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polyurethane: towards a greener and sustainable production route

A review on vegetable oil-based non isocyanate

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The transition from conventional polyurethane (PU) to non isocyanate polyurethane (NIPU) is driven mainly by safety concerns, environmental considerations, and sustainability issues associated with the current PU technology. NIPU has emerged as a promising alternative, addressing limitations related to traditional PU production. There has been increasing interest in bio-based NIPU aligning with the aspiration for green materials and processes. One important biomass resource for the development of bio-based NIPU is vegetable oil, an abundant, renewable, and relatively low cost feedstock. As such, this review aims to provide insight into the progression of NIPU derived from vegetable oils. This article highlights the synthetic and green approach to NIPU production, emphasizing the method involving the polyaddition reaction of cyclic carbonates and amines. The review includes case studies on vegetable oil-based NIPU and perspectives on their properties. Further, discussions on the potential applications and commercial importance of PU and NIPU are included. Finally, we offer perspectives on possible research directions and the future prospects of NIPU, contributing to the ongoing evolution of PU technology.

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1 Introduction

Polyurethane has become an essential part of the modern lifestyle. PU was first discovered in the 1930s by Otto Bayer while looking to replace natural rubber during World War II.1 Currently, PU is an important class of polymers, along with polyethylene, polypropylene, polyvinyl chloride, and polyethylene terephthalate. The global scale production of PU in 2020 exceeded USD 70.67 billion and is expected to reach USD 94.59 billion by 2029. This remarkable scale is due to the wide range of applications of PU in diverse fields. It is mainly used as flexible and rigid foams, resins, adhesives, sealants, and coatings,2 commonly found in biomedical applications, textiles, the building, construction and automotive industries, and many others. The outstanding advantages of PU are its durability, toughness, and exceptional chemical resistance. Also, PU possesses durability comparable to metals, yet the elasticity of rubber, making it a versatile alternative to metals, plastics, and rubber in various engineering and commodity applications.³ Despite the many advantages of PU, the downside of this

In view of the recent trend towards green chemistry, the development of bio-based PU has been mainly directed towards the preparation of bio-based polyol, bio-based isocyanate, biomass chain extender and isocyanate-free polyurethane.⁵ In particular, isocyanate-free PU or commonly known as NIPU is gaining research interest due to the immense benefits this route has to offer. With the absence of isocyanate, NIPU production is deemed to be safe, have reduced environmental impact, lower volatile organic compounds emissions, and potential for cost savings. NIPU can be synthesized via a catalyst-free and solventfree under moderate reaction conditions. This approach is considered atom-efficient and generates no by-products.6 Presently, there is a surge of interest in producing bio-based NIPU where the materials used are derived from renewable biomass sources. Different types of building blocks can be produced from renewable feedstock. In particular, vegetable oils present vast opportunities for this purpose, as they can be transformed into many novel products.7 The rationale for using vegetable oils in the production of NIPU is related to their wide availability, renewable nature, eco-friendliness, and the potential to reduce toxicity and health concerns.

polymer relates to the use of toxic and moisture-sensitive isocyanate in the manufacturing process. The involvement of this hazardous component renders PU production extremely toxic and dangerous.⁴ In addition to the risks associated with isocyanate, the majority of PU available today was derived from non-renewable petroleum-based chemicals. These challenges motivate researchers and industries to find greener and more sustainable alternative routes to produce PU.

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Considering the significant growth of studies using vegetable oils as the starting materials to produce NIPU, this work aims to convey the current state of the art in this field. To the best of our knowledge, even though there are some reviews focusing on bio-based NIPU, there is no work specifically focusing on vegetable oil-based NIPU in detail. Therefore, this review could serve as a foundational framework and complementary to the existing literature.

2 Insight into PU and NIPU

Polyurethane is a polymer joined by the urethane linkage comprising of alternative isocyanate (mono-, di-, polyisocyanate) unit, referred to as the hard segment, and the alcohols that possess two or more hydroxyl groups (from here on refers to as polyols) as the soft segments. In the structure of PU, the hard segment contributes to the stiffness and rigidity of the polymer, whereas the soft segment is responsible for the elasticity or flexibility properties.8 It is worth to mention that the ratio of the hard segment to the soft segment affects the structure, physical, mechanical, and thermal properties of PU. An increasing amount of isocyanate leads to a more rigid and hydrophobic PU, while excess amount of polyol produces softer and more hydrophilic PU.9 Therefore, the ratio of polyol to isocyanate plays a vital role in designing PU products. In addition to polyol and isocyanate, additives are often incorporated to aid the reaction or to achieve specific requirements of end applications. Frequently used additives during PU processing include catalysts, surfactants, blowing agents, plasticizers, pigments, chain extenders, fillers, flame retardants, smoke retardants, and stabilizers. The functions of each materials and additive are presented in Table 1.10

PU can be produced as a thermosetting or thermoplastic, either rigid and hard or flexible and soft material, depending on the specific choice of isocyanate and polyol. In general, isocyanate is characterized by the highly reactive isocyanate (R-N=C=O) group and this reagent is available commercially in liquid or powder form. The R group can be aliphatic, cycloaliphatic, polycyclic, or aromatic in nature. Comparatively, aromatic isocyanate is more reactive than other types, as the negative charge gets delocalized into the R. Diisocyanate which possesses two reactive isocyanate groups in the structure, is the

 Table 1
 Materials used in PU production and their functions

Material/additive	Function
Isocyanate	Hard segment, responsible for rigidity in PU
Polyol	Soft segment, responsible for flexibility in PU
Catalyst	Speed up the reaction, enhance reactivity
Surfactant	Improve dispersion of isocyanate and polyol
Blowing agent	Aid the production of PU foams
Plasticizer	Reduce hardness
Pigment	Produce coloured PU
Chain extender	Structural modification of PU, mechanical support
Filler	Reduce the cost, improve mechanical properties
Flame retardant	Reduce flammability
Smoke retardant	Reduce smoke generation when material is burnt
Stabilizer	Protect PU from damage caused by heat, light, etc.

most widely used due to its high reactivity.¹¹ In the case of polyols, they can be broadly divided into monomeric and polymeric polyols. Monomeric polyols are low molecular weight polyols such as glycerol, ethylene glycol, propylene glycol, and others. They can be further categorized as diols, triols, tetraols, *etc.* depending on the number of hydroxyl groups they possess. Meanwhile, polymeric polyols refer to high molecular weight compounds.¹² Fig. 1 presents the general reaction of polyol and isocyanate to produce PU, highlighting the urethane linkage. Some common example of isocyanates and starter polyol are included.

PU is considered to be physiologically inert and safe for many applications. However, the isocyanate monomers are not and they can pose health risks. Isocyanates, being highly reactive chemicals, may cause irritation to the skin, eyes, and respiratory tracts upon exposure. Therefore, elaborate safety procedures are necessary during handling, transport, and storage. In addition to this, the manufacturing of isocyanates involves the use of hazardous and toxic phosgene, a deadly gas used as the starting material. Moreover, isocyanate can contribute to air pollution, specifically in the form of volatile organic compounds released during the production process, impacting air quality and potentially causing environmental issues.13 Another concerns in the production of PU is their moisture sensitive nature of isocyanates. They have strong reactivity with water and can react rapidly upon contact. This reaction can caused premature curing and often results in the formation of unusable by-products.14 Furthermore, as for the vast majority of polymers, the precursors of PU, polyols and isocyanates mainly derived from chemical intermediates based on the non-renewable petroleum or natural gas resources.

To address these challenges, efforts have been directed towards the synthesis of bio-based PU. In this context, various studies have explored the production of bio-based polyol originated from renewable resources such as vegetable oil, 15 fatty acid,16 fatty acid methyl esters,17 crude glycerol,18 wood,19 crop residues,20 lignin,21 and protein feedstock.22 While bio-based polyols are readily available in the market and welldocumented in research, options for bio-based isocyanates are significantly limited. Despite that, commercial isocyanates with high renewable content are available including those based on fatty acids or amino acids. The main challenges in developing bio-based isocyanates are related to high production costs, low efficiency, reactivity limitations, and subpar technical performance.9 Consequently, the majority of bio-based PUs are only partially derived from renewable resources. Typically, these PUs are synthesized using bio-based polyols combined with petrochemical-based isocyanates.23 Currently, well-known companies like DuPont, BASF, BioAmber, Myriant, General Mills Co. Henkel Corporation are producing bio-based raw materials for PU synthesis.24

The next progressive step towards greener PU production is through the non isocyanate route. Various ways have been established to produce NIPU which will be discussed in Section 3, with the most common methods being the reaction of cyclic carbonate and amine groups. Through this method, NIPU enable to eliminate the need for isocyanates or phosgene

Fig. 1 Schematic representation of polyurethane production with some common example of isocyanates and starter polyol.

derivatives and contribute to carbon dioxide sequestration as cyclic carbonates are commonly prepared by carbon dioxide fixation of epoxies.13 Without the use of isocyanates, NIPU synthesis is considered reliable in terms of safety and environmental impact. Other than being moisture insensitive, the production of NIPU has little-to-no use of those volatile organic compounds.25 Though promising, some technical challenges somehow hinder the commercialization of NIPU such as slow polymerization, long reaction time, subpar mechanical properties, and difficulty to synthesize high molecular weight product.26,27 Various factors affects the production of high molecular weight products, including the concentration and ratio of monomers, synthesis conditions i.e. temperature, pressure and reaction time, the present and types of catalysts,

conventional PU.

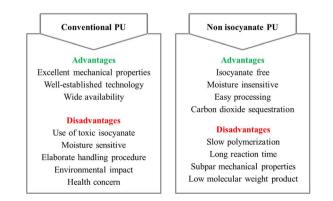


Fig. 2 Comparison of conventional PU and non isocyanate PU

and polymerization mechanism. Nevertheless, there are tradeoffs between the advantages and disadvantages of NIPU and conventional PU as depicted in Fig. 2. From this point of view, PU and NIPU differ significantly in synthesis methods, safety considerations, and environmental impact. While PU is already well-established, NIPU serves as a promising alternative with potential environmental benefits. As NIPU technology continues to evolve, it is expected to offer improvement in properties and production techniques to match or even surpass

3 Production routes of NIPU

The pioneer work on NIPU was reported by Dyer and Scott in 1957. In this study, they proposed the synthesis of urethanes and cyclic urea by reacting ethylene carbonate and various diamines to produce β-hydroxycarbamate. These compounds were subsequently polycondensed through a transesterification reaction.28 This early study laid the foundation for the development of NIPU and opened up new possibilities in polymer synthesis. NIPU based on cyclic carbonate-amine chemistry has since emerged as a significant area of study and is extensively explored for various applications. To date, there are four main approaches to synthesize NIPU including polycondensation, ring opening polymerization, rearrangement and polyaddition reaction29 as presented in Fig. 3. All the four methods are briefly described in this section.

Polycondensation reaction is one of the approaches used in NIPU synthesis, involving the condensation of suitable

Fig. 3 Common synthesis routes of NIPU. Adapted from ref. 2 with permission from MDPI, copyright 2023.

precursors to form the desired polymeric structure. Several routes can be used for this method such as the reaction between polychloroformate and polyamine, polycarbamate and polyol, polycarbamoyl chloride and polyol, and the interaction between polycarbonate and polyamine. However, the precursors for these routes are synthesized from toxic phosgene or its derivatives. Hence, there is not much difference in the production process of isocyanate-based polyurethane and NIPUs using polycondensation.³⁰ Additionally, the process releases side products such as alcohol or HCl, posing significant limitation in industrial applications. Other significant drawbacks of polycondensation include the need for a catalyst, extended reaction time, the necessity for purification of end products, and the formation of low molecular weight compounds.³¹

Ring-opening polymerization is another method for synthesizing NIPU, where a cyclic monomer such as aziridine or cyclic carbamate, undergoes a ring-opening reaction to form the polymer chain. Although this process does not produce any undesirable side-products, it is often performed at high temperature. Further, the precursors, cyclic carbamate are commonly derived from phosgene and aziridine is toxic remains significant issues for NIPU synthesis through the ringopening pathway.³² Rearrangement reaction can also be utilized in NIPU synthesis. The literature has reported the synthesis of NIPU through the rearrangement of acyl azides (Curtius rearrangement), carboxamides (Hoffman rearrangement), or hydroxamic azides (Lossen rearrangement).33 In these reactions, isocyanates are generated in situ, allowing the synthesis of polyurethane. However, it is worth mentioning that these routes still involve the use of isocyanates, albeit produced in situ. Furthermore, the reactants involved such as azides, carboxamides, and hydroxamic azides, are highly hazardous

substances, which add to the importance of handling them with care.

Lastly, NIPU can be synthesized through the polyaddition pathway by reacting cyclic carbonates with amines. By far, this is the most general approach used due to its inherent advantages. Polyaddition reaction offer flexibility choosing reactants, allowing for a wide range of compounds to be utilized. This enables the customization of NIPU properties for specific applications. NIPUs obtained through the polyaddition of cyclic carbonates and diamines or polyamines are extensively studied in the literature, particularly concerning their mechanical properties. Furthermore, the synthesis of these NIPUs is more economically advantageous from an industrial standpoint, as it eliminates the need for solvents, catalysts, and the absence of by-products.34 Table 2 presents the comparison of synthesis methods of NIPU. It is crucial to carefully evaluate the benefits and limitations associated with each method prior to NIPU synthesis. This review focuses on the production of NIPU mostly but not limited to using cyclic carbonate with amine group through the polyaddition reaction. Further details on cyclic carbonates and types of amine will be discussed in the following

Ghasemlou *et al.* proposed a mechanism for the NIPU synthesis involving a tetrahedral intermediate. Fig. 4 depicts the reaction mechanism between five-membered cyclic carbonates and amines. Initially, the amine group attacks the carbonyl group in the cyclic carbonate leading to the formation of a tetrahedral intermediate. The next step involves the deprotonation of the tetrahedral intermediate, along with the elimination of hydrogen ions resulting from the attack of another amine group. Then, the presence of nitrogen atom induces the cleavage of the carbon–oxygen bond, leading to the formation of

Table 2 Comparison on synthesis methods of NIPU

Polycondensation	Ring-opening	Rearrangement	Polyaddition
- Use solvent and catalyst	- Absence of side-products	- Isocyanates are generated <i>in situ</i> during reaction	- Can be conducted with solvent- and catalyst-free method
- Longer reaction time	- Performed at high temperature	- Reactants are highly hazardous substances	- Absence of by-products
 Formation of low molecular weight by-products during reaction Required purification of obtained polymer 	- Toxic precursors		- Purification is not needed

Fig. 4 Proposed reaction mechanism between cyclic carbonates and amines to produce NIPU. Adapted from ref. 4 with permission from Elsevier, copyright 2019.

