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Divinylglycol, a glycerol-based monomer: valorization, properties and applications

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Abstract

In the context of the development of bio-refineries, glycerol and its derivatives are coproducts of the oleochemistry for which new valorization routes must be found. In this study, the polymerizability of divinylglycol (DVG), a symmetrical C-6 glycerol derivative which bears a vicinal diol and two vinyl functions was studied. The reactivity of the hydroxyl and vinyl functions of DVG through polycondensation and polyaddition reactions was investigated. In a first step, the synthesis of polyesters was carried out by reaction of DVG with various biosourced diesters. In a second route, DVG was polymerized through its vinyl functions by ADMET and thiol-ene addition. Finally, three-dimensional epoxy-amine networks were prepared from a series of diamines and bis-epoxidized DVG, the latter being prepared by oxidation of the DVG double bonds. These different polymerization reactions showed that DVG double bonds were more reactive than the alcohol ones and that a panel of original polymers could be obtained from this bio-sourced synthon.

Introduction

Polymers are ubiquitous in our daily lives; the automotive, packaging, health and textile sectors, to name just a few of them, are meaningful. With the depletion of fossil resources and environmental concerns, the search for more sustainable solutions is becoming a necessity. Since the beginning of the 21st century, bio-sourced chemistry has been widely expending with the valorization of biomass, which is an abundant source of carbon

structures.¹ In this regard, the concept of a biorefinery was created with the objective of producing feed, food, fuels and molecule platforms for industry.² Vegetable oils are the second renewable resource used after ligno-cellulosic biomass.³ Vegetable oils are inexpensive, non-toxic and have a large potential as precursors of bio-based polymers. These oils are composed of triglycerides, which after reaction with an alcohol, water or a base will produce three molecules of fatty acids (or fatty esters) and a molecule of glycerol. Fatty acids are widely studied as precursors of thermoplastic or thermosetting polymers.^{4,5} Glycerol is a simple, non-toxic and versatile molecule that provides access to a wide range of high-value bio-based molecules.⁶

Recently, Len and coll. developed an eco-responsible synthesis of divinylglycol (DVG).⁷ Divinylglycol (DVG) also called 1,5-hexadiene-3,4-diol is a symmetrical C-6 monomer which bears a vicinal diol and two vinyl functions. It can be synthesized from mannitol^{8–10} or tartaric acid,¹¹ but this requires protection/deprotection steps and a rather low yield is obtained (20 to 52%). A third route to synthesize DVG is possible by a pinacol coupling of acrolein with a yield of 90% (Scheme 1).^{12,13} Acrolein can be synthesized from glycerol but this molecule is toxic; Len and coll. then proposed a direct synthesis of DVG from glycerol.⁷



Scheme 1: Synthetic routes to divinylglycol

Divinylglycol is used in the total synthesis of several analogues of natural molecules.^{14–16} As polymer precursor, DVG is actually used as cross-linker for thermosets; Noveon[®] and

Polycarbophil from Lubrizol.^{17–19} However, literature data about the reactivity of DVG and the physico-chemical properties of the polymeric materials are very scarce.^{20–26}

In this study, the reactivity of DVG with respect to chain-growth and step-growth polymerizations was investigated through the synthesis of linear or cross-linked polymers from the two types of available functions of DVG, the hydroxyl and unsaturation functions, respectively.

I. DVG polymerization via hydroxyl reactivity: example of polyester

The synthesis of DVG-based polyesters by polytransesterification was first investigated. DVG has two secondary alcohol functions, known to be of lower reactivity than that of primary alcohols, in particular in the esterification or transesterification reactions.^{27,28} Examples of polyester syntheses from short secondary diols can be find in the literature, in particular 2,3-butanediol which can be obtained by fermentation of glycerol. In the fifties, Watson and coll.²⁹ have described the synthesis of oligoesters from 2,3-butanediol exhibiting molar masses between 450 and 2,600 g/mol. More recently, bio-sourced polyesters based on 2,5-furandicarboxylic acid and 2,3-butanediol have also been synthesized.³⁰ Although the formation of rings was demonstrated, polyesters of relatively high molar masses up to 7,000 g/mol were obtained depending on the nature of the catalyst used. Avérous and coll.³¹ recently studied the esterification kinetics between 2,3-butanediol and 1,4-butanediol with adipic acid and described the synthesis of (co)polyesters from this mixture of diols.³²

In this study, the polyesters were prepared in bulk by copolymerization of DVG (mixture of (±) and meso) with dimethylsuccinate (DMSu) or dimethylsebacate (DMSe), as bio-sourced diesters (Scheme 2). Several catalysts were tested such as 1,5,7-triazabicyclo[4.4.0]dec-5-ene (TBD), sodium methanolate (MeONa), titanium butoxide (Ti(OBu)₄) and titanium isopropoxide (Ti($O^{i}Pr$)₄), which are commonly used in polyester synthesis.^{33,34}



Scheme 2: Polytransesterification reaction of DVG with DMSu (x = 1) or DMSe (x = 4) These experiments show that only oligomers are formed whatever the diester and the catalysts used (Table 1). The apparent molar masses measured by SEC do not exceed 1,840 g/mol for polyesters obtained with DMSe and 1,200 g/mol with DMSu in the presence of TBD, which was the most effective catalyst. Titanium catalysts lead to low conversion of hydroxyl functions of DVG (23 to 50%). These last results can be explained by a possible formation of a complex between the diol and the titanium, which prevents the reaction to take place as already reported by Gau and coll..³⁵

