in recent years. A recent example was the development of machine learning techniques for the large-scale binary classification of solvents and antisolvents for polymers, thus categorizing solvents into similar categories as HSPs. Specifically, Chandrasekaran et al.\textsuperscript{660} developed a deep neural network to classify 24 common solvents for over 4500 homopolymers with an accuracy of 93.8%. This example demonstrates the feasibility of applying data-centric methods to rapidly screen potential polymer–solvent combinations for selective dissolution processes, although more data will be required to extend the solvent library.

The preceding techniques primarily focus on qualitatively classifying solvents and antisolvents for target polymers. However, the development of dissolution/precipitation processes requires quantitative analysis of solubilities as a function of temperature in order to assess the feasibility and economics of potential industrial-scale processes. To address this challenge, Zhou et al.\textsuperscript{661} developed a computational approach for quantitatively predicting temperature-dependent polymer solubilities using molecular-scale models.\textsuperscript{661} This approach employs molecular dynamics simulations to sample polymer conformations in representative solvents. These conformations are used as input to the CONductor-like Screening MOdel for Real Solvents (COSMO-RS), a model that used quantum mechanical calculations and statistical thermodynamics methods to predict polymer solubilities through solid–liquid equilibrium calculations.\textsuperscript{610} After initial polymer conformations are available, temperature-dependent solubilities can be obtained rapidly (~minutes). This approach was applied to screen over 500 solvent combinations for the selective dissolution of PE and EVOH and extended to develop a STRAP process for a commercial multilayer film with four polymer components.\textsuperscript{609,610,661}

8.6 Techno-economic analysis of STRAP

A published techno-economic analysis of the STRAP process has demonstrated that plastics recovered via this technology can be sold at prices that are comparable to virgin resins.\textsuperscript{610} Due to STRAP being a new technology, the size of the process required to achieve a competitive sell price is important, as large capital costs can make scaling up to an industrial level difficult. At a feed rate of 3000 tonnes per year, plastic recycled by the STRAP process can be sold at a minimum selling price (MSP) of $2.18 per kg, while delivering a discounted return on investment (DROI) of 10% to investors over a 20 year lifetime. The MSP can be further reduced through optimization of STRAP design parameters, as shown in Fig. 25.\textsuperscript{609}

A key feature that drives economic performance is solvent selection. A solvent with a high affinity for a target polymer can decrease the dissolution time and solvent/polymer ratio which can in turn decrease the filter equipment cost by reducing its size. A low-boiling solvent/antisolvent pair with a high relative volatility or solvent that allows for temperature-controlled precipitation can reduce the size of or eliminate the need for distillation columns.\textsuperscript{609} Using less solvent, an easily separable solvent/antisolvent system, or eliminating the need for an antisolvent then reduces the quality and quantity of steam required. This illustrates that efficient solvent screening and selection is essential for establishing economic as well as technical feasibility of STRAP.

Fig. 26 shows that STRAP is far from the point of diminishing returns at a processing capacity of 3000 tonnes per year. These data indicate that there will be a significant economic incentive to scale STRAP up to larger sizes if the process is shown to be effective at smaller processing capacities. At a feed rate of 15 000 tonnes per year, the recycled plastic could be sold for $0.95 per kg to achieve a DROI of 10% or deliver a
DROI of approximately 30% if the selling price of $2.18 per kg is maintained while requiring about twice as much capital investment. This analysis demonstrates that the process has significant economic potential and can be made even more competitive via process optimization and solvent selection. More broadly, this analysis, along with the commercial demonstrations described previously, supports the economic viability of dissolution-based recycling techniques.

9. Chemical recycling of polyester to monomers

9.1 Chemical recycling of PET

The commercial uses of PET span a wide variety of industrial sectors, from fiber for textiles to resin for beverage bottles. In 2021, global PET consumption totaled 89.3 million metric tons (MMT) (Fig. 27). It is the fourth most consumed plastic in the world after PE, PP, and PVC. There are many reviews in the literature summarizing recent progress in chemical plastics recycling, as well as several which focus solely on PET. Chemical recycling of PET is an area of intense focus in the literature, and a comprehensive review of every report in this field is thus not attempted here. This section summarizes each strategy for PET depolymerization, with a focus on important and practical advances and their potential for industrial application. This review is organized into sections based on the approach to PET depolymerization:
aerobic oxidation, hydrolysis, alcoholysis, glycolysis, aminolysis, hydrogenolysis, and enzymatic depolymerization (Fig. 28). Current commercial activity in chemical PET depolymerization is included at the end of this section, along with comments on the economics of these processes.

**9.1.1 Aerobic oxidation.** Aerobic oxidation of PET involves heating the polymer to high temperatures in the presence of oxygen, and sometimes a catalyst or promoter. In 1984, Jabarin and Lofgren studied the thermal oxidation of PET and showed that it was an exothermic reaction with an activation energy lower than the thermal decomposition of PET under vacuum. At all temperatures and drying conditions studied, increased decomposition of PET was observed under air as compared to under nitrogen. However, specific products of the reaction were not identified and PET degradation was measured only by the weight loss of the sample.

Partenheimer received a patent in 2005 for the oxidation of a wide variety of polymers using a range of simple metal salt promoters. PET sourced from blue film, recycled resin, and shirts were depolymerized as 4–20 wt% solutions in acetic acid under 1000 psia air pressure at 150–205 °C. A combination of Co/Mn/Br/Zr in varying ratios was used as a catalyst, and the effect of adding co-oxidants such as toluene and p-xylene was also investigated (Fig. 29). TPA yields of up to 100% were reached, and more efficient conversion was observed at higher temperatures, longer reaction times, higher initial PET concentrations, and in the presence of either co-oxidant. Partenheimer further published that a 56% yield of TPA could be isolated from the oxidation of PET at 190 °C in a water/acetic acid mixture using a similar Co/Mn/Br/Zr catalyst. The TPA yield increased to 100% in the presence of toluene as a co-oxidant. This chemistry was based on an extensive review published by Partenheimer about a decade earlier on metal/bromide autoxidation of hydrocarbons.

**9.1.2 Hydrolysis.** Hydrolytic PET depolymerization involves the reaction of the polymer in either a basic, neutral, or acidic aqueous solution. The products of PET hydrolysis are TPA and EG. One major disadvantage of hydrolysis is the large volumes of inorganic salt and aqueous waste that are generated, along with the high corrosivity of the basic and acidic systems. It has been estimated that, on an industrial scale, 20–50 L of aqueous solution would be required to hydrolyze 1 kg of post-
However, it is also more tolerant of post-consumer waste contaminants and other polymers than many chemical recycling strategies. Several studies are included in this section that utilize post-consumer PET sources.668,680

9.1.2.1 Basic hydrolysis. Basic hydrolysis is typically carried out in a 4–20 wt% solution of NaOH or KOH in water. Reaction temperatures average around 200 °C and pressures are usually 15–20 bar (Fig. 30a). The disodium or dipotassium terephthalate product must be protonated with acid after the reaction, generating stoichiometric inorganic salt waste.681 LCA performed on this reaction demonstrated that decreasing the volume of water required for product purification is critical to achieve lower GHG emissions for this recycling process compared to PET incineration.679 Yoshioka and coworkers evaluated a system in which PET was hydrolyzed in concentrated NaOH at 250 °C and 5 MPa O2 pressure for 5 hours. The disodium terephthalate was not oxidized under these conditions and was obtained in quantitative yields. However, EG was oxidized to oxalic acid in roughly 70% yield, generating a value-added product that could be sold to further improve the economics of this recycling strategy.682

A phase transfer catalyst such as a quaternary ammonium salt can be used to lower the temperatures and pressures required for alkaline hydrolysis. Kosmidis and coworkers683 used this strategy to isolate TPA at high purity using aqueous NaOH and catalytic trioctylmethylammonium bromide to depolymerize post-consumer PET soda bottles under atmospheric pressure and at temperatures as low as 70 °C. Similarly, microwave irradiation has been used to lower the required temperature of basic PET hydrolysis.684 Khalaf and Hasan685 isolated TPA from post-consumer soda bottles in almost quantitative yields after only 1 hour of irradiation in the presence of tetrabutylammonium iodide as a phase transfer catalyst. The catalyst could be recovered from the reaction mixture after product extraction, which improves the potential practicality of this method.

Hu et al.686 found that using a mixed solvent system of an ether and an alcohol enabled complete depolymerization of PET with NaOH in 40 minutes at only 60 °C. Pure TPA and EG were easily isolated from the reaction mixture. In 2017, Essadarn687 received a patent for the room temperature hydrolysis of PET. The system consisted of a nonpolar solvent such as CH2Cl2, which swelled the polymer, and an alcoholic solution of NaOH to hydrolyze the PET. High purity TPA could be isolated using this method in up to quantitative yields. Ügdüler and coworkers679 used ethanol as a cosolvent in the
9.1.2.2 Neutral hydrolysis. Neutral hydrolysis is in many ways preferable to basic or acidic strategies, as the solution will not require specialized reactor materials to prevent corrosion and stoichiometric salts are not required or formed in the reaction. However, this also means that impurities and additives present in the PET remain in the recycled monomer, which is not the case in TPA generated from acidic and basic hydrolysis.\(^{680,681}\) Additionally, high temperatures and pressures are required for the reaction to run at an appreciable rate. The temperatures used are 200–300 °C, and often in the upper half of this range so the reaction runs in the melt phase of PET. Pressures of 10–40 bar are typically used, along with a large excess of water or steam (Fig. 30b). The large volume of water required leads to very dilute solutions of EG in the product mixture, which makes isolation more difficult and costly. The reaction proceeds in about 1 hour at 275 °C, and it has been proposed that the TPA formed in the reaction can catalyze the hydrolysis, rendering the reaction autocatalytic.\(^{681,688}\)

Campanelli and coworkers\(^{689}\) investigated possible catalytic effects in the neutral hydrolysis of PET but found that zinc salts exhibit only a minute accelerating effect on the reaction. Full conversions were only reached at temperatures above 250 °C. Xylene was utilized as a cosolvent in neutral hydrolysis, which reduced the required temperature, pressure, and volume of water in the reaction. Additionally, this strategy enabled the efficient isolation of EG as a concentrated solution in the organic layer.\(^{690}\)

Zhang et al.\(^{691}\) used a dual phase-transfer catalyst \([{(CH_3)_3N(C_{16}H_{33})_2)[PW_{12}O_{40}]}\) in the neutral hydrolysis of post-consumer bottle PET at 145 °C for 3 hours to obtain TPA yields of up to 93%. The catalyst allowed the reaction to be run at much lower temperatures and could be separated from the products and recycled at least three times without any observed decrease in efficiency. Stanica-Ezeanu and Matei\(^{692}\) found marine water to be an excellent solvent for neutral PET hydrolysis due to the presence of various metal ions that can act as catalysts. They used simple salts commonly found in the ocean, including NaCl, KHCO\(_3\), and CaCl\(_2\), as catalysts in the depolymerization of PET at 200 °C and 40 bar to obtain TPA in up to 85% isolated yield. Yields were increased at milder conditions when more than one salt was added or when marine water was used as a catalyst.