NIPU. This initiation of bond breakage is primarily attributed to the strong electron-withdrawing effect exerted by the nitrogen atom. The reaction between cyclic carbonate and amines involving a five-membered cyclic carbonate may lead to the formation of two isomers. The isomers contain a urethane moiety and a hydroxyl group in their backbone. The only difference between these isomers is the type of hydroxyl group present, whether a primary or secondary hydroxyl group. The formation of these isomers is dependent on the structure of intermediate compounds involved in the reaction.⁴

4 Reactants for preparing NIPU through polyaddition reaction

4.1 Cyclic carbonates

Cyclic carbonates (CC) are a class of versatile compounds that have garnered attention in both academic and industrial areas due to their attractive attributes in the context of green chemistry and sustainability, leading to the surge of interest in their synthesis and applications. In terms of properties, CC are relatively non-toxic, exhibit low volatility, and possess thermal and mechanical stability. They can be synthesized from a variety of relatively inexpensive starting materials. Owing to these advantages, cyclic carbonates have found diverse applications in various fields. Apart from their used as monomers in polymer synthesis, they can serve as reactants or green solvents in chemical reactions.35 Besides, CC are used in the formulation of electrolytes for energy storage devices like lithium ion batteries.36 Other than that, they have been used in industrial lubricants and as environmentally benign intermediates for the preparation of a wide variety of organic chemical products.37 There are several types of cyclic carbonates that can be classified based on their size and substitution patterns such as 5-, 6-, 7and 8-membered ring cyclic carbonates (5CC, 6CC, 7CC and

8CC), with the most common being the 5CC and 6CC. It is worth mentioning that the synthesis of 5CC is much easier as the 6CC have higher reactivity and lower stability.³⁸ Therefore, this review will be mainly focusing on the use of 5CC to synthesize NIPU. Fig. 5 depicts the structure of commonly produced 5CC.

The first work described on cyclic carbonates was conducted by Carothers et al. in the early 1930s. 39,40 Since then, the studies on cyclic carbonates have evolved, and they have now become commercially available with various methods, utilizing different precursors for their synthesis.41 Traditionally, the synthesis of cyclic carbonates relied on a chemical reaction between diols and phosgene or its derivatives. This method offered high yields due to the high reactivity of phosgene, albeit requiring additional purification steps. However, due to the toxicity associated with phosgene and its derivatives, alternative approaches have been developed.42 Fig. 6 displays some common routes to prepare cyclic carbonates using different starting materials. The most commonly studied and effective methods remain the cycloaddition reaction of carbon dioxide and epoxides. This reaction is favoured because it utilizes carbon dioxide, CO2 a renewable, non-toxic and abundant reactant, besides being relatively facile and straightforward. The preparation of CC can also be viewed as an efficient and practical method for CO₂ fixation and consumption. Remarkably, it can be performed efficiently without the need for a solvent. The reaction is also

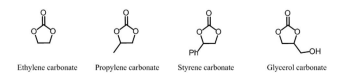


Fig. 5 Structure of commonly produced five-membered ring cyclic carbonates.

Fig. 6 Synthesis routes of cyclic carbonates with various precursors. Adapted from ref. 38 with permission from MDPI, copyright 2020.

thermodynamically favourable as the high free energy of epoxides counterbalance the high thermodynamic stability of CO_2 .

The industrial production of cyclic carbonates primarily utilized propylene carbonate and ethylene carbonate, derived from propylene oxide and ethylene oxide, respectively. These two epoxides are obtained from non-renewable petroleum derived chemicals. Other issues that raise concerns are about the toxicity and volatility of these starting materials. To address this issue, ongoing research is exploring various approaches aimed at achieving safe and sustainable production of renewable cyclic carbonates. One focus is on establishing alternative methods for the industrial production of epoxides from biobased materials. In view of this, various types of renewable resources can be used such as terpene, vanillin, tannin, lignin derivatives, glycerol, isosorbide, and vegetable oils.4 Among these sources, vegetable oils present as promising option as they are readily available and abundant. The presence of unsaturation sites in vegetable oils provides a wide range of possibilities for chemical modifications44 such as epoxidation, acrylation, carbonation, malenization, click chemistry and various other reactions. A detailed discussion on carbonated vegetable oils will be presented in Section 6 of this review.

4.2 Amine curing agents

The properties of NIPU can be tailored made based on the selection of amine used during the synthesis stage. The choice of amine can affect the molecular weight, crosslink density, and degree of branching of the polymer, which in turn dictates the mechanical, thermal, and other related properties of the resulting NIPU. The properties of NIPU can also be tuned by adjusting the ratio of amine to cyclic carbonate used in the synthesis.24 A higher amine-to-cyclic carbonate ratio can lead to higher crosslinking density and improved mechanical properties, whereas a lower ratio may result to a more flexible and lower stiffness polymer. Various types of amine have been investigated such as diamine, triamine or polyamine with aliphatic, cycloaliphatic, or aromatic structure. It is important to note that, in NIPU synthesis, the polyamine precursor must possess at least two amine groups within its structure. This is crucial for increasing reactivity towards cyclic carbonates and

achieving a high conversion rate.⁴ Primary amines are typically more reactive than secondary or tertiary amines due to their higher nucleophilicity. Diamines usually produced linear polymers, while polyamines can lead to branched or crosslinked polymers. The use of cycloaliphatic and aromatic amines can improve the chemical, mechanical, and thermal stability of the resulting polymer.⁴⁵

Currently, there is progressing research and development to produced bio-based amines from renewable resources such as biomass, agricultural waste, or other sustainable feedstock. Similarly to cyclic carbonates, the development of bio-based amines is driven by the desire to reduce reliance on fossil fuels and to mitigate the environmental impact associated with traditional amine production methods. The available literature suggests that amino acid, vegetable oil derivatives, sugar derivatives, lignin, and proteins can serve as feedstock for the production of amines.46 Overall, regardless of the source, the selection of amine plays a crucial role in determining the properties of NIPU, and careful consideration of the amine group and its ratio to cyclic carbonate is necessary to tailor the material properties and meet specific application requirements. Fig. 7 displays some examples of amine curing agents that has been used to prepare NIPU.

5 Properties of NIPU

The properties of NIPU can be tailor-made to meet specific requirements of the final products. The abundant options of starting materials and pathways ensure the diversity of NIPU produced.47 The properties of NIPU can be customized by selecting appropriate precursors and incorporating additives. This flexibility in properties customization allows for the optimization of NIPU for different applications. NIPU has been reported to demonstrate superior thermal stability, resistance to water and resistance to solvents compared to conventional PU.48 The presence of pendent hydroxyl groups in the NIPU structure make it capable of forming intermolecular hydrogen bonds with urethane carbonyl groups. These intramolecular hydrogen bonds lead to the decrease of the susceptibility of urethane linkages to hydrolysis. Furthermore, NIPU demonstrate enhanced thermal stability as they lack thermally labile allophanate and biuret groups that are typically present in the chemical structure of conventional PU.49 Meanwhile the mechanical properties of NIPU are significantly influenced by processing conditions, the carbonate/amine ratio, and presence of fillers. In Table 3, a comparison of isocyanate-based PU (both synthetic and bio-based) and NIPU is presented. Particularly, the mechanical properties of polyurethane vary based on the specific types of PU produced. Some interesting properties of NIPU will be briefly discussed in this section, such as selfhealing, shape memory effects, and reprocessibility.

5.1 Self-healing

The concept of self-healing materials has gained attention in recent years as a way to develop repairable materials that can last longer, required less maintenance, and reduced need for

4,4-diaminodiphenyl methane (DDM)

Fig. 7 Example of amine curing agents used to synthesize NIPU

4.4-diaminodiphenyldisulfide (DDS)

Table 3 Mechanical properties of isocyanate and non isocyanate-based PU

	Mechanical properties		
Sample	Tensile strength	Elongation	Ref.
Isocyanate-based PU			
Synthetic PU	11.0 MPa	440%	50
Palm oil-PU foam	60-115 kPa	38-80%	51
Rapeseed oil-PU foam	82–115 kPa	78-154%	52
NIPU			
Soybean oil_NIPU film	0.35-1.35 MPa	2.2-4.5%	53
Soybean oil_NIPU	1.7-6.2 MPa	165-306%	54
membranes			
Soybean oil_NIPU film	0.3-11.1 MPa	3-433%	55
Soybean oil_NIPU film	1.8-6.2 MPa	152-309%	56
Canola oil_NIPU film	5-8 MPa	103-192%	57
Jatropha oil_NIPU film	4-17 MPa	30-230%	58
Linseed oil_NIPU film	2-18 MPa	1-310%	59
Soybean oil_NIPU film	7.2-10.1 MPa	80-97%	60
Sunflower oil_NIPU film	0.9-5.2 MPa	100-240%	61
Linseed oil_NIPU film	11.9-18.9 MPa	26.0-57.1%	26
Linseed oil_NIPU film	5.0-9.5 MPa	2148-327.3%	62
Soybean oil_NIPU film	0.45-0.97 MPa	36.7-92.0%	63

replacements. Generally, self-healing polymers (SHP) can be classified based on their mechanism and working chemistry as presented in Fig. 8.⁶⁴ The SHP by mechanism involves either intrinsic or extrinsic repair of the materials. Intrinsic SHP refers to polymers that can restore their own structural integrity without relying on external assistance, whereas extrinsic SHP

requires external agent pre-embedded in the polymer matrix to trigger them to repair the damage. In terms of working chemistry, SHP can be categorized into autonomous and non-autonomous. Autonomous polymers can recover their shape automatically or respond to damage and recover without relying on external stimuli to initiate the healing process. Meanwhile, non-autonomous self-healing polymers require external stimuli to heal the damage surface. Without the presence of stimuli, the healing process does not occur. ⁶⁵⁻⁶⁷

Diaminodiphenylsulphone (DDPS)

Self-healing NIPU has been investigated and reported by several studies. Raut et al. described the preparation of selfhealable hydrophobic NIPU based on carbonylimidazoleamine reaction and dynamic Diels-Alder click reaction employing furan-maleimide cycloaddition. The produced NIPU exhibits excellent hydrophobic and self-healing properties.⁶⁸ In one study, Hu et al. produced degradable and self-healable NIPU by copolymerizing bis(6-membered cyclic carbonate) and amino-terminated liquid nitrile rubber. The report suggested that NIPU indeed possesses self-healing characteristic with a healing efficiency of more than 88%. They suggest the mechanism is due to the internal transesterification exchange between the hydroxyl and carbamate groups.⁶⁹ Further, Wang et al. studied the self-healing properties of NIPU from the reaction of bio-based diglycerol bis(cyclic carbonate) with furan methylamine and trimethylolpropane tris(poly[propylene glycol], amine terminated) ether or tris(2-aminoethyl) amine under mild conditions, followed by the reaction with bismaleimide. The highest healing efficiency of NIPU was obtained at 80%.70 In other study, they prepared soft self-healing NIPU via

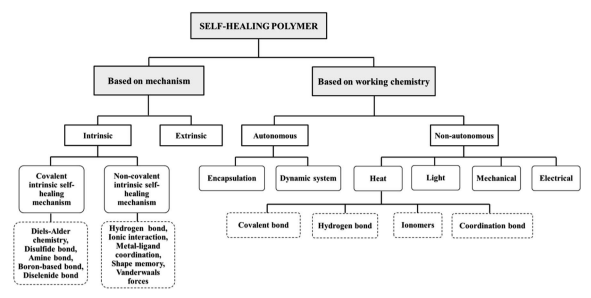


Fig. 8 General classification of self-healing polymer. Adapted from ref. 64 with permission from Wiley, copyright 2022.

Diels-Alder reaction between furan terminated NIPU and unsaturated polyester. The obtained NIPU achieved 100% self-healing efficiency.⁶⁶

5.2 Shape memory effects

The concepts of self-healing and shape memory effects of polymers are interconnected. While self-healing polymers repair damage by restoring their molecular structure, shape memory effects (SME) refer to the ability of polymers to remember their original shape and recover it after being deformed, typically when subjected to a specific stimulus, such as heat, light, electricity, and chemical induction. SME is one approach to facilitate healing of materials. Shape memory is the consequence of molecular structure, reversible mobility changes, conformational entropy, and programming.65 To enable shape memory in a polymer, two essential structural elements are required which are; (a) permanent net-points or junctions, and (b) reversible molecular switching segments. The permanent net-points or junctions serve as stable anchor points within the polymer network that maintain the original shape of the material. They are responsible for holding the polymer chains in place and preserving the memory of the permanent shape. Meanwhile, reversible molecular switching segments allow the polymer chains to undergo reversible conformational changes when subjected to a stimulus. The reversible switching enables the transition between the temporary shape and the permanent shape. These combinations allow the polymer to exhibit shape memory behaviour.71

Several studies have reported the shape memory effects of NIPU. For instance, Yin *et al.* synthesized semicrystalline NIPU with tunable triple shape memory properties by introducing behenic acid into the system. They were able to obtained NIPU with a shape recovery rate close to 100%.⁷² Another study on triple shape memory NIPU was reported by Schimpf *et al.* NIPU was obtained by curing pentaerythritol-based cyclic carbonates

with hyperbranched, semi crystalline polyamidoamines with the subsequent addition of behenic amides. The production of nanocrystalline with programmable shapes can be attributed to the crystallization of the behenic amide side chains.73 Meanwhile, another study on lignin/NIPU hybrid polymers were synthesized by Zhang et al. using bis(6-membered cyclic carbonate) and polyetheramine. The lignin/NIPU hybrid polymers exhibited excellent shape memory effects due to the transcarbamoylation reactions in their dynamic covalent network triggered by high temperature.74 Another study based on lignin derived NIPU was presented by Sternberg and Pilla. In their study, Kraft lignin was functionalized with glycerol carbonate and subsequently reacted with Priamine 1074 curing agent. The cured NIPU samples were subjected to shape memory effect test by heating them at 105 °C for 10 minutes. A semicircular deformation was induced as the samples cooled down to room temperature. The produced lignin-based NIPU exhibits the ability to retain the deformation after cooling to room temperature and shows reversibility by returning to the pre-deformed shape upon re-heating.75 Further, Li et al. synthesized a five-membered bicyclic carbonate by reacting meso-erythritol with carbon dioxide. The as-synthesized carbonate was reacted with Jeffamine T-403, a trifunctional polyetheramine curing agent to produce thermosets NIPU. This NIPU was then incorporated with poly(2-oxo-1,3-dioxolane-4-yl) methyl methacrylate -grafted Fe₃O₄ nanoparticles. It was found that the nanocomposites retained excellent shape memory properties, and this shape memory behaviour could be triggered by the photothermal effect of Fe₃O₄ nanoparticles.⁷⁶

5.3 Reprocessibility

Reprocessability is an important property of polymers that enables recycling, reshaping, and reuse of these materials. Reprocessability refers to the ability of a polymer to undergo reprocessing without significant degradation or loss of

properties. Reprocessable polymers are highly sought after as they can contribute to achieving sustainability goals, waste reduction, and resource efficiency. One example of study on reprocessibility of NIPU was conducted by Chen et al. They synthesized NIPU using poly(propylene glycol) dicyclocarbonate and tris(2-aminoethyl)amine in the presence of 4-(dimethylamino)pyridine catalyst. The reprocessing ability of NIPU was performed at 140 °C for 2 hours, and the resulting NIPU elastomer showed full property recovery after multiple reprocessing steps. This study revealed that the rearrangement of NIPU networks during reprocessing involves both dissociative reversible cyclic carbonate aminolysis and associative transcarbamoylation exchange reactions. Through these reactions, NIPU can be designed to be intrinsically reprocessable with full property recovery.⁷⁷ A subsequent study on NIPU reprocessibility was conducted by Hu et al. They synthesized NIPU through the reaction of carbonated sovbean oil with a difunctional amine with 4-(dimethylamino)pyridine catalyst. This study achieved a complete recovery of NIPU network after multiple melt-state recycling steps under relatively mild reprocessing conditions. In addition to the reversible cyclic carbonate aminolysis and transcarbamoylation exchange reactions, the networks were found to undergo a third dynamic chemistry through a transesterification exchange reaction.⁶⁹ Recently, Purwanto et al. described a rapid synthesis of reprocessable self-blowing NIPU foams through the competitive reactions of cyclic carbonate aminolysis and thioldecarboxylation. They able to achieve a significant reduction in foaming production time, from 20 hours to 30 minutes when the gelling reaction conducted at 80 °C. The NIPU foams were capable to be melt-reprocessed into well-consolidated films for multiple cycles with a full recovery of cross-link density.78 Another recent study investigated the reprocessability of biobased non isocyanate polythiourethane (NIPTU) conducted by Chen et al. In this study, raw materials were obtained from biowastes i.e. five-membered cyclic dithiocarbonates (DTC); NC-514-DTC and Cyclo-DTC were derived from cashew nutshell liquid and rice husks. They employed JEFFAMINE T-403 as the

6 Case studies on vegetable oil derived NIPU

Vegetable oils are derived from plant sources such as seeds, fruits, nuts, and grains. They can be broadly classified as edible and non-edible oils based on their chemical content. Edible vegetable oils are derived from sources that are commonly used for direct human consumption as food intake, whereas non-edible oils are derived from sources that are not used for food purposes. Fig. 9 illustrates some examples of edible and non-edible vegetable oils available in the market and

curing agent. Due to its dynamic covalent thiourethane and

disulphide cross-links, this NIPTU demonstrated excellent

reprocessability with full recovery after multiple reprocessing

stages. 79 Based on the observations from the presented studies

so far, it can be deduced that NIPU has reprocessable proper-

ties, including those prepared from bio-based resources.