Entry	Diester	Catalyst	Conv (ersion %)	М_п (а) -1	Đ ^(a)	T _d 5%	T _g	T _f
		(10mol.%)	DVG	Diester	(g.mol ⁻)		(°C)	(°C)	(°C)
1		TBD	100	96	950	2	220	-39	-
2	DMSe	MeONa	100	100	1,840	2	260	-54	-
3	2	Ti(OBu)₄	41	23	500	1.2	97	-	-40, 0
4		Ti(O ⁱ Pr) ₄	50	50	700	1.6	120	-	-37, -16
5		TBD	100	90	1,200	1.2	199	-17	-
6	DMSu	MeONa	37	84	800	1.8	147	-47	-
7	DIVISU	Ti(OBu)₄	(b)	(b)	(b)	(b)	88	-	-
8		Ti(O ⁱ Pr) ₄	(b)	(b)	(b)	(b)	88	-	-

Table 1: Characteristics of polyesters synthesized with DVG and DMSe or DMSu as diester in presence of different catalysts

(a) Measured by SEC in THF, PS calibration, (b) Insoluble polymer

Nevertheless, the various NMR analyses confirm the presence of pendant double bonds along the polyester chains. A chemical shift from 5.81 to 5.74 ppm of the peak corresponding to the protons a' of the double bond of DVG is noticed and protons c' in alpha of the alcohols are shifted from 4 to 5.35 ppm, which confirms the reaction of the alcohol functions (Figure 1).



Figure 1: ¹H NMR in CDCl₃ of DVG, DMSe and polyesters synthesized (Entries 1-4, Table 1) ; catalyzed by (a) TBD, (b) MeONa, (c) Ti(OBu)₄ and (d) Ti(OⁱPr)₄

In the following, DVG was further used as a co-monomer (co-diol) with 1,3-propanediol (1,3-PD) or 1,12-dodecanediol (1,12-DD) in the course of polyester synthesis; DMSu or DMSe were kept as diesters and TBD as catalyst (Figure 2). Data are gathered in Table A1 in appendix.



Figure 2: Co-monomers used for polytransesterification of DVG

Results obtained are similar with the two diesters used; incorporation of DVG is difficult in both cases and do not exceed 67 and 58% with DMSu and DMSe, respectively- Moreover the incorporation of DVG leads to a decrease of the polymers molecular masses. The copolyesters synthesized with 1,12-DD are semi-crystalline with T_g around room temperature and several melting temperatures, attesting for a heterogeneous composition of the chains. With the short co-diol, 1,3-PD, the copolyesters formed are all amorphous with a T_g between -52 and -40°C.

In conclusion, the synthesis of high molar mass polyesters from DVG revealed impossible. Nevertheless, the advantage of this monomer is the provision of pendant double bonds in the copolymers, allowing a potential post-functionalization that was not addressed in this study.

II. DVG polymerization via double bond reactivity

a) ADMET polymerization of DVG

Acyclic diene metathesis (ADMET) polymerization follows a step-growth mechanism to produce linear polyalcenamers from α, ω -dienes.³⁶ ADMET proceeds through a transalkylidenation reaction with the release of ethylene, which can be removed by applying vacuum or a constant flow of an inert gas to obtain high conversions and high molecular weight polymers.³⁷



Scheme 3: ADMET polymerization of DVG

ADMET polymerization enables the synthesis of polyalcenamers with well-defined architectures and various functions (alcohol, halogen, amine) giving them unique properties.^{38,39} The reactivity of DVG and derivatives through ADMET polymerization was thus investigated. Knowing that DVG is a short diene with alcohol functions, the possibility of the negative neighboring group effect (NNGE) with catalysts was analyzed. Wagener and coll.⁴⁰ showed that close proximity of functional groups strongly reduces the monomer reactivity, presumably due to complexation of the heteroatom's nonbonded electrons with the metal center. In addition, at higher temperatures, the DVG may be evaporated ($T_b = 198$ °C under 1 atm).

The polymerization of DVG by ADMET (Scheme 3) was tested with five different catalysts, i.e. Schrock catalyst, first and second generation Grubbs catalysts (G1 and G2) and Hoveyda-Grubbs catalysts (HG1 and HG2). The reaction was carried out at 35°C, without solvent and under dynamic vacuum to remove the ethylene formed and shift the equilibrium of the reaction (Table 2).

Condition	Catalyst	Quantity of catalyst (mol.%)	p ^(a) (%)	Mn ^(b) (g/mol)	Đ ^(b)	T _g (°C)	T _d ^{5%} (°C)	Xn
	Schrock	1	-	-	-	-	-	-
	G1	1	-	-	-	-	-	-
Bulk	G2	1	26	780	1.2	-49	74	1.35
24h, 35°C	HG1	1	-	-	-	-	-	-
Under vacuum	HG2	1	65	300	1.5	-21	126	2.9
	HG2	5	68	350	1.1	-10	77	3.1
	HG2	0.5	75	890	1.2	36	122	4

Table 2: Characteristics of poly(DVG) synthesized by ADMET polymerization with differentcatalysts

(a) Conversion of vinyl group calculated by ¹H NMR, (b) Measured by SEC in DMF, LiBr, PS calibration

As expected, no reaction takes place with the Schrock catalyst which is presumably deactivated by the alcohol functions of DVG. Polymerization also does not occur with G1 and HG1, although these are more robust and polar-resistant than Schrock's catalyst. With the 2nd generation catalysts, an oligomerization reaction is observed. In the case of G2 catalyst, however, the double bond conversion of DVG remains very low (26%), probably because of

the low polymerization temperature; indeed, G2 is truly active above 45°C but its activity is 10 times slower that Hoveyda-Grubbs catalyst.^{41,42} On the other hand, the reaction is more favorable with HG2, as evidenced by the release of ethylene observable from the addition of the catalyst. Even if this value is still too low, a 65% conversion to poly(DVG) is reached when HG2 is set to 1mol.%. Other tests at HG2 concentrations of 0.5 and 5 mol.% respectively were carried out but, in all cases, the conversion remains of the same order of magnitude and therefore still insufficient to achieve "correct" molar masses.

