Advanced Polymer Nanocomposites with Tailored Morphologies for High Voltage Insulating Systems

by

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DEDICATION

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To my husband, Mohamed ZEMZEM, who has been a constant source of support for me.

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NANOCOMPOSITES DE POLYMÈRES À MORPHOLOGIES CONTRÔLÉES POUR DES APPLICATIONS D'ISOLATION HAUTE TENSION

Emna HELAL

RÉSUMÉ

Cette thèse rapporte différentes approches pour préparer des matériaux d'isolation nanostructurés qui bénéficient d'une dispersion contrôlée de nanoparticules dans l'objectif d'optimiser leurs propriétés. Ces approches se basent sur l'utilisation de matrices polymères multi-phases, telles que les copolymères à blocs et les mélanges de polymères, comme gabarit pour guider la dispersion des nanoparticules. Deux types de nanoparticules, organiquement modifiées, ont été utilisés: l'oxyde de zinc (ZnO) sous forme de sphères et l'argile Montmorillonite sous forme de plaquettes. En plus, un copolymère à blocs a été sélectionné comme gabarit, à savoir: le polystyrène-*b*-poly(éthylène-*co*-butylène)-*b*-polystyrène (SEBS) constitué de deux blocs de polystyrène (PS) sous forme de domaines nanométriques et d'un bloc élastomère de poly(éthylène-*co*-butylène) (PEB).

Des nanocomposites SEBS/argile et SEBS/ZnO ayant des morphologies variées ont été fabriqués. En particulier, l'orientation des nanodomaines du bloc PS et des plaquettes d'argile dans ces nanocomposites a été contrôlée en ayant recours à différentes méthodes de fabrication. Les morphologies obtenues varient de l'isotropie totale à l'orientation totale suivant une seule direction, en passant par des morphologies intermédiaires partiellement orientées. La distribution spatiale des deux types de nanoparticules ainsi que leurs affinités respectives aux blocs PS et PEB ont été également contrôlées par la présence ou non de groupements fonctionnels d'anhydride maléique (MA) greffés sur le bloc élastomère PEB. Les propriétés diélectriques, rhéologiques, thermiques et mécaniques de ces nanocomposites ont été caractérisées en corrélation avec leurs morphologies.

Dans les nanocomposites SEBS/argile, l'incorporation des argiles a réduit la mobilité des chaînes du bloc PEB situées à l'interface polymère/nanoparticule. Une transition vitreuse, plus élevée que celle du bloc PEB, a été attribuée à ces chaînes de la zone interfaciale. De plus, l'orientation et la localisation des argiles ont affecté la mobilité de ces chaînes et en conséquence les propriétés d'ingénierie. Plus précisément, les nanocomposites SEBS/argile ayant une morphologie partiellement alignés ont assuré la combinaison optimale de résistance diélectrique et mécanique.

Dans les nanocomposites SEBS/ZnO et en présence du MA, la dispersion des nanoparticules et leur affinité au bloc PEB ont été nettement améliorées. Par conséquence, un réseau entre les nanosphères de ZnO et les chaînes de polymère a été formé. Ce réseau a été associé à une amélioration de la conductivité thermique et une remarquable augmentation de la résistance à l'érosion de surface par décharges couronne; le volume érodé étant réduit de 90% à seulement 5 wt% de ZnO. Dans la dernière partie du projet, des nanocomposites SEBS/ZnO sélectionnés ont été mélangés avec du polyéthylène (PE) pour préparer des nanocomposites hybrides PE/SEBS/ZnO candidats pour l'isolation haute tension. La résistance à l'érosion de surface et la flexibilité mécanique des nanocomposites PE/SEBS/ZnO ont été améliorées par rapport aux nanocomposites de référence PE/ZnO. Cette augmentation a été associée à l'amélioration de la dispersion des nanoparticules de ZnO et à leur localisation sélective dans SEBS et à l'interface entre PE et SEBS.

Mots clés: isolation haute tension, nanocomposite de polymère, nanodiélectrique, dispersion contrôlée de nanoparticules, copolymère à blocs, mélange de polymères...

ADVANCED POLYMER NANOCOMPOSITES WITH TAILORED MORPHOLOGIES FOR HIGH VOLTAGE INSULATING SYSTEMS

Emna HELAL

ABSTRACT

This thesis reports different approaches to prepare a new generation of nanostructured insulating materials featuring controlled nanoparticles dispersion, using block copolymers and polymer blends as template matrices. Two types of nanoparticles, both organically modified, were used: zinc oxide (ZnO) and Montmorillonite clay. In addition, polystyrene-*b*-poly(ethylene-*co*-butylene)-*b*-polystyrene (SEBS) triblock copolymer, composed of two polystyrene (PS) endblocks in the form of well-ordered nanodomains and one poly(ethylene-*co*-butylene) (PEB) elastomer middle block, was selected as a template matrix.

SEBS/clay and SEBS/ZnO nanocomposites featuring different configurations of PS domains and clay platelets, namely: isotropic, partially oriented and oriented morphologies were achieved by varying the processing techniques. Besides, the spatial distribution of clay platelets and ZnO nanospheres and their affinities to either PS block or PEB block were tuned by the presence or not of maleic anhydride (MA) graft attached to PEB block. In particular, the dispersion of both types of nanoparticles was considerably improved in the presence of MA. Dielectric, rheological, thermal and mechanical properties of these nanocomposites were characterized in correlation with their morphologies.

In SEBS/clay nanocomposites, it was found that the incorporation of clay induced slower dynamics of PEB chains located in the interfacial region. A new interfacial glass transition (T_{gi}), higher than the glass transition (T_g) of bulk PEB, was attributed to these interfacial chains. Furthermore, the orientation and location of clay affected the interfacial dynamics: the highest T_{gi} temperatures were related to samples with lower alignment degree and preferential location of clay in PEB phase. Functional properties were also affected by the orientation. To be more specific, SEBS/clay nanocomposites with totally aligned clay platelets and PS domains were simultaneously the most efficient in improving the breakdown strength up to 45% and the less efficient in improving the best combination of dielectric breakdown strength and mechanical strength.

In SEBS/ZnO nanocomposites, the improved dispersion and affinity to PEB block, achieved in the presence of MA, induced the formation of networks between ZnO nanoparticles and SEBS chains. This behavior was accompanied by an increase of thermal conductivity and excellent improvement of the resistance to surface erosion: eroded volume reduced by 90% at only 5wt% ZnO. In the last part of the project, selected SEBS/ZnO nanocomposites were mixed with polyethylene (PE) to prepare blend nanocomposites as new candidates for HV insulation. Although the overall dielectric performance of unfilled PE/SEBS blend was reduced compared to neat PE, PE/SEBS/ZnO blend nanocomposites featured higher resistance to surface erosion and mechanical flexibility compared to conventional PE/ZnO nanocomposites. This improvement was correlated with the improved dispersion of ZnO nanoparticles in PE/SEBS/ZnO compared to PE/ZnO nanocomposites and their selective localization in SEBS phase and potentially at the interfaces between PE and SEBS.

Keywords: High voltage insulation, polymer nanocomposite, nanodielectric, tailored nanoparticles dispersion, block copolymer, polymer blend...

TABLE OF CONTENTS

Page

INTRO	DUCTION	N	3
CHAP	TER 1	NANOCOMPOSITES WITH TUNED MORPHOLOGY FOR HV	
		INSULATION	13
1.1	Definition	of nanocomposites and nanodielectrics	13
	1.1.1	Polymer nanocomposites definition	13
	1.1.2	Definition of nanodielectrics	14
	1.1.3	Theories and models regarding the interfacial region	15
		1.1.3.1 Interlayer model	16
		1.1.3.2 Multi-core model	18
	1.1.4	Nanoparticles vs. Microparticles in polymer based insulations	19
	1.1.5	Review of nanoparticles used in HV insulation systems	22
		1.1.5.1 Metal oxide nanoparticles	22
		1.1.5.2 Nitride and Carbide nanoparticles	24
		1.1.5.3 Anisotropic nanoparticles	24
	1.1.6	Parameters affecting the role of nanoparticles in HV	
		insulating materials	25
		1.1.6.1 Effect of nanoparticles size	25
		1.1.6.2 Effect of nanoparticles shape and orientation	26
		1.1.6.3 Effect of nanoparticles dispersion	29
1.2	Control of	morphology: application to nanodielectrics	30
	1.2.1	Block copolymer nanocomposites	30
	1.2.2	Polymer blend nanocomposites	33
1.3	Review of	the materials	36
	1.3.1	Polyethylene	36
	1.3.2	SEBS thermoplastic elastomer	37
	1.3.3	Zinc Oxide nanoparticles	39
	1.3.4	Organically modified Montmorillonite Clay	40
CHAP	TER 2	ARTICLE 1: STYRENIC BLOCK COPOLYMER-BASED	
		NANOCOMPOSITES: IMPLICATIONS OF	
		NANOSTRUCTURATION AND NANOFILLER TAILORED	
		DISPERSION ON THE DIELECTRIC PROPERTIES	43
2.1	Introductio	on	44
2.2	Materials	and experimental characterization	46
	2.2.1	Experimental strategy	46
	2.2.2	Materials and methods	47
2.3	Results an	d discussion	50
	2.3.1	Morphology	50
		2.3.1.1 State of dispersion and location of nanoclay	50
		2.3.1.2 Amount of nanoparticle surface area	52

		2.3.1.1 Nanoclay orientation	55
		2.3.1.2 SEBS morphology	56
		2.3.1.3 Stability of SEBS morphology	57
	2.3.2	Dielectric properties	61
	2.3.3	AC short-term breakdown strength	70
	2.3.4	Summary of results	74
2.4	Conclusio	on	75
CHAJ	PTER 3	ARTICLE 2: INTERFACIAL MOLECULAR DYNAMICS OF	
		STYRENIC BLOCK COPOLYMER-BASED NANOCOMPOSITE	S
		WITH CONTROLLED SPATIAL DISTRIBUTION	77
3.1	Introduct	ion	78
3.2	Materials	and methods	81
3.3	Character	rization	86
	3.3.1	Small Angle X-ray Scattering	86
	3.3.2	Transmission Electron Microscopy	86
	3.3.3	Broadband dielectric spectroscopy	86
	3.3.4	Dynamic mechanical analysis	87
3.4	Results an	nd discussion	87
	3.4.1	Morphology	87
		3.4.1.1 Totally isotropic morphologies	87
		3.4.1.2 Totally aligned morphologies	90
		3.4.1.3 Partially aligned morphologies	90
	3.4.2	Dielectric properties	97
		3.4.2.1 Effect of orientation on dielectric behavior	102
		3.4.2.2 Effect of styrene content and diameter of PS cylinders	105
		3.4.2.3 Effect of nanoparticles location and intercalation degree	109
	3.4.3	Dynamic mechanical analysis	111
3.5	Concludi	ng remarks	119
CHAI	TER 4	ARTICLE 3: THERMOPLASTIC ELASTOMER	
		NANOCOMPOSITES WITH CONTROLLED NANOPARTICLES	
		DISPERSION FOR HV INSULATION SYSTEMS:	
		CORRELATION BETWEEN RHEOLOGICAL, THERMAL,	
		ELECTRICAL AND DIELECTRIC PROPERTIES	123
4.1	Introduct	ion	124
4.2	Experime	ntal section	127
	4.2.1	Materials	127
	4.2.2	Processing	128
	4.2.3	Characterization	129
4.3	Results an	nd discussion	132
	4.3.1	Dispersion and morphology	132
	4.3.2	Rheological properties	137
	4.3.3	Low-field dielectric characterization	142
		4.3.3.1 Complex dielectric permittivity	142

	4.3.4	High-field dielectric characterization	151
		4.3.4.1 Short-term breakdown strength	151
		4.3.4.1 Resistance to surface erosion by partial discharges	153
		4.3.4.1 DC electrical conductivity: dependence on nanoparticles	
		networks	155
	4.3.5	Thermal conductivity	159
4.4	Concludin	ig remarks	161
CHAP	TER 5	ARTICLE 4: EVALUATION OF	
		POLYETHYLENE/THERMOPLASTIC ELASTOMER/ZINC	
		OXIDE BLEND NANOCOMPOSITES FOR HIGH VOLTAGE	
		INSULATION APPLICATIONS	163
5.1	Introducti	on	164
5.2	Experimen	ntal	166
	5.2.1	Materials	166
	5.2.2	Processing	167
	5.2.3	Characterization	167
5.3	Results an	d discussion	170
	5.3.1	Scanning electron microscopy (SEM)	170
	5.3.2	Thermodynamic prediction of ZnO nanoparticles localization:	
		Wetting coefficient	174
	5.3.3	Rheological properties	177
	5.3.4	Complex dielectric permittivity	181
	5.3.5	AC short-term breakdown strength	185
	5.3.6	Resistance to surface erosion by partial discharges	
	537	Mechanical properties	189
54	Concludin	g remarks	192
	e chier dan		
CHAP	TER 6	ARTICLE 5: TUNING THE MECHANICAL AND DIELECTRIC	
		PROPERTIES OF CLAY-CONTAINING THERMOPLASTIC	
		ELASTOMER NANOCOMPOSITES	195
6.1	Introducti	on	196
6.2	Processing	and characterization	199
	6.2.1	Materials	199
	6.2.2	Processing	199
	6.2.3	Summary of nanocomposites morphologies	200
6.3	Characteri	ization	
	6.3.1	Mechanical properties	203
	6.3.2	AC short-term breakdown strength	203
6.4	Results an	d discussion	
	641	Mechanical properties	203
	642	AC short-term Breakdown strength	208
6.5	Conclusio	n	
	2011010010		
CHAP	TER 7	DISCUSSION	215
7.1	Nanocom	posites with tailored morphologies	215

XVIII

7.2	SEBS/Cla	y nanocomposites	
7.3	SEBS/ZnO	D nanocomposites	
7.4	PE/SEBS/	ZnO blend nanocomposites	
CONC	CLUSION		
RECO	MMENDA	TIONS	
ANNI	EX I	SUPPORTING ELECTRONIC INFORMATION FOR ARTICLE 1	
ANNE	EX II	SUPPORTING ELECTRONIC INFORMATION FOR ARTICLE 2	241
ANNE	EX III	SUPPORTING ELECTRONIC INFORMATION FOR ARTICLE 3	
APPE	NDIX VIT	Α	
LIST	OF BIBLIC	GRAPHICAL REFERENCES	

LIST OF TABLES

Table 2-1	Properties of Cloisite 20A grade	48
Table 2-2	Summary of SEBS/clay nanocomposites with different structures	60
Table 2-3	Designation of the nanocomposites	61
Table 2-4	Dielectric strengths of SEBS, SEBS_MA and their nanocomposites	73
Table 2-5	Summary of different effects of SEBS/nanoclays morphology on the complex dielectric permittivity and breakdown strength	75
Table 3-1	Properties of SEBS-13, SEBS-20 and SEBS-30	82
Table 3-2	Nomenclature of nanocomposites	85
Table 3-3	Nomenclature, morphology description and morphology sketch of the nanocomposites	96
Table 3-4	VFT fitting parameters corresponding to SEBS-20 based nanocomposites.	.104
Table 3-5	Dielectric strength of SEBS-20 based nanocomposites at 60 °C and 80 °C	.105
Table 3-6	VFT fitting parameters corresponding to SEBS-13 and SEBS-30 based nanocomposites.	.109
Table 3-7	VFT fitting parameters corresponding to SEBS-30-MA-20A-oriented nanocomposite	.110
Table 3-8	Dielectric strength corresponding to SEBS-30-20A vs. SEBS-30-MA-20A nanocomposites prepared by extrusion	.110
Table 3-9	Tgs of both PS and PEB blocks in different samples of SEBS based nanocomposites	.114
Table 3-10	Interfacial Tgi, bound fraction and interfacial layer thickness in SEBS-30 based nanocomposites	.117
Table 3-11	Average dimensions of clay tactoids based on TEM quantitative analysis published by Carastan et al. (Carastan et al., 2014)	.117

v	v
А	А

Table 4-1	Values of real part of complex conductivity at 10 ⁻¹ Hz	.147
Table 4-2	Dissipation factor (tan $\delta = \epsilon''/\epsilon'$) at 66 Hz for different nanocomposites	.150
Table 5-1	Nomenclature and composition of PE/ZnO and PE/SEBS-MA/ZnO nanocomposites	.167
Table 5-2	Total surface tension and Dispersive and Polar components for polymers and ZnO nanoparticles estimated at 160 °C	.176
Table 5-3	Estimated interfacial tension and wetting coefficient values	.177
Table 6-1	Nomenclature and morphology description of SEBS-20 and SEBS-13 based nanocomposites	.202

LIST OF FIGURES

Figure 0-1	Main steps of the PhD project	9
Figure 1-1	Illustration of size effect on interfacial volume: increasing dominance of interfacial areas with decreasing filler sizes	14
Figure 1-2	Polymer composite with an interfacial layer between the filler and the matrix	17
Figure 1-3	Conductive water layer surrounding nanoparticles embedded in a polymer matrix: (a) separate water layers (b) overlapping water shells.	18
Figure 1-4	Multi-core model	19
Figure 1-5	Effect of nanoparticles on the flow of electrical charge carriers in nanocomposites	22
Figure 1-6	Relative impacts of selected nanoparticles on the breakdown strength of polymer insulation matrices	23
Figure 1-7	DC breakdown strength of BN/Epoxy as function of filler size	25
Figure 1-8	Effect of the nanofillers alignment on: (a) the Displacement-Electric field (D-E) loops and (b) the breakdown strength of polyethylene	27
Figure 1-9	Thermal conductivity of BNNT/ PVA nanocomposite as function of the orientation	28
Figure 1-10	TEM images of: a) P3HT/ZnO nanocomposite and b) P3Ht-b-PEO/ZnO nanocomposite	31
Figure 1-11	Morphology of polyethylene: amorphous and crystalline regions	37
Figure 1-12	Schematic description of SEBS block copolymer: (a) without maleic anhydride (MA) and (b) with MA graft attached to PEB block	38
Figure 1-13	Crystal structure of phyllosilicates	41
Figure 1-14	Layered clay structures in nanocomposites	42

XXII

Figure 2-1	TEM images of respectively: (a) SEBS_5wt% nanoclay, (b) SEBS_MA_2.5wt% nanoclay, (c) SEBS_MA_5wt% nanoclay and (d) SEBS_MA_7.5wt% nanoclay (Prepared by extrusion)	1
Figure 2-2	(a) Storage modulus E', (b) Loss modulus E'' and (c) Damping factor (tan δ) at 1 Hz of SEBS vs. SEBS_MA nanocomposites containing 5wt% nanoclay (prepared by extrusion)	2
Figure 2-3	TEM image quantitative analysis of oriented hexagonal SEBS/clay nanocomposites, prepared by extrusion, containing different nanoclay loadings: 2.5, 5 and 7.5 wt%	1
Figure 2-4	(a) Storage modulus E', (b) Loss modulus E'' and (c) Damping factor (tan δ) at 1 Hz of SEBS_MA nanocomposites containing 2.5, 5 and 7.5 wt% nanoclay (prepared by extrusion)	5
Figure 2-5	Radial plot of peak intensities of SEBS vs. SEBS_MA nanocomposites containing 5 wt% nanoclay (prepared by solution): (a) unannealed samples and (b) annealed samples	7
Figure 2-6	Radial plots of SAXS peak intensities as function of time at 200 °C of: (a) unannealed pure SEBS and (b) unannealed SEBS_5wt% nanocomposite prepared by solution)
Figure 2-7	Orientation of electric field in dielectric spectroscopy and AC short-term breakdown measurements: (a) anisotropic nanocomposites prepared by extrusion and (b) isotropic nanocomposites prepared by solution casting	3
Figure 2-8	 (a) Real and (b) imaginary parts of the complex permittivity at room temperature of oriented hexagonal SEBS (type I) and SEBS_MA (type II) nanocomposites containing 2.5, 5 and 7.5 wt% nanoclay	1
Figure 2-9	Real and imaginary parts of the complex dielectric permittivity in the temperature range from 25 to 95 °C of: Pure SEBS (a) and (b), SEBS_5wt%_int_ori_hex (c) and (d) and SEBS_MA_5wt%_exf_ori_hex (e) and (f)	5
Figure 2-10	Real (a) and Imaginary (b) parts of the complex permittivity at room temperature of oriented hexagonal vs. random hexagonal SEBS/clay nanocomposites	3
Figure 2-11	Real (a) and imaginary (b) parts of the complex permittivity of SEBS_5wt%_int_ran_hex nanocomposite with random hexagonal structure in the temperature range from 25 to 95 °C	9

