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Recycling of polyurethanes: where we are and where we are going

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Polyurethanes (PUs) represent a family of useful synthetic polymers (thermoplastic or thermosetting) obtained from diisocyanates and diols/polyols *via* polycondensation reactions. Within the circular economy concept and also considering the current need for limiting the environmental impact of plastics, several methods have been designed, assessed, and exploited for the recovery at the end-of-life of polyurethanes and for their recycling. Indeed, the processing of polyurethane wastes can be significantly beneficial not only from an ecological but also from an economic point of view. At present, feedstock (namely, glycolysis) and mechanical recycling are the two most important strategies to recover and recycle polyurethanes; notwithstanding, "biological recycling", an approach that exploits the biological degradation of the polymer, is gaining interest. This review aims to elucidate the recycling processes of both thermoplastic and thermosetting polyurethanes, providing the reader with some perspectives about their possible future developments.

Introduction

Versatility is undoubtedly one of the main characteristics exhibited by polyurethanes (PUs), an important "family" of thermoplastic and thermosetting polymers containing a remarkable number of urethane groups ($-\text{HN}-\text{COO}-$), derived from isocyanates and hydroxyl-containing compounds *via* the reactions presented in Fig. 1a. This reaction is exothermic and occurs spontaneously even at room temperature; nevertheless, one or more catalysts are commonly used to speed up the reaction. The hydroxyl-containing compound could be a diol, polyol, or water, which is used as a chemical blowing agent for producing polyurethane foams. Indeed, isocyanate reacts with water, forming an unstable substituted carbamic acid that decomposes into an amine species together with carbon dioxide, leading to the blowing of the polyurethane polymer (Fig. 1b). The as-obtained primary amines may react again with isocyanate groups giving a substituted urea (Fig. 1c). At room temperature, this reaction is faster than the urethane formation. Active hydrogen atoms of substituted urea compounds or allophanates (formed by the reaction of isocyanates with urethanes, Fig. 1d) may react again with isocyanates yielding a biuret (Fig. 1e). Isocyanates are highly reactive, so they can also react themselves leading to polymerization reactions; dimerization yields symmetric or asymmetric uretdiinediones;

trimerization leads to the formation of isocyanurate rings that are responsible for the high thermal stability of polyisocyanurate foams.

Isocyanates could be aliphatic or aromatic and bifunctional, however, their functionality could be higher than 3. For some specific applications, such as flexible foams, elastomers, and thermoplastic polyurethanes, prepolymeric isocyanate, *i.e.*, an

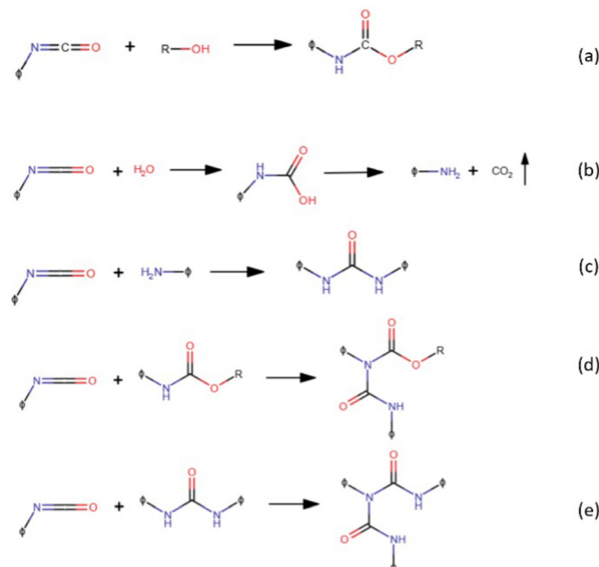


Fig. 1 Reaction between isocyanates and hydroxyl-containing compounds (a), water (b), amine (c), urethane (d), and urea (e).

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NCO-functional reaction product of aromatic or aliphatic isocyanates and polyols, is commonly used.¹ Similarly, a large variety of polyols are offered, but most of the polyols used fall into two major categories: hydroxyl-terminated or amino-terminated polyols. The structure of the polyols has a direct bearing on both processing and finished properties of the polyurethane polymer. In fact, the majority of the linkages found in polyurethanes are derived from the linkages found in the polyols.¹ Polyols differ from each other with regard to the molecular weight (up to 6500 Da), hydroxyl number (up to 1000 mg KOH per g) and functionality (up to 8). Polyols with a higher molecular weight and a lower hydroxyl number and functionality are used for elastomers, coatings and flexible foams, while the others are employed for elastoplastics, rigid coatings and rigid foams.¹

The wide availability of these co-reactants accounts for the possibility of tailoring the final properties of the obtained PUs, hence manufacturing from thermoplastic to thermosetting polymers, from flexible to semi-rigid and rigid polyurethanes (depending on the ratio between soft segments and hard segments²), which can be employed in different application sectors, including upholstered furniture (as flexible foams), wall and roof insulating rigid foams, thermoplastic footwear and medical devices, commercial refrigerators, adhesives and sealants, and coatings.³ A general classification scheme of PUs and some of their main current industrial applications are presented in Fig. 2 and Table 1. Besides, in EU27 + 3 countries, polyurethanes represent around 7.8% of all the produced plastic materials, which places them in 6th place in the EU27 + 3 plastics demand rank.⁴

The advantage of employing polyurethanes for replacing synthetic plastics (such as polystyrene, polyvinyl chloride, and synthetic rubbers) or natural polymers (*i.e.*, leather) is undeniable, also considering the possibility of using bio-sourced polyols, which reduces the environmental impact of PUs, making these polymer systems greener than the fossil-based counterparts.⁵ Further, waterborne polyurethane coatings take advantage of the use of water instead of potentially toxic organic solvents.⁶

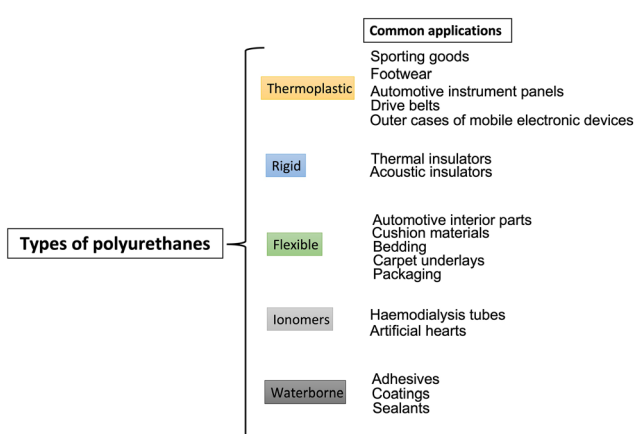


Fig. 2 Polyurethanes: classification of general uses.

Table 1 Main applications of polyurethanes and related production value

Sector	Main applications	Production (%)
Binders, adhesives, and sealants	Sealants, casting, assembling of wood boards, rubber or elastomeric flooring surfaces	10
Coatings	Bumpers & side panels of cars	14
Elastomers	Medical uses	8
Flexible foams	Mattresses, seating, cars	36
Rigid foams	Packaging, appliances, insulation boards	32

Among the most interesting peculiarities, apart from enhanced electrical or adhesive properties, PUs exhibit high durability (higher than polyvinyl chloride) and high resistance toward different solvents/compounds (namely, water, organic solvents, and oils).⁷ These characteristics justify the very extended lifetimes of several polyurethane-based products, parts and components; however, the extension of their life cycle may have an important environmental impact, due to the accumulation of post-consumer polyurethane garbage, also considering the wearing out over time of such PU-based products as shoes or clothes, or the rapid exchange of some others (like sports equipment, cars, and furniture) for diverse or enhanced counterparts. Therefore, as witnessed by the increased number of publications in peer-reviewed journals on this topic (Fig. 3), the end-of-life recovery and recycling of polyurethanes are currently of extreme importance, in order to avoid landfill confinement (though it is still the most common strategy to exploit, despite the decreasing disposal of landfill spaces⁸) or incineration (*i.e.*, energy recovery⁹), due to the toxicity of some products originating during the combustion process. Further, the recovery and recycling strategies of polyurethanes are motivated by the significant depletion of world reserves of such fossil fuels as crude oil, *i.e.*, the raw material, on which the chemistry of polyurethanes is substantially based.

In general, the recycling processes can be classified as follows:¹⁰

- Primary recycling: the recovered plastic is used in products with performance characteristics that are equivalent to

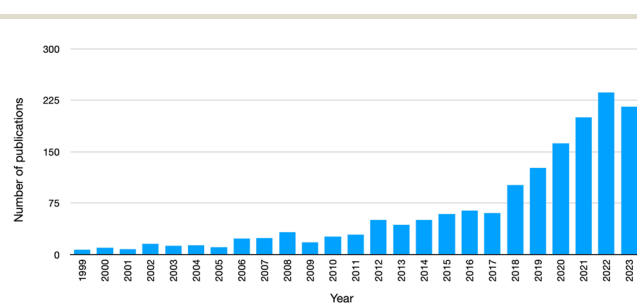


Fig. 3 Number of publications (from 1999 to 2023) in peer-reviewed journals, dealing with "recycling AND polyurethanes" (data collected from the Web of Science™ database, accessed on Oct 27, 2023).



those made using virgin plastics. This comprises closed-loop mechanical recycling.

- Secondary recycling: the recovered plastic is used in products that have less demanding performance requirements than the original application. This comprises open-loop mechanical recycling as well as physical recycling.

- Tertiary recycling: the waste plastic is used as the feedstock in a process that generates chemicals (*i.e.*, the monomers and/or other compounds) and fuel. For this reason, it is also called feedstock recycling and comprises both chemical and thermochemical recycling. Microbial degradation still falls under this category.

- Quaternary recycling: energy is recovered from waste plastics by incineration.

These recycling strategies have been applied to polyurethanes, and they are presented in Fig. 4. Each one shows several peculiarities that not always are very well known by both the academic and industrial communities, especially by the people who are not specifically familiar with this research topic. A comparison among them has been reported by Yang *et al.*,¹¹ and detailed explanations will be given in each specific section in the following.

At present, the main recovery and recycling routes for polyurethanes, which are the most preferred on a commercial/industrial scale, include mechanical, physical and feedstock recycling that is based on degradation and/or depolymerization reaction. For this latter, hydrolysis or glycolysis represent the recycling strategy that has been mainly exploited: in fact, it allows for the recovery of polyols that can directly be re-used for the manufacturing of new polyurethane systems (*i.e.*, regenerated polyurethanes), replacing up to 50% of virgin polyols with the recovered counterparts in polyurethane foam technology. The three bonds most susceptible to hydrolytic degradation are the ester, urea, and urethane. The ester reverts to the precursor acid and alcohol; this precursor acid further catalyzes ester hydrolysis, and thus the reaction becomes autocatalytic. Because of the autocatalysis of ester hydrolysis, this is the most prevalent hydrolytic degradation reaction. The urea bond can hydrolyze to form a carbamic acid and an amine. The carbamic acid normally is unstable and typically undergoes further reactions. The urethane linkage, although some-

what less susceptible, may undergo hydrolysis to yield a carbamic acid and the precursor alcohol.¹

In terms of thermo-chemical process, what is relevant is the thermal stability of the bonds within the polymeric chain. As explained before, there are several chemical groups in a polyurethane chain. The thermal degradation rate and onset are highly dependent on the specific group considered. The onset of allophanate dissociation is around 100–120 °C, while the dissociation temperature of the biuret linkage takes place around 115–125 °C. These reactions are dissociations and somewhat reversible. They revert to the urethane or urea, from which they were formed. The aromatic-based urethane bond begins its thermal dissociation around 180 °C, *i.e.*, before the thermal dissociation of the urea linkage (occurring at about 160–200 °C). The urethane can dissociate into the isocyanate and polyol from which it was formed. This reaction can also lead to the formation of amines and olefins and generates CO₂, which is lost as a gas.¹ The isocyanurate group is very stable and its decomposition starts around 300 °C.^{12,13}

While the ester is the weakest link in hydrolysis, now it is the ether that is the weakest link in thermooxidation. The order of stability of polyethers to thermooxidation is as follows: polytetra methylene ether glycol is more stable than poly(ethylene oxide) glycols that are, in turn, more stable than poly(propylene oxide) glycols.¹

In this context, the present review aims to provide the reader with an overview of the recycling processes of both thermoplastic and thermosetting polyurethanes, suggesting some perspectives about their possible future developments, also considering the strong current need for sustainability and low environmental impact of the designed recycling strategies.