Other conditions of reaction have been tested such as addition of a polar solvent (THF or water) or an additive (tetrachloro-1,4-benzoquinone or titanium butoxide) to avoid isomerization reaction or to prevent NNGE by hydroxyl screening respectively (Table A2). Better results were obtained when ADMET polymerization was performed in THF with tetrachloro-1,4-benzoquinone (5mol.%). A maximum of 80% of double bond conversion was obtained leading to the formation of oligomers with a degree of polymerization $X_n = 5$. These last polymerization attempts being carried out in diluted medium, the possibility of forming cycles is favored. MALDI-TOF analysis of the poly(DVG) synthesized in THF in the presence of HG2 as a catalyst demonstrates that no ring was formed and the repeat pattern of P(DVG), 86 g/mol, is clearly visible (Figure 3).



Figure 3: MALDI-TOF of poly(DVG) synthesized by ADMET polymerization with HG2 as catalyst

This first study showed that HG2 was the best catalyst for ADMET polymerization of DVG. Even though DVG exhibits an immediate reactivity with HG2, only oligomers were obtained with a maximum conversion of vinyl functions of 80%. Various reactions were tested by adding additives to the medium to suppress the isomerization reaction induced by the catalyst or the NNGE effect of the alcohol functions of DVG. Although the isomerization reaction was suppressed, similar conversions were calculated. The same results were observed by carrying out the reactions in diluted medium in THF. Again DVG revealed quite difficult to polymerize by ADMET.

In the following, DVG was copolymerized with hydrophobic undecyl undecenoate (UndU) (Figure 4) in order to increase the polymer molar mass and also to prepare copolymers exhibiting amphiphilic properties. Indeed polymers obtained by ADMET polymerization of DVG are water soluble. In a first stage, UndU was polymerized by ADMET at 80°C for 24h under dynamic vacuum using HG2 as a catalyst. Thanks to NMR analysis (Figure A1), calculation of the conversion of the vinyl functions of UndU (88%) allowed us determining an $X_n = 8.3$ (M_n NMR = 2,788 g/mol) while a value of 1,430 g/mol was measured by SEC in THF (Table 3). This difference between the molar mass values is due to the double bond isomerization reaction caused by the ruthenium catalyst (HG2) as confirmed by MALDI-TOF analysis of P(UndU) on which a distribution of the characteristic peaks of the isomerization at an interval of 14 g/mol are detected (Figure A2).



Figure 4: Structure of undecyl undecenoate (UndU)

The poly(UndU) formed was then used for the synthesis of multi-sequenced copolymers by ADMET copolymerization with DVG employed at different ratios (70, 50 and 30mol.%). The ¹H NMR analysis of the copolymers formed was carried out in CDCl₃/DMSO-d6 mixture (50/50 v/v) in order to identify the signal of the two moieties (Figure A3). Low conversions of vinylic functions were obtained in all cases; in the case of DVG/UndU mixture at a ratio 30/70mol.%, the conversion in terminal double bond was high (98%) but the signals corresponding to DVG were not detectable by NMR. For the other two copolymers, DVG signals could be observed and conversions of 24.5 and 36% were calculated for a DVG/UndU ratio of 70/30 and

50/50 mol.%, respectively. These conversions only lead to a degree of polymerization (X_n) less than 2 (Table 3).

	Ratio		p ^(a) (%)		Mn ^(b)	- (b)	ΔН	T _f	Td ^{5%}	
Product	DVG/UndU (mol.%)	DVG	UndU	Total	(g/mol)	שיייש	(J/g)	(°C)	(°C)	Xn
Poly(UndU)	0/100	-	-	88	1,430	1.6	96	34	227	8.3
	70/30	32	17	24.5	1,750	1.3	101	33	143	1.3
	50/50	45	26	36	1,870	1.4	60	34	151	1.5
UndU)	30/70	0	98	98	1,470	1.5	56	33	204	50

 Table 3: Characteristics of multi-sequenced polymers synthesized by ADMET

 polymerization from P(UndU) and DVG

(a) Conversion of vinyl group calculated by RMN ¹H, (b) Measured by SEC in THF, PS calibration

The self-assembly properties in water of the synthesized copolymers composed of hydrophobic (UndU) and hydrophilic (DVG) sequences, were evaluated by nanoprecipitation and analyzed by Dynamic Light Scattering (DLS). The copolymers were dissolved in 0.5 mL of THF at a C = 5 mg/mL and the solution added dropwise (every second) in 4.5 mL of water with a constant stirring of 250 rpm. THF was then evaporated at room temperature and a cloudy solution was obtained visually indicating the presence of dispersed particles (Figure 5). The solution was then filtered on a 0.8 μ m filter. DLS analysis was performed 3 times for each sample as reproducibility test.



Hydrophobicity





These analyses confirm that objects of the order of 200 to 500 nm are formed with polydispersity, PDI, of the order of 0.1 (Figure 5). These dispersions are stable for several months, except for the 50/50 copolymer that contains the largest objects. One should note that the P(UndU) do not lead to stable particle by this process indicating that the 30/70 copolymers contains a small fraction of DVG units.

In this part, multi-sequenced copolymers from DVG and UndU could be synthesized; DVG again shows a low reactivity by ADMET. However, these copolymers exhibit amphiphilic behaviors as confirmed by NMR and DLS analyses.

b) Polymerization of DVG by thiol-ene reaction

A series of aliphatic dithiols were tested in the course of thiol-ene addition to DVG following two procedures. The monomers were used at the 1:1 stoichiometry and the reaction conducted either at 80°C for 24 h or under UV (65 mW/cm² at 365 nm) for 15 min; AIBN and Irgacure 2659 were used as thermo- and photo-initiator, respectively (Figure 6).