Figure 2-12	Real (a) and imaginary (b) parts of the complex permittivity of SEBS_5wt%_int_ran_lam nanocomposite with random lamellar structure in the temperature range from 25 to 95 °C70
Figure 2-13	Weibull plot of breakdown strength of SEBS vs. SEBS/clay nanocomposites with oriented hexagonal structure
Figure 3-1	Illustrations showing: (a) sheet die extrusion, (b) film blowing with ratio R1= 1 and (c) film blowing with ratio R3= 3 (Coordinate system used as reference is indicated at the left of the figure)
Figure 3-2	Scattering intensity as function of scattering vector q at different temperatures of: (a) SEBS-20-solution and (b) SEB-20-20A-solution nanocomposite, (c) SEBS-13-solution and (d) SEBS-13-20A-solution nanocomposite
Figure 3-3	(a)-(b) 2D SAXS patterns of: (a) SEBS-20-film blowing R1 and (b) SEBS-20-20A-film blowing R1, (c)-(e) TEM images of SEBS 20-20A- film blowing R1 at 3 different magnifications showing imperfect alignment of PS cylinders and clay tactoids: (d) zoom on the section defined by black square in (c), (e) the white square indicates regions featuring perfect alignment of PS cylinders in the extrusion direction while dotted white rectangles indicate other directions of alignment (PS domains were stained with RuO4 (dark phase))
Figure 3-4	 (a)-(b) 2D SAXS patterns of: (a) SEBS-20-film blowing R3 and (b) SEBS-20-20A- film blowing R3 nanocomposite, (c)-(d) TEM images of SEBS-20-film blowing R3 at different magnifications (PS domains (darker phase) were stained with RuO4)
Figure 3-5	Imaginary part of the dielectric permittivity as a function of temperature of: (a) neat SEBS-20-partially oriented R3, (b) SEBS-20-20A-oriented, (c) SEBS-20-20A- partially oriented R1, (d) SEBS-20-20A-partially oriented R3, (e) SEBS-20-20A-isotropic and (f) an example of the fitting corresponding to SEBS-20-20A-partially oriented R3 at 75 °C (Different colors are used to denote the different contributions)
Figure 3-6	Temperature dependence of the relaxation times corresponding to SEBS-20 nanocomposites prepared by different processes: symbols correspond to relaxations times retrieved from the HN function and solid lines correspond to VFT fitting
Figure 3-7	Examples of fitting according to equation (A II.2) in annex II of respectively: (a) SEBS-13-20A-isotropic at 65 °C (ϵ " presented in log scale due to high contribution from charge fluctuation)

XXIV

	and (b) SEBS-30-20A-oriented at 70 °C (Different colors are used to denote the different contributions)	106
Figure 3-8	Temperature dependence of the relaxation times corresponding to SEBS-13 and SEBS-30 nanocomposites prepared by different processes: symbols correspond to relaxation times retrieved from the HN function and solid lines correspond to VFT fitting, plots corresponding to SEBS-20 nanocomposites are presented again for comparison	108
Figure 3-9	Temperature dependence of the relaxation times corresponding to SEBS-30-MA-20A-oriented nanocomposite: symbols correspond to relaxations times retrieved from the HN function and solid lines correspond to VFT fitting (plot corresponding to SEBS-30-20A-oriented nanocomposite is presented again for comparison)	111
Figure 3-10	tan δ of different SEBS-30 materials (prepared by sheet die extrusion) from -100 °C to 130 °C at 1 Hz	113
Figure 3-11	Fitted tan delta curves showing bulk Tgs of PS and PEB blocks as well as Tgi of the interfacial layer (an additional ADS term is considered for the increasing baseline):(a) SEBS-30-oriented and (b) SEBS-30-20A-oriented	115
Figure 3-12	Scheme of the interfacial layer covering the lateral side of clay particles: (a) First approximation: clay tactoid as a solid particle, (b) part of the interfacial layer is between clay galleries	118
Figure 4-1	Schematic description of SEBS block copolymer with MA group grafted to elastomer block ("Polystyrene-block-poly(ethylene-ran-butylene)-block-polystyrene-graft-maleic anhydride,")	128
Figure 4-2	Sketch of the set up for resistance to surface erosion test	131
Figure 4-3	SEM micrographs of: (a) SEBS-5 (arrows indicate the agglomerations) and (b) SEBS-MA-5	133
Figure 4-4	TEM micrographs of SEBS-MA-5 nanocomposite	133
Figure 4-5	FTIR spectra of SEBS-MA-ZnO nanocomposites: (a) in the absorption range 1500-2000 cm ⁻¹ and (b) in the absorption range 3000-3800 cm ⁻¹	135
Figure 4-6	Reaction scheme between maleic anhydride and the surface of the nanoparticles (R: alkyl ammonium salt)	136

Figure 4-7	AFM images of: (a) Neat SEBS-MA, (b) SEBS-MA-1 nanocomposite, (c) Zoom on the region indicated by the square in (b) (showing the same magnification as (a))
Figure 4-8	SAOS measurements of SEBS-ZnO vs. SEBS-MA-ZnO nanocomposites: (a)-(b) storage modulus G' (\bullet) and loss modulus G''(\blacksquare) and (c)-(d) complex viscosity modulus $ \eta^* $ 140
Figure 4-9	Time sweep test at 200 °C, 0.05 rad.s ⁻¹ and 0.5% strain141
Figure 4-10	Complex dielectric permittivity at room temperature of SEBS-MA-ZnO vs. SEBS-ZnO nanocomposites: (a) Real part of permittivity of SEBS-MA-ZnO nanocomposites, (b) Imaginary part of permittivity of SEBS-MA-ZnO nanocomposites, (c) Real part of permittivity of SEBS-ZnO nanocomposites and (d) Imaginary part of permittivity of SEBS-MA-ZnO nanocomposites
Figure 4-11	Imaginary part of the dielectric permittivity as functions of frequency and temperature of: (a) Neat SEBS-MA, (b) Neat SEBS,(c) SEBS-MA-5 nanocomposite and (d) SEBS-5 nanocomposite
Figure 4-12	Imaginary part of electric modulus as functions of frequency and temperature of: (a) SEBS-MA-20wt% ZnO nanocomposite and (b) SEBS-20wt% ZnO nanocomposite
Figure 4-13	Breakdown strength of: (a) SEBS-ZnO and (b) SEBS-MA-ZnO nanocomposites
Figure 4-14	Resistance to surface erosion by partial discharges155
Figure 4-15	 (a) current density vs. DC electric field plots of SEBS-MA-ZnO vs. SEBS-ZnO nanocomposites , (b) DC conductivity of SEBS-MA-ZnO vs. SEBS-ZnO nanocomposites as functions of ZnO loading (wt%) and electric field
Figure 4-16	Thermal conductivity of SEBS-ZnO vs. SEBS-MA-ZnO nanocomposites
Figure 5-1	SEM micrographs of PE/ZnO nanocomposites: (a) unfilled PE, (b) PE/1, (c) PE/5 and (d) PE/10171
Figure 5-2	SEM micrographs of samples treated by solvent extraction: (a) unfilled PE/SEBS-MA, (b) PE/SEBS-MA/1, (c) PE/SEBS-MA/5 and (d) PE/SEBS-MA/10172

XXVI

Figure 5-3	 HRSEM micrographs of PE/SEBS-MA/5 nanocomposite: (a) SEBS-MA not extracted, (b) SEBS-MA extracted, (c) Zoom inside PE region surrounded by blue continuous line in (b): poor ZnO phase, (d) Zoom inside SEBS-MA region surrounded by red dotted line in (b): poor ZnO phase, (d) Zoom inside SEBS-MA region surrounded by red dotted line in (b): rich ZnO phase (Colors are available in the online version) 	174
Figure 5-4	SAOS measurements of PE/ZnO vs. PE/SEBS-MA/ZnO nanocomposites: (a) storage modulus G' and (b) complex viscosity modulus $ \eta^* $ as function of angular frequency ω	180
Figure 5-5	Plots of imaginary permittivity ε'' as function of frequency measured at room temperature corresponding to: (a) PE/ZnO nanocomposites, (b) PE/SEBS-MA/ZnO nanocomposites and (c) comparison with SEBS-MA/ZnO master batches	183
Figure 5-6: Isothe	rmal frequency sweeps of the imaginary permittivity ε'' of PE/5 and PE/SEBS-MA/5 nanocomposites at different temperatures between 25 °C and 100 °C, performed before and after drying: (a) undried PE/5, (b) undried PE/SEBS-MA/5, (c) PE/5 dried and (d) PE/SEBS-MA/5 dried for 5 days at 65 °C	185
Figure 5-7	Weibull probability plots of: (a) PE/ZnO nanocomposites and (b) PE/SEBS-MA/ZnO nanocomposites	187
Figure 5-8	Resistance to surface erosion by partial discharges for PE/ZnO vs. PE/SEBS-MA/ZnO nanocomposites	189
Figure 5-9	Typical stress-strain curves of selected PE/ZnO vs. PE/SEBS-MA/ZnO materials	190
Figure 5-10	Mechanical properties of PE/ZnO vs. PE/SEBS-MA/ZnO nanocomposites: (a) Elastic Young's modulus E, (b) Elongation at break EB, (c) Ultimate tensile strength UTS and (d) Elongation at ultimate tensile strength EUTS (error bars stand for standard deviation)	192
Figure 6-1	Cartoons illustrating: sheet-die extrusion (a), films blowing extrusion using a tubular die with two different ratios: R1 (b) and R3 (c) (the coordinate system is used as a reference for the following characterizations and discussions)	200
Figure 6-2	Characteristic stress-strain plots of SEBS-20 based materials	207

Figure 6-3	Characteristic stress-strain plots of SEBS-13 based materials	207
Figure 6-4	Young's modulus of respectively SEBS-20 and SEBS-13 sets of materials	208
Figure 6-5	Breakdown strength of SEBS-20 based nanocomposites	212
Figure 6-6	Breakdown strength of SEBS-13 based nanocomposites	212

LIST OF ABREVIATIONS

AC	Alternating current
ADS equation	Asymmetric Double Sigmoid equation
AFM	Atomic force microscopy
Al ₂ O ₃	Aluminium oxide or Alumina
BDS	Broadband dielectric spectroscopy
BN	Boron nitride
BNNS	Boron nitride nanosheets
BNNT	Boron nitride nanotubes
CNT	Carbon nanotubes
DC	Direct current
DMA	Dynamic mechanical analysis
DSC	Differential scanning calorimetry
E	Elastic modulus
EB	Elongation at break
EUTS	Elongation at ultimate tensile strength
FTIR	Fourier transformed infrared spectroscopy
GO	Graphene oxide
HDPE	High density polyethylene
HRSEM	High resolution scanning electron microscopy
HV	High voltage
HVDC	High voltage direct current
LDPE	Low density polyethylene
MA	Maleic anhydride
MgO	Magnesium oxide
MMT	Montmorillonite
MWS polarization	Maxwell Wagner Sillars polarization
PE	Polyethylene
PEB	Poly(ethylene-co-butylene)

XXX

PEO	Polyethylene oxide
PMMA	Poly(methyl methacrylate)
POE	Polyolefin elastomer
POSS	Polyhedral Oligomeric Silsesquioxane
PP	Polypropylene
PS	Polystyrene
SAOS	Small amplitude oscillatory shear
SAXS	Small angle X-ray scattering
SBR	Styrene-Butadiene rubber
SEBS	Polystyrene-b-poly(ethylene-co-butylene)-b-polystyrene
SiC	Silicon Carbide
Si ₃ N ₄	Silicon Nitride
SiO ₂	Silicon dioxide or Silica
TEM	Transmission electron microscopy
TiO ₂	Titanium Oxide
UTS	Ultimate tensile strength
VFT equation	Vogel Fulcher Tammann equation
XLPE	Crosslinked Polyethylene
ZnO	Zinc Oxide

LIST OF SYMBOLS

F	Fragility index
G'	Storage modulus
G''	Loss modulus
t	Thickness
Т	Temperature
tanδ	Loss tangent or dissipation factor in BDS measurements (Chapters 4 and 5) or DMA measurements (Chapters 2 and 3)
Πg	Glass Transition
T_{gi}	Interfacial Glass Transition
Wa	Wetting coefficient
20A	Organically modified clay Cloisite 20A
δ	Surface tension
бр	Polar component of the surface tension
δd	Dispersive component of the surface tension
δ12	Interfacial tension
Δε	Dielectric strength
3	Complex dielectric permittivity
٤'	Real part of dielectric permittivity
ε''	Imaginary part of dielectric permittivity
η*	Complex viscosity
λ	Thermal conductivity
σ	Conductivity modulus
σ'	Real part of complex conductivity
τ	Relaxation time
ω	Angular frequency

INTRODUCTION

0.1 Context of the research

Existing electrical insulating polymers currently used in high voltage (HV) apparatus are subjected constantly to increasing electrical and thermal stresses due to growing operating voltages and power ratings required by the current market demand and the emergence of new techniques suitable for long-distance electric power transmission such as high voltage direct current (HVDC) cables (Astrom & Lescale, 2006; Ghorbani, Jeroense, Olsson, & Saltzer, 2014). For example, in the case of extruded HVDC cables, the maximum transmission voltage level is currently equal to 525 kV. By 2030, the objective is to increase this level up to 1 MV while maintaining the present cables geometry unchanged (Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016). To meet these pressing requirements, there has been a critical need for the development of new reliable and cost-effective HV insulating materials with improved dielectric and thermal performance. The ensuing research, during the last decades, led to the introduction of nanocomposites as a new conceptual design for dielectric materials used in HV insulation.

The new generation of nanocomposite dielectrics, known also as nanodielectrics (David & Fréchette, 2013; Michel F Fréchette, Trudeau, Alamdar, & Boily, 2004; Michel F Fréchette et al., 2010; Tanaka & Imai, 2013) featured several promising properties, albeit depending on the nanoparticles challenging dispersion, which unfortunately compromises their reliability and reproducibility (Balazs, Emrick, & Russell, 2006). In fact, inorganic nanoparticles typically form submicrometric and micrometric agglomerations, during nanocomposites processing, due to their incompatibility with the organic polymer hosts, which supress or reduce potential benefits associated with their nanoscopic dimensions. In addition to the agglomerations issue, the spatial distribution of nanoparticles as well as their orientation and selective location are considered as major challenges of morphology design that could affect drastically the functional properties of nanocomposites in general and those of nanodielectrics in specific, such as the breakdown strength, dielectric loss, electrical

conductivity, thermal conductivity and resistance to surface erosion, to name a few (David & Fréchette, 2013; Roy et al., 2005). Therefore, recently researchers began seeking the possibilities to design nanocomposites with prescribed morphologies that optimize their functional properties (Balazs et al., 2006; Richard A Vaia & Maguire, 2006).

Several techniques have been tested to avoid nanoparticles agglomerations, through the mediation of the interaction between the nanoparticles and polymer chains, such as surface functionalization of nanoparticles (Balazs et al., 2006; Huang et al., 2013; D. Ma et al., 2005; Polizos, Tomer, Manias, & Randall, 2010; Virtanen et al., 2014), in situ synthesis of nanoparticles (Dirix, Bastiaansen, Caseri, & Smith, 1999) or in situ polymerization of the polymer matrix (Zapata et al., 2011). Besides, some attempts to orient nanoparticles and tailor their spatial distribution were also performed (Richard A Vaia & Maguire, 2006). The current findings are partially successful in solving the issue of agglomerations. However, they are far from governing all different aspects involved within the concept of tailored dispersion. Moreover, a large-scale production based on the current techniques is often inappropriate due to several factors including increased complexity and cost, limited yield, poor efficiency and/or potential drawbacks induced by some modifiers on the dielectric properties.

A new approach is nowadays considered as a promising technique to overcome the hurdles related to the control of dispersion. It consists in using template matrices in order to guide the dispersion, the spatial distribution as well as the orientation of nanoparticles (Balazs et al., 2006). In particular, the template can be a multiphase polymer system with controllable morphology, such as block copolymers or immiscible polymer blends, which offers various possibilities to disperse nanoparticles based on its architecture and the affinity of the studied nanoparticles to one phase or another. Block copolymers are composed of covalently bonded blocks that exhibit nanoscale dimensions and self-assemble in different morphologies depending on several factors including the molecular weight of the blocks, their affinity and the ratio between them (Helfand, 1975; Sarkar & Alexandridis, 2015). Immiscible polymer

blends are mixtures of at least two polymers which cannot be homogenized. They feature several characteristic morphologies as well, of microscale dimensions though.

Owing to their ordered structures, these classes of materials might lead nanoparticles orientation, location and dispersion in the aim of optimizing the ultimate performance of the nanocomposites (Balazs et al., 2006; Carastan, Amurin, Craievich, do Carmo Gonçalves, & Demarquette, 2013; Carastan, Amurin, Craievich, Gonçalves, & Demarquette, 2014; Kao, Thorkelsson, Bai, Rancatore, & Xu, 2013; Park, Yoon, & Thomas, 2003; Sarkar & Alexandridis, 2015). Furthermore, they could exhibit useful physical properties depending on the choice of the blocks or constitutive phases of the blend. In particular, styrenic thermoplastic elastomers are block copolymers, composed of a combination of styrenic hard blocks and elastomeric soft blocks, which display excellent mechanical properties (Holden, Kricheldorf, & Quirk, 2004), electromechanical coupling (B. Kim et al., 2011) and good resistance to water treeing (Liu, Mhetar, & Freestone, 2011; Z. Ma, Jiang, Wang, & Yang, 2010). Among styrenic thermoplastic elastomers, polystyrene-b-poly(ethylene-co-butylene)*b*-polystyrene SEBS is an interesting candidate commercially available and widely used in several industries as a compatibilizer. It consists of two polystyrene PS endblocks dispersed in the form of cylinders, lamellae or spheres within a hydrogenated Polybutadiene midblock matrix, known as poly(ethylene-co-butylene) PEB. The latter has a structure equivalent to the structure of polyethylene PE in its amorphous form which ensures good compatibility with polyolefins (Agari, Ueda, & Nagai, 1993).

0.2 Motivation and Objectives

This PhD project is part of a bigger research project conducted in collaboration between École de Technologie Supérieure (ÉTS) and Hydro-Québec Research Institute (IREQ) and involving several graduate students, in the aim of producing novel nanodielectrics for HV insulation systems and tailoring their functional properties in correlation with their microstructures. Besides, a better understanding of the underlying structure-property relationships governing the dielectric and thermal performance of nanodielectrics is expected. Within this framework, the main goal of this thesis was to design advanced SEBS

block copolymer nanocomposites with tunable nanoparticles dispersion that take advantage from SEBS architecture as a template or guiding matrix (Carastan et al., 2013; Carastan et al., 2014). Two types of nanoparticles, respectively: clay nanoplatelets and Zinc oxide (ZnO) nanospheres featuring distinguished properties and shapes were used. The effects of their patterned arrangement in SEBS, going beyond single-particle dispersion limits, on the dielectric performance of the insulating matrix were evaluated. The resulting block copolymer nanocomposites were subsequently used as master batches and blended with polyethylene (PE), a polyolefin widely used in its crosslinked form as insulating material in the current state-of-the-art extruded HV cables. The dispersion and the selective location of the nanoparticles in these particular blends were emphasized. Ultimately, their dielectric properties were compared to those of PE as a reference insulating material.