Primary and secondary recycling of polyurethanes

Undoubtedly, primary and secondary (*i.e.*, mechanical and physical) recycling represents the easiest and most basic strategy to recycle PUs, apart from landfill confinement. It is cost-effective and the investment for process equipment is quite low, though many times the performance of the recycled products is decreased, thus reducing the market potential with a low economic benefit.

In these recycling processes, the materials have to undergo several processing steps first that are needed for reducing the original particle size to an appropriate level for being reprocessed in secondary manufacturing events. To this aim, polyurethane waste products (comprising old recycled components or production wastes – such as scrap parts and trimmings –) are converted into powders, pellets, or flakes, on the basis of the specific type of polymer that is recycled. Several methods can be successfully employed for fragmenting polyurethanes, namely: shredding, tearing, cutting, and grinding. It is worth noting that the average size of the fragmented particles is strictly related to the fragmentation technique: as an example, two-roll milling can be exploited for obtaining polyurethane

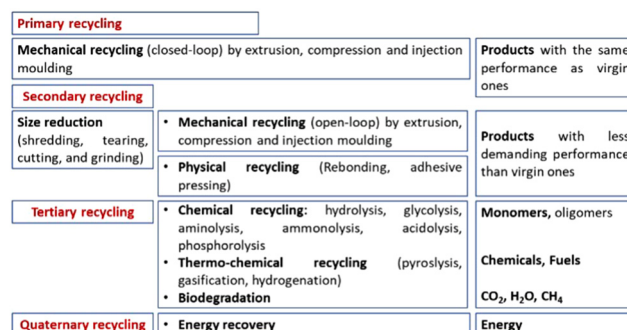


Fig. 4 Possible treatment and recycling processes for polyurethane wastes.



particles having an average size below 100–125 µm, which can be employed as a filler in newly processed polyurethanes. Conversely, polyurethane powders having a larger particle size, between 125 and 250 µm, can be obtained by knife cutting. Further, pellet mills can be exploited for preparing PU pellets.

The following steps depend on the nature of polyurethanes, since it shall be kept in mind that the recycling processes are quite different for thermoplastic or thermosetting polyurethanes. Since, by definition, thermoplastic polyurethanes (TPUs) can be melted, they can be mechanically recycled using the common method employed for thermoplastic polymers in general, *i.e.*, by extruding the recycled materials in the form of flakes or pellets, in the presence of some virgin polymers and/or compatibilizers.^{14–17}

Conversely, thermosetting PUs, at least highly crosslinked PUs such as foams, cannot be recycled in such way, but what is generally called “physical recycling” is the eligible method. Physical recycling implies the direct reuse polyurethane wastes, still in form of powders, flakes or small particles without chemical treatment,¹¹ and thus it can be considered among the secondary recycling methods. The PU particles are mixed with another component that will create a continuous matrix in which the particles are embedded. Generally, the use of heat and pressure is required to obtain the final recycled material. Physical recycling method is simple and convenient, with low cost, but there are still certain technical limitations. The performance of the physically recovered products is usually poor, so these are suitable for non-demanding applications only, thus limiting the market potential.¹¹ The different mechanical and physical recycling processes are described in more detail below.

Mechanical recycling by extrusion, compression and injection moulding

Thermoplastic polyurethanes (TPUs) can be reprocessed by means of extrusion, hot compression, or injection moulding, without the use of any binder and obtaining, at the same time, up to 100% recycled material. As mentioned before, the first step in the mechanical recycling is the fragmentation of PU wastes.

A general problem of mechanical recycling is related to the degradation of the polymer properties after each recycling cycle, because of the degradation caused by reprocessing (thermo-mechanical degradation) that sums to the degradation during lifetime (thermo-oxidative one).¹⁸ In each processing cycle, the material can undergo irreversible thermal and mechanical degradation, which leads to physical and chemical changes and a detriment of the final properties.¹⁹ Therefore, the material cannot be subjected to extensive reprocessing cycles,^{17,20} and the number of reprocessing cycles is generally limited to two or three.¹⁷ Sometimes, oligomers have been proposed as chain extenders or recycling aids which, when reacted in simple extrusion or injection moulding equipment with virgin, post-industrial recycle, or post-consumer recycled polyurethanes, effectively revert molecular weight degradation even at very small use levels.²⁰

The possibility of PU waste melting (and bonding in the melted state) is strongly affected by the degree of cross-linking, which increases in the sequence: elastomers < flexible foams < rigid foams. Thus, some reaction injection moulding (RIM) and reinforced RIM (RRIM) polyurethanes, which are strictly speaking thermoset polymers, thanks to their very low cross-linking degree can be still mechanically recycled.²¹ For example, polyurethane scraps (such as car bumpers at the end of their life; very often the PU wastes primarily derive from RIM polyurethanes) are reground into fine particles that are subsequently compressed at high temperatures (about 180 °C) and pressures (around 350 bar). In this way, it is possible to fabricate such rigid components as motor and pump covers; besides, the incorporation of glass fibers allows for obtaining strengthened parts for door panels and dashboard panels (containing about 6 wt% of reground PU and 15 wt% glass fibers).²² Further, it is possible to use polyurethane particles as fillers for other resins (as an example for polyester resins), taking advantage of the toughening effect exerted by the PU particles compared to mineral fillers. Finally, this technique exhibits two main drawbacks: the difficulty of managing the presence of dyes and pigments in the polyurethane wastes and the limits in the obtainment of finely ground particles.²³

When finely ground polyurethane particles cannot be obtained, it is possible to exploit the so-called structure reaction injection moulding (SRIM), employing up to 30 wt% of coarsely ground polymer scrap as an inner layer in a sandwiched assembly made of two glass fiber layers, which, in turn, are covered by a two-component PU resin (Fig. 5).

As assessed by Bayer,²⁴ it is also possible to exploit the injection moulding process to recycle relatively low crosslinked polyurethanes: for this purpose, polyurethane granules (size between 250 and 1000 µm) were employed for producing automotive parts through injection moulding carried out at around 180 °C and high shear compression (beyond 350 bar).

Further, Hulme and Goodhead²⁵ demonstrated the suitability of dual injection moulding processes for recycling and reusing thermoplastic and thermosetting polyurethane wastes and scraps: this technique is based on the concurrent injection moulding into a single mold of a skin (made of virgin polyurethanes) and a core (made of recycled polyurethanes), as shown in Fig. 6. Its main advantages refer to the possibility of obtaining parts and components with enhanced mechanical features and good aesthetical properties (with improved surface finish); besides, it is possible to reprocess colored polyurethanes (as materials for the core).

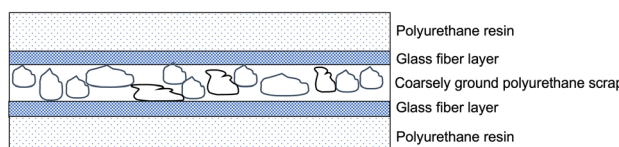


Fig. 5 Recycling of PU scraps through structure reaction injection moulding (SRIM).



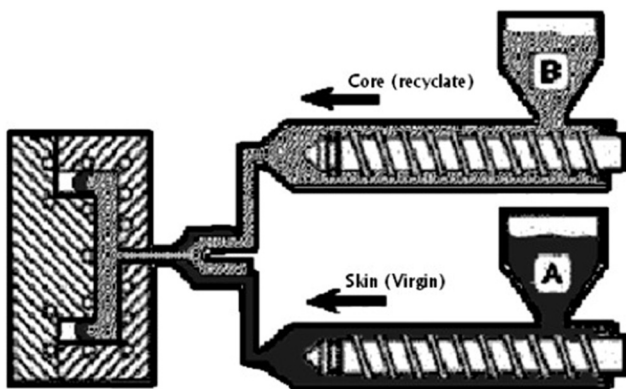


Fig. 6 Scheme of the recycling and reuse of polyurethanes by dual injection moulding: two injection machines concurrently inject recycled (B) and virgin (A) polyurethanes; the former is employed for the core of the injection molded part, while the latter, which exhibits good aesthetic properties, is used as an external layer (skin). Reproduced with permission from reference.²⁶ Copyright 2007 Elsevier.

Rebonding re-processing (physical recycling) with adhesives

Polyurethane flakes or granules (hence particles showing an average size of a few mm) are coated with an adhesive at 5–10 wt% loading (such as poly(phenylenemethylene isocyanate) or 4,4'-diphenylmethane diisocyanates), as shown in Fig. 7. Fig. 8 shows the schematic of the rebonding of a flexible PU foam, while Fig. 9 displays the distribution of the foam flakes within the adhesive.

Then, a hot water stream (usually at high temperatures, beyond 100 °C) or steam melts the polyurethane that subsequently undergoes compression (usually between 30 and 200 bar). The steam reacts with the prepolymer or binder, which leads to the formation of rebonded PU product. Usually, the squeezed foam flakes are fully saturated with steam, which permits maximum contact with the binder; generally, several minutes are required to saturate compressed PU foams with steam. The rate of reaction can be controlled using catalysts.²²

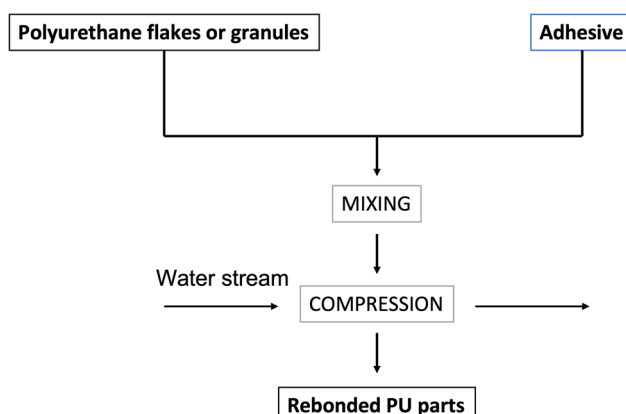


Fig. 7 Scheme of the rebonding process for polyurethane flakes or granules.

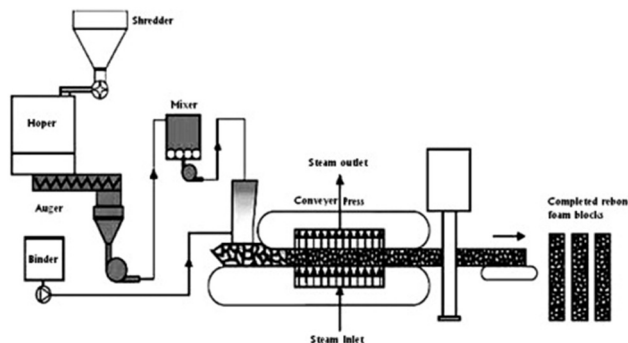


Fig. 8 Schematic of flexible foam re-bonding. Reproduced with permission from ref. 27. Copyright 2007 Elsevier.