Figure 6: Thiol-ene reaction of DVG and structure of dithiols and initiators used

Results obtained show that the photochemical initiation (procedure b) is more efficient than the thermal initiation (procedure a) (Table 4). Indeed, by photopolymerization, polymers of high molar masses (6,000 g/mol) were obtained in short time (15 min). In the case of polymerizations carried out under UV, no kinetic monitoring was carried out but the polymers were analyzed by ¹H NMR after 15 min of irradiation; conversions of double bonds from 93 to 96% could be calculated. The various reactions tested indicate that the chosen aliphatic dithiols have different reactivities with DVG. The best system is with HDT under photochemical activation (Table 4, entry 4); a total conversion is obtained in few hours.

Entry	Dithiol	Procedure	р (%)	Xn[Mn RMN]	M _n ^(c) (g.mol ⁻¹)	Đ ^(c)	T _d 5% (°C)	T _g (°C)	T _f (°C)
1	PDT	(a)	93	14.3[3375]	2,900	1.7	262	-26	-
2	ועם	(b)	93	14.3[3375]	7,400	1.5	277	-22	-
3		(a)	93	14.3[3775]	4,000	2	232	-14	73
4	וטח	(b)	96	33[8712]	6,600	1.6	279	-25	83
5	NDT	(a)	83	5.9[1800]	3,700	1.7	267	35	89
6	NUT	(b)	94	16.7[5100]	6,000	1.5	275	35	90

Table 4: Characteristics of polymers synthesized by thiol-ene reaction with DVG

(a) AIBN (5mol.%), 24h à 80°C, (b) Irgacure (0,05mol.%), 15min under UV (365 nm), (c) Measured by SEC DMF, LiBr, PS calibration.

NMR analysis also shows that the alcohol functions of DVG (characteristic signal at 4.4 ppm) are still present (Figure 7). Moreover, whatever the initiation procedure, NMR spectra of the polymers formed reveal the presence of multiplets at 0.9 ppm which are characteristic of methyl groups (-CH₃). This phenomenon indicates the possible addition of the radical on the most substituted carbon of the DVG double bond whereas anti-Markovnikov addition is generally expected.



Figure 7: ¹H NMR in DMSO of polymers synthesized with DVG and a) BTD, b) HDT or c) NDT by photopolymerization

In conclusion of this section, DVG exhibits a good reactivity towards thiol-ene reactions through thermal or photochemical initiation. The light-curing process enables to synthesize polymers in a simple and rapid manner with conversions of DVG double bonds higher than 93%, in 15 min. Although DVG is hydrophilic and provides hydroxyl functions, the polymers synthesized by thiol-ene addition from DVG and linear dithiols revealed insoluble in water.

Thiol-ene reaction provides access to DVG-based polymers with relatively high molar masses indicating that vinylic functions of DVG display a good reactivity. The synthesis of cross-linked materials by thiol-ene reaction was thus carried out in the presence of a symmetric tetrathiol (PTM), which have the advantage to be not odorous, and Irgacure 2659 as a photo-initiator (Figure 8). The thiol-ene reaction was conducted by mixing DVG and thiol monomer at different ratios. The initiator was then added (0.5 mol.%) and the mixture poured into a Teflon mold of 7.2×5.1cm and then cross-linked under UV (365 nm) and a power of 65 mW/cm² for 15 minutes to form films 2 mm thick.



Figure 8: Structure of tetrathiol and initiator used for the synthesis of DVG-based thermosets

Cross-linked materials by thiol-ene reaction were thus synthesized from DVG and tetrafunctional PTM. The thiol/double bond ratio was varied from 50% to 100% in order to measure the impact of the amount of PTM used on the cross-linking reaction and the properties of the networks obtained (Table 5).

PTM (%)	Swelling THF[H ₂	Soluble part O] (%)	T _d ^{5%} (°C)	T _g (°C)	Е' _{тб+зо°с} (MPa)	ρ (mol/dm ³)	Strain (MPa) ^(a)	Stress (%) ^(a)	Modulus (MPa) ^(a)
50	290[10]	39[15]	120	-33	0,15	0,057	0,1	99 ±7.8	0.13
70	191[7]	14[6]	170	-22	1,8	0.67	0.6 ± 0.1	35.8 ±1.4	1.96 ±0.3
90	163[6]	14[3]	285	-17	2,5	0.93	0.6	28 ±3.8	2.2 ±0.3
100	189[10]	8[3]	290	-16	3,5	1.3	1.2 ± 0.2	33.6 ±4.2	4.1 ±0.6

Table 5: Properties of networks synthesized by thiol-ene with DVG and PTM

(a) Average values

Swelling tests in THF were carried out for 24 hours. The swelling ratios vary from 189% to 290% in agreement with the crosslinking density, p. Similarly, the soluble fraction decreases with the percentage of PTM used, from 39% to 8%. These experiments confirm that a high content of tetrathiol (PTM) is required to obtain the formation of a 3D-network. Since DVG brings some hydrophilicity through its OH groups, swelling tests in water were also carried

out; the swelling ratios, around 10%, are lower in water in comparison to THF, but follow the same trend according to the PTM content. These results were confirmed by FTIR analysis of the films obtained (Figure A4); a characteristic signal of the double bond of DVG is observable at 3080 and 1640 cm⁻¹ and decrease with higher PTM concentrations demonstrating the almost complete crosslinking of the polymers.