9.1.2.3 Acidic hydrolysis. Several systems have been described for the acidic hydrolysis of PET in concentrated (>50 wt%) H\(_2\)SO\(_4\) solutions to generate high purity TPA in high yields at ambient pressures and relatively low temperatures (Fig. 30c). However, these reactions are highly impractical at scale due to the corrosivity of the solution, the need to recycle large volumes of acid, the generation of stoichiometric salt waste, and the difficulty in isolating EG from the reaction.\(^{672,681,693}\) Yoshioka and coworkers\(^{694}\) discovered that they could decrease the H\(_2\)SO\(_4\) concentration by about half and still achieve complete hydrolysis of PET to TPA in 5 hours at 150 °C. The acid was subsequently recovered by dialysis. The same group also reported that 7–13 M HNO\(_3\) completely hydrolyzed PET over about 2 days at temperatures below 100 °C. The EG was simultaneously oxidized to oxalic acid, a value-added product.\(^{695}\)

Liu et al.\(^{696}\) studied the hydrolysis of PET using the ionic liquid (IL) 1-butyl-3-methylimidazolium (Bmim) chloride as the solvent and an acid-functionalized IL (1-methyl-3-(3-sulfo-propyl)-imidazolium hydrogen sulfate) as a catalyst. After 4.5 hours at 170 °C, they isolated TPA in 88% yield, and were able to recycle the ILs at least 8 times with no decrease in yield. Yang and coworkers found that TPA itself was an effective catalyst for acidic PET hydrolysis. They achieved a 95.5% yield of high purity TPA after 3 hours at 220 °C. The only workup required was a simple filtration, and the hydrolysis efficiency was maintained over 8 consecutive reaction cycles.\(^{697}\)

9.1.3 Alcoholysis. Alcoholysis depolymerizes PET in the presence of alcohols to form an ester and EG (Fig. 31). Methanol and ethanol are the most common alcohols utilized in this reaction, but other alcohols can also be used.

9.1.3.1 Methanolation. Methanolysis of PET produces dimethylterephthalate (DMT) and EG. DMT can be used directly in transesterification polymerization to form PET or further hydrolyzed to TPA.\(^{572,680,698}\) However, methanolysis of post-consumer PET leads to high separation and purification costs due to the mixture of glycols, alcohols, and phthalate derivatives produced in the reaction.\(^{663}\) Methanolysis typically operates at temperatures of 180–280 °C and pressures of 20–40 bar. The high pressures ensure that the methanol remains a liquid during the reaction. Transesterification catalysts such as zinc acetate, magnesium acetate, cobalt acetate, or lead dioxide are

![Fig. 31 Typical reaction conditions and products for PET alcoholysis.](image-url)
typically used, and yields of DMT average 80–85%. This reaction can be run in continuous flow, but complex reactors are required to add feedstocks while maintaining the required high reaction pressures. In a typical continuous method, melted PET and preheated methanol are fed into an autoclave with a set residence time to complete the depolymerization, then the mixture is flowed to the bottom of a second pressurized reactor at a lower temperature so that high density impurities can be removed. After leaving the second autoclave the reaction pressure is reduced, the product mixture is further cooled in a mixer, and finally DMT is isolated by precipitation.672,680,699

Sako and coworkers700 reported in 1997 that PET could be completely depolymerized to DMT, EG, and some oligomers with no catalyst in only 30 minutes in supercritical methanol. This was achieved at temperatures above 300 °C and pressures of 110 bar. Above supercritical conditions, the pressure of the reaction was not found to have a significant effect on DMT yield, however the severe conditions make this process too cost intensive for large scale use and too difficult to adapt to continuous flow.580

Kurokawa et al.698 used the transesterification catalyst Al(OiPr)3 to promote the methanolysis of PET from soda bottles under more mild conditions. They achieved 88% DMT and 87% EG yields in 4:1 methanol:toluene at 200 °C and no added pressure. Improved yields in this solvent mixture compared to pure methanol indicate that the rate of methanolysis strongly depends on PET solubility. Hofmann and coworkers701 found that microwave irradiation facilitated the methanolysis of PET under even milder conditions using zinc acetate as a catalyst and CH2Cl2 as a cosolvent to improve solubility. In only 20 minutes at 160 °C, moderate to high yields of DMT were isolated from a variety of post-consumer PET sources, including a dish detergent bottle (46% yield), a soap dispenser with a label (70% yield), and a transparent food container (98% yield).

Pham and Cho702 reported that the inexpensive salt K2CO3 catalyzes PET methanolysis at room temperature. A 93% yield of DMT was obtained after 24 hours in a mixed solvent system of 1:1 methanol:CH2Cl2. Tanaka and coworkers703 developed a new strategy wherein they trapped the EG formed in the reaction as ethylene carbonate. This biased the depolymerization equilibrium towards the monomers and enabled the reaction to run in only 5 hours at ambient temperature. Various simple alkali salts were used as catalysts, and optimized yields of DMT exceeded 90% using 5 mol% LiOMe.

9.1.3.2 Ethanolysis. PET ethanolysis yields diethylterephthalate (DET) and EG. The strategy is similar to methanolysis in many ways but could be particularly advantageous in countries such as Brazil where ethanol from sugarcane is cheap and abundant.704

In 2006, it was reported that PET could be completely depolymerized to its monomers in 5 hours in supercritical ethanol. At 528 °C and pressures of 76–116 bar, DET yields of 65–98% were obtained using post-consumer PET from a variety of sources including green bottles, mechanically recycled fiber, and polyester string.705 Fávaro and coworkers704 studied the ethanolysis of multilayer food packaging containing PET, PE, and aluminum. The material was depolymerized over 2 hours in supercritical ethanol at 255 °C and 116 bar. High purity DET was obtained in 80% yield after precipitation, and metallic aluminum could also be isolated from the reaction. Lozano-Martínez and coworkers found that the product distribution from PET ethanolysis could be changed by altering the reaction conditions. Long reaction times in supercritical ethanol generated DET as the only product, while shorter reaction times at lower temperatures and pressures led to the formation of TPA.706

Nunes et al.707 reported that the addition of catalytic [Bmim][BF4] reduced the required reaction time for PET depolymerization in supercritical ethanol from 6 hours to only 45 minutes. Cobalt and nickel oxides have also been found to be effective catalysts for PET ethanolysis under supercritical conditions, and provide almost quantitative yields of DET. The required reaction time was only 90 minutes with low catalyst loadings.708

9.1.4 Glycolysis. Glycolysis is one of the most intensely studied routes for chemical PET depolymerization, and several systems have been commercially adopted. Glycolysis involves the transesterification of PET with excess glycol, most commonly EG, to generate the corresponding ester (bis-2-hydroxyethyterephthalate, BHET, when EG is used) and EG as products (Fig. 32). Many different classes of catalysts have been studied in this reaction.663,670,709 Without a catalyst, PET glycolysis requires temperatures of 200–240 °C and pressures of 2–6 bar, and high monomer yields are not obtained.881 In 1991, Chen and coworkers710 studied the relationship between reaction pressure and EG:PET ratio on depolymerization kinetics. Higher pressures were reported to enable faster reactions, and at higher EG concentrations BHET was in equilibrium with PET oligomers. At lower EG concentrations, no monomer was observed.

Metal acetates were first reported as PET glycolysis catalysts in 1989 and have been widely studied.709,711,712 Güçlü and coworkers713 used xylene to form a multiphase reaction in zinc acetate-catalyzed PET glycolysis with EG at temperatures of 170–245 °C. BHET was pulled into the xylene layer to bias the

![Fig. 32 Typical reaction conditions and products for PET glycolysis with EG.](image-url)
equilibrium of the depolymerization and prevent oligomer formation, providing monomer in up to 80% yield. Other diols such as butanediol and triethylene glycol have also been used in PET glycolysis with zinc acetate to generate monomers and dimers.\textsuperscript{714} Chen found that the optimal PET glycolysis temperature was 190 °C when using manganese acetate as a catalyst. Conversion of PET to BHET and dimer reached ∼100% after 1.5 hours.\textsuperscript{715} Ghaemy and Mossaddegh\textsuperscript{716} used various metal acetate catalysts in the glycolysis of PET in refluxing EG. Yields of BHET reached up to 85% from both fiber and flake PET.

Troev and coworkers\textsuperscript{717} demonstrated that light metal salts could also be used as glycolysis catalysts. They reported that titanium(w) phosphate catalyzes the depolymerization of PET to BHET in up to 97% selectivity at 190 °C with a reaction time of 2.5 hours. Fang et al.\textsuperscript{718} utilized polyoxometalates as catalysts for PET glycolysis under mild conditions. BHET was isolated in 85% yield after only 40 minutes at 190 °C and 0.018 mol% catalyst loading, with no loss in activity observed after the catalyst was recycled four times.