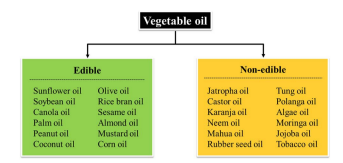


Fig. 9 Some examples of edible and non-edible vegetable oils.

ready for modification. While it is relevant to use edible oil as precursor for polymer production, there are valid concerns and criticisms regarding its potential impact on food security. In this context, the use of non-edible oils as a feedstock for polymer production is an attractive alternative to edible oils, providing a commercially viable and sustainable solution for the polymer industry.⁸¹

The next subsection presents a series of case studies on vegetable oil-based NIPU derived from both edible and non-edible oil that have been reported to this date. The case studies highlight the general characteristics of the oils, including their origin, fatty acid composition, and other related properties. They also offer insights into the preparation methods of NIPU, including the choice of curing agent, catalyst, and other additives, as well as their impact on the final properties of NIPU. Moreover, the studies probe into the properties and performance of NIPU and their suitability for various applications in diverse fields.

6.1 Soybean oil

Soybean oil (SBO) is extracted from the seeds of the soybean plant (*Glycine max*). Soybean originated from China and now has been cultivated in other parts of the world. Currently, SBO is the second largest source of vegetable oil after palm oil. South America currently accounts for over 50% of global soybean production worldwide. The oil is high in unsaturated fatty acids, comprising approximately 80% of oleic, linoleic, and linolenic acids. In addition to its use in food applications, SBO is also gained significant interest in scientific and industrial fields to produce bioplastics, coatings, paints, and lubricants. Other than these applications, the use of soybean oil as a raw material in the production of polyurethane also has been

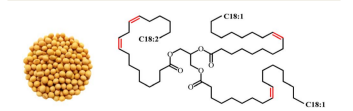


Fig. 10 Image of soybean seeds and the chemical structure of soybean oil. The image is obtained from ref. 85 with permission from MDPI, copyright 2023.

extensively studied in recent years. Fig. 10 shows the image of soybean seeds and the chemical structure of SBO.

Among various types of vegetable oil reported for NIPU production, soybean oil has been the most investigated so far. The earliest study on soybean oil-based NIPU was conducted by Tamami et al. to examine the effects of different amine groups on the physical properties of the resulted NIPU. Two diamine groups, ethylenediamine (EDA) and hexamethylenediamine (HMDA), were used along with one triamine group, tris(2aminoethyl)amine (TA). TA_NIPU display the highest T_g of 43 °C, the smallest strain at break (70%) and the lowest soluble fraction (6.8%) compared to EDA and HMDA. This can be attributed to the fact that TA has a tighter network than the diamine groups. HMDA_NIPU has the lowest $T_{\rm g}$ value of 18 °C and EDA_NIPU has 34 °C. The higher level of flexibility of HMDA moiety explains the lower $T_{\rm g}$ value of HMDA_NIPU compared to EDA_NIPU regardless being the same type of diamine, only differs in chain length.86

In another study, Javni et al. study the use of aromatic and cycloaliphatic diamines to increase the strength and rigidity of NIPU. The properties evaluated included mechanical, thermal, and swelling behaviour. The findings indicate that the mechanical properties depend on crosslinking density and the hydrogen bond between urethane, hydroxyl, and ester groups, which are controlled by the carbonate to amine structure and the amine ratio. The presence of rigid aromatic and cycloaliphatic structures increases intersegmental physical interactions, resulting in higher strength. Thermal analysis revealed that the T_g was affected by the carbonate to amine ratio. In regards to NIPU swelling in toluene, factors like the degree of crosslinking and solubility parameters of NIPU affected the swelling behaviour. Additionally, NIPU based on aromatic and cycloaliphatic diamines exhibited higher hydrophobicity and were less susceptible to water penetration compared to those based on aliphatic diamines.55

Lee and Deng described the synthesis of NIPU using soybean oil and lignin. Initially, the urethane monomer was prepared by the reaction of carbonated soybean oil (CSBO) with APTES. Following that, the monomer was reacted with various loading of lignin to produce polyurethane. In this case, lignin functions as the hard segment whereas polysiloxane linkages (-Si-O-Si-) act as the soft segments. Two different curing temperatures were used, which were drying at room temperature and curing at 60 °C. It was found that tensile strength enhanced with the increase of lignin content and curing at higher temperatures. This enhancement can be attributed to the rigidity of the lignin aromatic structure and lignin acting as the crosslinking agent. In contrast, curing at low temperature resulted in an increase in strain, as the lignin is predicted to blend with the polymer instead of acting as a crosslinking agent. This study proves that the mechanical properties of NIPU can be controlled by varying the temperature and lignin content.53

Poussard et al. prepared CSBO from epoxidized soybean oil (ESBO) under supercritical CO2 conditions at 120 °C and 100 bar. A complete conversion of ESBO to CSBO was achieved within a 9 hours reaction time. The use of supercritical CO2 conditions decreased the viscosity of soybean oil and facilitated

the coupling of CO2 with epoxides leading to complete conversion in a shorter time compared to lower pressure and higher temperature conditions. In parallel, bio-based amino-telehelic oligoamides were synthesized from a dimeric fatty acid and ethylenediamine. NIPU was prepared with short 1,4-butanediamine (BDA) and 1,5-pentanediamine (PDA), and with the higher molecular weight as-prepared oligoamide by meltcopolymerizion with ESBO. Thermal stability analysis of NIPU based on CSBO and BDA or PDA showed a relatively lower thermal stability compared to NIPU containing the bio-based oligoamide. Despite this, the thermal stability of all prepared NIPUs was higher than conventional polyurethane, which recorded around 310 °C. The improvement in stability was expected due to the hydroxyl functions in the urethane generating intra- and intermolecular hydrogen bonds.87

A study on chitosan-based non isocyanate polyurethane was reported by Das et al. In their study, CSBO was mixed with chitosan solutions at different concentration (1:1, 1:2, 1:3)weight ratio of CSBO to chitosan) to obtain a fully bio-based NIPU. Thermal analysis based on TGA revealed NIPU 1:3 has the highest thermal stability due its highly crosslinked polymer network structure. This study found out that increasing chitosan content improves the thermal stability of NIPU. Meanwhile, DSC analysis shows that NIPU samples exhibit two glass transition temperatures for he hard segment, $T_g(h)$ and soft segment, $T_g(s)$ of the polymer, indicating phase separation phenomena. The produced NIPU also exhibit amorphous nature as there is no endothermic or exothermic behaviour observed. In addition, this study also evaluated the chemical resistance and optical properties of the NIPU_1:3 in different chemical environments: acid, alkali, and neutral. The samples exhibited color changes as the pH of the environment changed, as displayed in Fig. 11. They also observed changes in the morphology of the NIPU films due to leaching of some components into the chemical media.88

Further, Doley et al. prepared a hybrid NIPU from CSBO and highly branched cyclic carbonate polyester (CHPE) reacted with polyamidoamine. Thermal analysis study by TGA showed that the hybrid NIPU has good thermal stability due to the presence of rigid aromatic rings, polar-polar interactions, and hydrogen bonding interactions. The presence of hydroxyl groups in the structure also contributes to the increased thermal stability. In addition, the chemical resistance of NIPU was tested in three media; HCl (10% aq.), NaOH (2% aq.), and water for 21 days. The results show that NIPU has satisfactory resistance towards water and acidic media due to the more crosslinked polymer

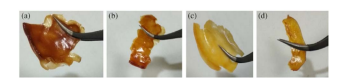


Fig. 11 Physical appearance of NIPU_1: 3 films tested for 48 hours in different media; (a) HCl, (b) NaCl, (c) NaOH, and (d) no chemical environment. Adapted from ref. 88 with permission from Wiley, copyright 2020

chain. However, NIPU's resistance to alkali media was poor due to the presence of hydrolysable ester bonds in both CSBO and CHPE. 60

Gholami et al. prepared wound dressing membranes from soybean oil-based NIPU containing azetidinium groups as an antibacterial agent. The article highlights the importance of using antibacterial wound dressings to reduce the population of microbes at the wounded site and eliminate the need for oral uptake of antibiotics that can be harmful. Two types of NIPU membranes were prepared; one with an internal secondary amine group, tetraethylenepentamine (TEPA_NIPU) and one without (HMDA_NIPU). The azetidinium containing crosslinked networks were obtained by the addition of epichlorohydrin (ECH) at 1:1 and 1:2 molar ratio of ECH to TEPA_NIPU. For the antibacterial assay, HMDA NIPU was used as a control. The antibacterial activity against Staphylococcus aureus (S. aureus) and Escherichia coli (E. coli) was monitored. Overall, the dressings showed excellent antibacterial properties against both bacteria and were non-cytotoxic to fibroblast. The NIPU dressings had desirable tensile properties with \sim 6 MPa tensile strength and ~200% elongation at break that remained under a hydrated state. The dressings were capable of maintaining a moist environment over mildly to moderately exuding wounds due to their ability to absorb (reaching up to 30% equilibrium water absorption) or release them as vapour (water vapour transmission rate of 390 to 710 g per m² per day for dry and wet dressings). Therefore, these dressings have the potential to promote the healing of injured skin tissue in a hygienic setting.54

Dong et al. design and prepared a self-healing CSBO derived NIPU incorporated with dynamic disulphide bonds. The NIPUs were prepared using different types of amine; bio-based dimeric fatty diamine PA650, isophorone diamine (IPDA), amino acidderived flexible disulphide-based cystamine (CA), and the premixture of these amines. The self-healing properties were adjusted by varying the amine ratio. The healing efficiency was estimated through the recovery of tensile strength, and it was observed that the self-healing capability of NIPU was improved with the addition of CA. In this case, the different diamines used have specific contribution to the properties of NIPU. The use of PA650 contributed to higher thermal stability and elasticity, while the addition of IPDA resulted in good tensile strength. The incorporation of CA significantly improved the self-healing efficiency by 45%. The efficiency was enhanced by one time when NIPU was healed at 80 °C for 4 hours. The good mobility and flexibility of NIPU networks formed by linear CA and bulky PA650 contribute to the improved healing ability. The proposed mechanism for self-healing of NIPU and the image of healed NIPU are presented in Fig. 12.56

Another study on self-healing and potential reprocessing of soybean oil-based NIPU was reported by Yang *et al.* In this work, CSBO was reacted with two types of amines; 4,4'-diaminodiphenylmethane (DDM) and 4,4'-diaminodiphenyldisulfide (DDS). In essence, disulfide bonds are one kind of dynamic covalent bond that can undergo bond exchange reactions under various stimuli (light, heat, ultrasound, *etc.*) making them attractive for materials to reprocess

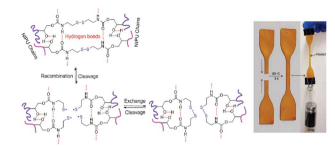


Fig. 12 Proposed mechanisms for self-healing NIPU and image of NIPU healed at 80 °C for 4 hours. Adapted from ref. 56 with permission from RSC, copyright 2020.

and heal under relatively mild conditions. The produced NIPU have good thermal stability and the tensile strength properties were improved by the introduction of disulphide bonds in the structure. Moreover, the disulphide bond was found to be effective in promoting self-healing properties. Fig. 13 shows the image of reprocessed ESO-DDS, CSBO-DDM and CSBO-DDS samples by using hot pressing method. The study revealed that the different network structures had a significant impact on the performance of the samples, and the dual dynamic covalent bond (CSBO-DDS) was more effective in accelerating molecular chain rearrangement than either single bond type (ESO-DDS or CSBO-DDM). Overall, the results suggest that the synergistic effect of disulphide bond exchange and carbamate bond exchange provides a feasibility study for designing NIPU with milder self-healing and reprocessing behaviours.⁸⁹

Liu *et al.* prepared a fully bio-based NIPU from CSBO and a terpene derivative with a dynamic network structure. In their work, 1,8-menthane diamine (MDA) derived from turpentine was used as the curing agent. The ratio of CSBO to MDA was varied using the amino/cyclic carbonate molar content with ratio of 1.2, 1.0 and 0.8, respectively. Through a stress relaxation study, the produced NIPU displayed dynamic covalent bond exchange. Additionally, the NIPU possesses self-healing, recyclability, reprocessing, and shape memory effect as can be observed in Fig. 14. In fact, the recycling process can be

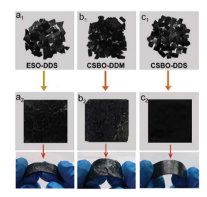


Fig. 13 Reprocessing of (a) single-disulphide ESO-DDS, (b) single carbamate CSBO-DDM, and (c) dual dynamic CSBO-DDS by hot pressing technique. The image obtained from ref. 89 with permission from RSC, copyright 2021.