0.3 Industrial impact

From an industrial point of view, the optimization of nanoparticles spatial distribution using a block copolymer template is expected to result in an optimized use of nanoparticles properties at lower loadings and to induce as a consequence an effective cost reduction. Hence, the method might be generalized to different types of nanoparticles and versatile processing techniques (solution casting, extrusion, injection...) to evaluate its scalability. Furthermore, the as-prepared nanocomposites are expected to exhibit good compatibility with polyolefins, good resistance to water treeing and excellent electromechanical coupling, which are intrinsic properties of the styrenic thermoplastic elastomer matrix. They might be used as final materials or as master batches to blend with polyolefins for larger spectra of dielectric applications including but not limited to HV insulation, capacitors and dielectric elastomer actuators.

0.4 Approach and methodology

This PhD project was divided in three main parts that are summarized in Figure 0-1.

- SEBS/Clay nanocomposites: In addition to their reduced cost, nanoclays were selected due to their anisotropy and confirmed electron scattering effect especially
upon exfoliation and orientation (David et al., 2013; Tomer, Polizos, Randall, & Manias, 2011). Moreover, successful orientation and localization of these nanoplatelets in SEBS was recently reported (Carastan et al., 2013; Carastan et al., 2014). In this step, the effect of nanoclays and PS domains orientations as well as the effect of nanoclays selective location on the dielectric performance of the nanocomposites were emphasized. Different morphologies, i.e. isotropic vs. oriented vs. partially oriented, were obtained through the variation of the processing technique and the ratio between the blocks of SEBS. The location of nanoclays either in the hard PS block or soft PEB block was tuned through the presence or absence of maleic anhydride (MA) group grafted onto the rubber block. The latter exhibits good affinity to the functional groups attached to the nanoparticles surface. Afterward, changes in the dielectric and mechanical properties depending on the configuration were evaluated. Moreover, the interfacial dynamics were investigated as function of morphology. The results of this part led to the publication of two articles (Chapter 2 and Chapter 3) and one conference paper. A third manuscript (submitted), related to this part, is also reported in Chapter 6.

- SEBS/ZnO nanocomposites: In a second step, nanocomposites of SEBS were prepared with a different type of nanoparticles: functionalized ZnO nanospheres that are commercially available. ZnO nanoparticles were selected for their relatively high thermal conductivity and excellent UV light shielding capability as well as their confirmed ability to reduce space charge accumulation in polymer matrices, which is a key requirement for HVDC cables (Huang, Jiang, & Tanaka, 2011; Mazzanti & Marzinotto, 2013; Tian, Lei, Wang, & Wang, 2011, 2012). Only isotropic morphology was investigated in this part but nanoparticles distribution and selective location was modulated using the MA graft. The effect of morphology on the dielectric, electrical and thermal properties was emphasized at this step. A correlation with rheological properties was also established. The findings of this part were reported in one submitted article (Chapter 4) and two conference papers.
- Blend of Polyethylene and SEBS/ZnO nanocomposites: In a third step, block copolymer nanocomposites containing ZnO nanoparticles studied in the previous step

were used as master batches and blended with polyethylene. These nanocomposites were selected due to their good potential for HV insulation and HVDC cables insulation in particular. The morphology of the resulting blend nanocomposites and the selective location of ZnO nanoparticles in this blend were investigated. The effect of the obtained morphologies on the dielectric and mechanical properties was evaluated. The main conclusions of this part were reported in one submitted article (Chapter 5) and two conference papers.

The microstructures of all the fabricated nanocomposites were investigated by means of Scanning Electron Microscopy, Transmission Electron Microscopy and Atomic Force Microscopy. The morphology of the block copolymer and the arrangements of both the block copolymer domains and the nanoparticles were assessed by means of Small Angle X-Ray Scattering. Besides, the dielectric performance was evaluated by means of: dielectric broadband spectroscopy, AC breakdown strength, resistance to surface erosion by partial discharges, electrical conductivity and thermal conductivity measurements. Tensile properties and linear viscoelastic properties were evaluated in some cases as well, to achieve better understanding of the correlation between the microstructure and dielectric properties.



Figure 0-1: Main steps of the PhD project

0.5 Organization of the PhD thesis

This thesis is divided in 7 chapters. Following the introduction, a brief literature review regarding nanodielectrics and the concept of tailored nanoparticles dispersion is presented. Then, in each one of the next 5 chapters, an article related to a part of the above-mentioned parts of the project is presented. Finally, in the last chapter, a general discussion is provided and followed by conclusions and recommendations for future work.

In chapter 2, the paper untitled "Styrenic block copolymer-based nanocomposites: implications of nanostructuration and nanofiller tailored dispersion on the dielectric properties" published in *Polymer* is presented. This article is based on results from the first part of the project regarding block copolymer nanocomposites containing clay nanoparticles. It emphasizes the effect of orientation of nanoparticles and morphology of the block copolymer on dielectric properties namely dielectric loss and short-term breakdown strength.

Chapter 3 presents the paper '' Interfacial molecular dynamics of styrenic block copolymerbased nanocomposites with controlled spatial distribution'' published in *Polymer* as well. This article is also related to the first part of this project. It highlights a correlation between the studied nanocomposite morphologies and the interfacial dynamics at nanoclay-block copolymer interfaces. An extended range of complex morphologies is achieved through the variation of the blocks proportions and the processing techniques as well as the use of MA graft.

In chapter 4, the paper untitled "Thermoplastic elastomer nanocomposites with controlled nanoparticles dispersion for HV insulation systems: correlation between rheological, thermal, electrical and dielectric properties" submitted to *European Polymer Journal* is presented. This study summarizes the results of the second part of this project related to block copolymer nanocomposites containing Zinc Oxide nanoparticles. In particular, a correlation between the dielectric, electrical, thermal and rheological properties and the formation of a network between ZnO nanoparticles and the block copolymer chains is established.

In chapter 5, the paper untitled "Evaluation of polyethylene/thermoplastic elastomer/zinc oxide blend nanocomposites for high voltage insulation applications" submitted to *Polymer* is presented. This paper is based on the conclusions found in the third part of this project dealing with the use of block copolymer nanocomposites with controlled morphology as a master batch to blend with polyethylene. It focuses on the potential of these blend nanocomposites with tuned nanoparticles dispersion for HV insulation applications.

In chapter 6, the study "Tuning mechanical and dielectric strength of clay-containing thermoplastic elastomer nanocomposites" is reported. The manuscript is submitted to *Polymer Engineering and Science*. This work is also based on the findings of the first part of the project. It presents an evaluation of the dielectric breakdown strength and mechanical strength of the nanocomposites with different morphologies studied in chapter 3.

In addition to the main chapters, three annexes are also included at the end of the thesis. They provide supplementary information regarding the articles reported respectively in chapters 2, 3 and 4.

For more details regarding the fundamentals of dielectric spectroscopy and the physical properties of solid insulations, good reviews can be found respectively in references (Kremer & Schönhals, 2012; C. C. Ku & Liepins, 1987) and references (Blythe & Bloor, 2005; Dissado & Fothergill, 1992; C. C. Ku & Liepins, 1987). For block copolymers (including thermoplastic elastomers) and polymer blends, respectively references (Holden et al., 2004; Mai & Eisenberg, 2012) and reference (Paul & Bucknall, 2000) can be consulted.

CHAPTER 1

NANOCOMPOSITES WITH TUNED MORPHOLOGY FOR HV INSULATION

Polymer nanocomposites have shown promise recently for the development of advanced materials intended for use in a large spectrum of applications including dielectric applications and HV insulation. Their established and foreseen superior properties are often correlated with the formation of polymer/nanoparticle interfacial layers featuring distinct behavior compared to the bulk polymer host.

In this chapter, the concept of polymer nanocomposites and nanodielectrics is reviewed, with a focus on the impact of the interfacial region. Besides, a summary of the most common types of nanoparticles investigated, during the last decades, in HV insulation is presented. The influence of several parameters such as the nanoparticles shape, size, orientation and quality of dispersion on the efficiency of their use in nanodielectrics is investigated as well. Then, the concept of controlled nanocomposites morphology is emphasized. In particular, the effect of tuning nanoparticles dispersion in polymer blends and block copolymers, used as template polymer matrices, on their dielectric performance is highlighted. Finally, a review of the polymers and nanoparticles studied in the present project is reported.

1.1 Definition of nanocomposites and nanodielectrics

1.1.1 Polymer nanocomposites definition

Polymer nanocomposites are polymeric materials containing particles, with at least one dimension below 100 nm. Polymer nanocomposites differ from conventional microcomposites in two major aspects:

- Particles size in the nanometer range: this feature results in significantly higher specific surface area, i.e. surface area per mass of a material, compared to similar masses of larger scale particles. To illustrate better the tremendous effect of

nanoparticles size on the specific surface area, one can compare the total surface area of a single solid cube of volume equal to 1 cm^3 with the same volume made up of 1 nanometer-sized cubes; each of them having a surface area equal to 6 nm^2 . The latter contains 10^{21} nanoscale cubes with a total surface area equal to 6000 m^2 compared to only 6 cm^2 for the single solid cube ("What is so special about the nanoscale?,").

- Small fraction of fillers compared to conventional composites: This can be also considered a consequence of the nanometric size. In fact, when incorporated in a polymer matrix, the large surface area of nanoparticles constitutes an interaction zone that comes into contact with the surrounding polymer and favorably affects its properties. As the particle size decreases, the specific surface area increases and the interfacial region becomes more prominent for the same amount of particles, as illustrated in Figure 1-1. As a consequence, lower loading of nanoparticles is needed, compared to microparticles, to achieve equivalent or higher properties.



Figure 1-1: Illustration of size effect on interfacial volume: increasing dominance of interfacial areas with decreasing filler sizes Taken from (Andritsch, Kochetov, Gebrekiros, Morshuis, & Smit, 2010)

1.1.2 Definition of nanodielectrics

A dielectric material is an electrical insulating material that has the ability to polarize in the presence of an electric field. When an electric field is applied, a dielectric responds by distorting slightly its charges from their equilibrium position and redistributing them in a

more organized form: positive charges are shifted toward the field and negative charges in the opposite direction. The resulting dipoles orient parallel to the electric field with opposite polarity and create an internal field that reduces the overall field within the dielectric. There are three basic groups of dielectrics: solid, fluid and gaseous dielectrics. In particular, electrically insulating polymers belong to the group of solid dielectrics. Comprehensive details regarding insulating polymers, their electrical properties and the polarization phenomena can be found in (Blythe & Bloor, 2005; Dissado & Fothergill, 1992; Kremer & Schönhals, 2012; C. C. Ku & Liepins, 1987).

The term nanometric dielectric or nanodielectric for short was proposed for the first time in 2001 by Fréchette et al. (M. F. Fréchette, Trudeau, Alamdari, & Boily, 2001) as a "multicomponent dielectric processing nanostructures, the presence of which lead to changes in one or several of its dielectric properties". This notion includes nanostructured ceramics and polymer nanocomposites. In particular, in last decades, an important and growing fraction of nanodielectrics has been developed based on the second group, namely polymer nanocomposites (M. F. Fréchette et al., 2001). It is now well established that the addition of specific nanoparticles to regular insulating polymers used as solid dielectrics can improve their performance in different aspects. Hence, the as-developed nanodielectrics have gained attraction in different dielectric applications including HV insulation, high energy density capacitors and dielectric elastomer actuators (Camargo, Satyanarayana, & Wypych, 2009; David & Fréchette, 2013; Michel F Fréchette et al., 2010; Pleşa, Noțingher, Schlögl, Sumereder, & Muhr, 2016; Tanaka, Montanari, & Mulhaupt, 2004). However, it is worth noting that the extent of the achieved improvement is strongly dependent on the properties of the interfacial area, which is actually the case for all nanocomposites. The interface region depends not only on the nanoparticles size and specific surface area but also on their dispersion/spatial distribution and their compatibility with the polymer host.

1.1.3 Theories and models regarding the interfacial region

Few properties of composite materials, such as the mechanical modulus, can be roughly estimated from the inherent properties of the polymer matrix and the added particles

respectively, using basic approaches such as the rule of mixtures (Bartczak, Argon, Cohen, & Weinberg, 1999; Selvin, Kuruvilla, & Sabu, 2004; Zare & Garmabi, 2014). However, the majority of them, including electrical and thermal properties, are more difficult to predict because they depend strongly on polymer/particles interactions in addition to the usual factors. In nanocomposites, this phenomenon is even more pronounced due to the presence of an important interfacial region. In this context, it was repeatedly observed that effective properties of nanocomposites are often far from the predictions of basic models and that the achieved/potential enhancements are rather governed by the properties of the interfacial region (Heid, 2015; Lewis, 2005; Nelson & Hu, 2005; Rittigstein, Priestley, Broadbelt, & Torkelson, 2007; Smith, Liang, Landry, Nelson, & Schadler, 2008; Zare & Garmabi, 2014). Therefore, several models have been proposed to describe the interaction zone in nanocomposites, on the basis of chemical and electrical analysis (Heid, 2015; Kremer & Schönhals, 2012; Pitsa & Danikas, 2011; Ioana Preda, Castellon, Frechette, & Agnel, 2014; Tanaka, Kozako, Fuse, & Ohki, 2005; Zou, Fothergill, & Rowe, 2007), in order to understand their electrical and thermal properties. These models based on different approaches and assumptions can be more or less accurate in describing the experimental results, depending on the property and the system studied. But, overall there is a general consensus on the great role of the interface region. One basic theoretical model that was initially developed is the interlayer model treating the case of a matrix-inclusion system with an interfacial layer (Kremer & Schönhals, 2012). Later, several models have been proposed such as the multi-core model proposed by Tanaka et al. (Tanaka et al., 2005) which suggests that the interfacial zone is divided in several layers featuring different physical properties. These two models will be reviewed briefly in the following sections.

1.1.3.1 Interlayer model

A polymer composite with an interfacial layer between the filler and the matrix, as illustrated in Figure 1-2, can be modeled using the asymmetrical effective medium approach. The analytical solution, also known as the interlayer model, can be used to describe several macroscopic properties including the complex dielectric permittivity (Kremer & Schönhals, 2012).



Figure 1-2: Polymer composite with an interfacial layer between the filler and the matrix Taken from (Kremer & Schönhals, 2012)

In the case of dielectrics and nanodielectrics, this model can be particularly interesting to describe the behavior of dielectrics and nanodielectrics containing water sensitive particles such as metal oxides. In fact, humidity absorption often leads to detrimental effects on the dielectric behavior of the polymer matrix. In particular, simultaneous increase of the dielectric constant and dielectric loss are commonly reported since water has relatively high dielectric constant and high electrical conductivity. The increase of the dielectric loss is dominant at low frequencies and attributed to quasi DC conductivity increase. This behavior is explained by the fact that absorbed water resides in the interfacial region leading to the formation of a conductive interlayer between the particles and the polymer matrix. In addition to the increase of conductivity and dielectric loss, this conductive interlayer gives rise to an interfacial loss process at low frequencies. Furthermore, in nanocomposites, water shells around the nanoparticles have higher probability to overlap with increasing concentrations as illustrated in Figure 1-3 due to the increased interfacial area and reduced interparticle distance, providing conduction paths for charge carriers (Kremer & Schönhals, 2012; Zou et al., 2007).



Figure 1-3: Conductive water layer surrounding nanoparticles embedded in a polymer matrix: (a) separate water layers (b) overlapping water shells Adapted from (Zou et al., 2007)

1.1.3.2 Multi-core model

According to this model, the interfacial layer is composed, on the basis of chemical and electrical analysis, of 3 different physical layers overlapped by an electric diffuse layer, as illustrated in Figure 1-4 (Tanaka et al., 2005). Each of these layers has a distinguished physical, chemical and/or electrical behavior, as following:

- **The bonded layer:** this first layer is tightly bonded to the surface of nanoparticles by coupling agents.
- **The bound layer:** this second layer consists of polymer chains strongly bound to the first layer and the surface of the nanoparticles. The thickness of the bound fraction, in the range from 2 to 9 nm, is increasing with the strength of polymer-nanoparticle interaction. The mobility of chains belonging to this region is particularly altered but the polymer conformation, local density and crystallinity are also subjected to changes, as compared to the bulk polymer.
- **The loose layer:** this third layer is loosely interacting with the second layer but still affected chemically by its behavior and expected to exhibit different behavior from the bulk. Its thickness is up to several tens of nanometers.

- The diffuse electric layer: Charge injection from nanoparticles to polymer matrix often occurs resulting in the formation of charge distribution layer. This diffuse layer is superposed to the 3 physical layers explained above.

Moreover, the overlapping of multiple interaction zones corresponding to several nanoparticles might result in collaborative effects and further complication of the resulting macroscopic behavior. Models treating this situation were also investigated (Ioana Preda et al., 2014).



Figure 1-4: Multi-core model Taken from (Tanaka et al., 2005)

1.1.4 Nanoparticles vs. Microparticles in polymer based insulations

The performance of electrically insulating polymers is reflected by their resistance to electrical degradation when placed in operational conditions. This degradation is a complex phenomenon that depends on a multitude of interdependent parameters such as dielectric losses, electrical conductivity, resistance to space charge accumulation as well as the thermal

and mechanical properties (Blythe & Bloor, 2005; Dissado & Fothergill, 1992). Although it is a long-term process, the susceptibility of polymers to electrical degradation might be reasonably evaluated by short-term measurements such as the short-term breakdown strength or mid-term measurements such as the resistance to surface erosion by partial discharges. Longer-term measurements such as voltage endurance and resistance to electrical treeing remain the most trustable tools but unfortunately time consuming (Tian et al., 2012). An exhaustive review of all the notions covered by the subject of electrical degradation of polymers might be found in (Blythe & Bloor, 2005; Dissado & Fothergill, 1992; C. C. Ku & Liepins, 1987).

To overcome the limitations faced by the existing polymer insulating materials, both microparticles and nanoparticles with desired intrinsic properties were investigated. Chronologically, microparticles raised interest first. In most of the cases, they resulted in enhanced heat dissipation and resistance to surface erosion (M. Frechette et al., 2012; Huang et al., 2011; Iyer, Gorur, & Krivda, 2012; Kochetov, Andritsch, Morshuis, & Smit, 2010; Tanaka, Matsuo, & Uchida, 2008), unfortunately at the cost of reduced dielectric and mechanical strengths (Fleming, Ammala, Casey, & Lang, 2008; Z. Li, Okamoto, Ohki, & Tanaka, 2010). In fact, inorganic microparticles incorporated in a polymer matrix result in the formation of defects and voids of micrometric size in the regions surrounding the particles. These defects constitute zones of reduced mechanical strength and increased electrical stress in addition to the distortion of the electric field which arises from the mismatch in permittivities or conductivities between the polymer and the inorganic filler. The increase of the electrical stress in these weak zones leads to initiation and propagation of electrical trees, i.e. interconnected channels which are generally filled with gas, through the polymer matrix and early dielectric failure (Blythe & Bloor, 2005; Danikas & Tanaka, 2009; Dissado & Fothergill, 1992; C. C. Ku & Liepins, 1987; Z. Li et al., 2010).