ILs have gained interest in the literature as catalysts for PET glycolysis due to the ease of separating the products from the reaction mixture, enabling facile catalyst recycling.\textsuperscript{719-721} H. Wang et al.\textsuperscript{719} first published the use of ILs as catalysts for PET glycolysis in 2009. They synthesized and tested several different ILs as catalysts and were able to achieve full conversion of PET to primarily BHET and some oligomers at 180 °C in 8 hours at atmospheric pressure. Higher reaction temperatures led to improved BHET selectivity. A subsequent study found that the IL could be recycled with no loss in activity.\textsuperscript{722} They then found that an iron-containing IL could catalyze the depolymerization at temperatures as low as 140 °C.\textsuperscript{723} Q. Wang\textsuperscript{720} and coworkers studied different metal-containing ILs as PET glycolysis catalysts. They achieved isolated yields of up to 80% high purity BHET after 75 minutes at 175 °C and atmospheric pressure. Other metallic ILs have been shown to be effective catalysts for this reaction under a variety of conditions.\textsuperscript{724} Yue et al.\textsuperscript{725} used basic ILs at catalysts in PET glycolysis to isolate BHET in yields of up to 71% after 2 hours at 190 °C.

Deep eutectic solvents (DESs) have also garnered recent interest as catalysts for PET glycolysis due to their low cost, simple synthesis, low toxicity, and other characteristics shared with ILs such as thermostability and tunability.\textsuperscript{726,727} Q. Wang and coworkers first used DESs as catalysts for PET glycolysis in 2015. Under optimized conditions of 170 °C and only 30 minutes, BHET was obtained in 83% yield. These results are similar to those obtained under supercritical conditions. The high activity is attributed to a synergistic effect between H-bonds and coordination bonds between the DES and EG. Sert et al.\textsuperscript{728} synthesized five different DESs to evaluate as PET glycolysis catalysts. The best catalyst tested was formed from potassium carbonate and EG, and reached isolated BHET yields of up to 88% after 2 hours at 180 °C.

A variety of different heterogeneous catalysts have also been used in the glycolysis of PET. Shukla and coworkers\textsuperscript{729} used β- and γ-zeolites as catalysts for PET glycolysis in 2008. BHET yields of over 60% were reached after 8 hours in refluxing EG. Al-Sabagh et al.\textsuperscript{730} used multiwalled carbon nanotubes doped with Fe₂O₄ as catalysts for PET glycolysis, and reported quantitative BHET yields after 2 hours at 190 °C. The catalyst was successfully recycled 8 times with no loss in activity. Veregue et al.\textsuperscript{731} used 3 nm cobalt nanoparticles as catalysts, and isolated pure BHET without additional water in 77% yield after 3 hours at 180 °C. The catalyst could be reused at least 5 times with no drop in activity. Laldinpuii and coworkers\textsuperscript{732} used bamboo leaf ash as a bio-waste derived catalyst for PET glycolysis. Several metal oxides and other salts were identified in the catalyst, which provided BHET in up to 83% yield after 3 hours at 190 °C. The catalyst could be reused four times with only a slight decrease in yield, and EG was also recovered from the reactions for reuse. Z. Wang et al.\textsuperscript{733} developed a colloidal catalyst based on graphitic carbon nitride for this reaction. 80% BHET yield was achieved in 30 minutes at 196 °C for five consecutive catalyst cycles. Several different post-consumer PET sources were successfully depolymerized to give BHET in ~80% yield, including green bottle flake, noise deadening foam, and bottle flake contaminated with PP.

IL catalysts have been tethered to heterogenous supports to enable even more facile catalyst recovery and product isolation. Najafi-Shoa\textsuperscript{734} and coworkers used a cobalt-containing IL supported on graphene to reach 95% isolated BHET yields after 3 hours at 190 °C and ambient pressure. The catalytic activity was maintained over five cycles. Al-Sabagh et al.\textsuperscript{735} supported iron-containing ILs on bentonite to reach a maximum 44% BHET yield. Similarly, Cano et al. supported iron-containing ILs on silica-coated, magnetic Fe₂O₄ nanoparticles. The recoverable catalyst gave quantitative BHET yields for an impressive twelve consecutive 24 hours cycles at 190 °C, and the magnetic particles greatly simplified reaction workup.\textsuperscript{736}

Organocatalytic PET glycolysis was first reported in 2011 by Fukushima et al.\textsuperscript{737} using the amine base 1,5,7-triazabicyclo[4.4.0]dec-5-ene (TBD). Isolated BHET yields of up to 78% were reached after 3.5 hours at 190 °C and atmospheric pressure, comparable to some of the most efficient metal-catalyzed reactions. A study of several other nitrogen-based catalysts led to the discovery that 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU) was an even more efficient catalyst when EG was used as the glycol. Mechanistic studies revealed that EG and other short chain diols can act as cocatalysts to activate the PET ester group and assist in depolymerization (Fig. 33).\textsuperscript{738} Jehanno and

![Fig. 33 Proposed mechanism of PET activation by the organic bases TBD or DBU and short chain alcohols. Longer chain alcohols cannot participate in this reaction.](image-url)
catalyst in PET glycolysis. 91% BHET was isolated after 2 hours at 180 °C. The catalyst could be recycled at least five times, and the monomer isolated from these reactions was used to synthesize new PET with similar thermal properties as commercial material.

Microwave irradiation is another growing strategy for facilitating PET glycolysis at milder conditions than conventional heating. Pingale and Shukla first published in 2008 that, under otherwise identical zinc acetate-catalyzed glycolysis conditions, microwave irradiation provided the same yield of BHET in 35 minutes that conventional heating did in 8 hours. This increased efficiency was confirmed by Chaudhary et al. Achilias et al. studied the microwave glycolysis of PET to oligomers using diethylene glycol. They found that complete degradation was achieved in only 5 minutes at 180 °C, compared to more than 4 hours using conventional heating. Saravari et al. glycolyzed PET bottles with propylene glycol under microwave irradiation with zinc acetate as a catalyst. Oligomers obtained from the reaction were reacted with linseed oil and toluene diisocyanate to obtain urethane oil with similar properties to commercially purchased material, showing the potential of this strategy for PET upcycling. Parrott received a patent for microwave-assisted PET glycolysis in 2020. BHET yields of up to 94% were achieved in only 5 minutes of irradiation at 250 °C with 0.1 wt% zinc acetate as a catalyst. A variety of simple salt catalysts were investigated, and zinc salts were generally the most effective.

9.1.5 Aminolysis. Aminolysis depolymerizes PET in the presence of amines to form terephthalamides and EG. This process is more thermodynamically favored than alcoholysis or glycolysis due to the stronger amide bonds formed in the products, and so less forcing conditions can be used (Fig. 34). The terephthalamide monomers can be used in a variety of upcycling applications, including antibacterial drugs, adhesion promoters, and as components for rigid polyurethane foams. Aminolysis was first reported in the 1960s using a variety of primary and secondary amines. In 2010, Soni and coworkers depolymerized PET to terephthalamides using hydrazine in an uncatalyzed reaction at ambient temperature and pressure. After 12 hours, the product was isolated and reacted with acryloyl chloride to provide a UV curable acrylic oligomer, demonstrating a potential application of this depolymerization technique to generate upcycled products. In 2013, Bhatnagar and coworkers working for the Indian Oil Corporation received a patent for aminolysis of virgin or post-consumer PET using (poly)amines to generate diamino mixtures with excellent bitumen anti-stripping properties. Reactions with quantitative product yield were run in xylene at 110–160 °C for 4 hours without a catalyst.

Son and Singh reported the aminolysis of PET in the presence of aqueous methylamine or ammonia using cetyl ammonium bromide as a catalyst in 2005. They found that the catalyst reduced the required time for complete degradation to N,N'-dimethylyterephthalamide to 45 days at 40 °C, while large PET flakes were still visible in the uncatalyzed reaction after the same time.

Ammonolysis of PET was much slower, and full depolymerization was not achieved under any tested conditions. In 2006, Shukla and Harad reported the aminolysis of PET using excess ethanolamine and simple catalysts including acetic acid, sodium acetate, and potassium sulfate. The product bis(2-hydroxy ethylene)terephthalamide (BHETA) was obtained in yields of up to 91% from post-consumer PET and 83% from bottle PET after 8 hours at reflux. Sodium acetate was found to give the highest yields in all reactions.

Tawfik and Eskander used dibutyltin oxide as a catalyst in the aminolysis of PET with ethanolamine at 190 °C and ambient pressure. Pure BHETA was obtained in yields of 49–62% after 1–4 hours. Chan and coworkers used zinc acetate as a catalyst to depolymerize post-consumer bottle PET in the presence of tri- and tetraamines. The reactions were complete in 30 minutes at 190 °C, and no purification of the polyamine products was necessary before the authors used them in crosslinking reactions with ethylene glycol diglycidyl ether to form hydrogels with potential uses in removing industrial dyes from water and other environmental and agricultural applications (Fig. 35).

Fukushima and coworkers used TBD as a catalyst in PET aminolysis with a variety of amines. The terephthalamide products were obtained in moderate to high yields at reaction temperatures of 110–190 °C in less than 24 hours. The monomers have potential applications as building blocks for high performance materials with desirable thermal and mechanical properties. The same group used this catalytic system to depolymerize PET with o-phenylenediamine and 2-amino phenol to generate bis-benzimidazole and bis-benzoazole, which can be used in the synthesis of medicines and high-performance polymers.

Microwave irradiation has also been studied in PET aminolysis. Pingale and Shukla reported the microwave-assisted aminolysis of PET using simple sodium salt catalysts and excess ethanolamine in 2009. BHETA yields of up to 90% were obtained in only 5 minutes under microwave irradiation, whereas similar yields required 8 hours using conventional heating. Cleaned post-consumer PET bottles and pristine PET

![Fig. 34 Typical reaction conditions and products for PET aminolysis.](Image)
fibers were both effectively depolymerized using this system. Achilias and coworkers utilized microwave heating to quantitatively depolymerize PET to BHETA in only 5 minutes at 260 °C without the addition of a catalyst. Hoang and Dang reported the uncatalyzed aminolysis of PET to bis(2-aminooethyl)terephthalamide and its oligomers \(\alpha,\omega\)-aminoligo(ethylene terephthalamide), but Sharma et al. found that the same reaction could be carried out in only 10 minutes at 250 °C using microwave irradiation. They then used the obtained amides to synthesize bis-benzoxazines, which can be used as advanced performance thermosets.