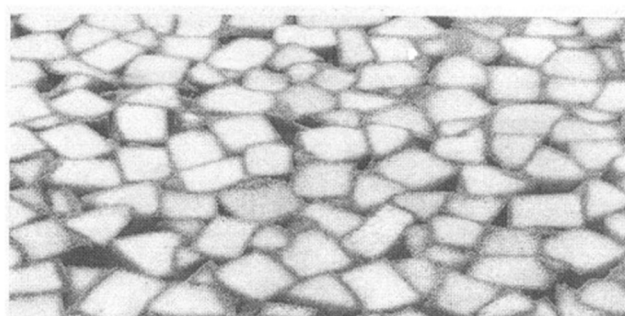


Fig. 9 Rebonded foam: homogeneous distribution of the flake-binder mixture. Reproduced with permission from ref. 27. Copyright 2007 Elsevier.

The binder is a crucial component in the production of rebonded PU foams.

This process allows for obtaining contoured parts that can be employed as covers for tires, athletic mats, car linings, and carpet underlays, among other uses.²⁷ Besides, it is possible to manufacture PU building panels exhibiting outstanding insulation features and high resistance to moisture, although characterized by quite a high density and, therefore, with some limits for market exploitation.²⁷

In a recent application of the rebonding strategy, Salino and Catai demonstrated the possibility of using polyurethane foam residues and recycled plasterboard sheet cores for acoustic applications.²⁸ In particular, they found that the recycled materials exhibited high porosity (about 47%) and interesting sound absorption coefficients over a wide range of frequencies (from 250 to 2000 Hz).

Chemical (tertiary) recycling of polyurethanes

Under suitable conditions, it is possible to make the polymerization of polyurethanes partially reversible: generally speaking, gradual depolymerization may occur when the polymer reacts with organic compounds bearing active hydrogen

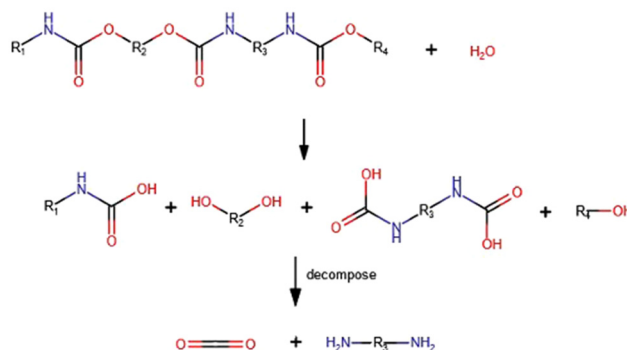
atoms, able to attack the polar groups of the polymer main chain. The most liable chemical groups, involved in the depolymerization process, have been reported in the “introduction” section. In general, depolymerization can be based on the cleavage/exchange of urethane bonds or depolymerization based on the cleavage/exchange of chemical bonds of the precursors used for PU production, such as the ester groups of the polyol.²⁹ This review will primarily focus on depolymerization reactions based on depolymerization involving urethane or urea groups.

The possibility of recovering monomers and oligomers from the polymer is strictly correlated with the adopted experimental conditions, namely: selected reactive organic compounds, temperature, pH, pressure, and atmosphere, as well as the type(s) of employed catalysts. After purification and distillation processes carried out on the attained depolymerized products, it is possible to recover monomers with high purity (namely, amines and polyols), which can be exploited for synthesizing not only polyurethanes but also other polymers (such as polyamides, polyesters, and polyureas). Based on the selected degradation pathway, it is possible to recover products with different functionalities and physicochemical features. The most performed processes for the chemical recycling of polyurethanes include hydrolysis, glycolysis, aminolysis, ammonolysis, acidolysis and phosphorolysis, which will be described in the following. It is noteworthy to mention that, compared to mechanical recycling, chemical recycling processes allow for the recovery of higher-value products; besides, although closed-loop recycling is possible, at the same time, these processes are more complicated than the mechanical ones. Further, selected polyurethane wastes are required and higher energy consumption is needed. Hence, the majority of these processes have not achieved the industrial and commercial scale. A full review of the chemolysis process that has gone beyond the limits of the laboratories, in pilot-plant or industrial scale, has been reported by Simón *et al.*³⁰ more attention is paid to glycolysis since it is by far the most implemented on a larger scale. In general, chemical recycling is applied to thermoset PUs, while for the thermoplastic ones, as well as for PUs with low crosslinking degrees (*e.g.*, RIM and RRIM), the primary and secondary recycling methods explained before are the preferred option.

Hydrolysis

Hydrolysis can be considered the first chemical technique developed for recycling foamed polyurethane wastes (both as post-consumer wastes and as production scraps). It refers to the reaction of the polymer with water, either in the liquid state or as vapor (steam),³¹ which gives rise to the formation of different products comprising carbon dioxide, polyols, and amine intermediates (see Scheme 1).

It should be noted that hydrolysis is almost always a side reaction to chemical recycling of polyurethanes, since any PU foam contains absorbed water (between 5 wt% in viscoelastic foams and 0.5 wt% in aged rigid foams). Water will promote side reactions giving undesired products, especially primary



Scheme 1 Hydrolysis of polyurethanes.

aromatic diamines, such as 4,4'-diaminodiphenylmethane (MDA) or 2,4- and 2,6-tolylene diamines, which are carcinogenic substances.³² This point is discussed in more detail in the “glycolysis” section.

Hydrolysis shows drawbacks, as it is carried out under high operating conditions, *i.e.*, high pressures (often between 15 and 50 atm) and temperatures (even 300–400 °C). It is worth noting that the polyol yield is strictly related to the severity of the adopted conditions. In this context, in a pioneering work, Campbell and Meluch³³ demonstrated the suitability of the hydrolysis reaction, carried out between 232 and 316 °C and at atmospheric pressure, for recovering about 20 wt% of polyols from a polyurethane foam, which was then reused for the synthesis of new polyurethane foams. Further, Gerlock and co-workers³¹ studied the hydrolysis of a poly(ether-urethane) foam between 190 and 230 °C: the reaction yielded high-quality polyols, CO₂, and toluene-diamines.

Although it has been reported several times in the scientific literature that hydrolysis can be performed using bases (such as sodium hydroxide) or acids (like hydrochloric acid), basic conditions seem to provide a more efficient process with higher conversions with respect to the acidic reaction medium. Further, acid catalysis combined with increasing hydrolysis temperatures was found to account for enhanced efficiency.³⁴

Then, sodium hydroxide was employed in combination with diaminotoluene, and their catalytic effects on the hydrolysis of post-use polyurethane foams were thoroughly investigated by Dai and co-workers.³⁵ In particular, the concurrent use of the two co-catalysts accounted for the decrease in both depolymerization activation energy and reaction temperature; however, the energy required for the overall process was very high and there were some difficulties in the purification of the resulting polyols.

In a further research effort, Nikje and co-workers³⁶ proposed the use of mixtures of binary (*i.e.*, water and glycerol) or ternary (*i.e.*, water, sorbitol, and glycerol) solutions for chemical recycling polyurethanes, either using a thermal treatment carried out at atmospheric pressure and 160–180 °C, or through microwave irradiation.³⁷ This way it was possible to exploit the synergistic effect provided by the combined action



of hydrolysis and glycolysis (in the so-called “hydroglycolysis”), employing milder conditions (temperatures around 200 °C and lithium hydroxide as a catalyst) with respect to hydrolysis only. The obtained decomposition products were employed for synthesizing new polyurethane foams.

Further, Goto³⁸ demonstrated the possibility of carrying out hydrolysis depolymerization reactions on polyurethanes by using sub- or super-critical water.

Actually, hydrolysis processes have only been scaled up to a pilot plant but, in no case, they have achieved the commercial scale due to the high energy input required for the reactor.³⁹ Moreover, the recent use of super and/or subcritical fluids (e.g., CO₂ and water) proposed in the literature⁴⁰ does not seem to be able to overcome this limitation.

Moreover, at the end of the hydrolysis process, as well as hydroglycolysis, it is necessary to completely separate the amines from polyols, if the amines are to be used to produce new isocyanates. The recovered polyols can be used in formulations for making a PU foam, preferably in admixtures with virgin polyethers.³² This separation step is quite complex since it requires solvent extraction, and this hinders the suitability of such a recycling method for industrial applications. Therefore, although the energy requirement of hydroglycolysis is lower than that of hydrolysis, the former is also not applied on an industrial scale for its separation and purification cost.

Glycolysis

Reacting polyurethanes (mainly rigid bulk materials, rigid and flexible foams) with diols at temperatures usually beyond 200 °C and under atmospheric pressure allows for depolymerizing these materials and recovering polyols or glycolysates (*i.e.*, oligomers embedding urethane groups and showing higher molecular weights compared to polyols), which, in turn, can be exploited for producing new polyurethanes. The reaction, called glycolysis (a general scheme is presented in Fig. 10) usually takes place with the glycol, in the presence of a catalyst (such as acetate inorganic salts, hydroxides of alkaline metals, amines, metallo-organic compounds – stannous octoate, dibutyltin dilaurate). It is very important to finely tune the reaction temperature, as the activity of the catalysts is not enough when the reaction is carried out below 180 °C. Conversely, when the glycolysis temperature exceeds 220 °C, undesired side reactions may occur, which lead to the formation of amines rather than polyols or glycolysates.

After completion, the glycolysis products are cooled down, filtered, and possibly mixed with virgin polyols to reformulate polyurethanes. The overall process requires about 8 h. Several parameters may affect the reaction yields, as well as the quality of the obtained recyclates, namely, temperature and reaction time, mass/molar ratio of polyurethane scraps to glycols, possible stirring of the glycolysis reactor, type and concentration of the employed catalyst, glycol type and size of the polyurethane scraps.^{41,42}

Glycolysis involves two main reactions, namely, the break of biuret and allophanate groups, and the transesterification of urea and urethane bonds. Further, when poly(ester-urethane)

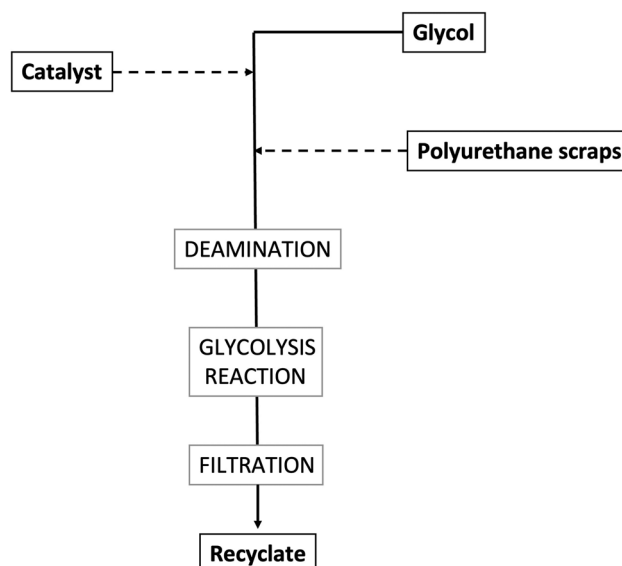


Fig. 10 Scheme of the glycolysis process for polyurethane wastes.

scraps are processed, esterification of the polyol-rich moieties (*i.e.*, of the soft segments of the polymers) may occur. Besides, it is very important to limit the presence of amines in the recyclates, as amine groups are more reactive than OH functionalities with isocyanates, leading to faster crosslinking phenomena in the new polyurethanes obtained from the recyclates. Moreover, aromatic amines, formed by PUs comprising aromatic isocyanates (*i.e.*, the largest part of PU) are carcinogenic and thus their content must be limited below 1000 ppm to avoid the need for labelling according to classification and labelling of chemical (CLP) regulation. Indeed, if the concentration of aromatic amines in a polyol exceeds this limit, the polyol can still be used but must be labelled, handled and transported accordingly, increasing the degree of complexity for the managing of the supply chain of recycled polyols for PU production.⁴³ Aromatic amines are formed through the transesterification of urea groups or by the side reaction of urethane with water that is always absorbed in PU wastes or by thermal degradation of urethane because of the high temperature required by the glycolysis process.