Progressively, nanoparticles replaced microparticles in composites in order to avoid the abovementioned issues. Given their excellent intrinsic properties, their nanoscale dimensions and their high interfacial area, nanoparticles were reported to improve the overall dielectric

performance without sacrificing a specific property (Andritsch, 2010; Camargo et al., 2009; Danikas & Tanaka, 2009; David & Fréchette, 2013; Michel F Fréchette et al., 2010; Heid, 2015; Pleşa et al., 2016; Tanaka et al., 2004). In fact, when nanoparticles are well dispersed, the density and the size of defects are reduced. In addition, the distortion of the electric field caused by the difference in conductivities or permittivities between the polymer and the inorganic filler is less pronounced due to the small size of nanoparticles leading to more homogeneous electric field distribution over the nanocomposite. Furthermore, nanoparticles act as barriers resisting the penetration and propagation of electrical trees in the bulk of the polymer insulation by the formation of traps to charge carriers (Danikas & Tanaka, 2009; Z. Li et al., 2010; Tian et al., 2011, 2012). As a consequence, they induce higher breakdown strength, resistance to surface erosion and resistance to electrical treeing while reducing space charge accumulation and maintaining low dielectric losses (Andritsch, 2010; Danikas & Tanaka, 2009; David & Fréchette, 2013; Michel F Fréchette et al., 2010; Heid, 2015; Z. Li et al., 2010; Pleşa et al., 2016; Tanaka et al., 2004; Tiemblo et al., 2008). In this context, Danikas et al. (Danikas & Tanaka, 2009) suggested that, in the presence of nanoparticles impeding the flow of electrical charge carriers, three possible directions of electrical paths can be considered as shown in Figure 1-5:

- Propagation of the electrical tree through the polymer without contacting the nanoparticles (Figure 1-5(a)).
- Propagation of the electrical tree through the polymer contacting the nanoparticles but without penetrating them (Figure 1-5(b)).
- Propagation of the electrical tree through the polymer contacting the nanoparticles and circumventing them; the tree structure will grow partially at the polymernanoparticle interface (Figure 1-5(c)).

In all these three cases, the electrical tree path is extended resulting in an increase of the time required to produce dielectric failure and ultimately an increase of the lifetime of the insulating material.



Figure 1-5: Effect of nanoparticles on the flow of electrical charge carriers in nanocomposites Replotted from (Danikas & Tanaka, 2009)

1.1.5 Review of nanoparticles used in HV insulation systems

Several nanoparticles, mostly of spherical shape, such as Alumina (Al₂O₃), Silica (SiO₂), Zinc Oxide (ZnO) and Titanium Oxide (TiO₂) were first studied as nanofillers for materials used in HV insulation systems. Later, an increasing interest has been given to anisotropic nanoparticles of 1D and 2D dimensions to achieve more advanced nanodielectrics, due to their outstanding properties compared to their isotropic pairs. In the following paragraphs, a review of the different types of nanoparticles that have been used in HV insulation is presented. Furthermore, a classification of these nanocharges considering their chemical nature, their geometry as well as their dielectric and thermal properties is adopted.

1.1.5.1 Metal oxide nanoparticles

Metal oxide nanoparticles have been extensively investigated in HV insulation materials. Being either insulating such as Al₂O₃, SiO₂ and magnesium oxide (MgO) or semiconductive such as TiO₂ and ZnO, these nanoparticles exhibit in general values of electrical conductivity higher than those of insulating polymers such as polyethylene. When added in low weight fractions (< 5wt%) to polymer matrices, some metal oxide nanoparticles, mainly SiO₂, MgO and ZnO were reported to improve the breakdown strength and voltage endurance and more recently to supress space charge accumulation in the insulation matrix. These effects are explained by the formation of trapping sites that reduce the mobility of charge carriers in the polymer matrix including electrons, holes and ionic species, which results in an overall reduction of the insulation conductivity (David & Fréchette, 2013; Fleming et al., 2008; Hayase et al., 2006; Ju et al., 2014; Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016; Tian et al., 2011, 2012; Tian et al., 2015). For instance, Figure 1-6 reports a summary of the achieved improvements of breakdown strength in various insulating polymer matrices induced by different concentrations of metal oxide nanoparticles (David & Fréchette, 2013).

Before their use in nanocomposites, metal oxide nanoparticles are often coated or functionalized with different groups such as silane coupling agents in order to reduce their hydrophilic character and enable their proper distribution in hydrophobic insulating polymer matrices (Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016).



Figure 1-6: Relative impacts of selected nanoparticles on the breakdown strength of polymer insulation matrices Taken from (David & Fréchette, 2013)

1.1.5.2 Nitride and Carbide nanoparticles

Several nitride and carbide nanoparticles such as silicon nitride (Si₃N₄), aluminium nitride (AlN), boron nitride (BN) and silicon carbide (SiC) have been investigated in HV insulation systems and HV accessories, particularly for their high thermal conductivities going up to few hundreds W.m⁻¹.K⁻¹ (Huang et al., 2011). Among them, BN presents the advantage of high thermal conductivity combined with low dielectric permittivity and excellent electrical resistivity and thermal stability. Due to these attractive properties, BN nanoparticles and nanoflakes have gained increasing interest recently (Golberg et al., 2010; Heid, 2015; Song et al., 2012; Zhi et al., 2009).

1.1.5.3 Anisotropic nanoparticles

Layered and elongated nanoparticles such as nanoclays (David et al., 2013; Tomer et al., 2011; B Zazoum, E David, & Anh Dung Ngô, 2014), BN nanotubes and nanosheets (Golberg et al., 2010; Heid, 2015; Song et al., 2012; Terao et al., 2010; Zhi et al., 2009) as well as graphene oxide (GO) and graphene (Fabiani, Mancinelli, Vanga-Bouanga, Fréchette, & Castellon, 2016; MF Fréchette, Vanga-Bouanga, Fabiani, Castellon, & Diaham, 2015; Gaska, Xu, Gubanski, & Kádár, 2017; Ghosh et al., 2008) have been also investigated in HV insulation. These anisotropic nanoparticles usually feature outstanding properties in preferential directions depending on their shape. In particular, nanoclays were reported to induce a beneficial effect on breakdown strength in addition to their support to mechanical strength while BN anisotropic nanoparticles input was mainly on the thermal conductivity and resistance to surface erosion. Regarding graphene oxide and graphene, it was demonstrated recently that their incorporation in small quantities (<0.1 wt%) to polymer matrix reduces space charge accumulation (Y. J. Kim et al., 2013) in addition to improvements in mechanical and thermal properties. More details regarding improvements induced by anisotropic particles, mainly nanoclays and BN are discussed in the following sections regarding the effects of nanoparticles shape and orientation on properties of nanocomposites candidates for HV insulation.

1.1.6 Parameters affecting the role of nanoparticles in HV insulating materials

Several parameters affect the efficiency of the role of nanoparticles in the ultimate functional properties of the nanocomposites: the size, the shape, the orientation and the state of dispersion (nanoscale dispersion vs. agglomerations). In the following sections, selected studies from the literature illustrate the effect of each parameter on several properties related to the performance of HV insulating materials.

1.1.6.1 Effect of nanoparticles size

As discussed previously, decrease of nanoparticles size induces an increase of the interfacial region fraction. In correlation with this effect, several dielectric properties such as the breakdown strength and resistance to surface erosion were reported to increase. An illustrative example was reported by Andritsch et al. (Andritsch et al., 2010) in their study of the dielectric behavior of epoxy nanocomposites containing BN nanoparticles with various diameters ranging from 70 nm to 5 μ m. In particular, they observed an increase of the DC breakdown strength with decreasing nanoparticles size. The maximum strength was achieved at 10 wt % BN (70 nm) and corresponded to 40 % increase compared to the neat epoxy (Figure 1-7). A similar trend was reported by Yang et al. (W. Yang et al., 2012) as well.



Figure 1-7: DC breakdown strength of BN/Epoxy as function of filler size Taken from Andritsch et al. (Andritsch et al., 2010)

Regarding the resistance to surface erosion, multiple studies concluded that polymer matrices filled with nanoparticles feature higher resilience compared to matrices filled with the same concentrations of microparticles or even higher (M. Frechette et al., 2012; Iyer et al., 2012; Tanaka et al., 2008). For example, Iyer et al. (Iyer et al., 2012) found out that epoxy matrix filled with 5wt% silica nanoparticles perform slightly better than epoxy matrix containing 65 wt% of silica microparticles. Taking into consideration that the degradation in the samples occurs exclusively through the polymer matrix which creates an erosion path growing between the particles of higher thermal stability, this behavior can be explained by the fact that nano-filled samples have longer erosion paths compared to micro-filled samples due to reduced interparticle distance and hence increased tortuosity.

Thermal conductivity as well is affected by the size of particles. In this context, a recent study performed by Kochetov et al. (Kochetov et al., 2010) on epoxy composites filled with BN particles of different sizes from 70 nm to 5 μ m has shown that thermal conductivity tends to increase with increasing nanoparticles loading but features an optimum as function of BN size at 0.5 μ m. This behavior is probable related to facilitated formation of a thermal network using submicron particles, as compared to either nanometric or micrometric particles. Few other studies reported size dependence of thermal conductivity on the filler size but no obvious trend was concluded (Reading, Vaughan, & Lewin, 2011).

1.1.6.2 Effect of nanoparticles shape and orientation

Some anisotropic nanoparticles such as nanotubes and nanoplatelets feature high aspect ratios and considerably larger interfacial areas compared to their isotropic pairs, in addition to the anisotropy of their intrinsic properties. Due to these features, the geometry and the orientation of nanoparticles incorporated in polymer insulations was found to affect their functional properties.

In this context, Tomer et al. (Tomer et al., 2011) studied the effect of alignment of nanoclays on the dielectric properties of an 80/20 LLDPE/ LDPE polymer blend. Films with oriented nanoclays were prepared in a twin-screw extruder whereas films with random distribution of

nanoparticles were obtained by hot-pressing a stack of 4 oriented films sequentially rotated by $\pi/4$. The orientation of nanoparticles didn't induce any particular effect on the low field dielectric response of the PE matrix. In fact, all the nanocomposites were found to exhibit similar losses that are approximately two orders of magnitude higher than the dielectric loss of unfilled PE. However, the group reported that the nanocomposites with randomly oriented fillers exhibit considerably higher losses when subjected to a high electric field (broader Displacement-Electric field D-E loop as reported in Figure 1-8(a)) as well as lower breakdown strength (Figure 1-8(b)), compared with nanocomposite containing oriented fillers. They attributed these results to the fact that oriented-fillers samples are expected to provide more ordered trapping sites and more efficient scattering for the injected charge, limiting its ability to cross the sample to the opposite electrode. This scattering effect is reduced for randomly dispersed fillers, even when they are in the form of high aspect ratio particles such as nanoclays.



Figure 1-8: Effect of the nanofillers alignment on: (a) the Displacement-Electric field (D-E) loops and (b) the breakdown strength of polyethylene Taken from (Tomer et al., 2011)

In the same context of orientation effect, Terao et al. (Terao et al., 2010) studied the effect of the alignement of boron nitride nanotubes (BNNT) by electrospinning on the thermal conductivity of BNNT/Polyvinyl alcohol (PVA) nanocomposites. They observed that in a general trend, the increase of thermal conductivity induced by BNNT incorporation is higher in the in-plane directions than along the thickness of the samples, regardless of the nanotubes

orientation as reported in Figure 1-9. Upon alignement of 10 wt% BNNT along one axis, the thermal conductivity value reaches 0.54 W.m⁻¹.K⁻¹ along the alignment axis and 0.27 W.m⁻¹.K⁻¹ along the other in-plane direction compared to 0.16 W.m⁻¹.K⁻¹ for the neat PVA. However, the values of thermal conductivity for the bioriented sample and the isotropic sample are respectively equal to 0.38 and 0.26 W.m⁻¹.K⁻¹. This increase along the alignment direction was attributed to the contribution of high axial thermal conductivity of BNNTs.

type of BNNT alignment	BNNT fraction		thermal conductivity (W/mE)		
		vol %is	X direction	J direction	2 direction
Y/	0	0	835	0.16	831
	4	0.45	0.17	0.17	0.13
	10	4.76	0.38	0.38	81.0
Y/	10	4.76	0.27	0.54 (0.59) ⁴	81.0
MA			0.10	0.10	0.10
(tait)	1	0,45	6.15	0.15 (0.17)	8.34
	10	4.76	0.26	0.26 (0.29)"	0.20

Figure 1-9: Thermal conductivity of BNNT/ PVA nanocomposite as function of the orientation Taken from (Terao et al., 2010)

Regarding the impact of nanoparticles geometry, a good illustration of the effect of nanoparticles shape on the dielectric behavior was reported in the work of Fabiani et al. (Fabiani, Montanari, & Testa, 2010). In this study, the dielectric response of Ethylene-vinyl-acetate (EVA) polymer matrix containing two clay particles featuring different aspect ratios was investigated. It was found out that the dielectric loss and the electrical conductivity were higher in the sample containing the nanofiller with the highest aspect ratio. This behavior was attributed to favored formation of a conductive path between the absorbed water shells surrounding the particles.

1.1.6.3 Effect of nanoparticles dispersion

The dispersion of nanoparticles is a key factor affecting the quality of the interfacial region and consequently the engineering properties. In the specific case of nanodielectrics, the thermal and dielectric properties are sensitive to the dispersion. Below, some illustrative examples from the literature regarding the effect of dispersion are provided. In particular, the engineering properties of samples containing agglomerated nanoparticles are compared to samples featuring improved dispersion, achieved either by specific processing techniques or by nanoparticles functionalization. For instance, Yang et al. (W. Yang et al., 2012) studied the dissipation factor and the short-term breakdown strength of an epoxy matrix containing respectively well dispersed nano ZnO particles vs. aggregated ZnO nanoparticles. The improved dispersion was achieved by adding high shear mixing and ultrasonication steps to the fabrication process. The group reported that the highest breakdown strength and the lowest dielectric loss were achieved in the nanocomposite featuring well dispersed nanoparticles due to more efficient scattering of charge carriers and reduced interfacial polarization. In the same context, a recent study performed by Zazoum et al. (Bouchaib Zazoum, Frechette, & David, 2016) investigated the dielectric loss of low density polyethylene (LDPE) containing respectively untreated titanium dioxide (TiO₂) nanoparticles and TiO₂ treated with Polyhedral Oligomeric Silsesquioxane (POSS) molecules. The surface modification with POSS resulted in improved nanoscale dispersion of TiO₂ nanoparticles that was confirmed by atomic force microscopy (AFM). As a result of this improved dispersion, a significant decrease of the dielectric loss was observed. Moreover, the dielectric strength of the interfacial relaxation observed in the untreated nanocomposite was reduced and the peak shifted toward lower frequencies indicating that most likely the effective electrical conductivity of the nanofillers was reduced after functionalization with POSS. As far as thermal conductivity and resistance to surface erosion are concerned, it was demonstrated that the dispersion quality might have an important impact as well. In recent publications by Heid et al. (Heid, Fréchette, & David, 2015, 2016), it was found out that 1 wt% of reactive POSS molecules incorporated in an epoxy matrix and evenly distributed at the nanoscale led to an increase of thermal conductivity by 20%. However, the sample containing non-reactive

POSS exhibited an increase of only 10%. Moreover, it was observed that the resistance to surface erosion of the sample containing reactive POSS was improved with increasing POSS loading while the opposite effect was observed in the sample containing non-reactive POSS due to the formation of agglomerations.

1.2 Control of morphology: application to nanodielectrics

As discussed in the previous sections, several parameters affect the performance of nanocomposites and specifically nanodielectrics, including the quality of dispersion and orientation. Recently, new approaches that might help improving the dispersion and more importantly controlling the selective localization and orientation of nanoparticles are being investigated. These approaches are based on the use of multi-phase polymer matrices as structure-guiding hosts to tailor nanoparticles dispersion, such as polymer blends and block copolymers which are self-assembled according to specific morphologies at micro and nanoscale respectively (Carastan et al., 2013; Carastan et al., 2014; Chipara, Artiaga, Lau, Chipara, & Hui, 2017; Kao et al., 2013; Yao Lin et al., 2005; Park et al., 2003; Sarkar & Alexandridis, 2015). In these multi-component systems, the control of dispersion and spatial distribution can be ensured based on the matrix ordered morphology and the affinity of the studied nanoparticles to one phase or another. In the last decades, several studies have investigated this approach for various applications including mainly microelectronics, polymer solar cells and optical sensors (As'habi et al., 2013; H. Choi et al., 2013; F. Li, Shi, Yuan, & Chen, 2013; Park et al., 2003; Sarkar & Alexandridis, 2015). However, only few of them addressed the field of dielectrics and more specifically HV electrical insulation. In the following sections, selected published studies on tailored dispersion of nanoparticles in block copolymers, polymer blends and polymer/block copolymer blends are presented with focus on dielectric applications.

1.2.1 Block copolymer nanocomposites

The control of nanoparticles dispersion in block copolymer hosts depends on few important criteria including the affinity between the selected nanoparticles and each phase of the block

copolymer, the nanostructure of the block copolymer as well as the geometry of the nanoparticles and the block copolymer nanodomains. In particular, some combinations of respectively nanoparticle geometries and block copolymer morphologies offer larger interfacial area (Bockstaller, Mickiewicz, & Thomas, 2005). Besides, the size of the nanoparticles, in comparison to the characteristic length scales of the block copolymer domains, might affect their selective localization and the block copolymer ordered structure, taking into account the thermodynamic constraints (Sarkar & Alexandridis, 2015).

Tailored dispersion of different nanoparticles, mainly of spherical shape, in block copolymers was reported in the literature (H. Choi et al., 2013; F. Li et al., 2013; T.-I. Yang & Kofinas, 2007). For instance, Li et al. (F. Li et al., 2013) studied the dispersion of ZnO nanoparticles in a poly(3-hexylthiophene)-b-poly(ethylene oxide) (P3HT-b-PEO) diblock copolymer matrix in comparison with ZnO dispersion in P3HT homopolymer matrix. The analysis of both types of nanocomposites processed in the same conditions by TEM revealed that fine and uniform dispersion of ZnO nanoparticles was achieved in the block copolymer while large clusters of ZnO nanoparticles were formed in P3HT matrix (Figure 1-10). Furthermore, the ZnO nanoparticles were preferentially located in the PEO nanometric domains. The group attributed this improved dispersion and selective localization to the favorable interaction between the oxygen atoms of the PEO backbone and ZnO nanoparticles.



Figure 1-10: TEM images of: a) P3HT/ZnO nanocomposite and b) P3Ht-b-PEO/ZnO nanocomposite Taken from Li et al. (F. Li et al., 2013)

Recently, controlled dispersion of anisotropic nanoparticles in block copolymers has been investigated as well (Carastan et al., 2014; Chipara et al., 2017; Peponi et al., 2009). As an illustration, the recent study carried out by Carastan et al. (Carastan et al., 2014) investigated the dispersion of clay nanoplatelets in polystyrene-b-poly(ethylene-co-butylene)-b-polystyrene (SEBS) triblock copolymer. The studied block copolymer features an hexagonally packed cylindrical morphology and is composed of two PS endblocks in the form of cylinders surrounded by an elastomer PEB midblock. The group reported successful orientation of nanoclays and PS nanodomains achieved using tape extrusion technique. Besides, clay nanoplatelets structure, i.e. exfoliated vs. intercalated, and their location in either PS block or PEB block were tuned through a maleic anhydride (MA) graft attached to the PEB rubber block.