The ammonolysis of PET involves the reaction of the monomer with excess ammonia to generate terephthalamide and EG as products. Terephthalamide can be further converted into platform chemicals \(p\)-xylenediamine or 1,4-bis(aminooethyl)cyclohexane. A patent from 1988 describes the ammonolysis of PET under 20 bar \(\text{NH}_3\) pressure at temperatures of 120–180 °C for 1–7 hours. A simple filtration and drying workup provided the product in >99% purity and >90% yield. The same patent describes the use of zinc acetate as a catalyst to facilitate the transformation at only 70 °C and provide terephthalamide in 87% yield.

9.1.6 Hydrogenolysis. Hydrogenolysis reductively depolymerizes PET under \(\text{H}_2\) pressure, typically forming 1,4-benzene-dimethanol (BDM) and EG, and sometimes providing deoxygenated products. BDM is a platform chemical that can be used to prepare polylactide-based thermoplastic elastomers, highly cross-linked polymers, and sulfonated aromatic resins. Organometallic catalysts have typically been used for these depolymerizations (Fig. 36).

![Fig. 35 Aminolysis of PET using a polyamine, followed by a crosslinking reaction to form a PET hydrogel.](image)

![Fig. 36 (A) Typical reaction conditions and products for PET hydrogenolysis and (B) organometallic catalysts used for PET hydrogenolysis by (I) Krall et al., (II) Fuentes et al., and (III) Westhues et al.](image)
In 2014, Krall and coworkers\textsuperscript{761} found that a ruthenium pincer complex developed by Milstein could be used as a catalyst in the hydrogenolysis of PET (Fig. 36B I). At 2 mol\% catalyst loading, 55 bar H\textsubscript{2}, and 160 °C in anisole/THF, quantitative conversion to BDM and EG was observed after 48 hours. Post-consumer PET soda bottles were used in these experiments without pretreatment. Fuentes \textit{et al.}\textsuperscript{762} found that a novel PNN pincer complex (Fig. 36B II) catalyzed PET hydrogenolysis with moderate BDM yields, but the catalyst was poisoned by the EG formed in the reaction. This precluded monomer yields higher than 73%. Westhues \textit{et al.}\textsuperscript{763} used a ruthenium triphos catalyst (Fig. 36B III) to depolymerize PET from a variety of sources, including water bottles, pillow filling, and a sports jersey. Reaction selectivity for BDM and EG were over 98% with most substrates, but relatively forcing conditions of 100 bar H\textsubscript{2} and 140 °C were required. An entire PET water bottle with PP cap and PE label was depolymerized without pretreatment. Quantitative conversion to BDM was observed, with no conversion of PP or PE. The other polymers could be filtered off and subsequently depolymerized.

Supported metal catalysts have also been used in PET hydrogenolysis. In 2020, Kratish \textit{et al.}\textsuperscript{764} selectively depolymerized PET to TPA and ethylene using a carbon-supported molybdenum-dioxo catalyst under only 1 bar H\textsubscript{2}. No solvent was required for the reaction, and TPA was obtained in 85% yield after 24 hours at 260 °C. The air and moisture stable catalyst could be recycled four times with no loss in activity, and the yield of TPA did not change when PET from a post-consumer beverage bottle was used as the substrate. Jing and coworkers\textsuperscript{765} used a Ru/Nb\textsubscript{2}O\textsubscript{5} catalyst to convert PET to benzene, toluene, and p-xylene in 84% total yield. The reactions were run in octane at 280 °C and 5 bar H\textsubscript{2} for 8 hours. The catalyst also successfully depolymerized a mixture of PET, polycarbonate (PC), PS, and polyphenylene oxide to a variety of aromatic monomers in 79% overall yields, showing the viability of this system in mixed waste streams. However, a 1 : 1 mass ratio of polymer to catalyst was required in this system. Wu \textit{et al.}\textsuperscript{766} prepared a N-doped carbon-supported bimetallic catalyst for the conversion of PET to TPA in 91% yield in 10 hours at 260 °C and 1 bar H\textsubscript{2}. No solvent was used in the reaction, and the catalyst could be reused at least six times with no observed decrease in activity. Almost the same yield of TPA was achieved with PET from bottles and with a mixture of PET and PP from bottles.

### 9.1.7 Biocatalysis

Given the ubiquity of natural ester-linked compounds in nature, including polymers such as cutin and suberin, enzyme catalysts have been pursued as another means to hydrolyze PET. These biocatalytic reactions have the advantage of enabling PET hydrolysis under mild conditions (30–75 °C and ambient pressure) in aqueous solutions to provide BHET, mono-2-hydroxyethylterephthalate (MHET), TPA, and EG with varying selectivity as products (Fig. 37). Many reviews have been written on this topic\textsuperscript{21,663,767–772} and it is a highly active area of research, thus we only discuss highlights here.

The first report of enzymatic hydrolysis of PET was in 2005 from Müller \textit{et al.}\textsuperscript{773} wherein they showed appreciable extents of PET conversion over 3 weeks at 55 °C using a cutinase enzyme from the thermophilic bacterium \textit{Thermobifida fusca}. This study prompted a search in genome and metagenome databases for additional cutinases capable of PET hydrolysis. In addition to the discovery and characterization of homologous enzymes from the genus \textit{Thermobifida} and closely related bacteria\textsuperscript{774–782}, Sulaiman \textit{et al.} reported the discovery of a leaf-branch compost cutinase (LCC) that has been subsequently engineered by Tournier \textit{et al.} for greater thermal stability and activity, enabling quantitative conversion of amorphous micronized PET substrates in 10 hours at 72 °C and enzyme loadings concomitant with industrial performance.\textsuperscript{783–785} Shirke \textit{et al.}\textsuperscript{786} demonstrated that the addition of glycosylation to the LCC enzyme \textit{via} expression in \textit{Pichia pastoris} results in greater stabilization of the enzyme, thus highlighting the potential for post-translational modifications as another means of enzyme stabilization and engineering for industrial applicability. A similar enzyme to LCC was recently reported by Sonnendecker \textit{et al.}\textsuperscript{787} that is able to achieve similarly high PET conversion with no substrate pretreatment. Additional studies to diversify the known PET hydrolytic enzyme suite will likely be enabled by computational methods and the massive number of genome and metagenome sequences that are continuously being reported.\textsuperscript{788,789}

The seminal discovery of \textit{Ideonella sakaiensis} by Yoshida and coworkers\textsuperscript{790} in 2016 highlighted that natural microbial systems are seemingly responding to the presence of PET in
the natural environment. In particular, *I. sakaiensis* secretes a two-enzyme system, comprising the enzymes PETase and MHETase, to fully depolymerize PET into TPA and EG via the intermediates BHET and MHET. This two-enzyme system has been extremely well studied to date, with structural biology techniques and many types of protein engineering and evolution approaches used to improve the stability and performance particularly of the PETase enzyme. The *I. sakaiensis* system also points to the potential utility of an enzyme cocktail for industrial application of an enzymatic recycling approach, as is common in industrial cellulose deconstruction, especially in this case to overcome product inhibition due to the buildup of MHET during enzymatic hydrolysis.

Singh et al. recently conducted a detailed techno-economic, energy, and GHG emissions analysis of enzymatic PET recycling using the patent literature and the work from Tournier et al. Therein, they estimated that with low-cost PET feedstocks, enzymatic recycling processes could achieve cost parity with virgin PET manufacturing at substantially reduced energy inputs and GHG emissions. This work directly highlighted two areas of prominence that merit further attention in process development for enzymatic PET recycling, which also may translate to other PET chemical recycling approaches discussed herein. Namely, it was estimated that approximately half of the energy input and GHG emissions arise from substrate pretreatment, which was modeled as thermal extrusion and cryo-grinding to yield micronized, amorphous PET. The development of enzymatic systems that can deconstruct crystalline substrates, which has been highlighted by studies as a major need for the field, would thus be a major step forward for this approach. Secondly, EG recovery from water was estimated to roughly equate to the other half of energy use and GHG emissions. Advanced approaches to separations for EG that operate in the condensed phase or employ reactive distillation may be of use to further improve this process and will likely be relevant for EG recovery in chemical solvolysis reactions as well.

9.2 Chemical recycling of polycarbonates and other polyester feedstocks

9.2.1 Polycarbonates. PC resins are generally distinguished by whether they have aliphatic or aromatic backbones. Aliphatic PC resins are not used as thermoplastics, while aromatic PCs are commonly used as engineering thermoplastics in electric and electronic equipment. Poly(BPA carbonate) (PBPAC) is the most commonly used PC and is the only PC resin that we will discuss in this section, though many chemical recycling techniques discussed previously for PET have been investigated on a laboratory scale for PBPAC, including pyrolysis, hydrolysis, alcoholysis, aminolysis, and hydrogenolysis (Fig. 38).

PBPAC pyrolysis produces a large amount of char due to the flame retardancy of the polymer. Uncatalyzed pyrolysis at 300–500 °C leads to the generation of BPA and phenolic compounds as the primary liquid products, leaving behind a residue which accounts for 20–30 wt% of the starting material. Metal chloride or oxide catalysts such as CuCl₂,
FeCl₃, ZnCl₂, SnCl₂, CaO, and MgO have been shown to lower the temperatures required for pyrolysis, narrow the product distribution, and decrease the amount of char waste formed in the reaction. Due to the hydrophobicity of PBPAC, hydrolysis of this polymer requires steam or supercritical conditions. At temperatures above 240 °C, BPA and CO₂ are the primary products of PBPAC hydrolysis. Alkaline earth metal oxides and hydroxides, alkali hydroxides and carbonates, and rare earth trihalides all catalyze PBPAC hydrolysis, enabling milder reaction conditions and higher BPA yields especially when an organic cosolvent such as 1,4-dioxane is used. Similar to hydrolysis of PET, liquid acid catalysts and microwave heating also facilitate this reaction.