It is worth underlining that it is possible to carry out glycolysis reactions without any catalyst, but with much longer reaction times; however, there is a need for higher temperatures, which may account for degradation phenomena involving the processed materials.

Concerning the effect of the mass/molar ratio of polyurethane scraps to glycols, the scientific literature clearly demonstrates that it is possible to perform the glycolysis process with a mass excess of either glycols⁴⁴ or polyurethane scraps.⁴⁵ Table 2 presents some examples of glycolysis processes performed with a mass excess of polyurethane scraps: the main advantage derived from the glycolysis with the mass excess of polyurethanes refers to the quite high homogeneity of the obtained products (single-phase), which do not need



Table 2 Some examples of glycolysis processes performed with a mass excess of polyurethane scraps

Type of polyurethane scrap	Glycol or diol employed	PU : glycol mass ratio	Catalyst	Reaction temperature (°C)	Ref.
Flexible PU foam	Diethyleneglycol	1 : 0.5	Potassium acetate	215–225	57
Flexible poly(ether-urethane) foam	Diethyleneglycol	1 : 0.9	Stannous octoate	179	58
Flexible PU foam	Ethyleneglycol	2 : 1; 5 : 1; 8 : 1, 10 : 1; 20 : 1	Potassium acetate	195	59
Flexible poly(ether-urethane) foam	1,4-Butanediol	2 : 1; 5 : 1; 8 : 1; 10 : 1	Potassium acetate	210–230	60
Flexible poly(ether-urethane) foam	Poly(ethylene glycol)	2 : 1	Potassium acetate	210–220	45
Flexible PU foam	1,6-Hexanediol	2 : 1; 4 : 1; 6 : 1; 8 : 1	Potassium acetate	230–245	50
Flexible PU foam	1,3-Propanediol, 1,4-butanediol, 1,5-pentanediol	10 : 1	Potassium acetate	190–250	61
PU elastomer	Ethyleneglycol	2 : 1; 5 : 1	Potassium acetate	170–180	62
Rigid PU foam	Diethylene glycol, ethyleneglycol, propyleneglycol	2 : 1	Sodium acetate	190	53

the purification and distillation of glycol excess, as for the process operating with a mass excess of glycol (split-phase). Moreover, the higher the PU/glycol ratio, the lower the investment cost for equipment because of the smaller reactor volume required for the same amount of scraps to be recycled.

Table 3 presents some examples of glycolysis processes carried out with a mass excess of glycols: this condition is adopted when the main target is the recovery of polyols that, after purification and distillation, can be exploited for synthesizing new polyurethane systems.⁴⁶

In general, the formation of one or separated phases depends, not only on the PU/glycol ratio, but also on the type of PU waste. The glycolysis of rigid foams generally provides a single homogeneous phase of high viscosity and high hydroxyl number, which can be reused, often mixed with virgin polyols, to make PU rigid foams again. Conversely, this product cannot be used to prepare flexible foams because of the too high hydroxyl number that is due to the high functionality and short molecular chain of these kinds of glycolysates. In case of flexible PU foams, because of their higher urea content, generally a split-phase product is obtained (Fig. 11): the upper phase contains the recovered original polyols, to be eventually

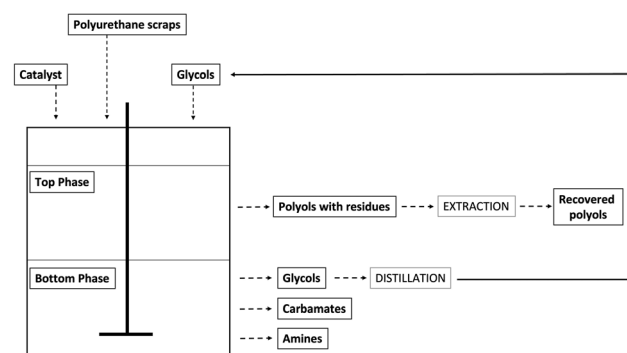


Fig. 11 Split-phase glycolysis.

further purified by means of distillation, while the bottom phase consists of the glycol excess, together with carbamates and aromatic amines (depending on the type of processed polyurethanes).

The main advantage of split-phase glycolysis with respect to single phase is related to the high quality of the recovered polyol;^{47–49} however, split-phase glycolysis processes present a

Table 3 Some examples of glycolysis processes performed with a mass excess of glycols

Type of polyurethane scrap	Glycol or diol employed	PU : glycol mass ratio	Catalyst(s)	Reaction temperature (°C)	Ref.
Flexible poly(ether-urethane) foam	Diethyleneglycol	1 : 1.5	K, Ca and Sn(II) octoates	189	63
Flexible poly(ether-urethane) foam	Diethyleneglycol	1 : 1.5	Diethanolamine, Ti butoxide, K and Ca octoate	189	47
Flexible poly(ether-urethane) foam	Diethyleneglycol	1 : 1.5	Sn(II) octoate	190	44
Flexible poly(ether-urethane) foam	Diethyleneglycol	1 : 1.3; 1 : 1.5;	Potassium octoate	190	48
Flexible poly(ether-urethane) foam	Ethyleneglycol, diethyleneglycol, propyleneglycol, dipropyleneglycol	1 : 2			
Flexible poly(ether-urethane) foam	Ethyleneglycol	1 : 1.5	Diethanolamine	189	49
Flexible PU foam	Poly(ethylene glycol)	1 : 2	Potassium acetate	195	59
Flexible poly(ether-urethane) foam	Ethyleneglycol/diethyleneglycol	1 : 2	Potassium acetate	210–220	45
Poly(ether-urethane) elastomer	Ethyleneglycol/diethyleneglycol	1 : 2	Li Acetate	170	64
Poly(ether-urethane-urea) fibers	Diethyleneglycol	1 : 3	Dibutyltin dilaurate	210	65
Rigid PU foam	Diethylene glycol, Dipropyleneglycol	1 : 1, 1 : 2, 1 : 3	Potassium acetate	220	54
Polyisocyanurate	Dipropyleneglycol	1 : 3, 2 : 3, 9 : 11	Potassium acetate, titanium (IV)-n-butoxide	200–230	66



great drawback relying on the huge molar excess of glycol required that causes an important increase in the operation costs, hence making almost impossible the implantation of industrial glycolysis plants.³⁰ Although split-phase situation is more common for flexible foams, the effective final product separation still depends on the PU/glycol ratio. For example, Datta *et al.*^{50,51} reported that the single-phase product is obtained from an elastic PU foam using hexamethylene glycol (HDO) when the foam/HDO weight ratio is up to 4/1; for higher ratios (6/1, 8/1, and 10/1), the reaction runs as 'split-phase' glycolysis.

Therefore, there is no general rule to define the usable weight ratio of glycol to PU.

Not only the weight but also the type of employed glycol greatly influences the glycolysis process. Dipropylene glycol and tetra ethylene glycol dissolved PUF in the shortest time among polypropylene glycols and polyethylene glycols (*e.g.*, DEG), respectively. PUF dissolution time was reduced to one-half for each 10 °C rise in the range of 170–200 °C. Moreover, PUF dissolution time was inversely proportional to the KOH (catalyst) concentration, while the dibutyltindilaurate concentration had less influence on the PUF dissolution time than the KOH concentration. Furthermore, smaller PUF particles dissolved in a shorter time. More specifically, the initial glycolysis conversion of PUF was proportional to the total surface area of PUF particles.⁵²

Moreover, the polyols produced from the reaction with DEG were more viscous than those from propylene (PG) or ethylene (EG) glycols due to the higher molecular weight of DEG and the slower reaction process.⁵³

The glycolysate final composition depends also on the catalyst used because of the different catalysts' selectivity toward transesterification and/or hydrolysis reactions. The use of potassium acetate (KOAc) is known to increase the amounts of amines in the glycolysis products of PU foams.⁵⁴ It has been shown⁴³ that aromatic methylenedianiline (MDA) content increases almost linearly with the reaction time and its formation is exponentially related to the KOAc concentration. An optimal KOAc shall be identified as a good trade-off between the reduced MDA concentration and the sufficient depolymerization degree to produce a recycled polyol with suitable (*i.e.*, not too high) viscosity. While a proper solution to reduce the free aromatic amine content for flexible PU foam wastes is available thanks to the implementation of an acidolysis process (as it will be explained later), for rigid PU wastes, the most effective solution to obtain products with a low amine content is adding chemical reagents to the glycolysis system, such as glycidyl ethers, hexamethylenetetramine, ethyl or methyl carbamate, propylene oxide and acetic anhydride or by alkoxylation of the final product.^{43,55,56}

Aminolysis

Aliphatic diamines or even polyamines bearing primary and secondary amine functionalities can be employed for the chemical decomposition of polyurethane scraps through transesterification (*i.e.*, aminolysis). The reaction kinetics is strictly



Scheme 2 Aminolysis of polyurethanes.

related to the basic character of the nitrogen of the amine groups: in particular, it is possible to perform the reaction under mild conditions (*i.e.*, low temperatures) when highly reactive amines are utilized. The general reaction is displayed in Scheme 2.

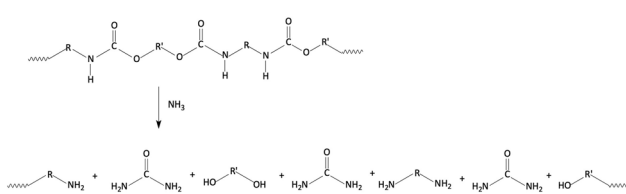
In one of the pioneering works, Xue and co-workers⁶⁷ investigated the aminolysis of rigid polyurethanes and polyisocyanurate foams carried out with different polyamines (namely, tetraethylenepentamine, triethylenetetramine, and diethylenetriamine); several experimental parameters were considered (*i.e.*, foam/polyamine ratio, the structure of the polyamine, and the reaction temperature). The resulting decomposed products were directly employed as curing agents for epoxy systems, particularly for designing new structural adhesives. The molecular weight of the polyamines turned out to remarkably affect the reaction kinetics: more specifically, low-molecular weight polyamines increased the degradation rates and the total amine number as well. Besides, the viscosity of the resulting decomposed products lowered with the increase in the reaction temperature and with the decrease in the foam/polyamine ratio.

Further, Kanaya and co-workers⁶⁸ demonstrated the feasibility of using alkanolamines for decomposing flexible polyurethane foams. More specifically, the reaction was carried out at 150 °C without employing any catalyst and obtaining two layered phases: the upper phase was rich in polyether polyol, while the lower consisted of 4,4'-methylene diphenylamine and some derivatives of alkanolamines. Despite its good potential, the aminolysis process is still on the research/lab scale.

Ammonolysis

The chemical decomposition of polyurethane scraps using ammonia or ammonium hydroxide is known as ammonolysis (Scheme 3).