As a conclusion of this part, 3D networks based on DVG and PTM were prepared. The latter have different mechanical properties whose properties vary with the degree of crosslinking. It was found also possible to vary the hydrophilicity of these materials by varying the thiol used.

III. DVG-derivate polymerization: example of epoxy-amine network

While DVG direct polymerization remains of low efficiency either using double bond or hydroxyl reactivity, the derivatization of DVG to more reactive function was investigated. At first, DVG was derivatized by oxidation of the double bonds in order to obtain a bis-epoxidized precursor (DAG). DVG was oxidized in dichloromethane during 24h in the presence of MCPBA in excess (Scheme 4). A maximum yield of 50% was obtained with incomplete conversion of DVG. DAG was then reacted with a series of diamines to prepare epoxy networks.



Scheme 4 : Epoxidation reaction of DVG with mCPBA

Syntheses with short alkyl diamine having primary or secondary amino groups (ethylene diamine, propylene diamine, isophorone diamine, diethylene triamine) lead to 'burned' materials; indeed, enthalpies of reaction values measured by DSC are quite high (422 to 647 J.g⁻¹) and the reaction starts at room temperature (Table A3). These preliminary investigations clearly demonstrated the high reactivity of DAG. To avoid this issue, diamines with long alkyl chains were selected, such as *Priamine*[®] 1075 which is a bio-sourced derivative from fatty acid and *Jeffamine ED600* and *ED900* which are oligo-ethers with molar masses equal to 678.8 and 972 g/mol, respectively. The curing reaction was followed by DSC. Enthalpy of reaction (Δ H), reaction start temperature ($T_{on set}$) and glass transition temperature (T_g) were determined by

DSC (Table 6). These analyzes reveal relatively low glass transition temperature values in agreement with the T_g of *Jeffamine* precursors, -39 and -45°C respectively. A T_g of 26°C was measured for the networks formed with *Priamine® 1075* which brings more rigidity. Formulation of networks was carried out from EEW (epoxy equivalent weight, g/eq.) value of DAG and from AHEW (amine hydrogen equivalent weight, g/eq.) value of the diamine. The curing of epoxy/amine formulation is theoretically performed with a ratio of 1 mole of epoxy group with 1 mole of active hydrogen from amine.

Diamina	£	AHEW	ΔΗ	T _{on set}	Tg
Diamine	T	(g.eq ⁻¹)	(J.g ⁻¹)	(°C)	(°C)
Priamine [®] 1075	4	136.17	433	36	26
Jeffamine ED600	4	169.7	353	62	-39
Jeffamine ED900	4	243	263	71	-45

Table 6: Characteristics of the reactivity of diamines with DAG

Networks obtained with these diamines display different properties. With *Priamine® 1075,* the curing is performed at room temperature during 24h. Swelling tests were made in THF and water (Table 7).

Diamine	Swelling (%)	Soluble part (%)	Td ^{5%} (%)	$T_{\alpha}(T_{g})$	E' _{Tα+30°C}	ρ (mal(dm ³)
	THF [H ₂ O]		(/0)	()	(IVIPa)	(moi/am)
Priamine® 1075	180 [20]	34 [2]	287	22(19)	6.2	1.6

Table 7: Characteristics of DAG/ Priamine® 1075 network

The network swelling was higher in THF (180%) and revealed a fairly large soluble fraction of 34%, demonstrating that the crosslinking was not complete. In water, due to its chemical structure, DAG provides hydroxyl functions and the opening of the epoxide ring generates new ones, which explains a certain affinity of this material with water. The proportion of soluble fraction remains nevertheless very low at 2%. Mechanical measurements by DMA

(Dynamic mechanical analysis) allowed us estimating a crosslink density of 1.6 mol/dm³ and measuring a phase transition, T α around 20°C.

With *Jeffamines*, networks were cured for 1h at 120°C and 2h at 140°C, the complete polymerization was confirmed by DSC (Table A4). The network prepared from *Jeffamine ED600* was similar to a viscous fluid while an aspect close to an elastomeric gum was obtained from *Jeffamine ED900*. Sticky materials were obtained so tack properties of these networks were measured by DMA (Figure A5); tack is measured as the force required to separate adhesive from a probe at the interface shortly after they have been brought into contact under a defined duration at a specific temperature (here, 10s at room temperature).

Diglycidyl ether of bisphenol-A (DGEBA) and isophorone diamine (IPDA) are classical components of epoxy-amine networks. One of the major issues with this type of material is the impossibility of degrading them in 'soft' conditions. With this purpose, DAG was incorporated into the DGEBA/IPDA network at different percentages (1, 5 and 10 wt.%) in order to provide 1,2-diol units within the network and then evaluate if the latter could be cleaved after treatment with a strong acid such as periodic acid (Table 8).

DAG (w.%)	Td ^{5%} (%)	Τ _α (Τ _g) (°C)	ρ (mol/dm ³)	Strain (MPa) ^(a)	Stress (%) ^(a)	Young modulus (MPa) ^(a)	Swelling	Soluble part
							THE [H]	20] (/8)
0	252	1/10/127)	10.6	74 +11	62+11	1337	2 [1]	~1
U	352	140(137)	10.0	74 111	0.5 ±1.1	±102	2[1]	< <u>1</u>
						1971		
1	341	135(122)	5.9	57 ±6.2	3.4 ±0.5	±165	4 [2]	<1
						1105		
5	312	151/120)	6.2	40 +14	3 / +1 3	1522	1 [2]	~1
5	542	131(123)	0.2	40 114	5.4 ±1.5	±17	4 [2]	1
						2105		
10	336	138, 157	6.6	56 ±12	3.9 ±0.8	+150	3 [2]	<1
						TT20		

Table 8: Properties of DGEBA / IPDA materials with addition of 1, 5 and 10wt.% of DAG

(a) Average values

This first study demonstrates that DAG could be used as a comonomer in DGEBA/IPDA mixtures without detrimental of the final thermomechanical properties. When DAG was used at 1 and 5wt.%, it is perfectly mixed within the DGEBA/IPDA system and the network so-formed remains homogeneous. However, at 10wt.%, a phase segregation is observed, as attested by the DMA analysis (Figure A6) that reveals the presence of two Tα values and by

DSC analysis, T_g was not detected (Table 8). IPDA has a similar reactivity with DGEBA and DAG ($\Delta H = 420 \text{ J/g}$) but the polymerization starts at room temperature with DAG instead of 88°C with DGEBA. Swelling tests in THF and water show that the level of soluble fraction is negligible (less than 1% in both cases), demonstrating that the networks so-formed are therefore completely cross-linked (Table 8).