To our knowledge, PBPAC alcoholysis was first reported with methanol, NaOH catalyst, and toluene cosolvent to swell the polymer and obtain BPA and methylene carbonate as products. The reaction has since been adapted to continuous flow, accelerated using microwave heating, and run under supercritical conditions. Alkali halide, heterogenious, ionic liquid, and organic base catalysts have also been employed to increase yield and selectivity for BPA under milder conditions. The aminolysis of PCs provides value-added ureas and BPA as products. The yield of monomeric products is increased when a diamine is used to form a stable cyclic urea. An ionic liquid/ZnO nanoparticle catalyst system provides excellent BPA and urea yields with a variety of mono- and diamines. Ammonolysis of PBPAC in an aqueous ammonia solution provides BPA and urea in high yields. Finally, many of the catalysts studied in PET hydrolysis also successfully depolymerize PCs under similar conditions to generate the corresponding alcohol monomer and methanol in high yields.

9.2.2 Other polyesters. Poly(butylene terephthalate) (PBT) (Fig. 39) can be used in similar applications as PET and also as an engineering thermoplastic in rigorous mechanical profile applications such as car bumpers. PBT resins have high strength and rigidity, low moisture absorption, and excellent electrical properties and chemical resistance. Due to their similar chemical structures, many of the PET depolymerization strategies discussed above also successfully depolymerize PBT.

PLA is a widely used, industrially compostable bioplastic derived from the fermentation of renewable feedstocks. PLA has good mechanical properties but a low glass transition temperature, narrow temperature-processing window, and undesirable brittleness. Depolymerization of PLA to the lactide monomer is not possible due to side reactions including elimination to acrylic acid and epimerization. The primary method used for the deconstruction of PLA on the lab scale is hydrolysis to lactic acid (Fig. 40). Additionally, some catalysts used in the hydrolysis of PET are capable of reductively depolymerizing PLA to form propylene glycol in high yields.

9.3 Commercial activity and economics

Much of the current commercial progress in PET chemical recycling is not captured in scientific publications, with patents and press announcements typically constituting the public information in this field. A recent review of advanced PET chemical recycling focuses primarily on the development of commercial processes. At the time of publication, commercial chemical PET recycling has not yet been realized. However, many commercialization projects exist using different depolymerization chemistries and subsequent downstream purification processes.

Mechanical recycling of PET is complementary to chemical recycling processes, and both are needed to achieve a high degree of circularity in PET applications. Chemical recycling feedstocks will comprise primarily colored and/or opaque plastic bottles and polyester-based textiles that are unsuitable for the high purity standards required in mechanical recycling. The availability of a high volume, low cost feedstock is critical for commercial process viability, which will require processing highly impure PET feedstocks to produce high purity, polymerization-grade monomers at scale. The quality of the output of rPET from chemical recycling is higher than that from mechanically recycled PET. This allows the rPET from chemical recycling facilities to be infinitely recyclable, whereas the rPET produced from mechanical recycling degrades during each cycle.

The commercial activity described below is in the research and development-to-pilot stage unless otherwise indicated. To our knowledge, hydrolysis, methanolysis, glycolysis, and enzymatic depolymerization are the current primary areas of commercial focus, but it is likely that additional technologies will soon emerge on the commercial landscape. The intention of the processes described here is the transformation of waste PET plastic and textiles to monomers for subsequent repolymerization to rPET with physical and mechanical properties identical to those of pristine material.

9.3.1 Hydrolysis. DePoly is a Swiss start-up that uses a room temperature basic hydrolysis process with TiO₂ or another metal oxide absorber in the presence of UV light.
The technology was originally developed at the Swiss Federal Institute of Technology Lausanne (EPFL). Tyton BioSciences, recently renamed Cric, is a start-up based in the US which uses alkaline water at high pressure and temperature, called hydrothermal processing, to hydrolyze PET fiber to TPA, EG, and products formed from the reaction of other materials such as cotton present in the textile waste stream. Gr3n, a Swiss-based start-up, announced the building of a demonstration plant in Italy in 2021. Their process features microwave-assisted hydrolysis that will be used to recycle both PET fiber and resin.

9.3.2 Methanolysis. Loop Industries, a publicly traded Canadian company, has signed multiple partnerships for a global rollout of their methanolysis technology. Using a patented process, the PET resin or fiber input is mixed with methanol, an organic swelling solvent, and an inorganic alkoxy catalyst. The reaction proceeds at a temperature below the boiling point of methanol and does not require added pressure. At the time of writing this review, a timeline on their website states that Loop Industries has raised over $116 million but as of yet does not have a commercial facility for their technology. Eastman Chemical has recently announced plans to build a $250 million, 100 kton per year PET methanolysis plant in Tennessee. They have been researching PET methanolysis for several decades, with patents in this field dating back to the early 1990s.

9.3.3 Glycolysis. There are many companies working to commercialize glycolysis processes with unique process concepts. JEPLAN, a Japanese start-up, uses a metal hydroxide catalyst and EG for PET glycolysis in a pilot plant in Japan. They utilize a two-step purification process involving crystallization and distillation to produce polymerization-quality BHET. Ioniqa, a spin-off from Eindhoven University, developed a process which features an imidazolyl base tethered to a magnetic iron oxide nanoparticle. The magnetic nature of the anchored catalyst facilitates simplified catalyst removal and recovery for reuse. Ioniqa has agreements with large consumer packaged goods companies in addition to PET producers and has recently announced a 10 kton per year facility. Researchers at IBM have taken a different path to enable the facile separation and reuse of a PET glycolysis catalyst. Their process is catalyzed by a volatile organic base that can be recovered by distillation after PET depolymerization is complete.

This technology, known as VolCat (Volatile Catalyst), is being scaled up in a recently announced joint venture with several commercial partners.

9.3.4 Biocatalysis. Commercial activity in biocatalytic PET depolymerization is highlighted by the publicly traded enzyme-catalyzed process from Carbios in France, who launched a demonstration plant in 2021. Carbios has announced partnerships including investors, end customers, enzyme producers, and process and plant designers to accelerate commercialization. Carbios has raised over $126 million to develop their technology, and they currently operate a 1.2 kton per year demonstration facility with goals to build a 40 kton per year facility by 2025.

10. Other catalytic approaches for plastics conversion

10.1 Hydrogenolysis with noble metals

Hydrogenolysis of plastics cleaves the carbon–carbon bonds in the presence of hydrogen converting the polymers into hydrocarbons (ranging from C1 to C50) typically with noble metal (e.g., Ru, Pt etc.) catalysts. In such processes, heterogeneous catalysts are used which enables simple separation and recycling from the reactants and products. Table 18 summarizes different technologies utilized in hydrogenolysis of waste plastics where plastics were converted in batch reactors. Ruthenium supported on metal oxides or carbon has been reported to catalyze the production of alkanes, aromatics, and liquid fuels from polyolefins (i.e., LDPE, HDPE, PP), PS, and PC. LDPE, HDPE, and PP can

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Polymer</th>
<th>Temperature (°C)</th>
<th>Pressure (Bar)</th>
<th>Time (h)</th>
<th>Polymer/catalyst (Mass)</th>
<th>Products</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ru/CeO3</td>
<td>LDPE, HDPE, PP</td>
<td>200–240</td>
<td>20–60</td>
<td>5–24</td>
<td>34</td>
<td>Liquid fuels, waxes (C5–C45)</td>
</tr>
<tr>
<td>Ru/TiO2</td>
<td>PP</td>
<td>250</td>
<td>30</td>
<td>8–16</td>
<td>20–40</td>
<td>Aromatic hydrocarbons</td>
</tr>
<tr>
<td>Ru/NbO3</td>
<td>PET, PS, PC</td>
<td>200–320</td>
<td>3–5</td>
<td>12–16</td>
<td>1–2</td>
<td>Liquid alkanes (C3–C13), light gases (C1–C6)</td>
</tr>
<tr>
<td>5Ru/C</td>
<td>PE, LDPE</td>
<td>200–225</td>
<td>20</td>
<td>16</td>
<td>25</td>
<td>Liquid alkanes (C5–C32), light gases (C1–C5)</td>
</tr>
<tr>
<td>5Ru/C</td>
<td>PP</td>
<td>250</td>
<td>35</td>
<td>8–24</td>
<td>14</td>
<td>Methane, light paraffins (C2–C11)</td>
</tr>
<tr>
<td>Ru/FAU</td>
<td>LDPE, PP</td>
<td>300–350</td>
<td>50</td>
<td>3</td>
<td>50</td>
<td>Lubricants, waxes, diesel (C4–C35)</td>
</tr>
<tr>
<td>Ru/WO3/ZrO2</td>
<td>LDPE</td>
<td>250</td>
<td>30</td>
<td>2</td>
<td>40</td>
<td>Lubricants, liquid fuels (C6–C38)</td>
</tr>
<tr>
<td>Ru/C</td>
<td>HDPE</td>
<td>220</td>
<td>30</td>
<td>1</td>
<td>2</td>
<td>Liquid fuels (C5–C22)</td>
</tr>
<tr>
<td>Pt/WO3/ZrO2 + zeolite</td>
<td>LDPE</td>
<td>250</td>
<td>30</td>
<td>2</td>
<td>10</td>
<td>Lubricants, waxes (Mw 200–1000 Da)</td>
</tr>
<tr>
<td>Pt/SrTiO3</td>
<td>PE</td>
<td>300</td>
<td>12</td>
<td>96</td>
<td>5</td>
<td>Fuels, lubricants (C8–C32)</td>
</tr>
<tr>
<td>SiO2/Pt/SiO2</td>
<td>HDPE</td>
<td>300</td>
<td>14</td>
<td>24</td>
<td>88</td>
<td>Lubricants (C5–C45)</td>
</tr>
<tr>
<td>Pt/C</td>
<td>PP</td>
<td>300</td>
<td>15</td>
<td>24</td>
<td>10</td>
<td>Lubricants (C5–C45)</td>
</tr>
</tbody>
</table>