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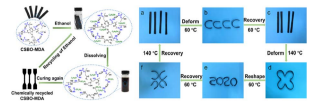


Fig. 14 Chemical recycling process and shape memory effect of CSBO/MDA_NIPU. (a) deformed at 60 °C and then cooled at 4 °C for a new temporary (b), recovered at 60 °C to the original rectangular shape (c), heated at 140 °C and then cooled at 4 °C (d), reshaped at 60 °C and then cooled at 4 °C shape (e), recovered at 60 °C to the permanent shape (f), and reconfigured at 140 °C to the original shape I. Images obtained from ref. 90 with permission from ACS, copyright 2021.

repeated multiple times with less impact on the properties of the recovered NIPU. The mechanical properties and thermal stability of NIPU are almost fully recovered (>85%) after several reprocessing cycles, indicating that the bond exchange repairs the network. Interestingly, this study produced NIPU with triple-shape memory due to the additional reversible phase transition.⁹⁰

6.2 Linseed oil

Linseed oil (LSO) is extracted from the seeds of flax plant (Linum usitatissimum). The terms "linseed" and "flaxseed" are often used interchangeably, depending on the geographic location and the context in which they are being used. The chemical composition of LSO is characterized by the presence of a high concentration of polyunsaturated fatty acids (73%) made up of linolenic acid and linoleic acid; 18% monounsaturated fatty acid (oleic acid); and 9% of saturated fatty acid (stearic acid).91,92 LSO is classified as a drying oil because when coated on a nonabsorbent substrate and being exposed to air, it dries promptly forming a hard, non-melting, and water insoluble film.93 The drying process occurs due to a polymerization reaction initiated by exposure to light and facilitated by the presence of atmospheric oxygen through a series of chemical reactions called autoxidation.94 These properties make LSO a popular ingredient in paints, varnishes, and wood finishes. The image of linseed and its oil chemical structure is displayed in Fig. 15.

Moritz and Rolf examined the effect of homogenous and heterogeneous catalyst for CLSO synthesis. TBAB was use as the homogeneous catalyst, whereas silica-supported 4-pyrrolidino-pyridinium iodide (SiO_2 -I) served as the heterogeneous catalyst.

C18:2

Fig. 15 Image of linseed and the chemical structure of linseed oil. The image adapted from ref. 95 with permission from MDPI, copyright 2020.

This study demonstrate the use of heterogeneous catalyst aids in simplifying the recovery process, however the activity is still not comparable to that of homogenous catalysts. At the same condition of 30 bar, TBAB catalyst shows complete conversion after 20 hours, whereas, SiO₂-I required 45 hours to reach completion. The produced CLSO was then cured with EDA, BDA and IPDA. The high carbonate content of CLSO led to high viscosity causing processing problems. Thus, CLSO was blended with CSBO to reduce the viscosity and increase the gel time. In this case, CSBO acted as a reactive diluent. Variations in carbonate content by varying the ratio of CLSO and CSBO significantly affected the thermal and mechanical properties of NIPU. They observed a high stiffness of NIPU was produced when the pure CLSO was cured with IPDA, increasing the glass transition temperature to 60 °C. Although the stiffness of NIPU has increased, there is a significant reduction in its ability to stretch without breaking, indicating severe brittleness. 59

Pouladi et al. conducted a study on the anti-corrosion performance of NIPU at various carbonate contents of CLSO. This work started with epoxidation, followed by carbonation to form cyclo-carbonate groups. During the carbonation stage, CLSO was prepared with three percentage of carbonate content which were 45% (CLSO45), 75% (CLSO75), and 100% (CLSO100) obtained at 24, 36 and 48 hours of reaction time, respectively. Each type of CLSO reacted with diethylenetriamine (DETA) to form NIPU. The results indicated an improvement in adhesion and hardness, and a reduction in flexibility with increasing carbonate content of CSLO. The increase in carbonate content contributed to a higher formation of hydroxyl groups in the NIPU structure due to the reaction of cyclo-carbonate and amine. This led to the improved adhesion strength and mechanical properties of the polymer. The corrosion study based on electrochemical impedance spectroscopy analysis revealed NIPU produced from CLSO75 exhibited superior anticorrosion properties, comparable to conventional PU. For CLSO100_NIPU, a high content of carbonate led to more formation of hydroxyl group upon reaction with amine and thus resulted to a decrease in corrosion resistance (Fig. 16).96

Zhang *et al.* reported a study on the preparation of waterborne non isocyanate polyurethane (NIWPU) from CLSO using different amine group 3,3'-diamino-*N*-methyldipropylamine (DMDPA), HMDA, *m*-xylylene diamine (*m*-XDA) and 2,2'-(ethylenedioxy)bis(ethylamine) (EDR) to produce cationic (CNIWPU), anionic (ANIWPU) and non-ionic (NNIWPU) polyurethane. The NIWPU dispersions exhibit excellent storage stability and



Fig. 16 NIPU foam produced from CLSO and HMDA. Image obtained from ref. 26 with permission from ACS, copyright 2021.

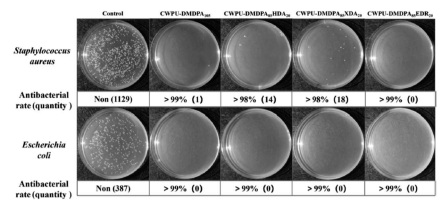


Fig. 17 Antibacterial study of cationic NIWPU against S. aureus and E. coli. The image obtained from ref. 62 with permission from Elsevier, copyright 2023.

tailorable properties. For instance, the anionic NIWPU exhibited good coating performance with the highest pencil hardness was recorded at 7H and crosshatch adhesion of 5B, whereas, the non-ionic NIWPUs showed a tailorable cloud points ranging from 60 to 80 °C. Meanwhile, the cationic NIWPU showcased good mechanical properties with tensile strength more than 9 MPa and elongation at break surpassing 200%. Notably, CNIWPU exhibited antibacterial properties against *Escherichia coli* and *Staphylococcus aureus* above 98%, as depicted in Fig. 17. Based on these findings, the authors suggested the potential use of these NIWPUs as antibacterial coatings, adhesives and soft elastomers.⁶²

6.3 Sunflower oil

Sunflower oil is extracted from the seeds of the sunflower plant (*Helianthus annus* L.). This plant is native in North America and has been cultivated around the world. The oil contains around 85% unsaturated and 15% saturated fatty acid, with 44–75% linoleic acids and 14–43% oleic acid content. In addition to its culinary uses, sunflower oil is also used in cosmetics and skincare products due to its moisturizing and nourishing properties. It is a common ingredient in a variety of products, including lotions, soaps, and hair care products. Sunflower oil also has been used to prepare polyurethane. A study synthesized PU films using sunflower oil-based polyol and diisocyanate with the addition of graphene oxide and reduced graphene oxide. Another study reported on TiO₂-nanocomposites coatings from sunflower oil-based amide diol as a soft segment. Additionally, Asare *et al.* prepared a polyol from sunflower oil, reacted it

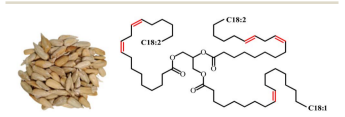


Fig. 18 Sunflower seeds image and the chemical structure of sunflower oil.

with methylene diphenyl diisocyanate to produced rigid polyurethane foams. ¹⁰² Fig. 18 displays the image of sunflower seeds and the chemical structure of the oil. ¹⁰³

In the case of NIPU, there have been a few studies on sunflower oil reported so far. In one study, Boyer et~al. synthesized terminal and internal epoxidized fatty acids diesters (TEFAD and IEFAD) from sunflower oil's oleic fatty acid methyl ester. TEFAD and IEFAD were then reacted with CO_2 to produce terminal and internal carbonated fatty acids diesters (TCFAD and ICFAD). Subsequently, NIPU was obtained by the polyaddition reaction of TCFAD and ICFAD with EDA and IPDA. The resulting NIPU exhibits molecular weights up to 13 500 g mol $^{-1}$ with a glass transition temperature (T_g) of -15 $^{\circ}$ C. This study reported that NIPU produced from IPDA resulted in the absence of the side reaction that typically occurs between amine and ester functions, which usually leads to the formation of amide linkages. 104

A fully bio-based NIPU was prepared using carbonated sunflower oil and novel bio-based amines synthesized form

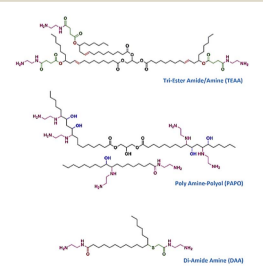


Fig. 19 The chemical structure of synthesized bio-based amines. Reuse from ref. 105 with permission from Elsevier, copyright 2018.

castor oil and oleic acid, as conducted by Farhadian *et al.* Three different amine groups were synthesized; tri-ester amide/amine (TEAA), polyamine-polyol (PAPO) and di-amide amine (DAA) as shown in Fig. 19. Then, carbonated sunflower oil was prepared by the reaction of epoxidized sunflower oil with CO₂ at atmospheric pressure. Subsequently, NIPU was obtained by the reaction of CSFO with the bio-based amine groups without a catalyst and solvent. Based on thermal and physical characterizations, bio-based NIPU exhibits excellent thermal stability, degradation resistance, and low water absorption. The thermal stability is enhanced is attributed to the presence of amide, ester and hydroxyl groups in their structure, as well as their comparatively higher molecular weight than short diamines.¹⁰⁵

Farid *et al.* prepared NIPU a flame-retardant coating from carbonated sunflower oil and 1,4-phenyldiamine with the addition of varied concentration of zirconia/silica (ZrO₂/SiO₂) nanoparticles. ZrO₂/SiO₂ was formed by an *in situ* reaction which binds ZrO₂ nanoparticles with SiO₂ nanospheres. Through thermal stability and flammability studies, the materials showed high resistance to flammability and demonstrated the ability to withstand direct contact with fire without ignition. The use of amine with aromatic properties significantly enhanced the flame retardancy of the produced NIPU and further improved by the addition of ZrO₂/SiO₂ nanoparticles. The authors reported that the optimum loading of ZrO₂/SiO₂ was up to 3% which give the best thermal, mechanical, and chemical resistance performance. At 5% ZrO₂/SiO₂, agglomeration was observed, impacting the properties of NIPU.

Doley and Dolui investigated the effects of different diamine groups on the properties of NIPU by using short chain diamine (EDA), long chain aliphatic diamine (DETA), and cyclo aliphatic diamine (EDA) for anti-corrosive coating. The findings indicate that the properties of NIPU were influenced by the diamines structure and the molar ratio of carbonate to amine used. Diamines with shorter chains resulted in a higher proportion of urethane and amide groups compared to diamines with longer aliphatic chains or cycloaliphatic diamines. In terms of coating properties, NIPU based on IPDA showed high tensile strength, hardness, thermal stability, gloss, and chemical resistance compared to NIPU based on EDA of DETA. Additionally, IPDA-based coating demonstrates the lowest corrosion rate and highest resistance due to its rigid structure.⁶¹

6.4 Castor oil

Castor oil is obtained from the seeds of castor plant (*Ricinus communis*) belonging to the Euphorbiaceae family. The main component of castor oil is ricinoleic (90%), an 18-carbon fatty acid with a hydroxyl group located on the 12th carbon, along with linoleic (4%), oleic (3%), palmitic (1%), stearic (1%), and linolenic (<1%) fatty acids. The oil has been used for medicinal and industrial purposes for centuries, including applications in lubricants, coatings, surfactants, printing inks, paints, *etc.*¹⁰⁷ Other than being non edible, the high content of ricinoleic acid made castor oil a viable option as a monomer in the production of non isocyanate polyurethane.¹⁰⁸ Fig. 20 displays the image of castor seeds and the chemical structure of castor oil.

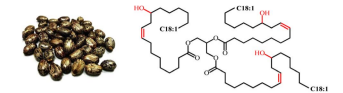


Fig. 20 Image of castor seeds and chemical structure of castor oil. The image was obtained from ref. 109 with permission from Elsevier, copyright 2020.

One of the earlier studies on NIPU from castor oil derivatives was reported by Pathak et al. The research commenced with the esterification of tris-2-hydroxy ethyl isocyanurate (THEIC) with dehydrated castor oil fatty acid (DCOFA) to produce THEICester of fatty acid (TEFA). Further, TEFA was epoxidized (ETEFA) and underwent a carbonation reaction (CTEFA). The produced CTEFA was subjected to reactions with three types of amine groups; diaminodiphenylsulphone (DDPS), HMDA and IPDA to produce NIPU. The type of amine used significantly influenced the properties exhibited by the NIPU coatings. In this case, the corrosion rates for HMDA_NIPU, IPDA_NIPU and DDPS_NIPU were recorded at 0.01599, 0.006516, and 0.0001989 mmpy, respectively. This finding proved that coating cured with aromatic and cyclo-aliphatic amines resulted in a better anticorrosion properties compared to coating cured with aliphatic amines. The aromatic hardener (DDPS) would impart rigidity and barrier protection to the coating, whereas the long aliphatic chain in the HMDA hardener produced a porous and permeable coating and thus decreasing the barrier properties.45

Following that, Rodrigues et al. prepared a simple on-pot procedure of non isocyanate poly(acyl) urethane using castor oil and urea in a presence of a catalyst. 110 Since castor oil contain large amount of ricinoleic acid, it can be directly utilized to prepare NIPU without prior modification. In this study, they assess the ideal reaction parameters to synthesize NIPU by varying the oil to urea ratios (1:2, 1:3, 1:4), synthesis temperature (50 °C to 100 °C), curing time (24, 48 and 72 hours), temperature (80 °C to 100 °C), type of catalyst (BF₃· OEt₂ (CH₃COO)₂Zn·2H₂O, SnCl₂·2H₂O), and percentage loading of the catalyst (5% and 10% w/w). They suggested 1:2 ratio of castor oil to urea is the best ratio for synthesis with 10% w/w of BF_3 · OEt_2 as the catalyst and the reaction proceeded at 70 °C. As for curing, the NIPU was well produced at 100 °C for 72 hours curing temperature and time. GPC analysis proved the formation of polymeric NIPU with an average molecular mass exceeding 10 000 g mol⁻¹ and a polydispersity index of 1.2. Results from DSC and DMA analyses indicated a relatively uniform chemical composition, while the polymeric chains exhibited varied sizes.110 This information is important for understanding the properties and potential applications of NIPU, as the size and distribution of its polymer chains can impact its strength, flexibility, and thermal stability.

6.5 Jatropha oil

Jatropha oil is derived from the seeds of Jatropha curcas plant. Jatropha curcas is belongs to the Euphorbiaceous family and is

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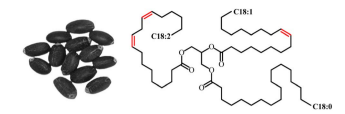


Fig. 21 Image of jatropha seeds and the chemical structure of jatropha oil. The image is obtained from ref. 119 with permission from Elsevier, copyright 2016.

native to Central America, but has been widely cultivated in other parts of the world for its oil. Jatropha seeds typically content 35–40% oil content, whereas the kernel can contain up to 35–40%. This makes jatropha a potentially valuable crop for producing many industrial products.¹¹¹ Jatropha oil contains about 79% unsaturated fatty acids and 21% saturated fatty acids, mostly oleic and linoleic acid. The presence of toxic and purgative compounds such as phorbol esters makes the oil non-edible and cannot be consumed without purification.¹¹² Thus, jatropha oil is commonly used for biodiesel,¹¹³ lubricants,¹¹⁴ cosmetics, and soap production,¹¹⁵ and other applications. Jatropha oil has also been used for polyurethane production such as for coatings,¹¹⁶ waterborne-PU,¹¹⁷ and polymer electrolytes.¹¹⁸ Fig. 21 shows the image of jatropha seeds and the oil chemical structure.