As already stated, the advantage of incorporating DAG in the DGEBA/IPDA network is to provide 1,2-diol units in the final network that can be cleaved. After 24h of reaction with periodic acid (1eq/DAG) in THF, the remaining polymer was oven-dried and weighed. The mass measured was nearly equal to the initial one indicating that the cleavage did not occur. However, the analysis by SEC of the soluble fraction revealed the presence of oligomers with molar masses between 700 and 6,000 g/mol (RI detection), also higher molar masses are detected by UV which could indicate the presence of aromatic rings resulting from DGEBA unit. These same samples were analyzed by FTIR; a signal at 1700 cm⁻¹ is identifiable and may correspond to the C=O elongation of the ketones or aldehydes formed after cleavage of the 1,2-diol moiety (Figure A7). This preliminary test gives some insights of polymer degradation and will be further investigated.

Conclusion

In conclusion, this study aims at investigating the polymerizability of DVG in various polymerization reactions for the development of original polymer materials. This work allowed us a better understanding of the reactivity of DVG and the access to potentially valuable oligomers and networks. However, this work also reveals the very low reactivity of DVG due to its very specific chemical structure, i.e the close vicinity of hydroxyl and unsaturation functions. Thiol-ene reaction was found the most realistic and efficient reaction to prepare linear polymer and cross-linked materials from DVG. These materials have pendant hydroxyl functions which can be post-functionalized in order to bring new properties and can be potentially degraded by cleavage of 1,2 diol moieties in acidic conditions.

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References

- (1) Okkerse, C.; Bekkum, H. van. From Fossil to Green. *Green Chem.* 1999, *1*, 107–114.
- (2) Marion, P.; Bernela, B.; Piccirilli, A.; Estrine, B.; Patouillard, N.; Guilbot, J.; Jerome, F. Sustainable Chemistry: How to Produce Better and More from Less? *Green Chem.* 2017, 19, 4973–4989.
- (3) Gallezot, P. Process Options for Converting Renewable Feedstocks to Bioproducts. *Green Chem.* 2007, *9*, 295–302.
- (4) Meier, M. A. R.; Metzger, J. O.; Schubert, U. S. Plant Oil Renewable Resources as Green Alternatives in Polymer Science. *Chem. Soc. Rev.* 2007, *36*, 1788–1802.
- (5) Maisonneuve, L.; Lebarbé, T.; Grau, E.; Cramail, H. Structure–properties Relationship of Fatty Acid-Based Thermoplastics as Synthetic Polymer Mimics. *Polym. Chem.* 2013, *4*, 5472–5517.
- (6) Bozell, J. J.; Petersen, G. R. Technology Development for the Production of Biobased Products from Biorefinery Carbohydrates—the US Department of Energy's "Top 10" Revisited. *Green Chem.* 2010, *12*, 539–554.
- (7) Sotto, N.; Cazorla, C.; Villette, C.; Billamboz, M.; Len, C. Toward the Sustainable Synthesis of Biosourced Divinylglycol from Glycerol. *ACS Sustain. Chem. Eng.* 2016, *4*, 6996–7003.
- Rao, R.; Mysorekar, S. V.; Gurjar, M. K.; Yadav, J. S. Synthesis of (3R,4R)-1,5-Hexadien-3,4-Diol and Its Unsymmetrical Derivatives : Application to (R)-(+)-α-Lipoic Acid. *Tetrahedron Lett.* 1987, *28*, 2183–2186.
- Yadav, J. S.; Mysorekar Pawar, S. V. S. M.; Gurjar, M. K. Synthesis of (3R,4R)-1,2-Divinylglycol and Its Unsymmetrical Derivatives: An Application to the Synthesis of R-(+)-(Alpha)-Lipoic Acid. J. Carbohydr. Chem. 1990, 9, 307–316.
- (10) Aravind, A.; Mohanty, S. K.; Pratap, T. V.; Baskaran, S. Regioselective Cleavage of the Bis-Benzylidene Acetal of D-Mannitol under Oxidative and Reductive Conditions: A New Approach to C2-Symmetric Chiral Ligands. *Tetrahedron Lett.* 2005, *46*, 2965–2968.
- (11) Michaelis, S.; Blechert, S. Total Synthesis of (+)-Phomopsolide C by Ring-Size Selective Ring-Closing Metathesis/Cross-Metathesis. *Org. Lett.* 2005, *7*, 5513–5516.
- (12) Braun, R. A. Preparation of 4,5-Dihydroöxepine and 1,2-Divinylethylene Oxide. *J. Org. Chem* 1963, *313228*, 1383–1384.
- (13) Billamboz, M.; Legeay, J.; Hapiot, F.; Monflier, E.; Len, C. Novel Strategy for the Bis-Butenolide Synthesis via Ring-Closing Metathesis. *Synthesis (Stuttg)*. 2011, 44, 137–143.
- (14) Burke, S. D.; Voight, E. A. Formal Synthesis of (+)-3-Deoxy-D-Glycero-D-Galacto-2-Nonulosonic Acid (KDN) via Desymmetrization by Ring-Closing Metathesis. *Org. Lett.* 2001, *3*, 237–240.