In the field of dielectric applications, few block copolymers gained attraction either for their nanostructure or for their functional properties such as their good electromechanical coupling (B. Kim et al., 2011). As far as the nanostructure is concerned, it was demonstrated that the breakdown strength of a block copolymer is highly sensitive to its ordered nanostructure. For example, polystyrene-b-poly(methyl methacrylate) (PS-b-PMMA) block copolymer with highly oriented lamellar morphology featured great enhancement of breakdown strength compared to unordered films prepared by solvent casting (Samant et al., 2016). The enhancement was attributed to the fact that the multiple interfaces between the lamellae block components act as barriers to the flow of electric charge carriers. This behavior has an important potential to design nanodielectrics with high energy density for instance. Styrenic block copolymers, as part of the category thermoplastic elastomers, were among those recently investigated within the scope of actuation properties. The main objective of their use was the design of sophisticated, light-weight and shape-flexible high dielectric permittivity polymer nanocomposites intended for service in embedded capacitors, piezoelectric and pyroelectric sensors, microwave communication devices and/or dielectric elastomer actuators (Kofod et al., 2011; Mc Carthy, Risse, Katekomol, & Kofod, 2009; Mi, Li, Turng, Sun, & Gong, 2014; Saleem, Thunga, Kollosche, Kessler, & Laflamme, 2014; Stoyanov, Kollosche, Risse, McCarthy, & Kofod, 2011; T.-I. Yang & Kofinas, 2007). In general, the two main approaches to achieve this type of nanocomposites, which are based on the use of either high dielectric permittivity ceramics or conductive particles (sub-percolation loadings), induce relatively high dielectric losses; a design limitation for the cited applications. Interestingly, through tailoring the size distribution of nanoparticles and their spatial distribution in block copolymers featuring highly periodic nanostructures, structural heterogeneities and agglomerations contributing to the increase of dielectric losses might be significantly reduced (T.-I. Yang & Kofinas, 2007).

In HV electrical insulation, the use of styrenic block copolymers is mainly limited to their role as compatibilizers to improve nanoparticles dispersion (Liang & Tjong, 2006) or as water treeing retardant agents (Liu et al., 2011; Z. Ma et al., 2010). However, their potential as nanostructured template matrices is not fully exploited yet.

1.2.2 Polymer blend nanocomposites

Recently, a need to replace commercial cross-linked polyethylene (XLPE) cables, currently on the market, has progressively emerged due to their vulnerability to space charge accumulation induced by the crosslinking by-products, which makes XLPE non-recyclable and unsuitable for increased HV ratings and especially HVDC cables (Fu, Chen, Dissado, & Fothergill, 2007; Ghorbani et al., 2014; Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016). Unfortunately, the crosslinking is required in current HV cable insulations to improve the mechanical and thermal stability of LDPE, which is the most common polymer insulation used in HV cables (Arora & Mosch, 2011).

In this context, polymer blends have been investigated since the last decades in order to improve the performance of LDPE and ultimately avoid crosslinking. In this context, Hosier et al. (I. Hosier, Vaughan, & Swingler, 1997) studied the effect of the presence of a dispersed high density polyethylene (HDPE) phase on the breakdown strength of an LDPE matrix. The prepared blends were subjected to thermal annealing at two different temperatures: 115 °C and at 124 °C. The group reported a clear improvement of the breakdown strength in blends

containing HDPE fractions above 7% and annealed at 115 °C, whereas, in those annealed at 124 °C, a slight improvement occurred only above 17% of HDPE. Below these limits, both types of blends featured a reduction in the breakdown strength compared to the pure LDPE. These behaviors were correlated with changes in spherulite structures upon blending HDPE with LDPE. In particular, it was concluded that morphologies based upon small well distributed HDPE-rich inclusions achieved at 124 °C (compact spherulites which did not interpenetrate) lead to lower breakdown strength while space-filling spherulites, achieved at 115 °C, cause the tree to adopt a more extended path or, in the extreme cases where they impinge, force the tree to penetrate the highly crystalline material leading to higher breakdown strength compared to the pure LDPE.

To improve further the properties of LDPE/HDPE blends, incorporation of nanoparticles has been investigated as well. For example, Zazoum et al. (B Zazoum et al., 2014) investigated the effect of organically modified nanoclays on the dielectric properties of a blend containing 80 wt% LDPE and 20 wt% HDPE. They observed that the incorporation of 5 wt% nanoclays in the blend induced a consistent increase of the breakdown strength equal to 17%. This improvement was extended further to 22% when polyethylene grafted maleic anhydride (PE-MA) was used as compatibilizer. This behavior was correlated with microscopy results confirming the achievement of better intercalated/exfoliated nanoclays structure in the compatibilized blend nanocomposites.

Recently, other alternatives have been investigated. In fact, high density polyethylene (HDPE) and especially Polypropylene (PP) feature better thermal and mechanical stability, compared to LDPE, due to their higher melting points (Arora & Mosch, 2011). Therefore, they might constitute potential candidates for recyclable HV insulation materials, considering the omitted need for crosslinking. However, these materials suffer from increased brittleness at low temperatures. To adapt better their properties to the purpose of recyclable HV insulations, blending was investigated again as one of the solutions. In particular, blends with thermoplastic elastomers such as ethylene-vinyl-acetate (EVA) and polyolefin elastomers were considered (I. Hosier, Vaughan, & Tseng, 2007; I. L. Hosier, Vaughan, & Swingler,

2010; Y. Zhou, He, Hu, Huang, & Jiang, 2015). It was found out that such blends feature considerably improved mechanical flexibility and occasionally improved electrical resistivity. Nevertheless, reduction of breakdown strength and increase of space charge accumulation, a key property for HVDC cable insulation, were unfortunately inevitable after blending. To cope with these drawbacks while taking advantages from the blends properties, several nanoparticles have been added, among them: ZnO and BN (Dong, Han, & Han, 2012; Du, Xu, Li, & Li, 2016; D.-L. Zhang et al., 2017). For instance, Du et al. (Du et al., 2016) reported that the addition of ZnO nanoparticles to PP/Polyolefin elastomer (POE) blends results in suppression of space charge accumulation, encountered in the unfilled blend, while maintaining the excellent flexibility ensured by the POE phase. This reduction of space charge accumulation is due to increased trap level density and consequently reduced charge injection upon the addition of ZnO nanoparticles. The maintained mechanical flexibility of the blend is more likely related to the fact that the majority of ZnO nanoparticles migrated to PP phase during the processing and consequently didn't affect the mechanical flexibility of POE phase. In the same line of research, another blend nanocomposite containing a thermoplastic elastomer was recently investigated by Zhang et al. (D.-L. Zhang et al., 2017). In this system, the main matrix is PP and the selected thermoplastic elastomer is polystyreneb-poly(ethylene-co-butylene)-b-polystyrene (SEBS) while the nanoparticles are Boron Nitride nanosheets (BNNS), known for their excellent thermal conductivity and electrical insulation. The proportions of PP and SEBS in the blend are respectively equal to 35 and 65 wt%. In this blend, BNNS were selectively located in SEBS phase. At 3wt% loading, these high aspect ratio nanoparticles simultaneously formed a thermal network inside the SEBS phase and induced a morphology change in the blend from dispersed SEBS phase to cocontinuity, leading to a "double percolation" phenomenon as described by the authors. This selective location of BNNS and specific configuration of the blend induced great enhancement of thermal conductivity in addition to reduction of space charge accumulation and increase of breakdown strength. Another example of a successful controlled dispersion in a conventional HV insulating material induced by blending has been reported by Peng et al. (Peng, Xu, Li, Zhang, & Zheng, 2016). In this work, an epoxy resin was blended with an amphiphilic triblock copolymer composed of a poly(POSS) midblock and two

polycaprolactone (PCL) endblocks. The PCL block is miscible with epoxy while the poly(POSS) midblock self-assembles in the form of non-spherical domains of size 20-50 nm well distributed within the epoxy matrix. This periodic organization allowed a consistent reduction of the dielectric permittivity compared to unfilled epoxy combined with an enhancement of the surface hydrophobicity.

1.3 Review of the materials

In this section, a review of the polymers and nanoparticles that were investigated in this project is presented. In particular, the choice of cost-effective nanoparticles is emphasized.

1.3.1 Polyethylene

Polyethylene (PE) is a polycrystalline polymer belonging to the family of polyolefins. It features a simple hydrocarbon structure and can be obtained with different molecular weights and branching degrees, with the highest branching occurring in low density polyethylene (LDPE). Moreover, its semi-crystalline structure with crystalline/amorphous interfaces provides traps that reduce charge carriers mobility and the overall electrical conductivity (Figure 1-11). Therefore, it is already considered as a nanodielectric, even without the incorporation of nanometric fillers (Ieda, 1984; Lewis, 2014). In HV insulating materials industry, PE is attractive due to its excellent electrical resistivity, high breakdown strength and good processability by melt compounding techniques. Currently, cross-linked low density polyethylene (XLPE) is the most widely used form of PE in HV cable insulation. The crosslinking is particularly important to improve the thermal and mechanical stability of LDPE at increased temperatures as mentioned previously (Arora & Mosch, 2011; Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016). Unfortunately, it renders the insulating material unrecyclable and sensitive to space charge accumulation (Fu et al., 2007; Ghorbani et al., 2014). Compared to LDPE, high density polyethylene HDPE features higher thermal stability but has been less common in cable insulation due to its relatively high rigidity (Arora & Mosch, 2011). However, recently it has been investigated in several blends in the objective of designing insulating materials suitable for HV insulation without the need

of crosslinking (I. Hosier et al., 1997; I. Hosier et al., 2007; I. L. Hosier et al., 2010). In this project, HDPE was investigated in a blend with SEBS thermoplastic elastomer.



Figure 1-11: Morphology of polyethylene: amorphous and crystalline regions Taken from (Lewis, 2014)

1.3.2 SEBS thermoplastic elastomer

Polystyrene-b-poly(ethylene-co-butylene)-b-polystyrene (SEBS) is a triblock copolymer composed of two polystyrene (PS) endblocks and a poly(ethylene-co-butylene) (PEB) midblock that belongs to the family of styrenic thermoplastic elastomers. Its chemical formula is reported in Figure 1-12.



Figure 1-12: Schematic description of SEBS block copolymer: (a) without maleic anhydride (MA) ("Polystyrene-block-poly(ethylene-ran-butylene)-block-polystyrene,") and (b) with MA graft attached to PEB block ("Polystyrene-block-poly(ethylene-ran-butylene)-block-polystyrene-graft-maleic anhydride,")

Being a block copolymer, SEBS features self-assembled nanodomains that vary from lamellar to spherical depending mainly on the styrene fraction (Carastan et al., 2013; Carastan et al., 2014; Carastan, Demarquette, Vermogen, & Masenelli-Varlot, 2008). Moreover, it exhibits excellent mechanical properties combining both the thermoplastic and the elastomer behaviors (Balsamo et al., 2006; Holden et al., 2004). Besides, its hydrogenated polybutadiene midblock has a structure equivalent to the structure of PE in its amorphous form, which ensures a good compatibility with PE and polyolefins in general (Agari et al., 1993). Simultaneously, the relatively polar aromatic rings of PS block exhibit chemical affinity to several inorganic nanoparticles such as nanoclays and carbon nanotubes (CNT) (Carastan et al., 2014; Kuester, Barra, Ferreira Jr, Soares, & Demarquette, 2016). Therefore, SEBS and especially SEBS grafted maleic anhydride (MA) were recently widely used either as matrices or compatibilizers in blends with polyolefins to improve the dispersion of inorganic nanoparticles including nanoclays, metal oxides and CNT, and to compensate for toughness decrease induced by their incorporation in polymer matrices (Carastan et al., 2014; Kuester et al., 2016; Liang & Tjong, 2006).

In dielectric applications, SEBS has been also investigated, as mentioned in previous sections, mainly as a water treeing retardant agent for cable insulation (Liu et al., 2011; Z. Ma et al., 2010) or as a dielectric elastomer actuator (Kofod et al., 2011; Mc Carthy et al., 2009; Mi et al., 2014; Saleem et al., 2014; Stoyanov et al., 2011; T.-I. Yang & Kofinas, 2007). In addition to these attractive functional properties, it started recently to gain

attraction, among other thermoplastic elastomers, in the design of recyclable polyolefin/thermoplastic elastomer blends suitable for HV insulation and especially for HVDC cable insulation (D.-L. Zhang et al., 2017).

In this study, SEBS was investigated first as a template block copolymer matrix to evaluate the effect of controlled dispersion and spatial distribution of nanoparticles on the dielectric, electrical and thermal properties of nanodielectrics; an approach that has not yet been widely studied in this field. In particular, different grades of SEBS with various PS block fractions and subsequently various morphologies of the block copolymer were studied. In a second step, SEBS was investigated in a blend with HDPE where both its functional properties and nanoarchitecture are taken into account to design nanocomposites with improved dielectric performance intended for use in HV insulation.

1.3.3 Zinc Oxide nanoparticles

Zinc oxide (ZnO) is a large band gap semi-conductor material characterized by its nonlinear electrical conductivity (Hong, Schadler, Siegel, & Mårtensson, 2006; Varlow, Robertson, & Donnelly, 2007), relatively high thermal conductivity (around 60 W.m⁻¹.K⁻¹ compared to 0.33 W.m⁻¹.K⁻¹ for LDPE and 2000 W.m⁻¹.K⁻¹ for diamond according to (Huang et al., 2011)) and ability to shield UV light that can be released during partial discharges for instance (S. Chen, Huang, Peng, Wang, & Cheng, 2010). Similar to other metal oxides, ZnO nanoparticles were studied in polymer nanocomposites for insulation applications, mainly PE based (Fleming et al., 2008; Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016; Tian et al., 2011, 2012; Tian et al., 2015). As mentioned in paragraph 1.1.5.1, the undertaken research into the properties of these nanocomposites demonstrated the ability of ZnO nanoparticles to relieve local electrical stress concentration and consequently to homogenize electric field distribution, which hinders the growth of electrical trees within the nanocomposite. Besides, they were reported to increase traps density in the insulating matrix leading to decrease of charge carriers' mobility (Tian et al., 2011). In correlation with these features, significant reduction in space charge accumulation and improved resistance to surface erosion and electrical treeing were observed in ZnO-containing nanocomposites.

In addition to the functional properties, the geometry of ZnO nanoparticles can be tailored into a broad range of particle sizes and shapes using mainly chemical routes. Moreover, their surface can be modified to increase their hydrophobicity and ultimately improve their compatibility with organic polymer matrices. Both the processes of synthesis and surface functionalization of these nanoparticles are relatively simple and cost-effective (Amir M Pourrahimi et al., 2016). In this project, commercial ZnO nanoparticles of spherical shape and functionalized with an alkyl ammonium salt were used.

1.3.4 Organically modified Montmorillonite Clay

Smectite clays such as Montmorillonite are the inorganic particles most used in nanocomposites. They belong to the phyllosilicates family characterized by a 2:1 structure built of layers; each of them is composed of aluminium or magnesium hydroxide octahedral sheet sandwiched between two silicon oxide tetrahedral sheets, as illustrated in Figure 1-13. The layers are placed on the top of each other's forming stacks. Besides, Van Der Waals and weak electrostatic interactions occur between them and result in the creation of interlayer spaces or galleries where exchangeable cations reside. The thickness of the constitutive layers is around 1 nm while the lateral dimensions may vary from 30 nm to several microns leading to high aspect ratio and large surface area that dominate the interactions of these particles with polymers (Anadão, 2012; Choudalakis & Gotsis, 2009; Nguyen & Baird, 2006; Powell & Beall, 2006).



Figure 1-13: Crystal structure of phyllosilicates Taken from Thomas el al. (Thomas & Zaikov, 2008)

Layered clays are hydrophilic in their pristine state. Therefore, for non-polar polymer matrices, an organic treatment on the clay surface is needed to obtain satisfactory dispersion. The most commonly used organic treatment is based on quaternary ammonium salts, which can have a variety of chain lengths (Choudalakis & Gotsis, 2009; Powell & Beall, 2006). Depending on the polymer matrix, the selected organic modification and the processing method, two main types of clay structures can be obtained as illustrated in Figure 1-14: intercalated and exfoliated. In the case of untreated clays or inappropriate surface functionalization, the layers don't separate and the clay particles remain in the initial form of stacks or tactoids.



Figure 1-14: Layered clay structures in nanocomposites Taken from (Tronto, Bordonal, Naal, & Valim, 2013)

Nowadays, polymer clay nanocomposites are already used in many applications to enhance targeted properties such as mechanical strength, flame retardancy and gas-barrier properties (Nguyen & Baird, 2006). Besides, the existing processes to obtain organically modified clays are well developed and cost-effective.

In dielectric applications, organically modified clays were also studied and reported to enhance the performance of polymer insulations by increasing their breakdown strength and resistance to electrical treeing. In fact, their high aspect ratio and surface area induce interfaces with increased scattering and tortuosity that oppose the flow of charge carriers (David et al., 2013; Tomer et al., 2011; B Zazoum et al., 2014). In particular, the orientation of clays was observed to increase the tortuosity effect and consequently to improve further the dielectric strength (Tomer et al., 2011; B Zazoum et al., 2014). However, this effect is still not well quantified/optimized. In this project, organically modified Montmorillonite clay treated with quaternary ammonium salt were used and the effect of their tailored distribution and alignment was emphasized.
CHAPTER 2

ARTICLE 1: STYRENIC BLOCK COPOLYMER-BASED NANOCOMPOSITES: IMPLICATIONS OF NANOSTRUCTURATION AND NANOFILLER TAILORED DISPERSION ON THE DIELECTRIC PROPERTIES

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Abstract

In this work, the effect of controlling the morphology on the dielectric properties of triblock copolymers and their clay-containing nanocomposites was evaluated. Two different copolymers: polystyrene-b-poly(ethylene-co-butylene)-b-polystyrene (SEBS) and SEBS grafted with maleic anhydride were used for that purpose. Morphologies with different degrees of intercalation, exfoliation and orientation were obtained and tested. At the highest state of dispersion, achieved at a clay loading equal to 5wt%, 50% of clay nanoplatelets were individually dispersed and located within the PEB soft domain of the block copolymer and a maximum of interfacial polarization and a minimum of dynamic mechanical damping factor were respectively exhibited. When the nanoclays were oriented, the dielectric loss due to nanoclays conductivity contribution was reduced up to 2 orders of magnitude at high

temperatures and low frequencies and the AC short-term breakdown strength increased up to 45%.

Keywords: block copolymer; tailored dispersion; dielectric properties...

2.1 Introduction

"Nanodielectrics" or nanocomposite dielectrics are a new generation of dielectric materials containing fillers that have at least one dimension less than 100 nm (Michel F Fréchette et al., 2010). This class of materials is gaining a lot of interest aiming at developing dielectrics with distinct properties originating from the intrinsic properties of the nanofillers and mostly from the important interfacial region introduced due to the huge surface area of the nanofillers (Lewis, 2004; Roy et al., 2005). Several publications over the last decade proved the efficiency of nanofillers at improving several aspects related to the dielectric performance of insulating polymers in applications such as cable insulation (David & Fréchette, 2013; M. Fréchette, 2009), high energy storage capacitors (Dang, Yuan, Yao, & Liao, 2013) and dielectric elastomer actuators (McCarthy et al., 2012). In these studies, different types of nanofillers and polymers have been tested.

In terms of nanofillers, nanoclays were the most common nanoparticles used due to their relatively low cost. More specifically, organically-modified nanoclays have been the nanoparticles of choice since their surface modification made them more compatible with organic polymer media (Carastan & Demarquette, 2007; Lei, Hoa, & Ton-That, 2006). These nanoparticles have proved their positive role in improving the mechanical properties of polymers and in few studies in improving their dielectric properties. In effect, it was demonstrated that nanoclay addition and orientation has an influence on the high field properties of polyolefins such as the short-term breakdown strength (David et al., 2013; Liao et al., 2013a; Liao, Li, Bai, Yang, & Gu, 2014; Tomer et al., 2011). In this context, David et al. (David et al., 2013) reported an increase up to 15% of the breakdown strength of low density polyethylene (LDPE) matrix filled with a low content of nanoclay equal to 3wt%. This improvement was combined with an increase of the dielectric loss by roughly two orders of magnitude. They attributed the increase of the breakdown strength to the exfoliated

structure of nanoclays and the high dielectric loss to the high conductivity along the clay nanoplatelets. Also, Tomer et al. (Tomer et al., 2011) reported an increase of the breakdown strength by 20% upon the alignment of nanoparticles in LDPE containing 6 wt% nanoclay whereas they observed a deterioration in the breakdown strength and consequently in the maximum recoverable energy when nanoclays are randomly oriented. The authors reported as well an increase of the dielectric loss by two orders of magnitude in both types of nanocomposites with either random or aligned nanoclays.