The reaction, as aminolysis, is a transesterification that originates such decomposition products as polyols (usually suitable for synthesizing rigid polyurethane foams), amines, and urea, due to the higher nucleophilic character of



Scheme 3 Ammonolysis of polyurethanes.



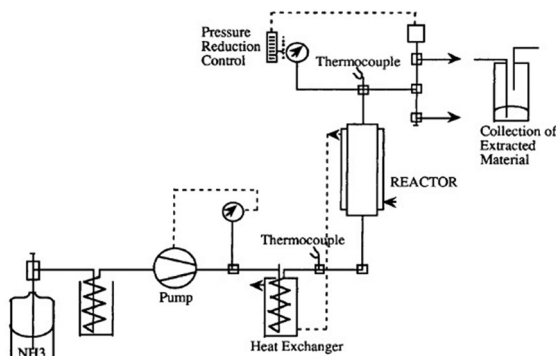


Fig. 12 Schematic of the ammonolysis apparatus working with supercritical ammonia. Reprinted with permission from ref. 69. Copyright 1992 Wiley.

ammonia with respect to H_2O . Besides, the reaction kinetics increases with the increase in the ammonia-to-polyurethane mass ratio.

In addition to the “standard” ammonolysis, Lentz and Mormann⁶⁹ succeeded in developing a particular ammonolysis process (Fig. 12), suitable for the feedstock recycling of rigid polyurethane elastomers and flexible foams, working with supercritical ammonia. The mass ratio of ammonia to polyurethane scrap was fixed at 1/1 and the reaction was performed at 139 °C, and 140 atm, for 2 h.

Acidolysis

Despite the fact that the first investigation on the acidolysis of PUs dated back to 1992, only recently acidolysis has started to emerge as a promising method for the chemical recycling of PUs. It is a method that converts the PU into a polyalcohol, *i.e.*, a polyol, using acids,^{70,71} possibly in the presence of a polyol, used as a solvent. Organic or inorganic acids can be successfully exploited for acidolysis recycling of polyurethane scraps: in particular, inorganic acids (such as HCl) give rise to the formation of polyols, CO_2 , and amine salts. Conversely, the reaction of organic acids (usually dicarboxylic structures) with the urethane groups accounts for the formation of amide groups. This is one of the main advantages of the use of organic acids, *i.e.*, the absence or the very low amount of aromatic amines in the final product. Acidolysis is particularly convenient for obtaining a recycled polyol with a very low hydroxyl value, suitable for flexible PU applications, while, for rigid PU applications, since a high hydroxyl value is needed, glycolysis is the most convenient choice with respect to acidolysis.⁴³

Besides, two main types of organic acids can be employed, namely, saturated (such as adipic acid or succinic acid) and unsaturated (like fumaric or maleic acid) dicarboxylic acids. The saturated acids allow for carrying out the chemical decomposition reactions under milder conditions (*i.e.*, low temperatures – around 60 °C – and short reaction times), giving rise to the formation of oligomers functionalized with amines, amides, urea compounds, or -OH groups. Conversely,

the use of unsaturated dicarboxylic acids accounts for the formation of oligomers bearing double bonds, suitable for the formulation of adhesives.⁷² Though organic acids work as reagents and catalysts as well, He *et al.*⁷³ proposed zinc acetate to speed up the process, without negatively affecting the quality of the recovered polyol.

Gama *et al.*⁷⁴ studied the acidolysis of the flexible PU foam by the design of experiment (DOE) method using succinic, phthalic, or adipic dicarboxylic acids (DA). They analyzed the process temperature in the range of 190–200 °C, with the PU/DA ratio between 4 and 5 and reaction time between 4.5 and 5.5 hours. They found that the temperature and PU/DA ratio are the input variables with a higher influence on the hydroxyl number of the recovered polyol, although its variation is quite limited by changing the reaction conditions and it is generally about 50 mg KOH per g, which is suitable for the production of flexible foams.

Phosphorolysis

Using esters of phosphoric or phosphonic acid allows for the decomposition of polyurethane scraps, by exploiting phosphorolysis reactions. Troev and co-workers have exploited this reaction for the chemical decomposition of both flexible foams and microcellular elastomers, as witnessed by their significant contribution to the scientific literature.^{75–78} Phosphorolysis reactions were performed between 140 and 180 °C, according to the type of polyurethane scrap and P-containing ester; dimethyl phosphonate, diethyl phosphonate, triethyl phosphate, and tris(1-methyl-2-chloroethyl) phosphate were successfully employed. Because of the presence of phosphorus, the recovered products (usually a mixture of P-containing oligo-urethanes) were utilized for synthesizing novel flame-retarded polyurethanes.

Thermo-chemical (tertiary) recycling of polyurethanes

Thermo-chemical recycling of polyurethane wastes was carried out to recover chemical products, energy, and fuels, among others. The main strategies that can be successfully exploited are pyrolysis, gasification, and hydrogenation, as described in the following paragraphs.

Pyrolysis of polyurethane scraps

Generally speaking, this process refers to the thermal decomposition of long macromolecular chains, with the formation of lower molecular weight products; the reaction is usually performed at high pressures and in inert gas. The main decomposition products comprise gases, oils, and ashes.

Specifically referring to polyurethane wastes, their pyrolysis occurs according to a two-step process: the first step, taking place between 100 and 300 °C, refers to the thermal decomposition of polyol-rich moieties (about 50% of the initial mass is lost), while the second step (occurring between 300 and



800 °C) involves the isocyanate parts of the polymer; the final ash content is usually below 3 wt%.⁷⁹

As reported by Cullis and Hirschler,⁸⁰ the thermal decomposition of polyurethanes may involve three different possible dissociation pathways of the urethane bond, namely:

- dissociation into olefin and carbamic acid
- dissociation into alcohol and isocyanate
- dissociation of carbamic acid into secondary amines and carbon dioxide.

All these reactions usually take place between 200 and 300 °C.

As observed by Takamoto and co-workers,⁸¹ the pyrolysis of reaction injection molded polyurethane scraps carried out beyond 450 °C normally originates 10–45 wt% oils (extremely viscous and prone to solidify over time), 5–25 wt% char and more than 40 wt% gases. Further, the use of either the char produced by the pyrolysis or activated carbon, during a secondary pyrolysis step, accounted for the enhanced amount and quality of the resulting pyrolyzed products, especially of the liquid fraction (oils), which showed lowered viscosity.

Moroi and co-workers^{82,83} highlighted the possible catalytic effect exerted by metal salts (namely, FeCl_3 , NaCl , MnCl_2 , CoCl_2 , CuCl_2 , and CrCl_3) on the pyrolysis of poly(ester-urethane)s. In particular, copper and chromium salts accounted for the best catalytic effect, favoring the occurrence of thermal decomposition at lower temperatures, but, at the same time, stabilizing the intermediate pyrolysis products. Conversely, all the other metal salts, despite an improvement in the thermal decomposition of the polymer, were not able to stabilize the intermediate degradation products.

Gasification

Gasification is a very exothermic reaction that is performed at high temperatures (ranging from 1200 to 1500 °C) and pressures (20–80 bar), in an oxygen (air) atmosphere; it produces heat, syngas (*i.e.*, synthetic gas, a gaseous mixture containing large amounts of hydrogen and carbon monoxide) and ashes. Further, it really takes a few seconds to achieve very high conversions of fuels (around 98–99%). Fig. 13 shows the general scheme of the gasification process.

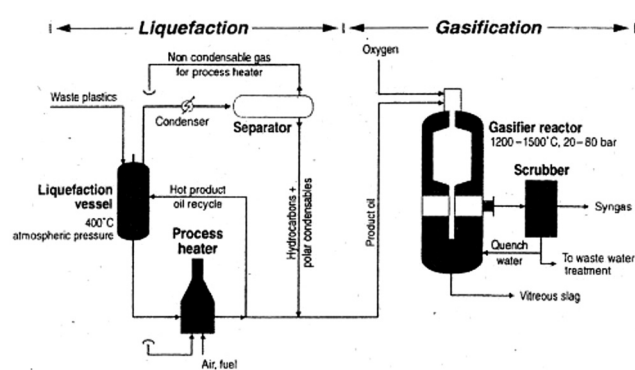


Fig. 13 Schematic of the liquefaction/gasification process. Reprinted with permission from ref. 27. Copyright 2007 Elsevier.

The obtained syngas can be employed to synthesize isocyanates and polyetherols, suitable for producing new polyurethanes.⁸⁴

Hydrogenation

To recover both liquid and gaseous products from polyurethane wastes, it is possible to perform hydrogenation processes, combining heat with high-pressure H_2 . Therefore, this process is in between pyrolysis (heat treatment) and gasification (treatment with high-pressure gas). Compared to simple pyrolysis, hydrogenation allows for obtaining purer recyclates. Similarly to other depolymerization processes, hydrogenation has not been implemented on a large scale yet.

Biodegradation of polyurethanes

It is possible to exploit such living microorganisms as bacteria and fungi to biodegrade polyurethane wastes, including post-consumer materials:^{29,85–87} because of the interaction with the microorganisms, either under aerobic or anaerobic conditions, the macromolecular chains are shortened, and the molecular weight of the polymer significantly decreases. Further, according to the adopted experimental conditions, it is possible to achieve the full mineralization of the biodegraded polymer. Biodegradation takes place according to three successive steps: break of the macromolecular chains and formation of oligomeric structures, further degradation to low-molecular weight species, and final conversion to CO_2 and H_2O (under aerobic conditions) and CH_4 (under anaerobic conditions). Compared to thermochemical degradation, biodegradation is more environmentally friendly, as it does not need the use of specific impacting reagents and takes place at room temperature (*i.e.*, no heat must be provided to the degrading system). Three different types of biodegradation can be considered, namely, fungal, bacterial, and enzymatic degradation; each of them usually suits better specific types of polyurethanes. As an example, fungal biodegradation is preferred to biodegrade polyether polyurethane foams, as the mycelium of fungi can easily penetrate the foam cells; conversely, bacterial degradation is ideal to biodegrade polyurethane coatings, as bacteria can effortlessly exploit the smooth surface of these coatings for the biofilm formation (Table 4).

Besides, it is worth highlighting that biodegradation is affected by several experimental factors including the cross-linking degree of the polyurethane substrate, its crystallinity, and molecular orientation as well. In particular, because of the easier accessibility, the amorphous parts of the polyurethane wastes are generally more prone to biodegrade with respect to their crystalline counterparts.⁸⁸

As far as enzymatic degradation is considered, many of the employable enzymes belong to the class of hydrolases (such as proteases, elastase, esterases, and ureases, see Table 5) and are suitable for polyurethane coatings and thermoplastic bulk polymers. Further, despite the quite high potential of biodegradation, the research is still at the lab scale, also consider-

Table 4 Some examples of fungal and bacterial degradation of polyurethanes

Polyurethane	Fungi	Bacteria
Thermoplastic polyester polyurethane	<i>Penicillium</i> sp. ⁸⁹ <i>Alternaria</i> sp. ⁸⁹	<i>Bacillus</i> sp. ⁹⁰ <i>Comamonas acidovorans</i> ⁹¹ <i>Staphylococcus epidermidis</i> ⁹² <i>Alyciphilus</i> sp. ⁹³ <i>Pseudomonas aeruginosa</i> ⁹⁴ <i>Bacillus subtilis</i> ⁹⁷ <i>Pseudomonas putida</i> ⁹⁸
Thermoplastic polyether polyurethane		
Polyester polyurethane foam		
Polyester polyurethane coating	<i>Penicillium crysogenum</i> ⁹⁵ <i>Cladosporium</i> sp. ⁹⁶	
Polyether polyurethane foam	<i>Alternaria</i> sp. ⁹⁹ <i>Cladosporium herbarum</i> ¹⁰⁰	

Table 5 Enzymatic degradation of polyurethanes

Polyurethane	Enzyme	Ref.
Thermoplastic polyester polyurethane	Lipase	101
	Esterase	102
	Protease	103
Thermoplastic polyether polyurethane	Esterase	103
	Protease	104
	Chymotrypsin	105
Thermoplastic poly(ester urea) polyurethane	Lipase	106
	Cholesterol esterase	107
	Elastase	108
Thermoplastic poly(ether urea) polyurethane	Papain	109
	Lipase	110
Polyester polyurethane coating		

ing that it usually requires a lot of time and research efforts to obtain reliable data.