Haniffa et al. conducted an optimization study on the carbonation reaction of epoxidized jatropha oil (EJO) using various parameters such as temperature and pressure. Besides, they prepared carbonated alkyd resin (CAR) from jatropha oil. The carbonated EJO (CJO) was blended with CAR at varied weight ratios (1:1, 1:2, 1:3, 1:4) and reacted with 1,3-diaminopropane (DM) and IPDA to produce NIPU. Then, the samples were assessed based on their thermomechanical properties and chemical resistance. The analysis revealed optimal condition at 120 °C, 2 MPa of CO₂ pressure and 30 hours reaction time, achieving 99% and 97% carbonate conversion for CJO and CAR, respectively. The blend exhibited improved thermal, mechanical, and chemical resistance compared to single CJO and CAR when cured with DM and IPDA. It was observed that NIPU based on CJO/CAR(1:3)_IPDA showed better solvent and chemical resistance compared to CJO/CAR(1:3)_DM. Other than the amine content, the blending of CJO and CAR contributed to the enhancement of the interfacial surface of NIPU.58

Fig. 22 Proposed mechanisms of NIPU synthesis using CJO and APTES in the presence of LiCl catalyst. Adapted from ref. 120 with permission from Tech Science Press, copyright 2023.

Another work on jatropha oil-based NIPU was reported by Bhakri *et al.* for wood composite applications. The CJO was reacted with 3-aminopropyltriethoxisylane (APTES) which known for its excellent amine activity with cyclic carbonate. The presence of silane in APTES allows for the formation of siloxane (–Si–O–Si–) bonds between molecules, producing a more hydrophobic PU. The addition of a Lewis catalyst like LiCl influenced the reaction speed by activating the cyclic carbonate while leaving the amine unaffected. They proposed mechanism of NIPU synthesis as depicted in Fig. 22.¹²⁰

6.6 Rubber seed oil

Rubber seed oil (RSO) is a non-edible oil extracted from rubber seeds, a by-product of the natural rubber tree (*Hevea brasiliensis*). Crude rubber seed oil cannot be used as food resource without refining due to the presence of toxic cyanogenic glycoside compounds. The oil is rich in unsaturated fatty acids with 77–82% made up of oleic, linoleic, and linolenic acid and less content of saturated acids for around 17–20%. So is primarily utilized in non-food purposes such as biodiesels, las been reported for the production of various types of polyurethane such as waterborne-PU, foams, the mage of rubber seeds and the chemical structure for rubber seed oil.

The current research on rubber seed oil and NIPU is limited with only two studies available to date by Raden and co-workers. In their work, a preliminary study was conducted using epoxidized linoleic acid (ELA) extracted from RSO. After converting ELA to carbonated linoleic acid (CLA), they observed increment of viscosity from 705 cP to 1471 cP. The presence of cyclic carbonate groups led to an increase in viscosity due to the enhanced intermolecular interactions involving the polar oxygen atoms in the cyclic carbonate groups. The CLA was then reacted with different percentage of EDA from 5, 10, 15, and 20 w/w% for the preparation of NIPU. An increase in EDA loading resulted in higher viscosity due to high degree of crosslinking in NIPU and a decrease in tack free time, implying a shorter curing time. 132 In their recent study, cyclic carbonate from epoxidized rubber seed oil (ERSO) was synthesized using a low-pressure method. This study focused more into the synthesis part of ERSO at which they investigate the effects of different reaction temperatures and ratios of hydrogen peroxide and formic acid on the properties of ERSO. However, there was less emphasis on the carbonated RSO and NIPU in the discussion. 133

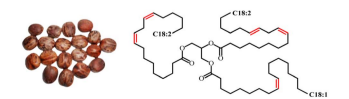


Fig. 23 Image of rubber seeds and chemical structure of rubber seed oil. The image is obtained from ref. 131 with permission from Elsevier, copyright 2017.

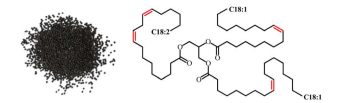


Fig. 24 Image of *Brassica napus* seeds and chemical structure of the oil. The figure is adapted from ref. 137 with permission from MDPI, copyright 2021.

6.7 Canola oil

Canola oil is an edible oil derived from a specific variety of rapeseed selectively bred to have low levels of toxic glucosinolates and erucic acid compounds that can be harmful when consumed in large quantities. 134 This rapeseed variety is known as "canola" in Canada and the United States, whereas in some other parts of the world, it is known as "rapeseed". This plant belongs to the Brassicaceae (Cruciferae) family that comprise thousands of species. Specifically, canola varieties could belong to Brassica rapa, Brassica napus or Brassica juncea. Canola oil is among important commodity oil globally and it is widely used as cooking oil. Canola oil contains high degree of monounsaturated fatty acids and very low in saturated fatty acids. 135 Canola oil consists of approximately 60% oleic acid (C18:1), 20% linoleic acid (C18:2), and 10% α -linolenic acid (C18:3). In comparison to other vegetable oils, canola oil has the lowest saturated fatty acid contain with 7% or less. Canola oil is considered very healthy oil owing to its fatty acid composition and the ideal 2:1 omega-6/omega-3 ratio.136 Other than being used for food purpose, canola is also being used for non-food products including biofuels, lubricants, and soaps. Fig. 24 displays the image of the seeds and the chemical structure of canola oil.

Research has been conducted to explore the possibility of utilizing canola oil as the raw material for the preparation of polyurethane for diverse applications such as coatings, adhesives, foams, foams, and elastomers. These studies focused on converting canola/rapeseed oil into polyols and subsequently reacting them with isocyanate groups to produce PU that meets the criteria for their proposed applications. In the case of NIPU, only one study was reported, conducted by Malik

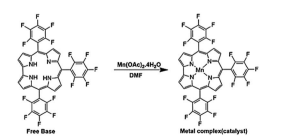


Fig. 25 Reaction scheme for the synthesis of 5,10,15-tris(penta-fluorophenyl)corrolato-manganese(III) complex catalyst. Adapted from ref. 57 with permission from Wiley, copyright 2017.

and Kaur. They used 5,10,15-tris(pentafluorophenyl)corrolatomanganese(III) complex as a novel catalyst for the preparation of carbonated canola oil (CCO) taking place under mild conditions. Fig. 25 shows the reaction scheme to synthesis the metal complex catalyst. The results indicate that the new catalyst can significantly reduce the reaction time to one-quarter of that required by conventional catalysts. The CCO was further reacted with four types of diamines; EDA, HMDA, IPDA, and paraphenyldiamine (PPDA) at a 1:1 molar ratio, cured at 80 °C for 8 hours to produce NIPU as the end product. Based on thermal analysis, all types of NIPU exhibited a relatively similar decomposition pattern and remained thermally stable up to 200 °C. On the other hand, the structure of diamine affected the tensile strength of NIPU. This study revealed NIPU with diamine having shorter carbon chain length i.e. EDA (8 MPa) demonstrated higher strength compare to HMDA (6 MPa), whereas the swelling behaviour of EDA NIPU was observed to be less than HMDA NIPU.57

6.8 Summary of vegetable oil-based NIPU

Table 4 provides a summary of reported studies on NIPUs derived from various types of vegetable oils. The literature indicates a consistent pattern of research in the production of NIPU from both edible and non-edible oils. Soybean oil stands out as the most reported study so far. This has to do with its high unsaturation value and wide availability. All the oils used contain high unsaturated value that can be exploited for chemical modification. On the side note, palm oil has not been explored for NIPU applications despite being the largest vegetable oil produced globally. This could be related to the high saturated fatty acid content of palm oil compared to other oils used for NIPU synthesis. It is known that saturated fatty acids have stable chemical structures which make them less reactive compared to unsaturated fatty acids. While palm oil may require additional processing steps for chemical modification, it can still be subjected to chemical transformations for NIPU production. Research has investigated various types of amine groups in the form of aliphatic, cycloaliphatic and aromatic amines to produce NIPU. Additionally, apart from using biobased cyclic carbonate, bio-based amines have also been utilized to produce fully bio-based NIPU. Based on the literature available so far, reactions were conducted with and without the presence of catalysts and solvents, depending on the nature of the experiments. Solvents typically used to reduce the viscosity of the carbonated resins, whereas catalysts are employed to accelerate the reaction rate, allowing more efficient polymerization. Catalysts are commonly used in the preparation of carbonated vegetable oils. Notably, most of studies excluded the use of catalysts in the aminolysis reactions.

7 Potential applications of NIPU

NIPU shares similarities with conventional polyurethane and offers a wide range of applications owing to its versatile properties and environmental advantages. There is a surge of interest to utilize NIPU in many areas, including those currently

Table 4 Summary of reported studies on vegetable oil-based non-isocyanate polyurethane

No.	Vegetable oil	Amine group	Synthesis methods	Observation	Applications	References
1	Soybean oil	EDA, HMDA and TA	(1) Carbonation (CSBO)Catalyst: TBAB(2) Aminolysis (NIPU)Catalyst: noneSolvent: none	NIPU made from triamine (TA) has the highest $T_{\rm g}$ and level of stress, smallest strain at break and lowest soluble fraction due to its tighter network structure	Not tested	86
2	Soybean oil	<i>m</i> -XDA, <i>p</i> -XDA and IPDA	(1) Carbonation (CSBO) Catalyst: TBAB (2) Aminolysis (NIPU) Catalyst: none Solvent: none	The three NIPUs produced were elastomeric PUs and the T_g was dominantly affected by the amine-to-cyclic carbonate ratio, while the impact of the amine's structure was relatively minor	Not tested	55
3	Soybean oil	APTES	(1) Carbonation (CSBO) Catalyst: TBAB (2) Aminolysis (NIPU) Catalyst: LiCl Solvent: THF (3) Addition of lignin	Tensile strength of NIPU increased with the addition of lignin by contributing to the rigidity of polymer	Not tested	53
4	Soybean oil	BDA, PDA and oligoamide	(1) Synthesis of oligoamide (2) Carbonation (CSBO) Catalyst: TBAB (3) Aminolysis (NIPU) Catalyst: none Solvent: none	NIPU produced with short diamines exhibit low mechanical properties, whereas NIPU derived from oligoamide shows better mechanical properties	Not tested	87
5	Soybean oil	PA650, IPDA, and cystamine	(1) Carbonation (CSBO) Catalyst: TBAB (2) Aminolysis (NIPU) Catalyst: none Solvent: none	The self-healing properties of produced NIPU were contributed by the intrinsic hydrogen bonds existed in the matrix and enhanced by the addition of cysteine-derived disulphide-containing cystamine	Not tested	56
6	Soybean oil	Chitosan	(1) Epoxidation (ESBO) (2) Carbonation (CSBO) Catalyst: TBAB (3) Aminolysis (NIPU) Catalyst: LiCl Solvent: THF	NIPU produced using chitosan as the amine source has good chemical and thermal stability	Not tested	88
7	Soybean oil	Polyamidoamine	(1) Epoxidation (ESBO) (2) Carbonation (CSBO) Catalyst: TBAB (3) Synthesis of CHPE (4) Aminolysis (NIPU) Catalyst: none Solvent: THF	NIPU exhibited good mechanical properties and thermal stability due to the increased in rigidity of the polymeric chain and dense polymer network with CSBO/ CHPE blend	Not tested	60
8	Soybean oil	DDS and DDM	Aminolysis (NIPU) Catalyst: none Solvent: none	The self-healing properties of NIPU under UV irradiation enhanced with the presence of disulphide bonds (DDS)	Not tested	89
9	Soybean oil	DDM	(1) Epoxidation (ESBO)(2) Carbonation (CSBO)(3) Aminolysis (NIPU)	NIPU has shape memory effect and reprocessing ability	Not tested	142
10	Soybean oil	MDA	(1) Carbonation (CSBO) catalyst: TBAI, 1-ascorbic acid (2) Aminolysis (NIPU) Catalyst: none Solvent: none	NIPU has shape memory properties, self-healing ability, chemical recyclability and physical reprocessing due to the dynamic covalent bond exchange	Not tested	90

Table 4 (Contd.)

No.	Vegetable oil	Amine group	Synthesis methods	Observation	Applications	References
11	Soybean oil	TEPA and HMDA	(1) Carbonation (CSBO) Catalyst: TBAB, CaCl ₂ (2) Aminolysis (NIPU) Catalyst: None Solvent: DMF (3) Addition of azetidinium	The NIPU wound dressing membranes possess good mechanical properties, showed no toxicity against fibroblasts and high antibacterial activity against <i>E. coli</i> and <i>S. aureus</i>	Antibacterial wound dressing membranes	54
12	Soybean oil	APTES and DETA	 (1) Carbonation (CSBO) Catalyst: TBAB (2) Aminolysis (NIPU) Catalyst: none Solvent: DMF, THF (3) Addition of lignin 	The addition of lignin causes a reduction in the adhesive properties in terms of peel force performance of NIPU	Adhesive	143
13	Soybean oil	Priamine 1075	(1) Carbonation (CSBO) (2) Aminolysis (NIPU)	The combination of reactive carbonates with epoxides reduced the occurrence of unwanted side reactions such as ester aminolysis and intramolecular cyclization	Not tested	144
14	Soybean oil	ТЕРА	(1) Synthesis of tetraniline (TANI) (2) Carbonation (CSBO) Catalyst: TBAB (3) Aminolysis (NIPU) Catalyst: none Solvent: DMF	Observation: Addition of TANI improve mechanical and anti- corrosion properties of NIPU	Anti- corrosion coating	63
15	Linseed oil	Phenalkalmine	(1) Carbonation (CLSO) Catalyst: TBAB (2) Aminolysis (NIPU)	Real-time FTIR spectra were utilized to measure the <i>in situ</i> decay of the carbonate groups and the appearance of C=O group of the urethane linkages	Not tested	145
16	Linseed oil	EDA, BDA and IPDA	(1) Carbonation (CLSO) Catalyst: TBAB (2) Aminolysis (NIPU) Catalyst: none Solvent: none	NIPU produced has a high dimensional stability and stiffness due to high carbonate functionality of CLSO	Not tested	59
17	Linseed oil	DETA	(1) Epoxidation (ELSO) (2) Carbonation (CLSO) Catalyst: TBAB (3) Aminolysis (NIPU) Catalyst: none Solvent: none	Adhesion and hardness properties of NIPU improved with the increase in carbonation number of the CLSO	Anti- corrosion coating	96
18	Linseed oil	BDA, PDA, ODA and HMDA	(1) Epoxidation (ELSO) (2) Carbonation (CLSO) Catalyst: TBAB (3) Aminolysis (NIPU) Catalyst: none Solvent: none	CLSO showed a high reactivity and this can shorten curing time for NIPU synthesis, however viscosity increases with the carbonated group content can be problematic to achieve a homogenous mixture	Foam	26
19	Linseed oil	m-XDA, HMDA	(1) Carbonation (CLSO)(2) Aminolysis (NIPU)(3) Computational approach	Density functional theory calculation revealed the reactivity of epoxy groups at different sites of the ELSO, whereas the reaction conditions for CLSO and NIPU were optimized to achieve high atom efficiency	Not tested	146

Table 4 (Contd.)

Table 4 (Contd.)