In terms of polymers, styrenic block copolymers constitute interesting candidates which represent an important part of the current market of thermoplastic elastomers. In fact, these materials exhibit attractive mechanical properties due to their combination of a soft elastomer phase and a hard polystyrene phase. Furthermore, as block copolymers, this class of materials presents the advantage of being self-assembled at the nanoscale in unique morphologies such as lamellar, cylindrical and spherical hard domains distributed within the elastomer domains. The shape of these ordered structures depends on the chemical composition, molar mass of the blocks, affinity between the blocks and processing method (Mai & Eisenberg, 2012). In the field of nanocomposites, this ability to control their spatial organization makes styrenic block copolymers attractive as promising template matrices for selective dispersion of nanoparticles with competitive mechanical properties (Bockstaller, Lapetnikov, Margel, & Thomas, 2003; Bockstaller et al., 2005; Chiu, Kim, Kramer, & Pine, 2005). In this context, a great number of recent publications were related to block copolymer-based nanocomposites with tailored dispersion of nanofillers. Among them, the studies performed by Carastan et al. (Carastan et al., 2014; Carastan, Vermogen, Masenelli-Varlot, & Demarquette, 2010) show that selective dispersion and orientation of nanoclays were successfully achieved in a triblock copolymer. The published researches treated various applications such as polymer solar cells (H. Choi et al., 2013; F. Li et al., 2013) and block copolymer electrolytes (Hur & Bae, 2015). They demonstrated that the orientation as well as the location of nanoparticles seems to affect the resulting properties. To the best of our knowledge, only few of these studies addressed the field of dielectric applications (H. Chen, Hassan, Peddini, & Mauritz, 2011; Mc Carthy et al., 2009; Vo, Anastasiadis, & Giannelis, 2011) in spite of the fact that thermoplastic

elastomers could represent promising candidates for dielectric elastomer actuators (Mc Carthy et al., 2009) or styrenic block copolymer/polyolefin blends for high voltage insulation (Liu et al., 2011).

In this paper, the dielectric properties of polystyrene-b-poly(ethylene-co-butylene)-bpolystyrene (SEBS), an interesting thermoplastic elastomer widely used (Arevalillo, Muñoz, Santamaría, Fraga, & Barrio, 2008; Carastan et al., 2013; Carastan et al., 2008; S. Choi, Lee, & Han, 2004; Ganguly, Bhowmick, & Li, 2008; Jeong et al., 2003), and its nanocomposites containing clay nanoparticles were investigated as function of various parameters related at times to the nanofillers and other times to the SEBS structure. In this context, the effect of clay dispersion (intercalated vs. exfoliated), clay location (within the hard or soft phase), clay concentration and clay orientation were evaluated. Besides, the effect of SEBS morphology was studied. In particular, two types of SEBS structures, namely a hexagonal structure where polystyrene cylinders are distributed within the rubber phase and a lamellar structure formed by alternating polystyrene and rubber layers, were evaluated. The structures of SEBS nanocomposites studied were characterized by different tools such as TEM, SAXS and DMA. The complex dielectric permittivity and breakdown strength of these nanocomposites were also evaluated and correlated to the different structures.

2.2 Materials and experimental characterization

2.2.1 Experimental strategy

In the nanocomposites studied in this paper, four microstructural aspects have been controlled, by varying the processing technique, in order to highlight their effects on the dielectric properties:

 Dispersion and location of nanoclays: two different grades of SEBS, one of them modified with a maleic anhydride (MA) group grafted to its rubber phase and the other without any modification, were used in order to tailor the location and state of exfoliation of nanoclays. In the pure SEBS, intercalated nanoclays located within the polystyrene (PS) domain were achieved while in the SEBS modified with maleic anhydride, exfoliated nanoclays located at the interface between the PS and the rubber domains were obtained. In fact, the grafted group has the greatest affinity to nanoclays and consequently attracts them to the interface (Carastan et al., 2014).

- Amount of surface area of nanoparticles: three different nanoclay concentrations were studied: 2.5, 5 and 7.5%.
- Orientation of nanoclays: two different techniques were used to control the orientation of the PS domain and consequently the orientation of nanofillers: extrusion process vs. solution casting. In particular, oriented nanoclays have been obtained by extrusion while random nanoclay distribution has been achieved in the nanocomposites prepared by solution casting (Carastan et al., 2014).
- Structures of SEBS: An oriented hexagonal structure, where oriented PS cylinders were dispersed within the rubber phase, was obtained by the extrusion process. Furthermore, the solution casting process was adapted to achieve respectively random hexagonal structure vs. random lamellar structure. The hexagonal structure was obtained when the solution casting process was accompanied by a specific thermal treatment while the lamellar structure was obtained when no thermal treatment was applied.

The details of experimental procedures to reach those morphologies are explained in the following section.

2.2.2 Materials and methods

Two grades of triblock copolymer SEBS donated by Kraton were used: G1652 and FG1901. The G1652 contains 30 wt % of polystyrene PS endblocks dispersed in the form of cylinders within a hydrogenated polybutadiene PEB midblock matrix. Its density is equal to 0.915 g/cm³. The FG1901 has the same composition with a maleic anhydride MA group grafted to the PEB midblock. Following, the G1652 and the FG1901 will be respectively referred to as SEBS and SEBS_MA. Montmorillonite clay grade Cloisite 20A from Southern Clay (Gonzales, USA), modified with dimethyl di (hydrogenated tallow) quaternary ammonium

salt was used as nanofiller for the nanocomposites preparation. More details regarding the modification and size of clay particles are provided in Table 2-1. All the materials studied in this paper were used as received.

Grade	OrganicCationcationicconcentrationmodifier a(meq/100g clay)		Weight loss at ignition (%)	Typical dry particle size (d50)	Basal spacing ^b (nm)
Cloisite 20A	CH3 1 CH3 - N'- HT 1 HT	CH, 1 N'- HT 95 1 HT		<10 µm	2.39
^a where HT is hydrogenated tallow (~65% C18; ~30% C16; ~5% C14) ^b Values according to reference [18]					

Table 2-1: Properties of Cloisite 20A grade

The nanocomposites were prepared by either melt compounding or solution casting method. The second method was combined in some cases with a thermal treatment to tune the structure of SEBS. The samples prepared by melt mixing were obtained according to a previous study published by Carastan et al. (Carastan et al., 2014). In a typical procedure, nanoclay was mixed with SEBS in a twin screw extruder equipped with a sheet die, at a temperature of 220 °C and a speed of 100 rpm. Further details on the processing steps and characterization by SAXS and TEM are available in the cited reference (Carastan et al., 2014). In the case of the solution casting process, the SEBS powder and the nanoparticles were mixed in toluene by magnetic stirring at 60 °C. The mixture was subsequently poured into a Petri dish and dried at room temperature for several days. The films obtained at this step had a random lamellar structure that will be confirmed in the characterization section. To obtain the random hexagonal structure, these samples were subsequently subjected to heat treatment to induce a microdomain transformation. This was performed in four steps: 20 min at 60 °C, 20 min at 100 °C, 20 min at 150 °C and 10 min at 200 °C. This transformation was enabled by the presence of toluene since the microdomain structure is strongly dependent on the solvent and annealing conditions (Hur & Bae, 2015). The progressive increase of temperature during the thermal treatment was applied to avoid the degradation of the polymer.

The morphology of the as-obtained nanocomposites was characterized by small angle X-ray scattering (SAXS) and transmission electron microscopy (TEM). The SAXS patterns were obtained using the synchrotron source of the National Synchrotron Light Laboratory (LNLS), Campinas, Brazil. The wavelength used in these measurements was 1.488 Å and the sample-to-detector distance was either 950 mm or 1125 mm. The TEM was performed with a Carl Zeiss CEM 902 transmission electron microscope for the nanocomposite containing 5 wt% nanoclay and with a Hitachi HD2700 field emission scanning transmission electron microscope (FE-STEM) for the nanocomposites containing 2.5 and 7.5 wt% nanoclay. Prior to observations, ultrathin sections of few tens of nanometers were cryo-cut using an ultramicrotome at -100 °C and deposited on copper grids. Some grids were also stained during 30 minutes with ruthenium tetroxide RuO4 vapor in order to determine the block copolymer morphology and to evaluate the location of clay nanoparticles.

Subsequently, the viscoelastic properties of all the nanocomposites were evaluated in order to confirm the different results obtained by microscopy and SAXS. In particular, the dynamic mechanical analysis (DMA) was done using a Q800 TA Instruments analyzer. Tensile mode was selected and rectangular specimens of 20 mm \times 6.5 mm x 1 mm were used. The measurements were performed from 20 °C to 140 °C. The test conditions were as follows: rate of heating equal to 3 °C/min, resonant frequency equal to 1 Hz and amplitude equal to 0.2 mm. The tensile storage modulus (E'), loss modulus (E'') and damping factor (tan δ) were evaluated.

Finally, in terms of dielectric properties, the complex dielectric permittivity of the nanocomposites was evaluated using a Novocontrol broadband spectrometer. Specimens of 20 mm in diameter were placed between two parallel brass plated electrodes. Measurements swept through a frequency range from 10^{-2} Hz up to $3*10^{5}$ Hz at a temperature range varying from 25 °C to 95 °C. The AC short-term breakdown strength of the samples was measured using ball-type electrodes of diameter 4 mm in a dielectric oil environment. The size of test specimens was approximately 3 cm x 3 cm. For each sample, 15 specimens were considered to calculate the dielectric strength using Weibull distribution, except for the extruded SEBS

nanocomposite containing 2.5 wt% nanoclay, where only 10 spots have been taken into account. A voltage ramp of 2 kV/s at a frequency of 60 Hz was applied until breakdown occurred. All measurements were done at a temperature equal to 23 °C. Since the thickness of the samples was not uniform, all the measurements were corrected to equal the breakdown strength of a 100 μ m thick film using the power law relationship between the breakdown field and the film thickness in order to obtain comparable results (Takala et al., 2010).

2.3 Results and discussion

2.3.1 Morphology

As mentioned above, four microstructural aspects of the nanocomposites were controlled: state of dispersion and location of nanoclays, amount of surface area of nanoparticles, orientation of nanoclays and structure of SEBS. These four aspects are reviewed below:

2.3.1.1 State of dispersion and location of nanoclay

In the nanocomposites produced by extrusion, both intercalated and exfoliated nanoclay structures were obtained depending on the use of SEBS or SEBS_MA as the matrix: nanoclay particles were intercalated and located within PS cylinders when SEBS was used while they were partially exfoliated and located in the PEB block when SEBS_MA was the matrix. The exfoliated vs. intercalated nanoclay structures can be observed respectively in Figure 2-1(a) and Figure 2-1(b)-(d). More details regarding the location of nanoclays in SEBS phases are available in reference (Carastan et al., 2014) and in the supporting file (annex I).



Figure 2-1: TEM images of respectively: (a) SEBS_5wt% nanoclay, (b) SEBS_MA_2.5wt% nanoclay, (c) SEBS_MA_5wt% nanoclay and (d) SEBS_MA_7.5wt% nanoclay (Prepared by extrusion)

DMA analysis was performed on the samples in order to confirm the difference in the state of clay dispersion, i.e. exfoliated vs. intercalated, as observed by TEM, between the nanocomposites prepared with SEBS and SEBS MA. Plots of storage modulus E', loss modulus E" and damping factor as functions of temperature are presented in Figure 2-2(a), Figure 2-2(b) and Figure 2-2(c) respectively. According to Figure 2-2(c), a relaxation peak around 104 °C was detected in tan δ curve for the pure SEBS. This peak corresponds to the glass transition temperature of the PS domains. The position of the damping peak was shifted by approximately 5 °C toward higher temperatures in the nanocomposites, indicating an increase of the PS glass transition temperature. Besides, the intensity of the damping peak decreased significantly in the nanocomposites. Both the increase of PS glass transition temperature and the decrease of the damping peak could be attributed to the reduction of polymer chain mobility within PS phase due to the presence of nanoclays. In particular, the lowest peak measured in SEBS MA 5% exfoliated nanocomposite may be considered as an indication of the efficient contribution of exfoliated clay nanoplatelets obtained in SEBS MA compared to intercalated nanoplatelets obtained in pure SEBS in blocking the segmental chain movement of PS cylinders and consequently decreasing the degree of disorder within the PS domain. These DMA results confirm the TEM observation shown in Figure 2-1.



Figure 2-2: (a) Storage modulus E', (b) Loss modulus E'' and (c) Damping factor (tan δ) at 1 Hz of SEBS vs. SEBS_MA nanocomposites containing 5wt% nanoclay (prepared by extrusion)

2.3.1.2 Amount of nanoparticle surface area

In addition to the nanocomposites prepared with 5 wt% nanoclay, two other loadings have been considered to highlight the effect of the interfacial region on the dielectric properties: 2.5 and 7.5 wt%. The quality of dispersion in these nanocomposites prepared by extrusion was assessed by image quantitative analysis involving TEM observations carried out at different scales following the procedure reported in Carastan et al. (Carastan et al., 2010) and Vermogen et al. (Vermogen et al., 2005). For that, a set of images taken at 3 magnifications was considered: 1 image at 20k, 4 images at 50k and 10 images at 100k. The thicknesses of

all the tactoids present in each image were estimated using image J software. The number of the nanoplatelets contained in each tactoid was then deduced. Once this step completed, the tactoids were classified in 6 different classes ranging from individual nanoplatelets to large tactoids containing more than 50 nanoplatelets. The final proportions were deduced taking into account a ponderation factor of the covered area corresponding to each magnification. More details regarding the quantitative analysis procedure are available in the supporting file (annex I) and references (Carastan et al., 2010; Vermogen et al., 2005).

Figure 2-3 shows the proportions of tactoid classes corresponding to the nanocomposites containing 2.5 wt% and 7.5 wt% of nanoparticles. The values corresponding to the nanocomposite containing 5 wt% of nanoparticles published in reference (Carastan et al., 2014) are also reported for comparison. It can be seen from Figure 2-3 that the 5 wt% nanocomposite has the highest percentage of individual layers and the best state of exfoliation. Almost 50% of its clay tactoids are individually dispersed in form of nanoplatelets and 97% of its tactoids contain less than 5 nanoplatelets according to the quantitative analysis. However, in the case of the nanocomposites containing 2.5 wt% nanoclay, the majority of nanoparticles (56%) were dispersed in form of small tactoids containing 3 to 5 layers and in the case of the nanocomposite containing 7.5 wt% nanoclay, the majority of the tactoids (51.5%) contained from 5 to 15 layers. A rough relative estimate of the number of tactoids and consequently the amount of interfaces in the nanocomposites containing respectively 2.5, 5 and 7.5 wt% nanoclay is: 52 vs. 350 vs. 137. This estimate is calculated based on the assumption that 1wt% clay tactoids should contain 100 individual nanoplatelets for example. The number of tactoids is deduced from this assumption by the following equation:

$$X = \sum_{i=1}^{6} \frac{x^{i*N}}{x_i}$$
(2.1)

Where:

N: the assumed total number of nanoplatelets;

xi: fraction of nanoplatelets in class i;

Xi: average number of platelets in class i;

X: total number of tactoids and consequently of interfaces.



Figure 2-3: TEM image quantitative analysis of oriented hexagonal SEBS/clay nanocomposites, prepared by extrusion, containing different nanoclay loadings: 2.5, 5 and 7.5 wt% (values for the 5 wt% nanocomposite are reported according to (Carastan et al., 2014))

Storage modulus, loss modulus and damping factor corresponding to SEBS_MA nanocomposites containing the 3 studied clay loadings are presented in Figure 2-4. In particular, Figure 2-4(c) shows that the highest glass transition temperature of the PS phase was exhibited by pure SEBS_MA (111 °C). However, all the nanocomposites have lower glass transition temperatures ranging from 107.5 °C for SEBS_MA_7.5% to 109 °C SEBS_MA_5%, which could be considered as a small difference. This remarkable increase of PS Tg in pure SEBS-MA could be related to the presence of MA. Furthermore, Figure 2-4(b) shows a remarkable decrease of the molecular relaxation peak of SEBS in the vicinity of PS glass transition upon addition of nanoclay (indicated by the arrow), which results in a decrease of the damping peak intensity as observed in Figure 2-4(c) and previously in Figure 2-2(c). Although the reduction of the damping peak occurred with increasing nanoclay loading, it can be seen that at 7.5 wt% nanoclay, only a slight additional decrease was induced compared to 5 wt% loading. This decrease is more likely attributed to the decrease of the loss modulus since no additional improvement of the storage modulus was observed at this loading (Figure 2-4(a)). This behavior may indicate that no further exfoliated

nanoparticles could be obtained at nanoclay loadings higher than 5 wt%. The present behavior is consistent with the result of quantitative analysis stating that the maximum exfoliation corresponds to 5 wt% nanoclay.



Figure 2-4: (a) Storage modulus E', (b) Loss modulus E'' and (c) Damping factor (tan δ) at 1 Hz of SEBS_MA nanocomposites containing 2.5, 5 and 7.5 wt% nanoclay (prepared by extrusion)

2.3.1.1 Nanoclay orientation

In order to characterize the orientation of nanoclays and PS micro-domains in the samples, SAXS analyses were performed. The SAXS diffraction patterns revealed that PS cylinders and clay nanoparticles were oriented in the extrusion direction for the extruded samples (see supporting file annex I). The TEM images presented in Figure 2-1(b)-(d) corresponding to the exfoliated samples clearly show evidence of this alignment while the TEM image

presented in Figure 2-1(a) and corresponding to the intercalated sample is not as conclusive. However, one should keep in mind that TEM images only probe a small area whereas SAXS present a more global analysis of the sample.

In the case of samples obtained by solution casting, for both the annealed and unannealed samples, the SAXS patterns revealed only concentric circles indicative of isotropic samples presenting random orientation of both PS domains and clay tactoids (see supporting file annex I).

2.3.1.2 SEBS morphology

The structure of extruded nanocomposites was studied in reference (Carastan et al., 2014). The authors reported that a hexagonal cylindrical morphology was present in both pure SEBS and the nanocomposites. The case of samples prepared by solution will be treated in this section. Figure 2-5(a) and Figure 2-5(b) present the radial plots of scattering intensity as function of scattering vector q of respectively unannealed and annealed samples. In Figure 2-5(a), corresponding to the unannealed samples, the relative q-position of Bragg peaks with the sequence 1:2:3 characteristic of lamellar structure was detected. This non-equilibrium morphology of alternating lamellae of PS and PEB domains was expected and attributed to the solvent effect since toluene dissolves better PS blocks compared to PEB blocks (Jeong et al., 2003), which favors the formation of PS lamellae and prevents their separation into cylindrical domains. In Figure 2-5(b) corresponding to annealed samples, the detected relative q-position of the peaks followed the sequence $1:\sqrt{3}:\sqrt{7}$, characteristic of hexagonal structure. This thermally induced order-to-order transition from lamellar to a more stable hexagonal cylindrical structure was reported in the literature as well when the annealing is performed at temperatures considerably higher than the PS glass transition temperature (Jeong et al., 2003). Furthermore, upon the addition of nanoclay, the third peak was broadened or disappeared from the spectra of both annealed and unannealed samples. This fact may indicate that the presence of nanoclay resulted in less well ordered SEBS structures compared to pure SEBS samples.