Table 18 Hydrogenolysis of plastics into various hydrocarbons in batch reactors

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be converted to liquid fuel (C5–C21) and lubricants/waxes (C22–C45) at low temperature and low H₂ pressure (i.e., 200–250 °C, 20–30 bar) on metal oxides-supported Ru (e.g., Ru/TiO₂, Ru/CeO₂). Over multifunctional Ru/Nb₂O₅, monocyclic aromatics can be selectively produced from single or mixed aromatic plastics (i.e., PET, PC, PS, polyphenylene oxide) at 200–320 °C in the presence of hydrogen and solvent (e.g., water, octane etc.). Ru hydrogenation sites with low coordination numbers (i.e., 5–6) prevent the over-hydrogenation of the aromatics, while the surface Lewis acid and Bronsted acid sites on Nb₂O₅ species act in concert to selectively absorb, activate C–O bonds, and cleave C–C bonds of the plastics. The support can be adjusted to treat different plastics and obtain varied products. Over Ru/WO₃/ZrO₂, LDPE can be converted into higher molecular weight fuels and wax/lubricant base-oils at 250 °C and 50 bar H₂. While on Ru/FAU, methane (>97%) can be produced from PE, PP, and PS under 50 bar H₂ at 300–350 °C. Ru nanoparticles can also be supported on carbon (Ru/C) to catalyze the hydrogenolysis of LDPE, HDPE, and PP towards liquid n-alkanes and alkane gases (C₁–C₆) under mild conditions (200–250 °C, 20–50 bar H₂). In the liquid phase (water, n-hexane etc.), around 90 wt% of HDPE can be converted to liquid hydrocarbons (C₈–C₃₈) within 1 h on Ru/C catalysts at 220 °C in the presence of H₂ (30 bar). Over platinum supported on strontium titanate or mesoporous shell/Pt/silica at 250–300 °C in the presence of H₂ (10–17 bar). Moreover, platinum can be supported on carbon which catalyzes the conversion of PP into liquid hydrocarbons (C5–C45) or on WO₃/ZrO₂ and mixed with zeolites which catalyzes the production of liquid fuels and short-chain hydrocarbons from PE, PP or PS. The synergistic effects between platinum and supports (carbon/metal oxides) or co-catalysts (zeolites) are vital in these processes. Additionally, the hydrogenolysis of plastics can also be carried out over Al₂O₃ or SiO₂ supported transition metals (Cr, Ni, Mo, Co, or Fe), micro-mesoporous zeolites, and other catalysts which are not discussed here.

10.2 Functionalization and reactive extrusion

Another strategy to valorize plastics is the addition of other chemical groups (e.g., aromatic ring, halogen, carboxylate, carbon–carbon unsaturated bond etc.) to these polymers via chemical methods (e.g., oxidation, halogenation, esterification, metathesis) as shown in Fig. 41. This approach is typically performed by reactive extrusion in an extruder. The products include polymers with new properties, liquid fuels, waxes, and platform molecules. The C–H bonds in PE (HDPE, LDPE, LLDPE) can be oxidized to ketone or hydroxyl groups, which generates functionalized polymers with unique physical properties (e.g., strong adhesion, ability to be painted with common paint etc.). A high-valent ruthenium-oxo catalyst (polyfluorinated ruth-
nium porphyrin) can be added with 2,6-dichloropyridine
N-oxide to form esters from PE at 120 °C. The same complex
can catalyze the oxidation of polyisobutene to o xo-pol y isobu-
tene (Fig. 41B) where the catalysts, reaction conditions, and
oxidants are vital in selectively oxidizing the methylene posi-
tions and avoiding chain cleavage.\textsuperscript{884} Reactions besides oxi-
dation can be utilized to functionalize the plastics. Kanbur
et al. reported that the conversion of polyolefins (PE, PP) to
shorter aliphatic alky laluminium species can be achieved \textit{via}
β-alkyl elimination and heterobimetallic alkyl metathesis
using organozirconium catalysts and aluminium reagents at
200 °C.\textsuperscript{885} The obtained shorter aliphatic alky laluminium
species can be further converted into alkyl alcohols, acids,
halides, and hydrocarbons with a quenching process. Cross-
alkane metathesis can convert PE into liquid fuels and waxes
(short-chain linear alkanes) at 175 °C.\textsuperscript{881} This process consists
of three steps: (1) the dehydrogenation of PE and cofed light
alkane; (2) the metathesis of olefins; (3) the hydrogenation
of the obtained olefins. An iridium complex can catalyze steps 1
and 3 while \( \text{Re}_2\text{O}_7/\gamma\text{-Al}_2\text{O}_3 \) catalyzes the olefin metathesis.
The catalysts are compatible with various polymer additives and
product distribution can be manipulated by the catalyst struc-
ture and reaction time.

Functionalized polymers can also be produced by dehydro-
genation, sulfonylation, amination, and vitrimerization. Zinc
sulfide is active in PP dehydrogenation\textsuperscript{886} while an iridium
complex can catalyze the dehydration of PE and poly(1-hexane)
in the \( p\)-xylene and norbornene mixture.\textsuperscript{887} For sulfonylation,
it has been reported that PS can be sulfonated by chlorosulfo-
nic acid, fuming/concentrated sulfuric acid, sulfur trioxide,
and acetyl sulfate.\textsuperscript{888–891} Tian \textit{et al.} reported that Ru/TiO\textsubscript{2}
can catalyze the amination of ether groups in PLA in ammonia
solution at 140 °C (Fig. 41C).\textsuperscript{883} Alane can be produced by
this process with high selectivity in the absence of hydrogen.
In most of the cases, other chemicals (e.g., light alkanes,
water, halogens) need to be cofed for the functionalization of
the waste plastics. However, the cofed species is not necessary
if the carbon and hydrogen in the polymers can be utilized.
One approach is to partially depolymerize the plastics. Long-
chain alky laramatics (ca. C\textsubscript{30}) can be produced from PE via
controlled depolymerization on Pt/\( \gamma\text{-Al}_2\text{O}_3 \) without solvent or
\( \text{H}_2 \) at 280 °C.\textsuperscript{882} This process couples exothermic hydrog enoly-
sis and endothermic aromatization which enables simulta-
naneous C–C cleavage, ring closure, and dehydrogenation
under mild conditions (280 °C). Vitrimerization is a process
converting polymers to vitrimers, which behave like silic a,
\textsuperscript{892} are covalent networks and can change their topology by ther-
mally activated bond-exchange reactions. Vitrimers can be pro-
duced from both thermoplastics (e.g., PB\textsubscript{T}\textsuperscript{893}) and thermoset
plastics (e.g. polyurethane\textsuperscript{894}). The key step in vitrimerization
is introducing covalent adaptable networks \textit{via} addition of cat-
alysts or cross-linkers. Although catalyst is not necessary for
vitrimerization, the properties of a catalyst-containing vitrimer
can be efficiently tuned.\textsuperscript{893,895}

Reactive extrusion is a commercially practiced approach to
install functionality of plastics.\textsuperscript{896} One example is the radical-
mediated addition of maleic anhydride to PE and PP.\textsuperscript{897} One
key step in reactive extrusion is the production of a pair of radicals
which can be achieved through heating or photolysis of
the reagent. One of the radicals cleaves a C–H bond in plastics
and a chain transfer step can functionalize plastics (Fig. 41D).
Thiocarbonyl amide reagent has been utilized to functionalize
branched polyolefins.\textsuperscript{898} The authors demonstrated that this
metal-free C–H functionalization of isotactic PP can be per-
formed on a decagram scale forming polymers with improved
adhesion to polar substrates. Recently, another study demon-
strated that an ideal reagent (e.g., \( O\)-alkeny lhydroxamate) enables
slow chain transfer kinetics which provides various
functionality to the backbone of plastics by other radical
sources.\textsuperscript{899}

The functionalization of waste polymers is being commer-
cialized (e.g., Novoloop\textsuperscript{900}). The company can recycle waste
LDPE, HDPE, and LLDPE. Those PE wastes are shredded,
cleaned, and fed into a patented, accelerated thermal oxidative
decomposition (ATOD) reactor (Fig. 42).\textsuperscript{900,901} The treated PE is
then heated to a temperature range from 60 to 200 °C, and oxi-
dized into dicarboxylic acid monomers by a group of oxidation
agents including oxygen, nitrous oxide, nitric oxide, and
aqueous nitric acid.\textsuperscript{962} The oxidized monomers are used to
produce thermoplastic polyurethane (TPU).