Energy recovery (quaternary recycling) from polyurethanes

Energy recovery (through incineration or combustion processes) is exploited only when polyurethane wastes and scraps are not suitable for any other recycling strategy; besides, it is very appropriate when PU-based products at the end of life are contaminated. The main difference between incineration and combustion is that the former is an incomplete combustion that may result in the production of toxic gases when not properly carried out.

Besides, two main advantages can be highlighted: energy recovery allows for reducing the volume of the wastes that are landfill confined by up to 99%; then, the produced energy is comparable to that which originated from coal combustion (the average calorific value of polyurethanes is around 25–30 MJ kg⁻¹) and it is marginally lower than that gathered from fuel combustion.¹¹¹ Conversely, the process is not feasible when flame-retarded polyurethane wastes or scraps have to be recycled, because of the presence of flame-retardant additives having a high environmental impact. Further, another issue arising from energy recovery refers to the possible generation of toxic and harmful gases, including nitrogen oxides, CO, and HCN.¹¹² This latter problem can successfully be solved by recirculating the exhaust gases that are generated during com-

bustion and further oxidizing the not-fully oxidized products with a secondary airflow.

Generally speaking, three main energy recovery approaches for polyurethane wastes have been considered so far, namely: co-combustion of polyurethanes with municipal solid wastes,¹¹¹ fluidized bed combustion,¹¹³ and two-stage incineration.⁹

Recent advances in polyurethane recycling

This paragraph aims to describe some recent examples that clearly indicate the direction taken for this research topic during the last years, especially toward the design and exploitation of strategies with low environmental impacts.

Zhang and co-workers¹¹⁴ succeeded in isolating a fungal strain (namely, *Cladosporium halotolerans*) from the deep sea, able to exploit a polyurethane foam (namely Impranil PU) as the only carbon source. In particular, as assessed by FTIR spectroscopy measurements, *Cladosporium halotolerans* accounted for the degradation of the foam up to 80% after 3 days of incubation at 28 °C (Fig. 14), through the breakage of C–N–H bonds and carbonyl groups. Further, gas chromatography-mass spectrometry measurements highlighted the formation of alkanes and polyols as degradation intermediates, due to the hydrolysis of urethane and ester bonds. Finally, urease and

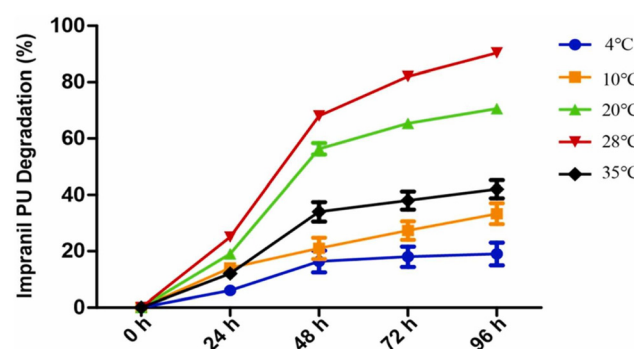


Fig. 14 Time course of the degradation of Impranil PUs by *Cladosporium halotolerans* 6UPA1 growing at different temperatures. Reprinted with permission from ref. 114. Copyright 2022 Elsevier.



esterase activities were identified in 7 days old cultures utilizing the polyurethane foam as the only carbon source.

Recently, Branson *et al.*¹¹⁵ demonstrated the suitability and effectiveness of urethanases, discovered from a metagenome library constructed from the soil, for recycling polyether-polyurethane foams through a chemo-enzymatic process. In particular, the urethanase was able not only to hydrolyze low-molecular weight dicarbamates resulting from chemical glycolysis of the foam but also to recover the polyether-polyols derived from chemocatalytic glycolysis.

Jeong and co-workers¹¹⁶ synthesized bio-polyurethane foams by reacting toluene diisocyanate with sucrose and propolis. A foaming rate as high as 1440% was observed for the polyurethane formulation containing 15 wt% of sucrose and 20 wt% of propolis. Besides, the presence of the sugar accounted for a high enzymatic degradability (using invertase), while the incorporation of the propolis was responsible for high antibacterial features, with about 94% bacteria reduction.

Recently, Xu and Hong¹¹⁷ nicely reviewed the potential of thermoplastic biodegradable polyurethanes for tissue repair and regeneration (Fig. 15), highlighting the possibility of synthesizing them starting from such aliphatic diisocyanates as 1,4-butane diisocyanate, 1,6-hexamethylene diisocyanate, and lysine-based diisocyanate, and exploiting their nontoxic biodegradation products (namely, 1,4-butanediamine, 1,6-hexanediamine, and lysine, respectively).

Pfohl and co-workers¹¹⁸ thoroughly investigated the biodegradation in compost (monitoring the carbon dioxide evolution) of (thermoplastic) polyurethanes differing with regard to their hard-to-soft segment ratios, the type of polymer backbone (aromatic or aliphatic), the presence of specific additives

(*i.e.*, hydrolysis stabilizers), and the possible degree of cross-linking. The increase in both the degree of crosslinking and the hard segments' content turned out to lower the rate and extent of the biodegradation rate (the maximum value achieved was about 72%). Besides, the incorporation of a hydrolysis stabilizer accounted for a decrease in the polyurethane fragmentation, without significantly impacting the conversion of the polyurethane carbon into carbon dioxide.

Espinosa *et al.*¹¹⁹ demonstrated the possibility of employing a soil bacterium, namely *Pseudomonas* sp. as an effective strain for biodegrading both a polyurethane-diol solution (*i.e.*, a polyurethane oligomer) and 2,4-diaminotoluene, a very common precursor in the synthesis of polyurethanes and, at the same time, a putative degradation intermediate.

Then, the possibility of designing renewable low viscosity polyester-polyols obtained from sebacic acid, azelaic acid and bio-based 1,3-propanediol, suitable for the synthesis of biodegradable thermoplastic polyurethanes was successfully demonstrated by Rajput and co-workers.¹²⁰ The resulting polyurethanes showed remarkable biodegradation under compost environmental conditions, with an as high as 57% decrease in molecular weight after 9 weeks of biodegradation, and up to 97% biodegradation over 120 days, as assessed by respirometric analyses.

Very recently, Liu *et al.*¹²¹ isolated 20 strains (namely, 9 fungi and 11 bacteria) and assessed their suitability for biodegrading a poly(1,4-butylene adipate)-based polyurethane. Among the different tested strains, *Cladosporium* sp. P7 was found to be the most efficient for biodegrading the polymer: in particular, it was found that both ester and urethane bonds were cleaved by this strain, leading to the formation of six metabolites comprising adipic acid, 1,4-butanediol, and 4,4'-methylenedianiline. Besides, after 28 days of cultivation, about 32 and 44% of the polyurethanes were converted into soluble small molecules when employed as the sole carbon source or in a medium with other co-carbon sources, respectively.

Concerning some recent outcomes in the mechanical and chemical recycling of polyurethanes, new strategies based on mechanochemistry have been reported in the literature. Mechanochemistry refers to the phenomenon of chemical reactions of substances caused by the input of mechanical energy (such as crushing in a ball mill or shearing). Guo and co-workers¹²² performed high-shear grinding on flexible polyurethane foam scraps by means of two-roll milling; the resulting powder was employed for partially replacing polyols in new polyurethane foams. In fact, the high shear grinding accounted for the activation of the powder, because of the appearance of several hydroxyl groups on its surface. In particular, the powder obtained by performing 7 milling cycles carried out at room temperature showed an average size of around 98 μm , and successfully replaced 15 wt% of the polyols in the preparation of the new polyurethane foams, whose microstructure and density were similar to those of the pristine polyurethane scraps. He and co-workers¹²³ proposed a mechano-chemical reaction for recycling rigid polyurethane insulation board, employing a self-made pulverizer. The

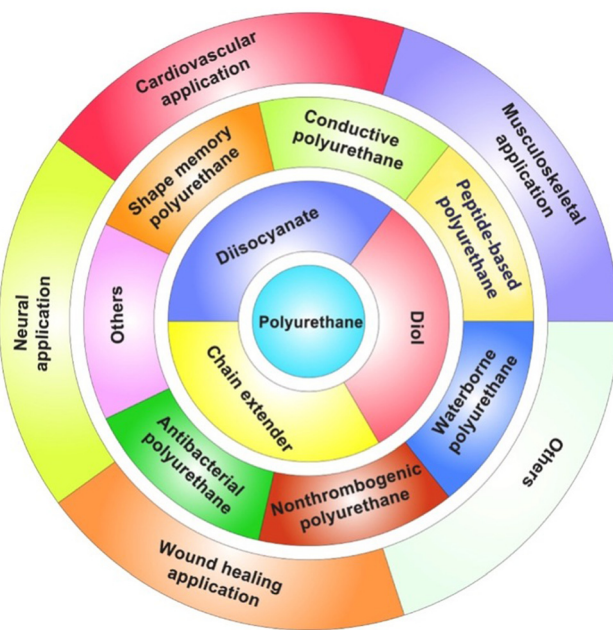


Fig. 15 Biodegradable polyurethanes for tissue repair and regeneration. Reprinted from ref. 117 under CC BY-NC-ND 4.0 License.



accumulation of mechanical energy during the process was responsible for the concentration of the impact energy on individual macromolecular chain segments, hence determining the break of their chemical bonds. The crushing parameters (feed amount – 90 g, crushing speed – 500 rpm, and crushing time – 30 min) were optimized; finally, the recovered polyurethane powder (exhibiting high activity of functional groups and reduced the cross-linking density as well) was exploited for producing composite panels with polypropylene.