Figure 2-5: Radial plot of peak intensities of SEBS vs. SEBS_MA nanocomposites containing 5 wt% nanoclay (prepared by solution): (a) unannealed samples and (b) annealed samples

2.3.1.3 Stability of SEBS morphology

The morphological stability of unannealed nanocomposites prepared by solution was checked by analysing the relative peak positions in radial SAXS plots as a function of time at a temperature high enough to induce transition from lamellar to cylindrical morphology, selected equal to 200 °C in our case. This study was performed in order to determine whether the lamellar morphology in unannealed nanocomposites changed or not after pressing the samples at 200 °C, which was a necessary step to prepare specimens with comparable thicknesses for the dielectric breakdown measurements which will be discussed later. The plots corresponding to pure SEBS and SEBS nanocomposite with 5 wt% nanoclay are presented in Figure 2-6(a) and Figure 2-6(b). They show respectively the relative peak positions of pure SEBS and SEBS 5% clay nanocomposites as functions of time. Peaks corresponding to the diffraction of clay particles are observed as well in Figure 2-6(b). It could be seen that the first and the second peaks relative to lamellar structure in the radial plots changed of positions while the third peak was transformed in two small peaks (indicated by two arrows in the figures), after 5 minutes in the pure SEBS and after 10 minutes in the SEBS_5% clay nanocomposite. These new peaks correspond to the $(\sqrt{7}, \sqrt{9})$ peaks in radial plots of annealed samples presented in Figure 2-5(b). As a consequence, it could be concluded from this observation that a possible transition from lamellar to cylindrical morphology could have occurred during the molding process. It is also worth mentioning that the two small peaks are less apparent in the nanocomposite sample; which was also observed in the case of annealed nanocomposites (Figure 2-5(b)). Therefore, the initial morphologies of annealed and unannealed samples were also investigated by TEM. More details are provided in Table 2-2 and the supporting file reported in annex I.



Figure 2-6: Radial plots of SAXS peak intensities as function of time at 200 °C of: (a) unannealed pure SEBS and (b) unannealed SEBS_5wt% nanocomposite prepared by solution

Table 2-2 presents a summary of all the morphologies studied in this work. In particular, nanocomposites of type I and II prepared by extrusion exhibit an oriented hexagonal structure where both PS cylinders and nanoclays were oriented in the direction of extrusion while nanocomposites of type III and IV, prepared by solution casting method, exhibit both an isotropic structure where PS domains are distributed randomly within a PEB matrix. In type III nanocomposites, the hexagonal structure was induced due to thermal treatment while in type IV, the lamellar structure was dominating according to SAXS plots presented in Figure 2-5(a). However, it is worth mentioning that both lamellar and hexagonal morphologies could co-exist in the unannealed nanocomposites as indicated by the TEM image corresponding to type IV presented in Table 2-2. In particular, the cylindrical morphology is observed surrounding the nanoclays which may indicate that these particles favor the transition from lamellar morphology to more stable hexagonal morphology despite the fact that the samples were not subjected to thermal treatment (more details are available in the supporting file (annex I)). This observation is also consistent with the assumption of less ordered structure and the absence of the third peak in radial SAXS plots presented in Figure 2-5 and Figure 2-6.

Table 2-2: Summary of SEBS/clay nanocomposites with different structures (Red, yellow and green domains in the sketches illustrate respectively nanoclays, PEB phase and PS phase, in some sketches the yellow domains are removed only for the sake of simplicity)

	Processing		TEM	Morphology	Sketch
I	Extrusion	Figure al., 20 (Leice & Den	6(d) in (Carastan et 08) and Figure 3(b) in G. Amurin, Carastan, narquette, 2012)	 Hexagonal cylindrical SEBS structure Oriented PS cylinders and nanoclays Intercalated nanoclays crossing PS cylinders 	
п	Extrusion			 Hexagonal SEBS_MA structure Oriented PS cylinders and nanoclays Partially exfoliated nanoclays located within the PEB domain 	- Correction
III	Solution Casting + Thermal treatment	441.10		 Hexagonal SEBS structure Groups of PS cylinders dispersed randomly in PEB matrix Nanoclays located within the PS cylinders or the PEB domain depending on the use of SEBS or SEBS_MA (see supporting file annex I) 	
IV	Solution Casting			 Mostly lamellar SEBS structure (see supporting file annex I) Localized cylindrical morphology surrounding nanoclay particles PS domains randomly dispersed in PEB matrix Nanoclays located within PS domain or PEB domain depending on the use of SEBS or SEBS_MA 	

For the sake of simplicity, in the following figures related to dielectric properties, all the nanocomposites will be designated as following: $SEBS_W_X_Y_Z$ or $SEBS_MA_W_X_Y_Z$ where W is the wt% of nanoclay, X is the type of dispersion, Y is the orientation and Z is the structure of SEBS, as indicated in Table 2-3.

Symbol	Parameter	Abbreviation
W	Weight%	2.5, 5 or 7.5 wt% of nanoclays
X	Dispersion	exf (exfoliated), int (intercalated)
Y	Orientation	ori (oriented), ran (random)
Z	SEBS structure	hex (hexagonal), lam (lamellar)

Table 2-3: Designation of the nanocomposites

2.3.2 Dielectric properties

The dielectric permittivity of an insulating material measures its ability to polarize in response to an applied electric field. It is expressed as follows (Kremer & Schönhals, 2012):

$$\hat{\varepsilon}(\omega) = \varepsilon'(\omega) - i\varepsilon''(\omega) \tag{2.2}$$

Where:

ω: Frequency;

 ε ': real part of the complex dielectric permittivity, which is related to the stored energy. It is also known as the dielectric constant;

 ε '': imaginary part of dielectric permittivity, which is related to the dissipated energy. It is also known as the dielectric loss.

The complex dielectric permittivities of all SEBS/Clay nanocomposites were evaluated as function of the nanoclays' state of exfoliation and orientation as well as the block copolymer structure. It is worth mentioning that the measurements were performed perpendicular to the direction of orientation in the case of extruded samples as indicated in Figure 2-7. Figure 2-8(a) and Figure 2-8(b) represent respectively the real and the imaginary part of the complex

permittivity measured at room temperature of intercalated SEBS/clay vs. exfoliated SEBS MA/clay nanocomposites prepared by extrusion with different nanoclay loadings: 2.5, 5 and 7.5 wt%. In all the nanocomposites, an increase of the dielectric constant accompanied with a simultaneous increase of the dielectric loss is observed at low frequencies. This increase is attributed to Maxwell/Wagner/Sillars polarization at the nanoclay/polymer interfaces (Kremer & Schönhals, 2012). In particular, the increase achieved in nanocomposites prepared from SEBS MA matrix is more pronounced compared to the nanocomposites prepared from SEBS matrix indicating that more nanoclay/polymer interfaces are present in the first type of nanocomposites. Besides, a broad relaxation peak corresponding to the interfacial polarization appeared in the curves of nanocomposites with exfoliated structure in the frequency range 0.1 to 1 Hz. This is consistent with the fact that a better state of exfoliation is achieved when SEBS MA is used. In order to highlight the effect of exfoliation vs. intercalation of nanoclays on the dielectric behavior, the dielectric permittivities of respectively pure SEBS. SEBS 5wt% int ori hex and SEBS MA 5wt% exf ori hex nanocomposites were scanned as functions of frequency in the temperature range: 25 to 95 °C. In this temperature range, no relaxation was observed for the unfilled SEBS block copolymer (Figure 2-9(a) and Figure 2-9(b)) since the material is essentially non polar and therefore the relaxation process that can be seen in Figure 2-4 is not dielectrically active. However, the oriented SEBS 5% nanocomposite with intercalated structure exhibits a first relaxation mode below the Tg of PS phase which is attributed to the interfacial polarization. As the temperature increases, the increase of the dielectric constant at low frequencies is more evident (Figure 2-9 (c)) and the relaxation peak is shifted to higher frequencies (Figure 2-9(d)). In the graphs corresponding to the exfoliated nanocomposite (Figure 2-9(e) and Figure 2-9(f)), the interfacial phenomena is more important and corresponds to a clearer step in the dielectric constant curve and to more intense interfacial relaxation peak. The relaxation peaks in Figure 2-9(d) and Figure 2-9(f) are indicated by the arrows. Furthermore, the increase of dielectric loss observed at low frequencies and high temperatures due to nanoclays conduction contribution is one order of magnitude less in the nanocomposite with exfoliated structure compared to the nanocomposite with intercalated structure (Figure 2-9(d) vs. Figure 2-9(f)). This behavior could be explained by the fact that the conductivity at the interfaces between intercalated clay nanoplatelets is higher than the conductivity at the interfaces between the polymer and exfoliated clay nanoplatelets.



Figure 2-7: Orientation of electric field in dielectric spectroscopy and AC short-term breakdown measurements: (a) anisotropic nanocomposites prepared by extrusion and (b) isotropic nanocomposites prepared by solution casting



Figure 2-8: (a) Real and (b) imaginary parts of the complex permittivity at room temperature of oriented hexagonal SEBS (type I) and SEBS_MA (type II) nanocomposites containing 2.5, 5 and 7.5 wt% nanoclay



Figure 2-9: Real and imaginary parts of the complex dielectric permittivity in the temperature range from 25 to 95 °C of: Pure SEBS (a) and (b), SEBS_5wt%_int_ori_hex (c) and (d) and SEBS_MA_5wt%_exf_ori_hex (e) and (f)

As far as the concentration of nanoclay is concerned, the nanocomposites containing 5 wt% nanoclay exhibit the highest permittivities compared to the pure SEBS, SEBS MA and the nanocomposites containing 2.5 wt% and 7.5 wt% nanoclay. The decline of the permittivity at 7.5 wt% indicates that concentrations above 5wt% may result in poor dispersion of nanoclays. This behavior was observed also in DMA results (Figure 2-4). The same trend of increase of the dielectric constant was observed in the dielectric loss of the nanocomposites, which confirms that the increase observed in the dielectric constant is due to conduction phenomena along the nanoclay/polymer interfaces which leads to both an interfacial polarization peak and some low frequency dispersion. These experimental results are consistent with the results of quantitative image analysis stating that a higher amount of interface is achieved when a 5 wt% nanoclay loading is used. A similar behavior was reported in the studies done by Pirani et al. (Pirani, Krishnamachari, & Hashaikeh, 2014) and Lam et al. (Lam et al., 2005) who observed that the highest improvements in the mechanical properties of the polymer matrices were achieved at optimum nanoclay loadings. They attributed the decline of the properties above these optimum loadings to the deterioration of the dispersion quality.

Figure 2-10(a) and Figure 2-10(b) show respectively the real and imaginary parts of the dielectric permittivities of random hexagonal SEBS nanocomposites vs. oriented hexagonal SEBS nanocomposites measured at room temperature. It can be seen that the orientation of nanoclays affects the dielectric response. In fact, both the real and imaginary parts of the dielectric permittivity of non-oriented nanocomposites are higher than the values corresponding to oriented nanocomposites. To illustrate better this increase, the complex dielectric permittivity of non-oriented hexagonal SEBS_5%_int_ran_hex nanocomposite as a function of increasing temperature from 25 °C to 95 °C, is presented in Figure 11. A more pronounced increase of the dielectric permittivity (Figure 2-11(a)) and the dielectric loss (Figure 2-9(c) and Figure 2-9(d)) is observed at high temperatures. For instance, at 0.1 Hz and 95 °C, respectively the dielectric constant and the dielectric loss of the oriented vs. random sample were: 3.4 vs. 14.4 and 1.3 vs. 30.9. This difference of dielectric behavior between the

anisotropic nanocomposite and the isotropic nanocomposite at low frequencies and especially at high temperatures could be explained by the fact that the increase of the dielectric constant and the dielectric loss is essentially due to the conductivity of clay tactoids which is an anisotropic property. In fact, the conductivity along clay platelets is largely higher than across them (David et al., 2013). Hence, the alignment of nanoclays along the direction of extrusion in the oriented nanocomposites may result in a decreased conductivity along the thickness of the nanocomposites samples and consequently lower dielectric permittivities and dielectric losses compared to the isotropic samples when measurements are done perpendicularly to the preferred orientation direction.



Figure 2-10: Real (a) and Imaginary (b) parts of the complex permittivity at room temperature of oriented hexagonal vs. random hexagonal SEBS/clay nanocomposites



Figure 2-11: Real (a) and imaginary (b) parts of the complex permittivity of SEBS_5wt%_int_ran_hex nanocomposite with random hexagonal structure in the temperature range from 25 to 95 °C

In order to study the effect of SEBS structure on the dielectric response, the complex permittivity of unannealed mostly lamellar SEBS 5% clay nanocomposite prepared by solution method was evaluated and compared to its corresponding nanocomposite with random hexagonal morphology. Figure 2-12(a) and Figure 2-12(b) show respectively the real and the imaginary parts of dielectric permittivity of SEBS 5wt% int ran lam nanocomposite in the temperature range 25 to 95 °C. The results indicate that the nanocomposite with dominant lamellar morphology exhibits largely higher dielectric loss compared to the nanocomposite with hexagonal structure (Figure 2-11(a) and Figure 2-11(b)). For example, at 0.1 Hz and 95 °C, the values of dielectric constant were 14.4 vs. 19.5 and the values of dielectric loss were 30.9 vs. 98.0 respectively in hexagonal vs. lamellar SEBS 5% clay nanocomposites. Prior to determining the origin of this difference, it is worth mentioning that both random hexagonal pure SEBS and random lamellar pure SEBS exhibit dielectric responses similar to the dielectric response of oriented cylindrical SEBS plotted in Figure 2-9(a) and Figure 2-9(b) (the results are not presented in this paper). As a consequence, the observed difference in the dielectric behavior of the nanocomposites is related to how the morphology of SEBS affects the orientation and dispersion of nanoclays.

One possible explanation is that the lamellar morphology may hinder the orientation of nanoclays more than cylindrical morphology.



Figure 2-12: Real (a) and imaginary (b) parts of the complex permittivity of SEBS_5wt%_int_ran_lam nanocomposite with random lamellar structure in the temperature range from 25 to 95 °C

2.3.3 AC short-term breakdown strength

The breakdown strength is equal to the maximum electric field that an insulating material can withstand without breaking down ("IEEE Guide for the Statistical Analysis of Electrical Insulation Breakdown Data," 2005). In this paper, the retrieved AC breakdown data were treated by means of a two-parameter Weibull distribution to assess the evolution of the breakdown strength of the tested nanocomposites. The expression of the two-parameter Weibull distribution is shown in equation (2.3). More details regarding the use of Weibull distribution for dielectric strength estimation could be found in IEEE 930 standard ("IEEE Guide for the Statistical Analysis of Electrical Insulation Breakdown Data," 2005).

$$P = 1 - \exp\left[-\frac{E}{E_0}\right]^{\beta}$$
(2.3)

Where:

P: is the cumulative probability of failure at an electric field less or equal to E;

 E_0 : is the scale parameter corresponding to the breakdown strength for which the cumulative failure probability is equal to 63.2%;

 β : is the shape parameter which measures the range of breakdown strengths. The larger β is the smaller is range for breakdown strengths.

The Weibull plots of the experimental data corresponding to oriented SEBS/clay nanocomposites are displayed in Figure 2-13 while the Weibull parameters of all the nanocomposites are summarized in Table 2-4. The electric field was perpendicular to the direction of preferred orientation in the anisotropic samples, for breakdown measurements as well (Figure 2-7). The average thicknesses of the nanocomposites and the relative humidity were also reported since they may affect the measurements. Regarding the oriented samples, a tremendous increase of the breakdown strength was achieved by the addition of nanoclays within the pure SEBS. The maximum increase corresponds to the intercalated SEBS_5wt% nanocomposite by 45%. At 7.5wt% loading of nanoclay, the breakdown strength starts to decrease again. This behavior is analogous to the behavior seen in dielectric spectroscopy and dynamic mechanical analysis.



Figure 2-13: Weibull plot of breakdown strength of SEBS vs. SEBS/clay nanocomposites with oriented hexagonal structure

In the oriented samples with exfoliated nanoclay structure, prepared with SEBS_MA as a matrix, the breakdown strength starts to decrease from 5wt% loading of nanoclay after an initial increase by 35% corresponding to 2.5wt% nanoclay loading. This behavior could be explained by the fact that in the presence of a large amount of exfoliated nanoclay in SEBS_MA_5wt%_exf_ori_hex nanocomposite as demonstrated by image quantitative analysis, a network starts to form in some regions of the polymer and consequently reduces the breakdown voltage. Furthermore, the breakdown values of these 3 nanocomposites prepared from SEBS_MA are relatively lower than the rest of the nanocomposites due to probably lower Young's modulus of SEBS_MA compared to pure SEBS induced by its less ordered structure. The increased relative humidity when the breakdown measurements were performed on these samples could also be responsible for this unexpected decrease.

In terms of isotropic samples, it could be seen from Table 2-4 that the dielectric breakdown strengths of the random hexagonal samples, either pure SEBS or SEBS_5% nanocomposites are higher than the values measured for the oriented samples prepared by extrusion. However, the nanoclays didn't induce any increase of the breakdown strength of the pure

matrix unlike the case of the oriented samples. Hence, it seems that the breakdown phenomena in these materials are mostly governed by the isotropic structure of SEBS. In the same context of the influence of SEBS structure, it could be also seen from Table 2-4 that the breakdown strength values of the samples with dominant lamellar morphology are higher than samples with cylindrical morphology. The presence of lamellar morphology could be responsible for this increase of the breakdown strength due to the mechanical strengthening of the structure. However, the breakdown measurements could be also overestimated after the thickness normalization. In fact, the lamellar samples are in average thicker compared to the rest of the samples and also compared to the normalization thickness used for correction.

Nanocomposite	Morphology	E ₀ (kV/mm)	В	Average thickness (μm)	Relative humidity (%)
SEBS_0	Intercalated Oriented Hexagonal (Type I in Table 2-2)	88	7.3	108	22
SEBS_2.5		123	5.9	96	22
SEBS_5		127	5.6	98	22
SEBS_7.5		120	3	129	22
SEBS_MA_2.5	Exfoliated Oriented Hexagonal	119	5.3	118	31
SEBS_MA_5		109	6.3	97	31
SEBS_MA_7.5	(Type II in Table 2-2)	113	3.8	120	31
SEBS_0	Intercalated Random	133	3.7	110	23
SEBS_5	Hexagonal (Type III in Table 2-2)	135	4.1	122	23
SEBS_0	Intercalated Random	147	3.4	155	23
SEBS_5	Lamellar (Type VI in Table 2-2)	144	6.1	171	23

Table 2-4: Dielectric strengths of SEBS, SEBS_MA and their nanocomposites

To conclude, the oriented nanoclays seem to be more efficient in scattering electrons transport and increasing the breakdown strength. The increase of the breakdown induced by these samples is the highest compared to the isotropic samples where nanoclays were randomly dispersed. However, in terms of SEBS structure effect, the orientation of PS cylinders in SEBS is more likely to produce a negative effect which was not observed in the isotropic structure. This effect could be attributed to the reduction of Young's modulus and stiffness along the thickness of the oriented samples. In fact, Wang et al. (Wang, Fujinami, Liu, Nakajima, & Nishi, 2010) estimated the Young's modulus of respectively PS and PEB blocks of an SEBS grade similar to the one used in this study to be equal to: 133.3±26 MPa and 25.6±9.2 MPa. Hence, the orientation of PS cylinders could significantly decrease the modulus along the thickness of the samples which was proved to be a key parameter controlling the breakdown strength in dielectric elastomers (Kollosche & Kofod, 2010; Kollosche et al., 2009). More precisely, it was reported that a consistent reduction in Young's modulus and stiffness could lead to electromechanical breakdown (Dissado & Fothergill, 1992). This effect of mechanical strength is further confirmed by the observed increase of the breakdown strength in samples containing lamellar morphology which normally exhibits stronger equivalent Young's modulus (compared to the cylindrical morphology) due to the arrangement of PS phase with high Young's modulus in form of lamellae.

2.3.4 Summary of results

The main effects of structural changes in the morphology of SEBS/nanoclays on the complex dielectric permittivity and ac the breakdown strength are summarized in Table 2-5.