- (15) Quinn, K. J.; Isaacs, A. K.; DeChristopher, B. A.; Szklarz, S. C.; Arvary, R. A. Asymmetric Total Synthesis of Rollicosin. *Org. Lett.* 2005, *7*, 1243–1245.
- (16) Marvin, C. C.; Voight, E. A.; Burke, S. D. Synthesis of (+)-Didemniserinolipid B via Ketalization/Ring-Closing Metathesis. *Org. Lett.* 2007, *9*, 5357–5359.
- (17) Robinson, J. R. Method of Moisturizing Tissue and Composition. EP 0429156 B1, 1994.
- (18) Ch'Ng, H. S.; Park, H.; Kelly, P.; Robinson, J. R. Bioadhesive Polymers as Platforms for Oral Controlled Drug Delivery II: Synthesis and Evaluation of Some Swelling, Water-Insoluble Bioadhesive Polymers. J. Pharm. Sci. 1985, 74, 399–405.
- (19) Trafton, P. E. L.; Crowley, A. P.; Stevens, R. L. Calcium Polycarbophil-Alginate Controlled Release Composition and Method. US 5110605 A, 1992.
- (20) Braun, R. A. Copolymers of a Vinyl Substituted Ethylene Glycol. US 3157623, 1964.
- (21) Stogryn, E. L.; Passannante, A. J. Preparation and Polymerization of 1,2-Divinyl Ethylene Oxide. US 3261819, 1966.
- (22) Bader, A.; Weiss, F. Hydroxy Alkylene Ethers of Divinyl Glycol. US 3714265, 1973.
- (23) Hoegger, E. F.; Werntz, J. H. Polyurethanes Produced from 1,2-Divinyl Ethylene Glycol. US 3376266, 1968.
- (24) Michelin & Cie. Improvements. Methods of Preparing Elastomers of the Polyurethane Type. US 3100759, 1963.
- (25) Boussu, G.; Neuville, L.; Massoubre, J.-M. Linear Polyurethane Elastomers. US 3219633, 1965.
- (26) Kuhar, G. A. Method for Making High Temperature Insulating Compositions and Said Compositions. US 3699023, 1972.
- (27) Brandner, J. D.; Birkmeier, R. L. Relative Esterifiability of the Primary and Secondary Hydroxyl Groups of Glycerol. *J. Am. Oil Chem. Soc.* 1960, *37*, 390–396.
- (28) Ramalinga, K.; Vijayalakshmi, P.; Kaimal, T. N. B. A Mild and Efficient Method for Esterification and Transesterification Catalyzed by Iodine. *Tetrahedron Lett.* 2002, *43*, 879–882.
- (29) Watson, R. W.; Grace, N. H.; Barnwell, J. L. Production and Properties of 2,3-Butanediol XXXVI. Linear Polyesters of 2,3-Butanediol. *Can. J. Res.* 1950, *28*, 652–659.
- (30) Gubbels, E.; Jasinska-Walc, L.; Koning, C. E. Synthesis and Characterization of Novel Renewable Polyesters Based on 2,5-Furandicarboxylic Acid and 2,3-Butanediol. *J. Polym. Sci. Part A Polym. Chem.* 2013, *51*, 890–898.
- (31) Debuissy, T.; Pollet, E.; Avérous, L. Synthesis of Potentially Biobased Copolyesters Based on Adipic Acid and Butanediols: Kinetic Study between 1,4- and 2,3-Butanediol and Their Influence on Crystallization and Thermal Properties. *Polymer (Guildf)*. 2016, 99, 204–213.
- (32) Hu, X.; Shen, X.; Huang, M.; Liu, C.; Geng, Y.; Wang, R.; Xu, R.; Qiao, H.; Zhang, L. Biodegradable Unsaturated Polyesters Containing 2,3-Butanediol for Engineering

Applications: Synthesis, Characterization and Performances. *Polymer (Guildf).* 2016, *84*, 343–354.

- (33) Lebarbé, T.; Maisonneuve, L.; Nga Nguyen, T. H.; Gadenne, B.; Alfos, C.; Cramail, H. Methyl 10-Undecenoate as a Raw Material for the Synthesis of Renewable Semi-Crystalline Polyesters and Poly(Ester-Amide)S. *Polym. Chem.* 2012, *3*, 2842.
- (34) Montaudo, G.; Rizzarelli, P. Synthesis and Enzymatic Degradation of Aliphatic Copolyesters. *Polym. Degrad. Stab.* 2000, *70*, 305–314.
- (35) You, J. S.; Shao, M. Y.; Gau, H. M. Enantioselective Addition of Diethylzinc to Aldehydes Catalyzed by Titanium(IV) Complexes of Diol Derivatives of D-Mannitol and the Spectroscopic Study of the Catalytic System. *Organometallics* 2000, *19*, 3368–3373.
- (36) Baughman, T. W.; Wagener, K. B. Recent Advances in ADMET Polymerization. *Adv. Polym. Sci.* 2005, *176*, 1–42.
- (37) Rybak, A.; Meier, M. a R. Acyclic Diene Metathesis with a Monomer from Renewable Resources: Control of Molecular Weight and One-Step Preparation of Block Copolymers. *ChemSusChem* 2008, *1*, 542–547.
- (38) Schulz, M. D.; Wagener, K. B. Precision Polymers through ADMET Polymerization. *Macromol. Chem. Phys.* 2014, *215*, 1936–1945.
- (39) Mutlu, H.; Montero de Espinosa, L.; Meier, M. a R. Acyclic Diene Metathesis: A Versatile Tool for the Construction of Defined Polymer Architectures. *Chem. Soc. Rev.* 2011, 40, 1404–1445.
- (40) Wagener, K.; Brzezinska, K. Kinetics of Acyclic Diene Metathesis (ADMET) Polymerization. Influence of the Negative Neighboring Group Effect. *Macromolecules* 1997, No. 30, 7363–7369.
- (41) Lehman, S. E.; Wagener, K. B. Comparison of the Kinetics of Acyclic Diene Metathesis Promoted by Grubbs Ruthenium Olefin Metathesis Catalysts. *Macromolecules* 2002, *35*, 48–53.
- (42) Courchay, F. C.; Sworen, J. C.; Wagener, K. B. Metathesis Activity and Stability of New Generation Ruthenium Polymerization Catalysts. *Macromolecules* 2003, *36*, 8231– 8239.