Plastic alloys are a type of functionalize plastic where
different plastic materials are functionalized to create a single
plastic phase.\textsuperscript{903,904} The properties of the plastic alloy can be
tuned by adding other materials (e.g., waste fibers) which
enables applications of plastic alloys in automotive, electrical
& electronics, construction, home appliances, and healthcare
equipment.\textsuperscript{905–908} The advantages of plastic alloying are short
reaction times, no need for solvents, and low operating/capital
costs compared with other plastic technologies like
extrusion.\textsuperscript{909}

Plastic alloying can introduce long chain branching of the
polymer backbone to improve the properties (e.g. melt
strength) of plastics. This approach has been used for functionalization of PP,\textsuperscript{905} PE,\textsuperscript{910} and polyhydroxybutyrate
(PHB).\textsuperscript{906} Long chain branching can be added using peroxides,
which can result in increases in melt elasticity.\textsuperscript{908} Long chain
branching can alter physical properties such as the crystallinity
and transition temperatures.\textsuperscript{911} The thermophysical properties
including crystallinity, melting behavior and tensile properties
of HDPE were shown to be influenced by di-tert butyl cumyl
peroxide induced cross-linking.\textsuperscript{910} Dicumyl peroxide (DCP)-
induced cross-linking in PE improves the melt strength and is
dependent on peroxide concentration.\textsuperscript{912} The lifetime of the
peroxide formed from DCP is relatively close to the residence
time of extrusion. The generally accepted process of peroxide
induced cross-linking of polymers follows three key steps:\textsuperscript{913}

1. The primary radical formation from the thermal
decomposition of peroxide,
\[ \text{RO} \rightarrow \text{OR} \rightarrow \text{2RO'} \] (Primary radicals)
2. H abstraction from polymer backbone by free radicals of
peroxide generated in step (1),
\[ \text{RO'} + \text{P (polymer)} \rightarrow \text{P'} \] (polymer radical) + ROH
(3) the bimolecular radical recombination of polymer radicals from step (2) to generate C–C cross-links.

\[ 2P^* \rightarrow P-P \text{ (crosslinking)} \]

There is limited information on the chemical position(s) associated with cross-linked molecular chains to form a gel network structure. Long chain branching increases the plastic molar mass, polydispersity, melt strength and strain hardening in PP.° There is one key step in plastic alloying is C–H activation which can be achieved by different routes.°

Plastic alloying has been explained by inter-linking and interfacial adhesion between different polymers. Wigotsky pointed out the importance of plastic alloying to the plastic industry because of the improved physical properties of the alloys.° The compatibilizing agents in plastics act as polymeric surfactants which lowers surface tension and promotes interfacial adhesion between polymer phases in plastic alloys.° In addition to this, PP functionalization (i.e., carboxylation) generates compatibilizers which can be used to produce plastic alloys.° Sakai reported a processing technology which can improve compatibility of plastic and produce plastic alloys using twin-screw extruders.° Asay et al. reported that PVC can be utilized to produce plastic alloys because of the varied degree of miscibility between PVC and a wide range of polymers.° Composite materials with improved properties can be obtained via addition of other materials. Kimura showed significant improvement in tensile strength by adding fillers to polyolefins.° Amor et al. studied the improvements of properties of PLA with biomass, demonstrating synergistic effect of small addition of PLA on the thermomechanical properties.° The above studies portray the importance of plastic alloying for the production of plastic and composite material products. However, these studies were all conducted with pure polymers by mixing two or three components. Aumnate et al. were the first to recognize that plastic waste could be also alloyed.° In their fundamental study, they simulated plastic wastes by mixing pure components of LDPE and HDPE with PP and showed the effects of the branching of PEs on mechanical properties and crystallization behavior.

Another plastic alloying approach is the “grafting onto” method which grafts the PHB polymer onto cellulose fibers through plastic alloying processing with the use of a small amount of DCP.° LDPE has also been shown to graft onto cellulose fibers, retaining the stiffness of cellulose and the flexibility of the polymer matrix.° When the peroxide is exposed to heat during extrusion, it will decompose into strong free radicals which tend to abstract H’s from the polymer and cellulose molecular chains and initiate the grafting between the two phases in composites.

10.3 Microwave-assisted conversion

Microwave radiation can be applied during catalysis to heat the reaction system, shorten reaction times, and enhance the kinetics of some chemical reactions.° In conventional thermal heating processes, the whole reaction system (catalysts, reactor, diluent, feedstock etc.) are heated while microwave radiation enables the electromagnetic energy to be absorbed evenly and efficiently at the microwave-adsorbing points (e.g., catalyst surface).° Furthermore, other researchers propose that the microwaves likely have intrinsic catalytic effects or assist in lowering the activation energy of the reactions.°

Microwave-assisted depolymerization and catalytic upgrading can be separated by the reactor system.° In microwave-assisted pyrolysis, LDPE is depolymerized into pyrolysis vapor and the microwave absorbent (e.g., SiC) enables even
heating of the feedstock which promotes the formation of light olefins. The products are then catalyzed by MgO to produce more valuable chemicals (i.e., aromatics, C12+, hydrocarbons) in a subsequent reactor. This process can be modified by changing the microwave absorbent to NiO and the catalyst in the separated reactor to HY zeolite which enhances the production of olefins in the pyrolysis zone and produces gasoline fuel with high yield as well as appropriate carbon number. Microwave-assisted conversion can also be performed with catalysts in the liquid phase. LDPE powder can be completely degraded in 1 h of microwave irradiation at 180 °C in the presence of dilute nitric acid solution (0.1 g mL\(^{-1}\)). In this process, water is used to absorb microwaves energy and nitric acid catalyzes the oxidation of LDPE towards dicarboxylic acids (e.g., malonic acid, succinic acid, glutaric acid etc.). Bäckström et al. pointed out that HDPE can be converted to dicarboxylic acids with a similar approach and the obtained products can then be utilized to synthesize plasticizers. Addition of the plasticizer to PLA improves the physical properties of the polymers. Much faster degradation can be achieved via microwave-assisted conversion combined with appropriate catalysts. Jie and coworkers reported that polymerized plastics (HDPE, LDPE, PP, PS) can be deconstructed into hydrogen and high-value carbon materials (e.g., carbon nanotubes) in 30–90 s. FeAlO\(_x\) plays a significant role in this process by both efficiently absorbing microwave energy, which initiates the physical heating process, and by catalyzing the depolymerization of the plastics.

Additionally, continuous processing and cofeeding of biomass can be applied to microwave-assisted conversion of plastics. Zhou et al. reported a continuous microwave-assisted pyrolysis system which can convert HDPE and PP to gasoline-range hydrocarbons rich in aromatic hydrocarbons. The system consists of microwave heating with a SiC mixing-ball-bed and a secondary catalyst bed with ZSM-5 and can process up to 10 kg of plastics an hour with 89.6% energy efficiency. Considering the oxygen deficiency of polyolefins, it can be used as a hydrogen source to be co-processed with hydrogen-deficient biomass via microwave heating, thus improving the quality of the bio-oil. Zeolites and metal oxides as well as the combination of these two materials have been reported to catalyze the production of hydrocarbons from a mixture of LDPE and lignin, where microwave-assisted heating is vital in achieving even and efficient internal heating.

10.4 Biocatalysis

This section will focus on the biocatalysis of other plastics as the application of biocatalysis for PET depolymerization has been discussed in Section 9. The biological breakdown of plastics by enzymes refers to an attack on water immiscible plastic polymers by appropriate microflora. Plastics are degraded by microbial flora’s enzymatic activity, which results in polymer chain fragmentation into monomers. Degradation is caused by the activity of enzymes secreted by microorganisms, which use hydrocarbons in the polymer backbone as their major carbon source. Enzymatic plastic degradation proceeds in two steps. The first step is the decomposition of polymers into smaller molecular compounds by attacking main chains of the polymer with abiotic and/or biotic reactions. The second step is the assimilation of the decomposed polymer (monomer, dimer, or oligomers) by means of microbials and its mineralization which creates CO\(_2\), H\(_2\)O, and CH\(_4\).

The degradation of HDPE by exposure to white rot fungus, Bjerkandera adusta TBB-03, in a lignocellulose substrate treatment has been demonstrated. TBB-03 is able to effectively improve laccase activity (211.4 U L\(^{-1}\)) in the presence of ash wood chips due to a mediator resulting from lignin decomposition. The mediator is oxidized by laccase and may play a role as a strong oxidant by producing radical intermediates, which results in the significant morphological change of HDPE.

Sakiriyachan and coworkers reported that thermophilic consortia of Aneurinibacillus and Brevibacillus sp. showed degradation potential for LDPE, HDPE and PP films and pellets. The percentage weight reduction for LDPE, HDPE and PP strips in the consortia of isolates at 50 °C in 140 days was calculated to be 58.21%, 46.6% and 56.3%, respectively. The report on the enzymatic degradation associated with PVC was discussed by Khatoon et al. Phanerochaete chrysosporium is able to degrade PVC film with 31% weight loss at 25 °C in the 4th week of the incubation period with the fungal strain. Furthermore, Sumathi et al. isolated a fungal species, Cochliobolus sp., for production of laccase, from plastic dumped soils and then determined the ability of the fungi to degrade low molecular weight PVC under lab conditions where the synthetic polymer served as the only carbon source for the Cochliobolus sp. They observed significant differences from the FTIR, GC-MS, and SEM results in between control and Cochliobolus sp. treated PVC.

11. Conclusions and outlook

Compared to other engineered materials (i.e., glass, metal, paper etc.), plastics are lightweight and can be easily manufactured into products for a variety of applications. Generation of plastic waste and the ratio of waste plastics in MSW have increased due to the growth of plastic production. Recycling of plastics is still far behind where it needs to be for a circular plastics economy, but improved technology is continuously being developed to efficiently increase the lifetime of plastics.

11.1 Collecting and sorting of waste plastics

The first step in any plastic recycling involves collecting and separating the plastics. There is an opportunity to divert more material from landfills through an upgrade in the recycling infrastructure. This can be achieved by an increase in the number of facilities, such as MRFs, or by improving process efficiency along the recycling process, such as improved sorting by adding more optical sorters. However, this will result in additional capital costs for MRFs. Education and outreach regarding recycling is also critical, as consumers need to...
learn the environmental, social, and economic benefits of recycling.

The operation of a MRF presents multiple challenges. One is the difficulty in finding personnel. Another challenge is the lack of predictability of the inbound stream, as each week, the received material can fluctuate according to holidays. For example, in the period of the year close to the Thanksgiving and Christmas holidays, there is an increase in food containers (from dinner parties) and fibers (ads from the shopping season). In terms of the materials collected, some MRFs view the presence of full-body shrink sleeve labels on PET bottles and aluminum cans as a challenge, as it affects the recyclability of two significant sources of income (PET and aluminum). Other challenges include the presence of films and flexible packaging, shredded paper, and packaging with mixed types of material (e.g., plastic bottles with metal closures).938

Increased contamination of the waste material stream is also an issue, especially with material collected from areas not used to having recycling programs, which often have less awareness of the waste sorting procedures. For example, lithium batteries are a growing concern due to the danger of starting fires, with 68 incidents reported in MRFs between 2013 and 2020.939,940 The challenges associated with the presence of contaminants in the bales could result in lower yields per bale, showing an opportunity to improve bale quality. Such measures would include an increase in sorting efficiency by adding more optical sorters, expansion of recycling education to reduce contamination in inbound streams, or even construction of new MRFs.