A further step in mechanochemical recycling is related to the use of covalent adaptable networks (CANs). The polymer materials with reversible CANs are described as vitrimers.¹²⁴

Under specific stimuli such as light and heat, the breaking, recombination and exchange of dynamic covalent bonds occur, which means that the crosslinked networks are no longer fixed, and the topological structures will change under external forces. Indeed, topological rearrangement reactions occur, which make the thermoset polymers flowable and thus allow for their reprocessing. Vitrimers have mechanical properties similar to common thermoset resins, but they can be remodeled during the bond exchange process, like thermoplastics since they can flow under shear stress. This bond exchange process needs a catalyst that can be inserted during the synthesis of new PU polymers but, to recycle an existing PU waste, the catalyst must be introduced into the PU just before reprocessing, *i.e.*, after reducing the size of the material by crushing and milling, among others. For example, Li *et al.*¹²⁵ studied the effects of different pretreatment methods on pulverized PU foams (PUF), after adding a dibutyltin dilaurate (DBTDL) catalyst, on the reprocessing properties of commercial rigid PUF. In particular, they considered direct pretreatment, swelling pretreatment, and ball milling pretreatment; they also tested multiple reprocessing sets by ball milling. The treated powders were placed in a mold and pressed into sheets for 30 min at a pressure of 30 MPa at 170 °C. They showed that, through strong mechanical forces, the ball milling pretreatment method could effectively reduce the size of PUF particles, increase the surface area, and generate new reactive groups *via* mechanochemical action, which effectively promoted urethane exchange and improved the reprocessing properties of PUF by final mechanical recycling. Bandegi *et al.*¹²⁶ proposed a similar approach, by using triazabicyclodecene (TBD) as a catalyst, and found that the permanent crosslinked structure of the PU thermoset foam can be converted into a dynamic network upon vitrimerization. The vitrimerized network retains high mechanical strength and can also be foamed by applying low pressures at high temperatures. Liu *et al.*¹²⁷ still used TBD to show that three-dimensional photo-printable resins with tunable material mechanical properties—which are superior to commercial high-performance counterparts—can be formulated with the addition of various network reforming additives, starting from waste PU foams. They thus showed that a more attractive approach with respect to chemical recycling is to partially disintegrate the network to a point that allows reworking. There are two notable benefits with this approach: (1) the reaction con-

ditions can be much milder and (2) the obtained mixture can be fully converted into a network without requiring purification. A more circular, direct and continuous foam-to-foam recycling method using twin screw-extrusion by leveraging the melt processability of PU CANs has been proposed by Kim *et al.*¹²⁸ by using zirconium(IV) acetylacetone as a replacement for toxic tin catalysts in PU CANs and azodicarbonamide as a chemical blowing agent. Their results indicate that waste PU foams can be directly recycled into a new foam while preserving its structural properties and chemical integrity. Bai and co-workers¹²⁹ exploited the combination of coordination and dynamic chemistry for facilitating the thermal and chemical recycling of polyurethanes. A general scheme is shown in Fig. 16. In particular, aluminum acetylacetone was employed as a crosslinker for the synthesis of the polyurethanes, which gave rise to the formation of thermally reversible bonds between isocyanate and acetylacetone groups. This way it was possible to provide the crosslinked polyurethanes with a worthy thermal remoulding capability at temperatures beyond 100 °C and, at the same time, allow for a simple chemical recycling route based on the pH-sensitive Al-acetylacetone coordination, using acetic acid for promoting acid hydrolysis. In fact, the coordination chemistry accounted for the break of the coordination bonds dissociating in aluminum acetylacetone and the reassociation of the same bonds after the removal of acetic acid through evaporation.

Disulfide as well as thiourethane, imine, urea, and boronic ester can be used to create CAN PUs.^{124,130}

Contrary to mechanical recycling, on which the research has not been focused so much during the last years, chemical recycling has highlighted significant outcomes.

In particular, Donadini and co-workers¹³¹ exploited a micro-wave-assisted glycolysis reaction catalyzed by monoethanolamine, stannous octoate, or potassium acetate, for recovering a polyol-like liquid product from rigid polyurethane foam wastes. The use of the microwave reactor accounted for a remarkable decrease in the reaction time (by about 94% with respect to conventional-heated processes) and significant energy saving (by about 45%). Finally, the replacement of

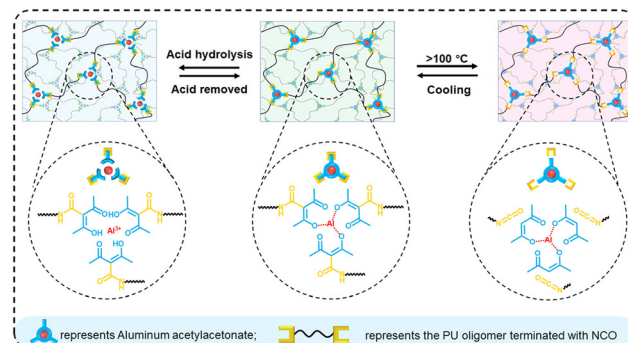


Fig. 16 Schematic of the thermal (*via* dynamic chemistry) and chemical (*via* coordination) recycling of polyurethanes. Reprinted with permission from ref. 129. Copyright 2022 ACS.



virgin polyol with the recovered counterpart allowed for a slight decrease in the thermal conductivity of the resulting re-foamed material, as well as for increasing its compression strength (by around 20%).

A microwave-assisted acidolysis reaction with adipic acid for recycling flexible polyurethane foams was proposed by Grdadolnik and co-workers,¹³² who demonstrated the feasibility of the process for the recovery of polyether polyols to be employed for the preparation of new polyurethanes. The molar ratio of adipic acid to urethane groups of polyurethane foam was set at 1.1 and the reaction was carried out at different temperatures (namely, 210, 220, and 230 °C) and reaction times (15, 30, and 40 min). The microwave reactor accounted for a significant decrease in the reaction time (below 1 h) compared to the acidolysis with conventional heating (several hours). It is worth noting that, with the increase in the amount of adipic acid, the content of toluendiamine (TDA) in the polyol decreased and the degree of degradation of the urethane groups increased, but at the expense of a higher degree of esterification of the hydroxyl groups of the polyol, resulting in the recovered polyols (RPs) being terminated with carboxyl groups to a greater extent. In other words, none of the reaction conditions employed resulted in the formation of a completely hydroxyl-functionalized polyol.

The results indicated that with the increase in the carboxyl functionality of the RPs in the formulation, the carboxyl-terminated polyol negatively affects the quality and performance of the flexible PUFs.

In a further research effort, Amundarain and co-workers¹³³ exploited the polyols, recovered from post-industrial rigid polyurethane foams by means of glycolysis reactions, for preparing new rigid foams. The recovered counterparts showed remarkably higher values for viscosity, acidity, and average molecular weight and more hydroxyl groups than those of commercial polyols: these findings were ascribed to both the residual ethylene glycol in the polyol after the purification of the glycolysate and in the presence of glycolysis by-products (such as amines and carbamates). Though it was not possible to incorporate more than 15 wt% of the recovered polyols into the formulation of the final polyurethanes, the latter showed enhanced tensile strength, likely due to a higher cross-linking of the foams containing the recycled polyols.

Pursuing this research, the same group¹³⁴ employed a glycolysis process for the recovery of polyols from rigid polyurethane foams either aromatic or aliphatic. Ethylene glycol or diethylene glycol was employed as a glycolysis reagent; sodium hydroxide, sodium acetate or diethanolamine was utilized as a catalyst. Unlike the glycolysis of the aromatic polyurethane, which achieved high conversions (about 90%) in about 2 h at 198 °C and without any catalyst, the presence of any catalyst was mandatory for the decomposition of the aliphatic polyurethane, leading to higher conversions (around 98%) under the same reaction conditions adopted for the aromatic counterpart. As assessed by gel permeation chromatography, the structure of the recovered polyols was similar to that of the pristine ones employed in the formulation of the initial poly-

urethanes, hence allowing for a partial replacement of "virgin" polyols with the recovered molecules (up to 15 wt%) in the preparation of new polyurethane systems.

Another interesting approach well matching the circular economy concept has been recently proposed by Sternberg and Pilla,¹³⁵ who investigated the feasibility of a high-pressure hydrolysis/hydroglycolysis recycling technique, for the recovery of lignin (exhibiting both improved solubility and lower hydroxyl contents) from a lignin-derived non-isocyanate foam and its reuse for the obtainment of second-generation polyurethane foams. The latter showed a mechanical behavior comparable to the foams made of virgin lignin, despite a slight decrease in the compressive strength (126 vs. 133 kPa, for the recycled lignin foams and the virgin lignin-containing counterparts, respectively), ascribed to a faintly lower cross-linking density due to the decreased hydroxyl content of the recycled lignin.

Yuan and co-workers¹³⁶ synthesized biodegradable and chemically recyclable polyurethanes using bio-based telechelic hydroxyl-terminated poly(γ -butyrolactone) diols as suitable precursors. Changing the molar mass of the diols allowed for obtaining polyurethanes with thermoplastic or elastomeric features. Finally, the authors demonstrated that simple heating of the polyurethanes in bulk at 170 °C for 2 h, in the presence of 4 wt% of stannous octoate as a catalyst, was enough for easily recovering highly pure γ -butyrolactone in about 99% yield.

Gausas *et al.*¹³⁷ synthesized an Mn-based complex for catalyzing the hydrogenative depolymerization of both real-life and end-of-life flexible polyurethane foams, using KOH, isopropyl alcohol, and H₂. This way, it was possible to deconstruct the foams into polyol and aniline fractions; the process was implemented on a gram-scale with a total of 90% mass recovery.

Gu and co-workers¹³⁸ employed four types of glycols (namely, butanediol, propylene glycol, diethylene glycol, and monodiethylene glycol) for degrading waste polyurethane elastomers. As an example, Fig. 17 shows the catalytic degradation mechanism of the polyurethane elastomer, using butanediol or diethylene glycol.

The recovered polyols were utilized for the preparation of regenerated polyurethane rigid foams showing physical, mechanical, and thermal features suitable for the envisaged applications. In particular, the foaming time of the regenerated polyurethane rigid foam was usually between 40 and 140 s. The compressive strength ranged from 0.131 to 0.176 MPa and the water absorption from 0.7265 to 1.9923%. Further, the apparent density of the regenerated foams was between 30 and 40 kg m⁻³, and the thermal conductivity ranged from 0.0151 to 0.0202 W m⁻¹ K⁻¹.

A very recent application of recyclable thermoplastic polyurethanes refers to the design of flexible strain sensors, suitable for detecting little changes in strain and pressure.¹³⁹ For this purpose, a thermoplastic polyurethane was mixed with graphene (at different loadings, ranging from 1 to 10 wt%) in a Haake internal mixer, working at 185 °C and 60 rpm for 2.5 min; the desired sheet geometry was obtained by sub-



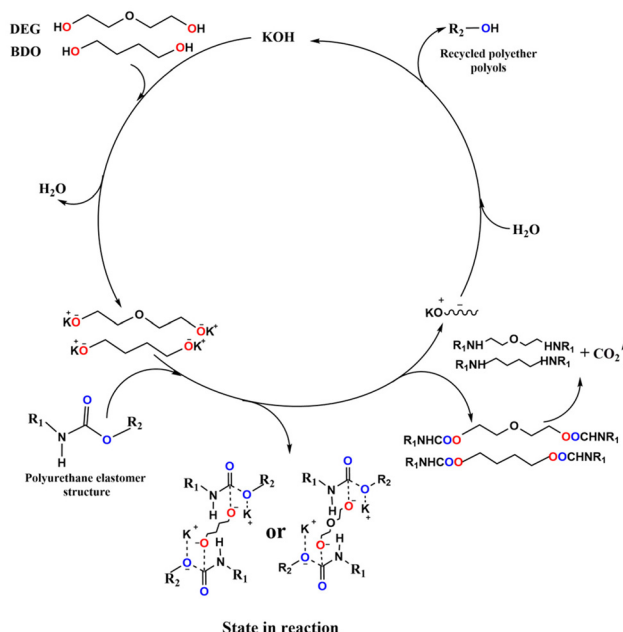


Fig. 17 Catalytic degradation mechanism of the polyurethane elastomer. Reprinted from ref. 138 under Creative Commons Attribution 4.0 International License.

sequent compression moulding. The optimum graphene loading for the envisaged application was 7 wt%: the related nanocomposite sensor exhibited good strain sensitivity, even after three mechanical recycling cycles, as displayed in Fig. 18.