No.	Vegetable oil	Amine group	Synthesis methods	Observation	Applications	References
29	Rubber seed oil	EDA	(1) Epoxidation (ELA) (2) Carbonation (CLA) Catalyst: TBAB (3) Aminolysis (NIPU) Catalyst: none Solvent: none	Increase in EDA loading resulted to increase in viscosity due to high degree of crosslink in NIPU and decrease in tack free time	Not tested	132
30	Rubber seed oil	EDA	(1) Epoxidation (ERSO)(2) Carbonation (CRSO)Catalyst: TBAB(3) Aminolysis (NIPU)Catalyst: noneSolvent: none	The optimum condition for epoxidation of RSO was 50 °C, and increasing molar ratio of hydrogen peroxide increased the oxirane number, whereas when molar ratio for formic acid increased, the opposite effect was observed	Not tested	133
31	Canola oil	EDA, HMDA, IPDA and PPDA	 (1) Synthesis of manganese-based catalyst (2) Epoxidation (ECO) (3) Carbonation (CCO) Catalyst: manganese-based (4) Aminolysis (NIPU) Catalyst: none Solvent: none 	The new catalyst can significantly decrease the carbonation reaction time to one-quarter of the time required by conventional catalysts	Not tested	57

served by traditional polyurethane as well as exploring new applications. NIPU finds application across various industries and sectors such as coatings, adhesives, packaging materials, insulation materials, biomedical applications, electronic components, and more. This section briefly reviews several applications where NIPU has been investigated, but not limited to those produced using vegetable oil-based feedstock.

7.1 Coatings

NIPU has found applications as coating materials due to its excellent adhesion, durability, and chemical resistance nature. It can be used in protective coatings for various surfaces such as metals, plastics, and woods. Various studies have already reported on the use of NIPU for coating applications. In a study by Zhang et al., NIPU coating was investigated for its high hardness, transparency, strong adhesion, and superior flexibility properties. The NIPU was prepared using epoxy-oligosiloxane nanoclusters and an amine curing agent. The mixture was casted onto different substrates i.e. glass, steel, ceramic, epoxy, aluminium alloy, tinplate, titanium, and polyethylene terephthalate. This study suggested that the strong adhesion between NIPU coating and the substrate results from the high density polar groups in the NIPU. Besides, the addition of amine-terminated low-surface-tension polydimethylsiloxane endowed the coating with exceptional anti-icing, self-cleaning, and anti-smudge abilities. This work able to achieve NIPU coating that exhibits a combination of the hardness properties of ceramic and the flexibility of polymers.147

Another research reports on NIPU coating using pentaery-thritol glycidyl ether-based cyclic carbonate (PGC)/single-end cyclic carbonate poly(dimethylsiloxane) (PDMS-C) and

ethylene imine polymer (PEI). In this case, the PDMS-C functions as a liquid-repelling agent to provide NIPU coatings with low surface energy, whereas PEI was used as a curing agent to achieve a high crosslinking density within the coating matrix. It was discovered that the produced NIPU coating displayed high repellency towards various liquids such as water, hexadecane, corn oil, pump oil, milk, fingerprint fluid, and cola. The transparent coating demonstrated excellent self-cleaning properties, allowing dust to slide off with water, leaving no residue of dust or water on the surface. The coatings also showed excellent resistance against ink deposition, maintaining anti-ink performance after 5000 abrasion cycles, making them suitable for various applications where ink smudging or deposition needs to be minimized.¹⁴⁸

Boisaubert et al. reported a study on non-isocyanate polyurethane acrylate coatings (NIPUAs) by photocrosslinking acrylate terminated NIPU oligomers (A-Ol) with reactive diluents. The A-Ol was obtained through a transurethane polycondensation pathway, followed by an acrylation reaction of the resulting hydroxyl chain-ends. This work investigates the effect of reactive diluents content and chemical structures on the properties of the coatings. Three types of diluents were used which were hexamethylene diol diacrylate (HDDA), urethane diol dimethacrylate (UDMA), and N-hexamethylene dihydroxyethyl carbamate diacrylate (BHECA). The different types of diluent significantly impact the thermal and mechanical properties of the NIPUA coating. The properties observed align with those of flexible PU currently being develop in the market. The UV-cured coatings are deemed feasible owing to their fast curing and low-energy consumption.149

Moreover, a series of bio-based NIPU coatings were synthesized by performing ring-opening polymerization of rosin-based

cyclic carbonate with amines. Rosin is natural resources derived from pine and conifer trees, available abundantly in nature. The NIPU was modified with epoxy and cyclic carbonatefunctionalized polyhedral oligomeric silsesquioxanes (POSS). The results demonstrated that the addition of POSS into the NIPU networks led to significant improvements in water tolerance, pencil hardness, and thermal stability of the NIPU/POSS coatings. 150 Another bio-based NIPU coating study was reported by Choong et al. with recyclable and intrinsic healing properties via three different types of healing mechanisms; thermo-healing, moisture-healing, and room temperature selfhealing. These desirable properties were achieved by the addition of bismaleimide cross-linker. Interestingly, the coating has remarkable moisture-healing property attributed to the presence of hydroxyl functionalities in the NIPU structure. The unique combination of bio-based resources, recyclable properties, and the presence of triple healing sites in their structure makes these materials highly promising for coating applications.151 Nevertheless, the exploration of vegetable oil-based NIPU has also been conducted and is currently being investigated for corrosion protection coatings, flame retardant composites coating, and others, as described in Section 6.

7.2 Adhesives

PU-based adhesives are commercially available in the market and are extensively used across various sectors such as packaging, automotive, construction, woodworking, and more. The demands of these adhesives continue to grow, driven by their excellent adhesion properties, flexibility and performance at low temperatures. This serves as a motivation for the continuous NIPU-based adhesives research to seek a green alternative to the conventional PU adhesives. In the case of vegetable oilbased NIPU, only one preliminary study so far has been reported for soybean oil-based adhesive143 as mentioned in Section 6.8. Despite the lack numbers of studies using vegetable oil, many works already have been conducted on other biobased resources. For instance, one study prepared low curing temperature tannin-based NIPU wood adhesives and evaluate its properties. In this work, commercial glycerol diglycidyl ether (GDE) was assesses as a promoter for the tannin-based NIPU adhesives. GDE facilitated the curing process at a reduced temperature and promoted the degree of crosslinking between the epoxy groups of GDE and amino groups present in the derived-products of NIPU. The inclusion of GDE in appropriate proportions was found to decrease the curing temperature, enhance bonding performance, and reduce the emission of harmful substances during the preparation of tannin-based NIPU adhesives. 152

Other adhesive from sucrose-based NIPU was synthesized by Xi *et al.* by the reaction of sucrose, dimethyl carbonate and hexamethylene diamine for bonding particleboard. A silane coupling agent was used as a crosslinking promoter to decrease the curing temperature. The results showed that the particleboards bonded with the sucrose-based NIPU adhesive at 230 °C exhibited excellent properties. However, the properties decreased as the press time was reduced. The coupling agent

significantly reduced the curing temperature of the adhesive, enabling good bonding even at lower press temperatures. This study confirmed the coupling of NIPU with silane produced improved results and made a suitable adhesive for particle-board, medium-density fibreboard, and other particulate wood panels.¹⁵³

A study by Gomez-Lopez *et al.* investigated the synergetic effect of dopamine and alkoxysilanes on the rheological and adhesive properties of NIPU. In this case, dopamine acted as an adhesive promoter, and aminopropyl trimethoxysilane functioned as a fast-curing promoter. They found out that adjusting the soft/hard ratios is crucial to achieving high lap-shear strength adhesion value. Furthermore, the addition of dopamine and silane compounds, maintained these adhesive properties even at high temperatures for optimal compositions. Adhesion tests on various substrates including polyamide, high-density polyethylene, poly(methyl methacrylate), oak wood, and aluminium showed the best performances on polar substrates, confirming the strong interactions between dopamine and hydroxyl groups of NIPU.¹⁵⁴

Another study on wood bio-based NIPU adhesives was performed by Chen *et al.* The preparation involves the reaction of soy protein isolate (SPI), dimethyl carbonate (DMC), and hexamethylene diamine. The reaction resulted in the formation of both linear and branched oligomers, which could further crosslink to form a hardened network. To evaluate the adhesive's performance, plywood panels were prepared and bonded using the SPI-NIPU adhesive. The dry strength of the adhesive met the requirements of relevant standards, indicating its suitability for interior grade plywood panels.¹⁵⁵

7.3 Foams

NIPU foams have gained interest in research and development due to their immense potential in diverse areas such as in packaging, furniture and bedding, automotive industries, etc. In the case of vegetable oil-based NIPU foams, only one study on linseed oil-based NIPU was reported.26 In contrast to the limited research on vegetable oil, other sources of bio-materials have been reported for NIPU foams production. One study constructed recyclable self-blown NIPU foams through an aminolysis approach. The method capitalizes on the divergent chemistries of amines with cyclic carbonates and thiolactone. By combining these reactants, a polymer network is created, and in situ generation of the blowing agent (CO2) occurs through the reaction of a thiol with a cyclic carbonate. This process results in the formation of multiple linkages including hydroxyurethane, thioethers, and amides within the polymer network. The foams produced exhibit open-cell morphology and can range from flexible to rigid depending on the formulation and reaction conditions. Fig. 26 shows the photographic images of produced NIPU foams at varied NAHcT content. This study reported that the foams can be easily repurposed into films or structural composites through thermal treatment, demonstrating their recyclability. Notably, both the formation and recycling of the foams can be achieved without the use of a catalyst.156

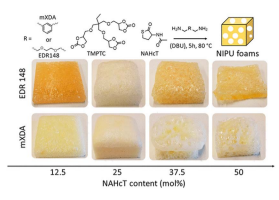


Fig. 26 The effect of NAHcT content on NIPU foaming. Reuse from ref. 156 with permission from ACS, copyright 2022.

Chen *et al.* described the preparation of condensed tanninglucose-based NIPU foams with improved fire retardancy. Tannin was added as a flame-retardant agent to improve the fire resistance property of glucose-NIPU. The weight ratios of tannin (T) to glucose ratios were varied from 0 g/40 g (NIPU_Tannin0), 10 g/30 g (NIPU_Tannin10), and 20 g/20 g (NIPU_Tannin20). The mixture was reacted with dimethyl carbonate, followed by addition of hexamethylenediamine in water. The visual appearance of the foams is presented in Fig. 27. Ignition combustion experiments indicated that tannin modified NIPU foams exhibit a longer burning time and higher residual weight compared to blank NIPU foam. Additionally, the addition of condensed tannin increased the limiting oxygen index from 17.5% to 25.5%. 157

Another study on flame-resistant rigid NIPU bio-foam based on chitosan and tannic acid was reported by Smith *et al.* The foams were prepared with the absence of solvents and catalysts. The foam's structure was achieved by incorporating glutaral-dehyde and four different carboxylic acids: malic acid, maleic acid, citric acid, and aconitic acid. The resulting foam samples were compared based on their morphology, thermal degradation, and flame resistance. This study observed that the properties of the foam varied depending on the carboxylic acid used. However, in all cases, the peak thermal degradation and peak heat release for NIPU foams were delayed by more than 100 °C compared to commercial rigid PU foam. Further, when subjected to a butane torch test, the NIPU foams exhibited significant improvements as shown in Fig. 28. The foams showed an 80% higher remaining mass and a 75% reduction in after-burn

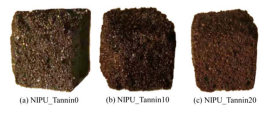


Fig. 27 Images of tannin–glucose-based NIPU foams. Reuse from ref. 157 with permission from Elsevier, copyright 2020.

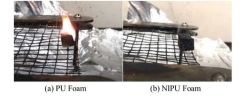


Fig. 28 Graphical images of commercial (a) PU foam and (b) NIPU foam 2 s after the removal of a butane torch flame. Reuse from ref. 158 with permission from MDPI, copyright 2022.

time compared to conventional PU. This finding highlights the superior fire-resistant properties of the bio-based NIPU foam which eliminates the hazards associated with traditional PU. ¹⁵⁸

7.4 Biomedical applications

The use of NIPU in biomedical applications has been investigated previously owing to its non-toxic and biocompatible properties. In the case of vegetable oil-based NIPU, Gholami and Yeganeh have prepared soybean-oil based NIPU containing azetidinium groups as antibacterial wound dressing membranes⁵⁴ as discussed in Section 6.1. A recent study was conducted on the fabrication of photosensitive NIPU acrylate (NIPUA) resin for 3D printing of customized biocompatible orthopedic surgical guides by Wang et al. Compared to commercial photosensitive resins, NIPUA exhibited higher thermal stability, hemocompatibility, superior biocompatibility to ME3T3-E1 bone cells and C1C12 muscle cells, and a nonimmunogenic effect towards macrophages. The NIPUA also did not caused inflammatory response during in vivo implantation unlike the severe response by commercial resins. Further, the transcriptome analysis revealed that NIPUA-treated MC3T3-E1 cells showed downregulation of genes associated with cell death and disruption of the cell cycle compared to those treated with commercial resin. On the other hand, there was an upregulation of genes related to autophagy and anti-tumor activity. To summarize, the NIPUA resin exhibited excellent mechanical and thermal properties as well as good biocompatibility towards bone cells, muscle cells, and macrophages suggests the suitability of this material for use in orthopedic surgical guides.¹⁵⁹

One work by Visser et al. reported on the electrospinning of NIPU by the transurethanization reaction of polycarbonate diols (PCLD) and 1,6-hexanedicarbamate. Different molecular weights of PCDL were utilized to optimize the electrospinnability of NIPU. Through scanning electron microscopy, they observed the electrospun NIPU formed fibrous scaffolds with submicron-sized fiber diameters. To assess the potential of these electrospun NIPU mats as biomimetic load-bearing pericardial substitutes in cardiac tissue engineering, their cytotoxicity was evaluated using primary human fibroblasts and a human epithelial cell line. Interestingly, the bare NIPU mats exhibited excellent cell adhesion without the need for further biofunctionalization, surpassing the performance of collagenfunctionalized NIPU mats. These results indicate that the NIPU mats possess significant potential for use in biomimetic scaffolds. 160 In another study, Aduba et al. prepared fibrous

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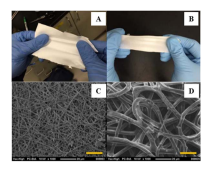


Fig. 29 Photographic images of electrospun and their scanning electron microscopy microscale images (A and C) NIPU, (B and D) TPU control. Reproduced from ref. 161 with permission from Wiley, copyright 2018.

mats NIPU using electrospinning method. Fig. 29 depicted the photographic images of (a) NIPU fibrous mats and (b) PU control and their scanning electron microscope images (c and d). The NIPU fibrous mats were found to be non-toxic and biocompatible as demonstrated by cell viability and attachment studies with human fibroblast. These finding suggest that the mats can be safely used in various biomedical applications including tissue engineering, drug delivery, and wound dressing technology.¹⁶¹

7.5 Food preservation

One interesting study was reported by Yang *et al.* on the use of bio-based NIPU in food preservation application. The NIPU was prepared from sweet potato residual (SPR) from the starch industry. The crude SPR was first converted into a liquid state, forming chain-extending homogenous molecules with active hydroxyl groups. Next the liquefied SPR underwent epoxidation and carbonation reactions to produce cyclic carbonates. The cyclic carbonate precursor was then reacted with different types of amines (EDA, HMDA, IPDA, DETA and TAEA). Among the five testing groups, TAEA_NIPU film exhibited the highest tensile strength (8.5 MPa) due to the strong curing and crosslinking properties of TAEA. The authors suggested that the produced NIPU derived from SPR would be safe for use as a food cling material. To evaluate its potential, the researchers atomized the