In developing countries, the waste pickers also need support. The main needs of waste pickers include improving their working conditions, learning to manage their family needs, receiving education and training, preventing violence and guidance to reduce teenage pregnancies, alcoholism and drug addiction, and training to “produce something” from the waste that gives them more income.941 Some actions that would lead to fair living wage waste picking jobs include: (a) develop strategies for precise censuses of waste picker communities and to have dynamic updates of this information; (b) integrate waste pickers in the recycling value chain; (c) harmonize recycling strategies with frontier technologies and train the waste pickers; (d) develop innovative business models for MSW management with a social perspective; (e) update and enforce the legal and regulatory framework of recycling value chains in compliance with human rights of waste pickers; (f) integrate and establish social programs for waste pickers, from public, private, local, and international organizations.

11.2 Mechanical recycling of waste plastics

Mechanical recycling works well for a very homogeneous plastic stream like HDPE milk bottles. However, contamination and other plastics decrease the quality of the plastics produced during mechanical recycling. Some plastics cannot undergo mechanical recycling or do not flow upon heating after cure (thermosets) as they cannot tolerate thermal treatment. Multi-layered plastic products (i.e., juice bottles) cannot be mechanically recycled. These multi-layered packaging materials are more challenging to recycle as each plastic in a different layer has a different melting temperature, and different plastics are often immiscible which makes them practically non-recyclable.942 Colored plastic products, for example PET containers with carbon black in them, are not detected by the sensors during the sorting stage of recycling and complicate the recycling process.943 Films produce a low quality product during mechanical recycling. To improve the quality of the products from mechanical recycling, more advanced separating techniques can be used to obtain feedstock with high a degree of homogeneity. In addition, additives can be added to the extruder to improve the quality of the resin.

11.3 Pyrolysis, liquefaction, and gasification of waste plastics

Pyrolysis, liquefaction and gasification have the advantage of being able to process mixtures of plastics and more easily dealing with contamination than mechanical recycling. Several plastic pyrolysis approaches are being commercialized. Catalytic pyrolysis is used to increase product selectivity.944

Despite continuing advancement of the technology, commercialization of plastics pyrolysis faces significant challenges which can be addressed by further research. Better understanding of the complex reaction mechanisms controlling depolymerization of plastics could improve selectivity to desired produced molecules. Novel approaches to process intensification will be important to overcome the combination of relatively modest chemical reaction rates and heat and mass transfer limitations that characterize many pyrolysis processes. Using process intensification to achieve economies of scale, determining optimal reactor and process conditions, and securing consistent feedstock streams will increase the profitability of pyrolysis plants. In addition to this, the reactors used in pyrolysis need to be well designed to enable efficient mass and heat transfer. A better understanding of approaches to convert the plastic pyrolysis oils into chemicals, plastics and fuels is needed as steam cracking of the oil has several challenges.

Liquefaction is another technology currently being commercialized. The major challenges of liquefaction are high costs related to the equipment and operation conditions. Supercritical HTL also increases reactor corrosion. Enabling a high feedstock mass loading in the solvent without significantly affecting liquefaction efficiency is also critical, as it is less efficient to heat a large amount of solvent and convert only a small amount of plastic waste. According to Licella, feedstock loading must be sufficiently high (i.e., 70%) for their CAT-HTR process to be economical. Developing an improved conversion strategy to produce higher quality products with minimal required upgrading is also essential. While lab studies indicate that feedstock compositions and reaction conditions strongly impact liquefaction product yield and quality, fewer studies discuss cross-interactions among different plastics. There is also a lack of information about the fate of contaminants and pollutants during liquefaction. HTL of polymers also promotes oxidative reactions, which become more
noticeable under supercritical conditions. The effect of oxygenated compounds on the direct use of the crude oil or during subsequent upgrading should be addressed in future studies.

Gasification of MSW shares similar challenges and opportunities as other fossil-based gasification technologies, primarily associated with the high capital cost of gasification. Converting synthetic gas to heat and power produces energy that competes with the declining costs of solar and wind power generation which are now less than $0.04 per kW per h.945 Pursuing higher-value chemicals such as ethylene which is valued at more than $1000 per metric ton946 is possible but requires several catalytic steps (methanol synthesis and methanol to olefins). The varying composition of waste plastic streams presents an additional operational and management challenge for recycling facilities. An incentive program tailored to reduce the risk of economic losses due to market price volatility could serve as a complement or alternative to environmental regulations. Waste reduction policies and incentive programs will likely spur the development of waste gasification facilities.947

11.4 Dissolution-based approaches for waste plastics

Dissolution-based plastic recycling approaches can have advantages over other recycling technologies as they produce the virgin resins. Proper solvent selection for specific types of plastic waste is the key to the feasibility of dissolution-based plastic recycling. Many solvent systems for common polymers have been disclosed in the literature and in patents by different companies. However, when considering more complicated plastic waste feedstocks, advanced thermodynamic computational tools are a means to expedite the development and design of these processes. The recycled materials obtained via dissolution/precipitation methods could be economically competitive with virgin resins, showing the capability of these approaches to reduce plastic waste. These techniques can also be applied to complex multicomponent input streams (e.g., multilayer films) which are challenging for current mechanical recycling technologies.

Several challenges remain in the design of dissolution-based recycling techniques that merit future study. The use of large amounts of solvent in these processes poses potential challenges associated with toxicity and energy consumption.631,948,949 To address these concerns, methods for solvent selection based on polymer solubility can be combined with techno-economic and life cycle analysis techniques to assess tradeoffs in solvent selection.38,47,634 Another challenge is the removal of potential contaminants (e.g., residual inks or retained solvent) from the recovered polymers.588,950

The kinetics of dissolution also could hinder dissolution-based recycling for large amounts of plastic waste.609,651 These challenges could be addressed through additional processing steps for contaminant removal or mechanical shredding prior to dissolution; such steps will require further investigation. Finally, there are opportunities to explore the integration of dissolution-based techniques with liquid-phase chemical recycling techniques.669,951 For example, selective dissolution could be used to separate target polymers from mixed plastic waste prior to their chemical depolymerization, thereby mimicking pretreatment methods utilized in biomass conversion. In this context, solvents could be selected to promote both polymer solubility and desired reaction outcomes.

11.5 Chemical recycling approaches for polyesters and other plastics

New chemical recycling processes centered around catalysis are emerging. PET can be depolymerized via many different routes generating chemicals to produce rPET or other commodities (Fig. 28). For the hydrogenolysis of waste plastics, the product distribution can be tuned by adjusting the residence time, catalyst to polymer mass ratio, temperature, and H2 pressure etc. Detailed reaction mechanisms for the hydrogenolysis of plastics should be elucidated to provide structure–function relationships at the molecular/atomic level which will provide insights in the rational design of new catalysts with improved efficiency or cheaper metal centers. The products from hydrogenolysis are mainly paraffins and light gases. Converting these back into plastics would involve dehydrogenation followed by steam cracking or another type of processing. The cost of H2 and safety issues related to its use are major concerns in the industrialization of such processes. Continuous flow of feedstocks with uniform properties might be challenging due to the solid state of waste plastics. It would be ideal to design techniques which enable continuous feedstock flow with high polymer/catalyst ratio and short reaction times. Furthermore, noble metal centers are likely to be poisoned by the impurities in the feedstock, which also needs to be considered in the development of catalytic materials and processes for waste plastics hydrogenolysis.

In the functionalization of plastics, product selectivity and atom efficiency are two challenges since side reactions (e.g., undesired chain cleavage, over-functionalized etc.) easily occur under reaction conditions and an excess of reactant is often required to provide the functional groups. The impurities in waste plastics likely lead to contaminated products or poison the catalysts, which should also be considered in developing these techniques. The removal of impurities in feedstocks needs significant effort in the sortation and pretreatment of waste plastics. Instead, virgin plastic resin can be utilized to manufacture some of the functionalized polymers. Besides, some of the processes need long reaction times (i.e., up to 4 days), which needs to be addressed in scaling up the techniques. Plastic alloying has several advantages compared to other recycling technologies. The key advantages include: (1) little sorting is needed prior to forming the plastic alloys; (2) every type of plastic (#1–#7) can be recycled via this approach; (3) less manpower is needed to operate a plastic alloy facility compared to a MRF and the secondary recycling facility. Compared to pyrolysis of waste plastics, the advantages are: (1) higher ratio of the mixed plastic waste stream can be recycled; (2) fewer numbers of steps are required which results in lower capital/operating costs; and (3) higher value materials than a plastic oil can be produced. More work on this technology is
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needed to understand if it can be applied to real plastic waste feedstocks, understand the relationships between the catalysts/additives and identify the properties of the plastic alloys, as well as the desirable properties of the plastic alloys.

The removal of impurities in the feedstock is also one of the challenges in microwave-assisted conversion. The impurities that can absorb microwave energy likely lead to varied local temperature, which then results in wide product distribution. Accurate parameters (i.e., microwave power, radiation intensity, magnetic permeability etc.) need to be standardized for the development of this technique. The capital cost and energy efficiency should also be considered in scaling up microwave-assisted techniques for plastic recycling. As for the biocatalytic processes, it is important to standardize and improve the efficiency of the isolation procedure for microbial species which contain plastic degrading enzymes so that pure breeding colonies of these microorganisms can be quickly screened. Enzyme production from pure colonies can be boosted by standardizing the initial step, which enhances the overall performance of biocatalytic plastic degradation.

Overall, enhancing plastic recycling requires focused efforts, particularly in chemical recycling technologies. Efficient approaches to process waste plastics need to be developed to decrease plastic waste and pollution.

Conflicts of interest

The authors declare conflict of interest.

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