Diester [Diol]	Ratio DVG/Diol	(Conversion (%)		Final ratio DVG/Diol (mol.%)	M _n ^(a) (g/mol)	Đ ^(a)	T _d 5% (°C)	T _g (°C)	T _f (°C)
	(mol.%)	DVG	Diol	Diester	(mol.%)	(8/		(0)	()	(0)
	0/100	-	98	88	-	2,000	1,4	140	-48	-
DMSu	30/70	58	100	88	20/80	1,200	1,3	181	-40	-
[1,3- PD]	50/50	59	96	91	43/57	620	1,3	137	-52	-
	70/30	74	95	64	67/33	650	1,3	132	-48	-
	0/100	-	98	100	-	3,600	1,9	281	-	75
DMSu	30/70	100	75	89	12/88	2,800	1,7	109	26	67
[1,12- וחס	50/50	21	92	77	49/51	970	1,3	78	17	55, 59
	70/30	40	99	72	59/41	430	1,6	95	10	-14, 35, 43
	0/100	-	100	80	-	1,900	1,3	166	nd	nd
DMSe	30/70	50	95	72	21/79	1,000	1,2	124	-44	15, 33
[1,3- PD]	50/50	40	95	43	29/71	630	1,5	132	-47	13, 29
,	70/30	77	99	56	50/50	630	1,4	124	-48	-13, -9, -4
	0/100	-	100	100	-	6,700	1,8	254	50	80
DMSe	30/70	nd	100	95	4/96	3,300 ^(b)	2 ^(b)	269	nd	76
[1,12- DD]	50/50	100	100	53	32/68	1,450 ^(b)	1,8 ^(b)	150	nd	31, 61, 66
551	70/30	100	100	63	58/42	1,000 ^(b)	1,7 ^(b)	158	nd	27, 47

Table A1: Characteristics of copolyesters synthesized from DVG and a co-diol (1,3-PD or 1,12-DD) with a diester (DMSu or DMSe), in the presence of TBD as a catalyst (2mol.%)

(a) Measured by SEC in THF, PS calibration

Conditions	HG2 quantity (mol%)	Additive (5mol%)	p ^(a) (%)	Mn ^(b) (g/mol)	Đ ^(b)	T _g (°C)	Td ^{5%} (°C)	Xn
Bulk	1	Chloranil	74	890	1,1	51	149	4
24h, 35°C	T	Ti(OBu)4	64,5	715	1,2	-32	81	2,8
H ₂ O	1	_	63 5	690	1 1	11	120	27
48h, 35°C	-		00,0	030	-,-		120	2,,
	0,5	-	76	840	1,2	18	90	4,2
THF	1	-	75	900	1,2	33	97	4
24h, 35°C	1	Chloranil	80	1,000	1,1	45	142	5
	5	Ti(OBu)₄	78	940	1,3	44	116	4,6

Table A2: ADMET polymerization of DVG with HG2 as a catalyst in the presence of asolvent or an additive

(a) Calculated by RMN ¹H, (b) Measured by SEC in THF, PS calibration



Figure A1: ¹H NMR spectra in CDCl₃ of UndU and Poly(UndU)





Figure A2: MALDI-TOF of Poly(UndU), matrice DTCB



Figure A3: ¹H NMR spectra of Poly(DVG-UndU) in DMSO, CDCl₃ and a mixture of DMSO/CDCl₃ (50/50 v/v), respectively



Figure A4: FTIR spectra of networks synthesized with DVG and PTM

Diamina	£		ΔΗ	T _{on set}	Tg
Diamine	I	AREW	(J.g ⁻¹)	(°C)	(°C)
EDA	4	15	524	30	86
PDA	4	18,5	639	30	69
DETA	5	20,63	647	31	73
IPDA	4	42,56	422	39	124
СНМА	4	35,56	597	34	86

Table A3: Characteristics of diamines used to formulated epoxide/amine networks

Diamine	Tď²% (%)	T _g (°C)	F _{max} (N)	W _{adh} (N/s)	t _{rupture} (s)	Gap _{rupt} (mm)
Jeffamine ED600	287	-30	0,2	0,2	30	30
Jeffamine ED900	289	-42	0,35	4,1	10	10,8

0,4---DAG/Jeffamine ED600--DAG/Jeffamine ED9000,30,20,1

Figure A5: Tack property analysis of networks synthesized with DAG and *Jeffamine ED600* or *ED900*

Table A4: Characteristics of networks synthesized with DAG and Jeffamines



Figure A6: Analysis of cross-linked epoxide/amine networks synthesized with DGEBA/DVG and IPDA. A) Young modulus, B)Tan delta, C) tensile tests



Figure A7: Analysis of soluble part after cleavage reaction, SEC in THF, PS calibration (left) and FTIR analysis (right)