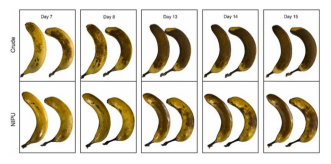


Fig. 30 Experimental study on preservation of bananas using SPR-based NIPU spray coating. Reproduced from ref. 162 with permission from Elsevier, copyright 2022.

initial liquid of SPR NIPU and spray coated the film onto the surface of banana peels to preserve freshness. The viscosity of the initial NIPU liquid was reported to be 6 Pa s⁻¹, indicating that it can be directly used for spray coating without further modifications. During the experiment, both treated and control groups were stored under similar conditions of 10–25 °C temperature, 30–65% humidity and in a ventilated dark place. After 7 days of storage, small black patches appeared on the surface of the banana peel for both treated and control groups. By the second week of storage, the control group began to spoil, whereas the banana peels in the coated with SPR-NIPU exhibited only slight darkening compared to 1 week after storage as displayed in Fig. 30. This indicates that SPR-NIPU can serve as a protective coating, effectively preserving the freshness of food. 162

8 Conclusions and future outlook

Significant progress has been made in NIPU development as outlined in this review. NIPU ability to avoid the use of isocyanates while maintaining desirable properties addresses both safety and environmental concerns associated with traditional PU production. The research and development in NIPU so far have demonstrated its potential across various applications. This is due to the wide range of properties NIPU such that it offers ample opportunities for alteration to meet diverse application requirements. Moreover, NIPU compatibility with bio-based feedstock further enhances its eco-friendliness and aligns with the growing emphasis on sustainable materials. In the case of vegetable oil-based NIPU, currently, there are still limited but continuous studies reported in this area. Both edible and non-edible vegetable oil has been utilized to prepare bio-based NIPU at which soybean oil being the most used source of oil so far. With ongoing research and development, it is likely to see an increasing number of reports on vegetable oilbased NIPU in the future. In addition, there are valid concerns related to the use of edible oil for non-food purpose applications. This has opened up the route to prepare NIPU using nonedible vegetable oil. Some studies already reported on the fully bio-based NIPU derived from bioresources. Overall, even though NIPU technology is promising, it is not yet attained the stage of industrial implementation. Presently, challenges related to synthesis issues and technical performance of NIPU hinders its commercialization. Specifically, issues like slow polymerization, extended reaction time, and technical performance pose obstacles in NIPU production. Addressing these challenges must involve a thorough consideration of various factors like the selection of monomer ratio, solvent, catalyst, and reaction parameters which collectively contribute to enhancing NIPU properties. Additionally, broader challenges like scalability, cost-effectiveness, and end-of-life considerations must be addressed for NIPU to be accepted as a suitable alternative to the traditional PU. To achieve this, the development of scalable synthesis route and the implementation of comprehensive cycle assessments are essential steps that will contribute to the competitiveness of NIPU. Furthermore, the industry is most likely to shift into NIPU only when they become

cost-competitive with the current PU technology. This is perhaps the direction of future efforts to materialize NIPU as commercial products.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 B. Bizet, B. Bizet, É. Grau, H. Cramail and J. M. Asua, Polym. Chem., 2020, 11, 3786-3799.
- 2 J. Catalá, I. Guerra, J. M. García-Vargas, M. J. Ramos, M. T. García and J. F. Rodríguez, Polymers, 2023, 15, 1589.
- 3 A. Das and P. Mahanwar, Adv. Ind. Eng. Polym. Res., 2020, 3,
- 4 M. Ghasemlou, F. Daver, E. P. Ivanova and B. Adhikari, Eur. Polym. J., 2019, 118, 668-684.
- 5 X. Ma, J. Chen, J. Zhu and N. Yan, Macromol. Rapid Commun., 2021, 42, 2000492.
- 6 M. Tryznowski, J. Izdebska-Podsiadły and Z. Żołek-Tryznowska, Prog. Org. Coat., 2017, 109, 55-60.
- 7 V. Ca, G. Lligadas, J. C. Ronda and M. Galia, Mater. Today, 2013, 16, 337-343.
- 8 A. Patti, F. Costa, M. Perrotti, D. Barbarino and D. Acierno, Materials, 2021, 14, 1951.
- 9 J. Niesiobędzka and J. Datta, Green Chem., 2023, 25, 2482-2504.
- 10 F. M. de Souza, P. K. Kahol and R. K. Gupta, in Polyurethane Chemistry: Renewable Polyols and Isocyanates, ACS Publications, 2021, pp. 1-24.
- 11 H. Choe and J. H. Kim, J. Ind. Eng. Chem., 2019, 69, 153-160.
- 12 Y. Li, X. Luo and S. Hu, Poyols and Polyurethanes from Vegetable Oils and Their Derivatives, in Bio-based Polyols and Polyurethane, Springer International Publishing, 2015, pp. 15-43, DOI: 10.1007/978-3-319-21539-6.
- 13 C. Pronoitis, M. Hakkarainen and K. Odelius, ACS Sustain. Chem. Eng., 2022, 10, 2522-2531.
- 14 M. A. Aristri, M. A. R. Lubis, S. M. Yadav, P. Antov, A. N. Papadopoulos, A. Pizzi, W. Fatriasari, M. Ismayati and A. H. Iswanto, Appl. Sci., 2021, 11, 4242.
- 15 G. dong Feng, Y. Ma, M. Zhang, P. you Jia, L. hong Hu, C. guo Liu and Y. hong Zhou, Prog. Org. Coat., 2019, 133, 267-275.
- 16 S. Laurichesse, C. Huillet and L. Avérous, Green Chem., 2014, 16, 3958-3970.
- 17 T. N. M. Tuan Ismail, N. A. Ibrahim, M. A. Mohd Noor, S. S. Hoong, K. D. P. Palam, S. K. Yeong, Z. Idris, C. M. Schiffman, I. Sendijarevic, E. Abd Malek, N. Zainuddin and V. Sedijarevic, J. Am. Oil Chem. Soc., 2018, 95, 509-523.

- 18 S. Cui, Z. Liu and Y. Li, Ind. Crops Prod., 2017, 108, 798-805.
- 19 R. Galhano, N. F. Acero, S. Matos, R. Carvalho, M. Vale, A. C. Marques, J. C. Bordado and M. M. Mateus, J. Polym. Environ., 2017, 1-10.
- 20 Q. Wang and N. Tuohedi, Sustainability, 2020, 12, 4214.
- 21 Y. Cao, Z. Liu, B. Zheng, R. Ou, Q. Fan, L. Li, C. Guo, T. Liu and Q. Wang, Composites, Part B, 2020, 200, 108295.
- 22 H. Tian, J. Wu and A. Xiang, J. Vinyl Addit. Technol., 2017, 24, 105-111.
- 23 R. Morales-Cerrada, R. Tavernier and S. Caillol, Polymers, 2021, 13, 1255.
- 24 K. Błażek and J. Datta, Crit. Rev. Environ. Sci. Technol., 2019, 49, 1-39.
- 25 P. M. Paraskar, M. S. Prabhudesai, V. M. Hatkar and R. D. Kulkarni, Prog. Org. Coat., 2021, 156, 106267.
- 26 T. Dong, E. Dheressa, M. Wiatrowski, A. P. Pereira, A. Zeller, L. M. L. Laurens and P. T. Pienkos, ACS Sustainable Chem. Eng., 2021, 9, 12858-12869.
- 27 T. Liu, X. Yang, S. Zhang, Q. Wang, N. Jiang and G. Wang, RSC Adv., 2022, 30167-30173.
- 28 E. Dyer and H. Scott, J. Am. Chem. Soc., 1957, 79, 672-675.
- 29 A. Z. Yu, R. A. Setien, J. M. Sahouani, J. Docken and D. C. Webster, J. Coat. Technol. Res., 2019, 16, 41-57.
- 30 Y. Suryawanshi, P. Sanap and V. Wani, Polym. Bull., 2019, 76, 3233-3246.
- 31 P. Stachak, I. Łukaszewska, E. Hebda and K. Pielichowski, Materials, 2021, 14, 3497.
- 32 A. Cornille, R. Auvergne, O. Figovsky, B. Boutevin and S. Caillol, Eur. Polym. J., 2017, 87, 535-552.
- 33 L. Filippi and M. A. R. Meier, Macromol. Rapid Commun., 2021, 42, 2000440.
- 34 J. Datta and M. Włoch, Polym. Bull., 2016, 73, 1459-1496.
- 35 N. Ismail, M. Essalhi, M. Rahmati, Z. Cui, M. Khayet and N. Tavajohi, Green Chem., 2021, 23, 2130-2147.
- 36 C. C. Su, M. He, R. Amine, Z. Chen, R. Sahore, N. Dietz Rago and K. Amine, Energy Storage Mater., 2019, 17, 284-292.
- 37 C. Claver, M. B. Yeamin, M. Reguero and A. M. Masdeu-Bultó, Green Chem., 2020, 22, 7665-7706.
- 38 E. J. C. Lopes, A. P. C. Ribeiro and L. M. D. R. S. Martins, Catalysts, 2020, 10, 479.
- 39 J. W. Hill and W. H. Carothers, J. Am. Chem. Soc., 1933, 55, 5031-5039.
- 40 E. W. Spanagel and W. H. Carothers, J. Am. Chem. Soc., 1935, 57, 929-934.
- 41 F. Camara, S. Benyahya, V. Besse, G. Boutevin, R. Auvergne, B. Boutevin and S. Caillol, Eur. Polym. J., 2014, 55, 17-26.
- 42 C. Carré, Y. Ecochard, S. Caillol and L. Avérous, ChemSusChem, 2019, 12, 3410-3430.
- 43 P. P. Pescarmona, Curr. Opin. Green Sustainable Chem., 2021, 29, 100457.
- 44 M. Rayung, M. M. Aung, S. C. Azhar, L. C. Abdullah, M. S. Su'ait, A. Ahmad and S. N. A. Md Jamil, Materials, 2020, 13, 838.
- 45 R. Pathak, M. Kathalewar, K. Wazarkar and A. Sabnis, Prog. Org. Coat., 2015, 89, 160-169.
- 46 V. Froidevaux, C. Negrell, S. Caillol, J. P. Pascault and B. Boutevin, Chem. Rev., 2016, 116, 14181-14224.

47 C. Zhang, K.-c. Huang, H. Wang and Q. Zhou, *Prog. Org. Coat.*, 2020, **148**, 105855.

- 48 B. Zhang, X. Yang, X. Lin, H. Shang, Q. Liu, H. Wang, S. Liu, X. Xu and F. Dong, ACS Sustainable Chem. Eng., 2023, 11, 6100–6113.
- 49 G. Rokicki, P. G. Parzuchowski and M. Mazurek, *Polym. Adv. Technol.*, 2015, **26**, 707–761.
- 50 E. Wondu, Z. Lule and J. Kim, Polymers, 2019, 11, 1103.
- 51 F. H. Yeoh, C. S. Lee, Y. B. Kang, S. F. Wong, S. F. Cheng and W. S. Ng, *Polymers*, 2020, **12**, 1842.
- 52 S. Dworakowska, D. Bogdal and A. Prociak, *Polymers*, 2012, 4, 1462–1477.
- 53 A. Lee and Y. Deng, Eur. Polym. J., 2015, 63, 67-73.
- 54 H. Gholami and H. Yeganeh, Eur. Polym. J., 2021, 142, 110142.
- 55 I. Javni, D. P. Hong and Z. S. Petrovic, *J. Appl. Polym. Sci.*, 2013, **128**, 566–571.
- 56 J. Dong, B. Liu, H. Ding, J. Shi, N. Liu, B. Dai and I. Kim, Polym. Chem., 2020, 11, 7524–7532.
- 57 M. Malik and R. Kaur, *Polym. Adv. Technol.*, 2017, 29, 1078– 1085.
- 58 M. A. C. M. Haniffa, Y. C. Ching, C. H. Chuah, Y. C. Kuan, D. S. Liu and N. S. Liou, *Polymers*, 2017, 9, 162.
- 59 B. Moritz and M. Rolf, Green Chem., 2012, 14, 483-489.
- 60 S. Doley, A. Bora, P. Saikia, S. Ahmed and S. K. Dolui, J. Polym. Res., 2021, 30–33, DOI: 10.1007/s10965-021-02485-2.
- 61 S. Doley and S. K. Dolui, Eur. Polym. J., 2018, 102, 161-168.
- 62 W. Zhang, T. Wang, Z. Zheng, R. L. Quirino, F. Xie, Y. Li and C. Zhang, *Chem. Eng. J.*, 2023, 452, 138965.
- 63 N. Dhore, E. Prasad, R. Narayan, C. R. K. Rao and A. Palanisamy, *Sustainable Chem.*, 2023, 4, 95–109.
- 64 M. Goyal, S. N. Agarwal and N. Bhatnagar, *J. Appl. Polym. Sci.*, 2022, **139**, e52816.
- 65 Z. Li, R. Yu and B. Guo, ACS Appl. Bio Mater., 2021, 4, 5926–5943.
- 66 D. Wang, S. Chen, J. Zhao and Z. Zhang, *Mater. Today Commun.*, 2020, 23, 101138.
- 67 F. Zhang, P. Ju, M. Pan, D. Zhang, Y. Huang, G. Li and X. Li, *Corros. Sci.*, 2018, **144**, 74–88.
- 68 S. K. Raut, P. K. Behera, T. S. Pal, P. Mondal, K. Naskar and N. K. Singha, *Chem. Commun.*, 2021, 57, 1149–1152.
- 69 S. Hu, X. Chen and J. M. Torkelson, ACS Sustain. Chem. Eng., 2019, 7, 10025–10034.
- 70 D. w. Wang, S. Chen, J. b. Zhao, Z. y. Zhang and J. y. Zhang, Polym. Eng. Sci., 2021, 61, 497–505.
- 71 C. C. Hornat and M. W. Urban, *Prog. Polym. Sci.*, 2020, **102**, 101208.
- 72 X. Yin, H. Liu, R. Lin, X. Liu, Z. Huang, J. Du, Y. Gu, X. Lin, W. Lin and G. Yi, J. Appl. Polym. Sci., 2023, 140, e53705.
- 73 V. Schimpf, B. Heck, G. n. Reiter and R. Mülhaupt, *Macromolecules*, 2017, **50**, 3598–3606.
- 74 T. Zhang, B. Xue, Q. Yan, Y. Yuan, J. Tan, Y. Guan, J. Wen, X. Li and W. Zhao, *Ind. Crops Prod.*, 2023, **199**, 116706.
- 75 J. Sternberg and S. Pilla, Green Chem., 2020, 22, 6922-6935.
- 76 L. Li, B. Zhao, H. Wang, Y. Gao, J. Hu and S. Zheng, *Compos. Sci. Technol.*, 2021, 215, 109009.