The scalability and cost-effectiveness of the chemical recycling process have been and are still very challenging. A

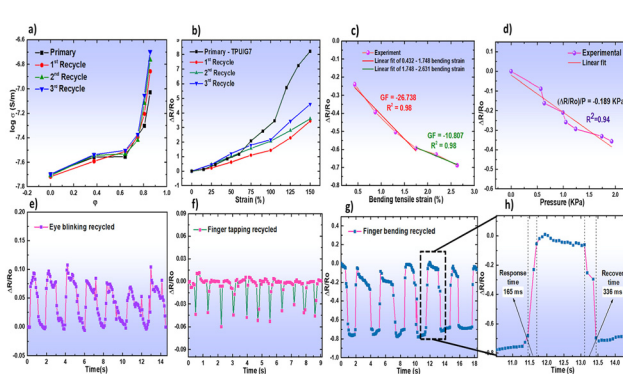


Fig. 18 Recyclability of thermoplastic polyurethane (TPU)/graphene nanocomposites: (a) log conductivity of TPU/graphene nanocomposites as a function of graphene content (φ) of each recycle; (b) tensile strain response of the strain sensor containing 7 wt% of graphene (sample coded as TPU/G7) for each recycling cycle; (c) relative resistance changes of the strain sensor as a function of the bending angle for the third recycled nanocomposite; (d) pressure response of the third recycled TPU/G7 nanocomposite. Human motion detection using the sensor made of the three-time recycled TPU/graphene nanocomposite: (e) eye blinking; (f) finger tapping; (g) finger bending. (h) Response and recovery time of the recycled sensor. Reprinted with permission from ref. 139. Copyright 2023 ACS.

potential solution to solve the high operating cost of split-phase glycolysis has been first proposed by Simón *et al.*,¹⁴⁰ who used crude glycerol, *i.e.*, a cheap by-product of biodiesel production, as a glycolyzing agent. It has been demonstrated that crude glycerol provides an upper phase with a lower content of by-products and transesterification agents than those in the case of using the best “common” option (*i.e.*, diethylene glycol (DEG)), as a consequence of its higher dielectric constant. Furthermore, as a result of this latter, a glycolysis bottom phase free of polyol, has been obtained thus increasing the net yield of the glycolysis process.

In a more recent work, a detailed evaluation of the economic feasibility of a split-phase glycolysis process (up to a pilot scale, Fig. 19) for the recycling of any kind of flexible polyurethane foam wastes, utilizing crude glycerol as cleavage agent, has been published by Del Amo and co-workers.¹⁴¹ In particular, a Net Present value of 1 464 555€, with an Internal Rate of Return of 27.99%, and a Payback Time between 4 and 5 years were estimated, hence confirming the economic viability of the proposed recycling process.

Zhang and co-workers¹⁴² prepared a series of fluorinated polyurethane rigid foams using fluorine-containing recycled polyols obtained from the degradation of waste polyurethane foams. The glycolysis reaction was carried out at 200 °C for 2 h, in the presence of ethylene glycol, diethylene glycol, and a fluorodiol synthesized on purpose. In particular, the incorporation of 8 wt% of fluorodiol in the waste polyurethane foam accounted for: (i) a significant increase in the compressive strength of the regenerated fluorinated rigid polyurethane foam (+ 43% with respect to that of the non-fluorinated regenerated counterpart), (ii) an enhanced waterproof behavior and

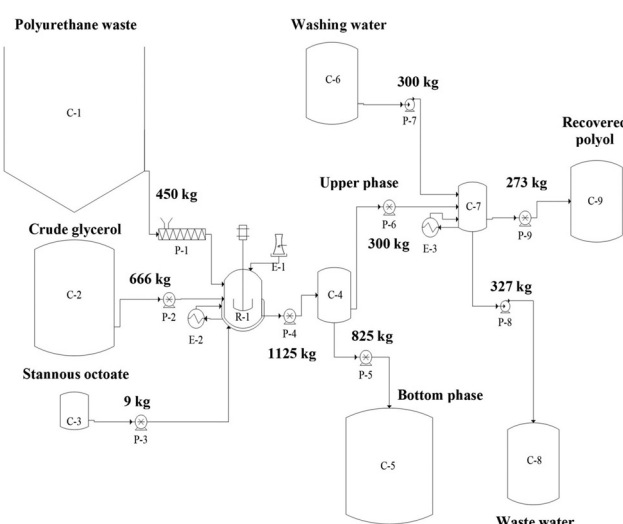


Fig. 19 Flow diagram of the split-phase glycolysis plant and mass balance per batch. Legend: C-1 = polyurethane silo; C-2, C-3, C-5, C-6, C-8 and C-9 = storage tanks; C-4 = decanter; C-7 = liquid-liquid extractor; R-1 = glycolysis reactor; E-1 = condenser; E-2 and E-3 = heat exchangers; P-1 = Archimedean screw; P-2, P-3, P-4, P-5, P-6 and P-9 = Gear pumps; P-7 and P-8 = centrifugal pumps. Reprinted from ref. 141 under Creative Commons Attribution 4.0 International License.

(iii) a decreased thermal conductivity (which was measured as low as $0.0227 \text{ W m}^{-1} \text{ K}^{-1}$).

A recent trend is also the “upcycling” of the polyurethane waste.^{143,144} The term ‘upcycling’ was first used to refer to any process that transforms by-products, undesired, unwanted or waste products into new materials of higher values.¹⁴⁵

Li *et al.*¹⁴⁶ upcycled TPU wastes into activated carbon through two steps of controllable carbonization and activation and demonstrated its excellent wastewater treatment performance.

Daniel *et al.*¹⁴⁷ showed the synthesis of iron–nitrogen–carbon (Fe–N–C) electrocatalysts for oxygen reduction reactions starting from a mix of waste polyethylene (PE) and polyurethane (PU) by adding FeCl_3 . In another work,¹⁴⁸ a PUR waste foam was converted into N-doped hierarchical porous carbon (NHPC) *via* an autogenic atmosphere pyrolysis (AAP)–KOH activation approach for electrochemical energy storage applications. The functional upcycling of the PU waste is also extensively reviewed in a recent paper,¹⁴⁹ where several strategies for the generation of new high-performance materials, especially high-capacity adsorbents for metal ions or oil–water separation, are proposed.

Another recent advance in the polyurethane field is the development of a non-isocyanate polyurethane (NIPU). These are a fairly new class of polyurethanes, which were initially discovered in 1957 by Dyer and Scott as a method to produce polyurethanes without using moisture-sensitive isocyanates. Because of the recent toxicity issues arisen around the use of isocyanates, they have recently attracted enormous attention. Advances in the synthesis of cyclic carbonates *via* the chemical insertion of CO_2 into epoxy resins have also contributed to push activity in NIPUs.¹⁵⁰ Although quite intense work dealing with NIPUs and several reviews have already appeared,^{151–154} only very few papers deal with the recycling of this new class of materials. Many of these latter rely on the use of dynamic covalent bonds (*i.e.*, use the covalent adaptable network concept) to guarantee the recyclability of such polymers, by heating and pressurizing, both in case of compact and foamed PUs.^{155–158} More specifically, they combine the recent trends in material synthesis (*i.e.*, NIPU) with recent trends in recycling (*i.e.*, vitrimers). There are very few examples dealing with the chemical recycling of NIPUs. Liu *et al.*¹⁵⁹ developed a series of isoeugenol-based NIPUs from bio-/CO₂-derived bis(6-membered cyclic carbonate) cured with amines, and showed that they can completely be degraded into pre-monomers by reacting them with ethanol in 30 wt% NaOH at 90 °C for 4 h, *i.e.*, under quite mild conditions. The pre-monomers can be separated by filtration with a high recovery ratio (90%) and used again, with CO₂, to produce the monomers. These recovered monomers were then exploited for preparing a NIPU that showed nearly the same mechanical properties of the virgin one. Despite a fairly limited number of papers on their chemical recycling, it can be expected that NIPUs can solve, at least partly, the “aromatic amine issue” in the chemical recycling process since aromatic amines cannot be formed. It can also be expected that NIPUs can help in closing the material loop,

with a good recycling efficiency under relatively mild degradation conditions.

Conclusions

Undoubtedly, the possibility of designing and effectively applying feasible and reliable recycling strategies to such fossil-derived polymers as polyurethanes has significantly reduced the general concern about their end-of-life and their difficult recyclability, while improving their overall image as recyclable materials with an important intrinsic added-value.

However, the recycling strategies that have been designed and exploited so far, or are being developed for polyurethane waste streams, have to face the great complexity of these latter, considering the wide range of chemical structures, molecular weights, degree of crystallinity, crosslinking density, and hard-to-soft segment ratios, among other parameters, all of which significantly affect the recycling processes and the properties of the final recycled materials and products. Therefore, it is not possible to identify an ideal, unique, feasible, reliable, and even scalable recycling technology that should be suitable for all polyurethanes, but it is necessary to develop specific recycling approaches that maximize the yields, purity, and exploitability of the resulting recycled products, trying, at the same time, to minimize the environmental impact and the overall energy consumption. As reported in the review, several recycling methods have been designed and successfully applied: among them, the most promising ones, also considering industrial exploitation, are mechanical recycling (through regrinding and compression moulding) and chemical recycling (mainly through glycolysis reactions). It is expected that these two recycling strategies will continue to be exploited in the future, as they are currently quite well-established and consolidated, even on an industrial scale. Conversely, almost all the other recycling strategies are still focused on lab-scale research investigation, despite the significant outcomes achieved up to now; it is, therefore, difficult to foresee their development and exploitation on a larger processing scale. Both regrinding and glycolysis are cost-effective and limit the impact on the environment; at variance, the energy recovery through pyrolysis, gasification, and two-stage combustion, despite the important reduction of the waste volumes that are landfill-confined and the high value of the recovered energy, are still limited by the quite complicate control of the emission of toxic and hazardous products.

Further, although biodegradation can be performed using mild conditions (*i.e.*, at room temperature and without the need for hazardous chemicals), it is still limited by the quite restricted number of suitable microorganisms and enzymes (indeed, the isolation of new microorganisms able to degrade polyurethanes is usually labor-intensive and time-consuming), and also by the biodegradation time that is usually high, and therefore has not expressed its full potential yet. To overcome these issues, the recently proposed metagenomic analysis of microbial population involved in the biodegradation of



polyurethanes^{160,161} could help elucidating the structure of the microbial community, mining the enzymes involved in the polymer degradation and predicting their biodegradation ability *in situ*.

A further issue that undoubtedly may slow down the progress in polyurethane recycling regards the diversified forms (*i.e.*, foams, bulk materials, and elastomers) and structures of the manufactured polyurethanes, which makes it more difficult to identify the most appropriate recycling strategy. In this context, a selective collection of polyurethane wastes could help a lot.

Further, it is essential to assess the influence of the polyurethane recyclates on the features (thermal and mechanical, among others) of the final products, into which recyclates are incorporated.

Finally, the development of reliable and feasible recycling strategies for polyurethane wastes and scraps demands cost-effectiveness and environmental friendliness: at present, these are still two challenging issues, especially when they are not considered individually. Besides, at present, the recovery and recycling of polyurethane wastes have no structured market; further, the cost and emission factors of the recovery processes have not been consolidated yet.¹⁶²

However, it is expected that, in the next few years, research on the recycling of end-of-life polyurethanes will make further progress, at least starting from the lab scale, leading to a significant improvement in the recycling processes (also including low environmental impact approaches) and their reliable use, hopefully on an industrial scale. These possible outcomes will certainly contribute to ameliorating the overall management of end-of-life polyurethanes, offering some new perspectives within the circular economy concept, to select which polyurethanes should progressively take part as valuable materials.

Conflicts of interest

There are no conflicts to declare.

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