- 77 X. Chen, L. Li, K. Jin and J. M. Torkelson, *Polym. Chem.*, 2017, **8**, 6349–6355.
- 78 N. S. Purwanto, Y. Chen, T. Wang and J. M. Torkelson, *Polymer*, 2023, 272, 125858.
- 79 Y. Chen, B. Chen and J. M. Torkelson, *Macromolecules*, 2023, 56, 3687–3702.
- 80 Y. Zhou, W. Zhao, Y. Lai, B. Zhang and D. Zhang, Front. Plant Sci., 2020, 11, 1315.
- 81 M. Rayung, M. Min, A. Ahmad, M. Sukor, L. Chuah, S. Nurul and A. Jamil, *Mater. Chem. Phys.*, 2019, 222, 110–117.
- 82 X. P. Song, M. C. Hansen, P. Potapov, B. Adusei, J. Pickering, M. Adami, A. Lima, V. Zalles, S. V. Stehman, C. M. Di Bella, M. C. Conde, E. J. Copati, L. B. Fernandes, A. Hernandez-Serna, S. M. Jantz, A. H. Pickens, S. Turubanova and A. Tyukavina, *Nat Sustainability*, 2021, 4, 784–792.
- 83 Q. Wu, Y. Hu, J. Tang, J. Zhang, C. Wang, Q. Shang, G. Feng, C. Liu, Y. Zhou and W. Lei, *ACS Sustainable Chem. Eng.*, 2018, **6**, 8340–8349.
- 84 M. Qi, Y. J. Xu, W. H. Rao, X. Luo, L. Chen and Y. Z. Wang, *RSC Adv.*, 2018, **8**, 26948–26958.
- 85 R. M. Nair, V. N. Boddepalli, M. R. Yan, V. Kumar, B. Gill, R. S. Pan, C. Wang, G. L. Hartman, R. Silva e Souza and P. Somta, *Plants*, 2023, 12, 609.
- 86 B. Tamami, S. Sohn and G. L. Wilkes, *J. Appl. Polym. Sci.*, 2004, **92**, 883–891.
- 87 L. Poussard, J. Mariage, B. Grignard, C. Detrembleur, C. Je, C. Calberg, B. Heinrichs, J. D. Winter, P. Gerbaux, J. Raquez, L. Bonnaud and P. Dubois, *Macromolecules*, 2016, 49, 2161–2171.
- 88 M. Das, B. Mandal and V. Katiyar, *J. Appl. Polym. Sci.*, 2020, 137, 1–12.
- 89 X. Yang, S. Wang, X. Liu, Z. Huang, X. Huang, X. Xu, H. Liu, D. Wang and S. Shang, *Green Chem.*, 2021, 23, 6349–6355.
- 90 X. Liu, X. Yang, S. Wang, S. Wang, Z. Wang, S. Liu, X. Xu, H. Liu and Z. Song, ACS Sustainable Chem. Eng., 2021, 9, 4175–4184.
- 91 X. Sun, Y. Wang, H. Li, J. Zhou, J. Han and C. Wei, *LWT*, 2021, **151**, 112137.
- 92 R. Turco, R. Tesser, V. Russo, T. Cogliano, M. Di Serio and E. Santacesaria, *Ind. Eng. Chem. Res.*, 2021, **60**, 16607–16618.
- 93 N. Nehchiri, S. Amiri and M. Radi, *Packag. Technol. Sci.*, 2021, 34, 283–295.
- 94 Q. Wang and G. W. Padua, J. Agric. Food Chem., 2005, 53, 3444-3448.
- 95 Á. Agüero, D. Lascano, D. Garcia-Sanoguera, O. Fenollar and S. Torres-Giner, *Sustainability*, 2020, 12, 652.
- 96 J. Pouladi, S. M. Mirabedini, H. E. Mohammadloo and N. G. Rad, *Eur. Polym. J.*, 2021, 153, 110502.
- 97 B. S. Adeleke and O. O. Babalola, *Food Sci. Nutr.*, 2020, **8**, 4666–4684.
- 98 M. R. Akkaya, J. Food Sci. Technol., 2018, 55, 2318-2325.
- 99 B. Behera, V. K. Singh, S. Kulanthaivel, M. K. Bhattacharya, K. Paramanik, I. Banerjee and K. Pal, *Eur. Polym. J.*, 2015, **64**, 253–264.
- 100 V. Suthar, M. A. Asare, F. M. d. Souza and R. K. Gupta, *Polymers*, 2022, **14**, 4974.

101 M. Alam, N. M. Alandis, F. Zafar, E. Sharmin and Y. M. Al-Mohammadi, *J. Macromol. Sci., Part A: Pure Appl. Chem.*, 2018, 55, 698–708.

RSC Advances

- 102 M. A. Asare, P. Kote, S. Chaudhary, F. M. d. Souza and R. K. Gupta, *Poolymers*, 2022, **14**, 5282.
- 103 S. Moqadam and M. Salami-Kalajahi, e-Polym., 2016, 16, 0152.
- 104 A. Boyer, E. Cloutet, T. Tassaing, B. Gadenne, C. Alfos and H. Cramail, *Green Chem.*, 2010, 12, 2205–2213.
- 105 A. Farhadian, A. Ahmadi, I. Omrani and A. Babaei, *Polym. Degrad. Stab.*, 2018, 155, 111–121.
- 106 M. E. Farid, M. A. El-Sockary, A. M. El-Saeed, A. I. Hashem, O. M. Abo Elenien, M. S. Selim and S. E. Park, *Mater. Res. Express*, 2019, 6, 065042.
- 107 I. Chakraborty and K. Chatterjee, *Biomacromolecules*, 2020, 21, 4639–4662.
- 108 A. F. Guzmán, D. A. Echeverri and L. A. Rios, *J. Chem. Technol. Biotechnol.*, 2017, **92**, 1104–1110.
- 109 V. Subramaniyan, in Nuts and Seeds in Health and Disease Prevention: Therapeutic Importance of Castor Seed Oil, ed.
 V. R. Preedy and R. R. Watson, Elsevier, 2nd edn, 2020, pp. 485–495, DOI: 10.1016/B978-0-12-818553-7.00034-6.
- 110 J. D. O. Rodrigues, C. K. Z. Andrade, R. L. Quirino and M. J. A. Sales, *Prog. Org. Coat.*, 2022, **162**, 106557.
- 111 J. Nisar, R. Razaq, M. Farooq, M. Iqbal, R. A. Khan, M. Sayed, A. Shah and I. u. Rahman, *Renewable Energy*, 2017, **101**, 111–119.
- 112 S. Saalah, L. C. Abdullah, M. M. Aung, M. Z. Salleh, D. R. A. Biak, M. Basri, E. R. Jusoh and S. Mamat, *Molecules*, 2017, 22, 1–17.
- 113 H. Acherki, A. Bouaid and J. M. Marchetti, *Biofuels, Bioprod. Biorefin.*, 2022, **16**, 219–227.
- 114 L. I. Farfan-Cabrera, E. A. Gallardo-Hernández, M. Gómez-Guarneros, J. Pérez-González and J. G. Godínez-Salcedo, Renewable Energy, 2020, 149, 1197–1204.
- 115 M. Shahinuzzaman, Z. Yaakob and M. Moniruzzaman, J. Cosmet. Dermatol., 2016, 15, 185–193.
- 116 N. H. Mudri, L. C. Abdullah, M. M. Aung, M. Z. Salleh, D. R. Awang Biak and M. Rayung, *Polymers*, 2020, 12, 1–17.
- 117 S. Saalah, L. C. Abdullah, M. M. Aung, M. Z. Salleh, D. R. A. Biak, M. Basri, E. R. Jusoh, S. Mamat and S. S. O. Al Edrus, *Polymers*, 2021, 13, 795.
- 118 M. Rayung, M. M. Aung, M. Sukor, L. C. Abdullah and A. Ahmad, *ACS Omega*, 2020, **5**, 14267–14274.
- 119 S. Dharma, H. H. Masjuki, H. C. Ong, A. H. Sebayang, A. S. Silitonga, F. Kusumo and T. M. I. Mahlia, *Energy Convers. Manage.*, 2016, **115**, 178–190.
- 120 S. Bhakri, M. Ghozali, E. Cahyono, E. Triwulandari, R. W. Kartika, N. N. Solihat, A. H. Iswanto, P. Antov, V. Savov, L. S. Hua, E. A. Agustiany, L. Kristak and W. Fatriasari, J. Renewable Mater., 2022, 1–19.
- 121 N. T. Thuy, V. M. Duc and N. T. Liem, *Vietnam J. Chem.*, 2018, **56**, 181–186.
- 122 C. F. Jisieike and E. Betiku, *Biocatal. Agric. Biotechnol.*, 2020, 24, 101522.
- 123 C. F. Uzoh, A. Nnuekwe, O. Onukwuli, S. Ofochebe and C. Ezekannagha, *J. Cleaner Prod.*, 2021, **280**, 124563.

- 124 J. Huang, J. Zhang, G. Zhu, X. Yu, Y. Hu, Q. Shang, J. Chen, L. Hu, Y. Zhou and C. Liu, *Prog. Org. Coat.*, 2021, **159**, 106391.
- 125 N. T. Thuy, B. X. Nam and V. M. Duc, *Vietnam J. Chem.*, 2019, 57, 735–740.
- 126 P. Chaikul, N. Lourith and M. Kanlayavattanakul, *Ind. Crops Prod.*, 2017, **108**, 56–62.
- 127 A. Saetung, A. Rungvichaniwat, P. Tsupphayakorn-ake, P. Bannob, T. Tulyapituk and N. Saetung, *J. Polym. Res.*, 2016, **23**, 64.
- 128 N. Saetung, S. Somjit, P. Thongkapsri, T. Tulyapitak and A. Saetung, *J. Polym. Res.*, 2016, 23, 58.
- 129 J. Hong, D. Radojčić, X. Q. Yang, X. Wan and Z. S. Petrović, *J. Appl. Polym. Sci.*, 2020, **173**, 48509.
- 130 J. Hong, X. Q. Yang, X. Wan, Z. Zheng and Z. S. Petrović, Polym. Int., 2017, 66, 126–132.
- 131 W. Roschat, T. Siritanon, B. Yoosuk, T. Sudyoadsuk and V. Promarak, *Renewable Energy*, 2017, **101**, 37–944.
- 132 A. H. Raden, F. M. Ahmad and S. Azemi, *Adv. Mater. Res.*, 2013, **812**, 73–79.
- 133 S. A. R. A. Raden, M. A. Faiza and A. Zuliahani, *Sains Malays.*, 2021, **50**, 2407–2417.
- 134 E. Cartea, A. De Haro-Bailón, G. Padilla, S. Obregón-Cano, M. Del Rio-Celestino and A. Ordás, *Foods*, 2019, **8**, 292.
- 135 S. Ghobadi, Z. Hassanzadeh-Rostami, F. Mohammadian, M. Zare and S. Faghih, J. Am. Coll. Nutr., 2019, 38, 185–196.
- 136 V. J. Barthet, in *Reference Module in Food Science: Canola: Overview*, Elsevier, 2016, pp. 1–5, DOI: 10.1016/b978-0-08-100596-5.00029-9.
- 137 N. Raboanatahiry, H. Li, L. Yu and M. Li, *Agronomy*, 2021, **11**, 1776.
- 138 M. Alam, M. Altaf and N. Ahmad, Polymers, 2021, 13, 3325.
- 139 X. Kong, G. Liu and J. M. Curtis, *Int. J. Adhes. Adhes.*, 2011, 31, 559–564.
- 140 M. Leszczyńska, E. Malewska, J. Ryszkowska, M. Kurańska, M. Gloc, M. K. Leszczyński and A. Prociak, *Materials*, 2021, 14, 1772.
- 141 K. Mizera, J. Ryszkowska, M. Kurańska and A. Prociak, *Polym. Bull.*, 2020, 77, 823–846.
- 142 X. Yang, C. Ren, X. Liu, P. Sun, X. Xu, H. Liu, M. Shen, S. Shang and Z. Song, *Mater. Chem. Front.*, 2021, 5, 6160–6170.
- 143 P. Naruebhorn, M. Seadan and S. Suttiruengwong, *Suan Sunandha Science and Technology Journal*, 2022, **9**, 8–14.
- 144 P. Helbling, F. Hermant, M. Pet, T. Tassain, T. Vidil and H. Cramail, *Polym. Chem.*, 2023, 14, 500–513.
- 145 A. R. Mahendran, N. Aust, G. Wuzella, U. Müller and A. Kandelbauer, *J. Polym. Environ.*, 2012, **20**, 926–931.
- 146 T. Wang, H. Deng, N. Li, F. Xie, H. Shi, M. Wu and C. Zhang, *Green Chem.*, 2022, 24, 8355–8366.
- 147 P. Zhang, G. Zhang, J. Pan, C. Ma and G. Zhang, ACS Appl. Mater. Interfaces, 2023, 15, 5998–6004.
- 148 C. Liu, J. Wu, X. Zhou, X. Zhou, Z. Wu and J. Qu, *Prog. Org. Coat.*, 2022, **163**, 106690.
- 149 P. Boisaubert, N. Kébir, A. S. Schuller and F. Burel, *Eur. Polym. J.*, 2020, **138**, 109961.

150 G. Liu, G. Wu, J. Chen and Z. Kong, *Prog. Org. Coat.*, 2016, **101**, 461–467.

- 151 P. S. Choong, N. X. Chong, E. K. Wai Tam, A. M. Seayad, J. Seayad and S. Jana, *ACS Macro Lett.*, 2021, **10**, 635–641.
- 152 X. Chen, A. Pizzi, E. Fredon, C. Gerardin, X. Zhou, B. Zhang and G. Du, *Int. J. Adhes. Adhes.*, 2022, 112, 103001.
- 153 X. Xi, Z. Wu, A. Pizzi, C. Gerardin, H. Lei, B. Zhang and G. Du, *Wood Sci. Technol.*, 2019, 53, 393–405.
- 154 A. Gomez-Lopez, B. Grignard, I. Calvo, C. Detrembleur and H. Sardon, *Macromol. Rapid Commun.*, 2021, 42, 2000538.
- 155 X. Chen, A. Pizzi, X. Xi, X. Zhou, E. Fredon and C. Gerardin, *J. Renewable Mater.*, 2021, **9**, 1045–1057.
- 156 F. Monie, B. Grignard and C. Detrembleur, *ACS Macro Lett.*, 2022, **11**, 236–242.

- 157 X. Chen, J. Li, X. Xi, A. Pizzi, X. Zhou, E. Fredon, G. Du and C. Gerardin, *Polym. Degrad. Stab.*, 2020, **175**, 109121.
- 158 D. L. Smith, D. Rodriguez-Melendez, S. M. Cotton, Y. Quan, Q. Wang and J. C. Grunlan, *Polymers*, 2022, **14**, 5019.
- 159 Y. Wang, Z. Zheng, J. L. Pathak, W. Feng, W. Wu, C. Yang, L. Wu and H. Zheng, *Int. J. Bioprint.*, 2023, 9, 80–93.
- 160 D. Visser, H. Bakhshi, K. Rogg, E. Fuhrmann, F. Wieland, K. Schenke-Layland, W. Meyer and H. Hartmann, *ACS Omega*, 2022, 7, 39772–39781.
- 161 D. C. Aduba, K. Zhang, A. Kanitkar, J. M. Sirrine, S. S. Verbridge and T. E. Long, J. Appl. Polym. Sci., 2018, 135, 46464.
- 162 Y. Yang, H. Cao, Y. Wang, J. Zhao, W. Ren, B. Wang, P. Qin, F. Chen, Y. Wang and D. Cai, *Ind. Crops Prod.*, 2022, 186, 115224