Morphology	Effect on dielectric properties		
Exfoliation vs. Intercalation	 Higher interfacial polarization in the exfoliated structures Relaxation peak in the dielectric loss of exfoliated structures Less improvement of the breakdown strength in the exfoliated structures. 		
Concentration of nanoparticles	 Highest dielectric constants and dielectric losses at 5wt% in both intercalated and exfoliated structures Highest breakdown strength at 5wt% in intercalated structure and at 2.5wt% in exfoliated structure 		
Orientation	 Oriented nanoclays Efficient scattering of electrons and increase of the breakdown strength by 45% Reduced dielectric losses at high temperatures Reduced breakdown strength of the pure SEBS due t reduction of mechanical strength perpendicular to th direction of orientation of PS cylinders Random nanoclays Negligible effect of random nanoclays on the breakdown strength High dielectric losses at high temperatures 		
Structure of SEBS	- Slightly higher breakdown strength in the lamellar structure due to the improvement of the mechanical strength		

 Table 2-5: Summary of different effects of SEBS/nanoclays morphology on the complex dielectric permittivity and breakdown strength

2.4 Conclusion

The dielectric properties and breakdown strength of SEBS/clay nanocomposites have been shown to be closely dependent on several structural properties such as the nanoclays' state of exfoliation, concentration and orientation as well as the structure of SEBS block copolymer. In particular, the exfoliation of nanoclays was responsible for the appearance of a relaxation peak caused by the interfacial polarization. Besides, the nanoclay orientation resulted in a significant reduction of the dielectric loss and a maximum increase of the breakdown strength by 45% compared to negligible or no increase induced by random nanoclays in isotropic nanocomposites. In these last nanocomposites, a beneficial effect from the random

distribution of PS domains in improving the mechanical strength and consequently the breakdown strength of the structure was rather seen and responsible for the increase of the breakdown strength of pure SEBS, especially in its lamellar structure compared to oriented SEBS. In all of the samples, it was demonstrated that a nanoclay loading of 2.5 to 5 wt% is an optimum content in terms of dispersion, number of nanoscale interfaces and dielectric performance. Finally, the results achieved so far in this study could be enhanced by studying the variation of the breakdown strength and dielectric loss in special morphologies where it is possible to take advantage of the orientation of nanoclays without degrading the mechanical strength of the SEBS matrix itself. An oriented lamellar morphology, a bi-oriented morphology or an isotropic morphology of PS cylinders with aligned nanoclays could represent interesting candidates in this context.

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CHAPTER 3

ARTICLE 2: INTERFACIAL MOLECULAR DYNAMICS OF STYRENIC BLOCK COPOLYMER-BASED NANOCOMPOSITES WITH CONTROLLED SPATIAL DISTRIBUTION

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Abstract

The dielectric properties of nanocomposites of polystyrene-*b*-poly(ethylene-*co*-butylene)-*b*-polystyrene (SEBS) triblock copolymers containing organically modified clay nanoparticles featuring controlled spatial orientation at the nanoscale: isotropic, totally oriented and partially oriented, have been investigated and correlated with the nanocomposite morphologies. A slow dielectric relaxation process attributed to elastomer chains with reduced mobility confined at nanoparticle/polymer interphase was observed in all the nanocomposites and was found to be dependent on the orientation of nanoclay and polystyrene (PS) domains, the location of clay tactoids as well as the PS block fraction. A dielectric "interfacial" glass transition temperature T_{gi} assigned to this characteristic

relaxation was estimated to occur at temperatures ranging between 6 °C and 35 °C depending on the nanocomposite, which is much higher than the bulk rubber phase glass transition temperature, normally lower than -40 °C for the studied block copolymers. Interestingly, the highest T_{gi} were associated with the nanocomposites featuring random or partial orientation and/or selective location of nanoparticles in the rubber phase.

Keywords: block copolymer nanocomposite, nanoclay, interphase, confinement, orientation, controlled dispersion, selective location

3.1 Introduction

In nanocomposite materials, controlled orientation of certain anisotropic nanoparticles such as nanoclay (Carastan et al., 2014; E Helal et al., 2015), carbon nanotubes (Goh, Ismail, & Ng, 2014) and recently boron nitride nanotubes and nanosheets (Z. Cui, Cao, Ma, Dobrynin, & Adamson, 2015; Terao et al., 2010) is very beneficial for a wide spectrum of applications requiring excellent mechanical, electrical and/or thermal properties. For example, in the case of mechanical reinforcement, the alignment of nanoclay was reported in several publications to induce an improved mechanical strength in the alignment direction (Galgali, Agarwal, & Lele, 2004). In the specific case of nanodielectrics, more efficient electron scattering and consequently higher breakdown strength perpendicular to the nanoparticles alignment direction were reported (David et al., 2013; Fillery et al., 2012; E Helal et al., 2015; Tomer et al., 2011). Simultaneously, nanoparticle alignment was shown to reduce dielectric losses in the direction perpendicular to the main plane of the aligned nanoparticles (E Helal et al., 2015). This controlled orientation can be obtained using a wide range of techniques (Goh et al., 2014) although spatial alignment of nanoparticles according to 2D and 3D patterns is still a challenging field of study (Richard A Vaia & Maguire, 2006).

In applications requiring tuned spatial distribution of nanoparticles, the use of block copolymers can be really an asset due to the different nanoscale morphologies these materials present (Hamley, 2001; Helfand, 1975; Ohta & Kawasaki, 1990; Sarkar & Alexandridis, 2015; Semenov, 1993). To probe the effect of tailored morphology and orientation of such
designed nanocomposites on their polymer-filler interactions and implicitly on their final performance, techniques such as broadband dielectric spectroscopy (BDS) are often required (Kremer & Schönhals, 2012).

In fact, the performance of polymer nanocomposites in general is governed by the interphase region (Roy et al., 2005; Leszek A Utracki, 2010) which consists mainly of a bound layer where the motion of macromolecular chains is strongly restricted affecting several properties including dielectric and mechanical properties (Hernández, Carretero-González, Verdejo, Ezquerra, & López-Manchado, 2010; Qu et al., 2011; Leszek A Utracki, 2010; Vo et al., 2011). The thickness and volume fraction of this interphase usually depend on the geometry of the nanoparticles and their compatibility with the polymer matrix (L. Chen et al., 2009; Klonos, Kyritsis, & Pissis, 2015; Yu Lin et al., 2015; Robertson & Rackaitis, 2011). It was estimated using both experimental techniques (L. Chen et al., 2009; Yu Lin et al., 2015) and molecular dynamics simulations (Gao, Liu, Zhang, & Cao, 2014; Ghanbari, Rahimi, & Dehghany, 2013). Furthermore, an additional glass transition corresponding to the interfacial polymer chains with restricted mobility was observed in some nanocomposites featuring strong attractive interfacial interactions (L. Chen et al., 2009; Y. Gao et al., 2014; Ghanbari et al., 2013; Hernández et al., 2010; Hernández, del Mar Bernal, Verdejo, Ezquerra, & López-Manchado, 2012; Holt et al., 2014; Klonos et al., 2015; Yu Lin et al., 2015; Robertson & Rackaitis, 2011; Tsagaropoulos & Eisenberg, 1995; Tsagaropoulos & Eisenburg, 1995; Vo et al., 2011) and predicted by modeling and simulations (Starr, Schroder, & Glotzer, 2002) as well, for relatively thick bound layers.

The reduced mobility of polymer chains in the interphase region of homopolymer-based nanocomposites has been well investigated during the last years, especially the effect of interaction strength between nanoparticles and the polymer matrix. However, to the best of our knowledge, there is only little literature regarding the interphase region in block copolymer based nanocomposite systems and specifically the effect of the orientation of nanoparticles and block copolymer nanodomains on polymer dynamics in this interphase region. In fact, in the case of nanocomposites prepared from multicomponent polymer

matrices, an additional degree of complexity is added to the system as the nanofillers can interact differently with the constitutive components (Sarkar & Alexandridis, 2015; Vo et al., 2011). These interactions are usually interdependent and hard to quantify separately. Few studies investigated the dielectric behavior of copolymers and its dependence on chemical and structural factors such as sulfonation (H. Chen et al., 2011) and compatibility with different nanofillers. In this context, Vo et al. (Vo et al., 2011) studied the dielectric behavior of styrene-butadiene rubber (SBR) random copolymer filled with three different nanoparticles: nanoclay, silica and carbon black. They reported a new relaxation mode for all three systems attributed to the segmental motion of rubber chains with reduced mobility at the polymer-nanoparticle interface. Moreover, they evaluated an interfacial glass transition temperature T_{gi} associated with this relaxation process. The highest T_{gi} was attributed to the soft strenger interaction and better compatibility compared to the two other fillers.

In this work, we investigated polystyrene-b-poly(ethylene-co-butylene)-b-polystyrene (SEBS) thermoplastic elastomer as the block copolymer matrix since it presents excellent features suitable for many applications, such as good mechanical properties (Balsamo et al., 2006; Holden et al., 2004), good resistance to water treeing (Liu et al., 2011; Z. Ma et al., 2010) as well as good electromechanical coupling (B. Kim et al., 2011). It is a symmetric triblock copolymer composed of two polystyrene (PS) end-blocks of the same length at the extremities and a poly (ethylene-co-butylene) (PEB) rubber mid-block. Organically modified nanoclays, known for their beneficial effect on mechanical and dielectric properties of polymer nanocomposites including breakdown strength, resistance to surface erosion and reduction of space charge accumulation (David et al., 2013; Fillery et al., 2012; MF Fréchette et al., 2008; Galgali et al., 2004; E Helal et al., 2015; Tomer et al., 2011; Leszek A Utracki, 2010), were added to the thermoplastic elastomer. Four sets of block copolymer nanocomposites containing three different weight fractions of PS phase (0.13, 0.20 and 0.30) were investigated in total. Depending on these ratios and using several fabrication processes, different morphologies were successfully prepared: isotropic vs. totally oriented vs. partially oriented polystyrene nanodomains and clay nanoparticles. Moreover, the degree of exfoliation and location of clay particles inside PS or PEB domains were tailored using a specific SEBS grade with a maleic anhydride (MA) grafted on the PEB block.

In a first step, the orientation of nanoclay and PS nanodomains in the different samples as well as the state of order of the block copolymer were fully characterized by SAXS and TEM. In a second step, the dielectric response was studied as a function of frequency and temperature in order to investigate the influence of the block copolymer tuned architecture on the polymer dynamics, which may affect implicitly the engineering properties such as dielectric losses, breakdown strength and mechanical stiffness, to name a few. The dielectric spectroscopy results were correlated to SAXS and TEM results in order to come up with a template of block copolymer nanodielectrics (M. F. Frechette, Trudeau, Alamdar, & Boily, 2004) with controllable morphology and properties suitable for different dielectric applications. In particular, a slower dielectric relaxation mode compared to the main relaxation responsible for the bulk glass transition of the rubbery phase was observed in all nanocomposites and attributed to the segmental motion of rubber chains with reduced mobility located at the polymer-nanoparticle interphase. An interfacial glass transition T_{gi} associated with this new relaxation mode was estimated and used to quantify the interaction strength between the nanoparticles and the polymer chains depending on the configuration, the styrene block content and the location of the nanoparticles in one block or another. Besides, this new dielectric relaxation process, attributed to rubber chains located at the interfacial layer, was also observed in the dynamic mechanical responses of samples prepared from SEBS grade containing 30 wt% of PS block. Finally, the thickness of the interfacial layer was estimated in the range 7-10 nm.

3.2 Materials and methods

Four grades of symmetric triblock copolymer SEBS donated by Kraton were used: G1643, G1645, G1652 and FG1901. These grades contain different fractions of PS block ranging from 13 to 30 wt%. Besides, the FG1901 grade contains 1.4-2 wt% of maleic anhydride (MA) group attached to its elastomeric PEB block. All the grades contain 0% of diblocks. More details regarding the physical properties and morphologies of these polymers are

reported in Table 3-1. Montmorillonite clay grade Cloisite 20A purchased from Southern Clay (Gonzales, USA) and modified with dimethyl di(hydrogenated tallow) quaternary ammonium salt, was used as nanofiller for the nanocomposites preparation. More details regarding the physical properties of the polymers as well as the modification and size of clay particles were provided in the previous studies published by co-authors in the same context of research (Leice G Amurin, Carastan, & Demarquette, 2016; Carastan et al., 2014; De Sousa Jr, Amurin, Demarquette, & Carastan, 2014; E Helal et al., 2015). All the materials studied in this paper were used as received.

Table 3-1: Properties of SEBS-13, SEBS-20 and SEBS-30

Block copolymer	Grade	PS wt%	MFI (g/10min)	ρ ^c (g/cm ³)	T _{OOT} d (°C)	Т _{орт} е (°С)
SEBS-20	G1643	20	18 ^a	0.9	always cylindrical	> 200
SEBS-13	G1645	13	40 ^b	0.9	between 150 and 160 (transition to spherical)	190- 200
SEBS-30	G1652	30	5 ^b	0.91	always cylindrical	>200
SEBS-30-MA	FG1901*	30	22 ^b	0.91	always cylindrical	>200

^a melt flow index measured at 230°C/2.16 Kg, ASTM D1238, provided by the supplier ^b melt flow index measured at 230°C/5 Kg, ASTM D1238, provided by the supplier ^c specific gravity, provided by the supplier

^d order to order transition (OOT) estimated from small amplitude oscillatory shear (SAOS) and small angle X-ray scattering (SAXS) measurements not presented in this manuscript.

^e order to disorder transition temperatures (ODT) estimated from small amplitude oscillatory shear (SAOS) and small angle X-ray scattering (SAXS) measurements not presented in this manuscript.

FG1901 grade contains 1.4-2 wt% of maleic anhydride (MA)

The nanocomposites were prepared by solvent casting, sheet die extrusion or film blowing extrusion to achieve different morphologies. In the case of solvent casting process, the SEBS powder and the nanoparticles were mixed in toluene by magnetic stirring at 60 °C. The mixture was subsequently poured into a Petri dish and left in open air under the fume hood

for several days until complete evaporation of solvent. They were subsequently dried in a vacuum oven and annealed according to specific temperature profiles selected depending on the styrene content, in order to reach equilibrium morphologies (Hadjichristidis, Hirao, Tezuka, & Du Prez, 2011). For SEBS-13 and SEBS-20, the sequence: 10 minutes at 50 °C, 30 minutes at 110 °C, 2 hours at 150 °C and 24 hours at 60 °C was used while for SEBS-30 and SEBS-30-MA, the sequence 20 min at 60 °C, 20 min at 100 °C, 20 min at 150 °C and 10 min at 200 °C was used as published in reference (E Helal et al., 2015). The samples prepared by sheet die extrusion were obtained according to references (Leice G Amurin et al., 2016; Carastan et al., 2014). In a typical procedure, clay nanoparticles were mixed with SEBS in a twin screw extruder equipped with a sheet die, at a temperature profile ranging from 160 to 190 °C for SEBS-20, equal to 150 °C for SEBS-13 and equal to 200 °C for SEBS-30 and SEBS-30-MA and a screw speed of 100 rpm. The films prepared from SEBS-20 by film blowing extrusion were processed in a first step following the same procedure of sheet die extrusion. Subsequently, the samples were processed in a single screw extruder using the same temperature profile and a screw speed equal to 30 rpm. The latter was connected to an annular die with controllable air pressure inside the tube in order to induce lateral elongation and promote biaxial orientation of the block copolymer. In this study, the resulting films were inflated with air at two blow-up ratios: R1=1 and R3=3 in order to initiate respectively uniaxial and biaxial orientation directions. More details regarding this processing method are available in reference (De Sousa Jr et al., 2014) published by coauthors. The sheet die extrusion and the film blowing extrusion processes are illustrated in Figure 3-1.



Figure 3-1: Illustrations showing: (a) sheet die extrusion, (b) film blowing with ratio R1= 1 and (c) film blowing with ratio R3= 3 (Coordinate system used as reference is indicated at the left of the figure)

The samples prepared by solvent casting were used without any other treatment for further characterization. The samples prepared by sheet die extrusion and film blowing extrusion were pressed for 2 min at 150 °C for SEBS-13 and SEBS-20 and at 200 °C for SEBS-30 and SEBS-30-MA, under 10 tons before further characterization, in order to have films with comparable thicknesses. The average thickness of all the films was around 550 μ m. Short time and relatively low temperatures were used to avoid possible changes in morphology or transition to disordered state (as what will be shown later in the manuscript). The list of samples and their nomenclature are presented in Table 3-2.

Polymer	Nanoparticles wt%	Porcessing method	Nomenclature
SEBS-20	0	Sheet die extrusion	SEBS-20-extrusion
SEBS-20	5	Sheet die extrusion	SEBS-20-20A-extrusion
SEBS-20	0	Solvent casting	SEBS-20-solution
SEBS-20	5	Solvent casting	SEBS-20-20A-solution
SEBS-20	0	Film blowing ratio R1	SEBS-20-film blowing R1
SEBS-20	5	Film blowing ratio R1	SEBS-20-20A-film blowing R1
SEBS-20	0	Film blowing ratio R3	SEBS-20-film blowing R3
SEBS-20	5	Film blowing ratio R3	SEBS-20-20A-film blowing R3
SEBS-13	0	Sheet die extrusion	SEBS-13-extrusion
SEBS-13	5	Sheet die extrusion	SEBS-13-20A-extrusion
SEBS-13	0	Solvent casting	SEBS-13-solution
SEBS-13	5	Solvent casting	SEBS-13-20A-solution
SEBS-30	5	Sheet die extrusion	SEBS-30-20A-extrusion
SEBS-30	5	Solvent casting	SEBS-30-20A-solution
SEBS-30-MA	5	Sheet die extrusion	SEBS-30-MA-20A-extrusion

Table 3-2: Nomenclature of nanocomposites

It is worth noting that the morphology and the dielectric response of the materials prepared from the SEBS-30 and SEBS-30-MA grades have been already reported in previous studies (Leice G Amurin et al., 2016; Carastan et al., 2014; E Helal et al., 2015). In particular, it was demonstrated, by TEM and XRD (Carastan et al., 2014), that in the presence of MA, clay particles are mostly exfoliated and located in the PEB phase while in its absence an intercalated clay structure is dominant and clay tactoids cross PS domains. At increasing clay loading up to 7.5wt%, both the average distance between (100) planes of the cylindrical structures and the diameter of PS cylinders increase (Leice G Amurin et al., 2016).

In terms of dielectric properties, it was shown that the alignment of PS cylinders results in an initial decrease of the breakdown strength that was compensated by the improvement induced through the alignment of clay tactoids. The highest increase was equal to 45% compared to unfilled and aligned SEBS-30, at 5wt% loading of clay. In this manuscript, the frequency-domain dielectric response of these samples will be investigated in terms of comparison.

3.3 Characterization

The morphology of the as-obtained nanocomposites was characterized by small angle X-ray scattering (SAXS) and transmission electron microscopy (TEM). Subsequently, the dielectric properties and the dynamic mechanical properties were characterized by means of respectively broadband dielectric spectroscopy (BDS) and dynamic mechanical analysis (DMA).

3.3.1 Small Angle X-ray Scattering

The SAXS patterns were obtained using Bruker NanoSTAR with 1.5 kV CuKa radiation. The sample-to-detector distance was 650 nm. Besides, bidimensional detectors were used in order to evaluate potential anisotropic features in the samples. The data were analyzed using Fit2D software, without any background subtraction.

3.3.2 Transmission Electron Microscopy

The TEM was performed with a JEOL 2100F microscope. Prior to observations, ultrathin sections of few tens of nanometers were cryo-cut using an ultra-microtome operated at -100 °C and deposited on copper grids. The grids were also stained during 30 minutes with ruthenium tetroxide RuO₄ vapor in order to determine the block copolymer morphology and to evaluate the location of clay nanoparticles.

3.3.3 Broadband dielectric spectroscopy

In terms of dielectric properties, the complex dielectric permittivity of the nanocomposites was measured using a Novocontrol broadband spectrometer. Specimens of 20 mm in diameter were placed between two parallel brass plated electrodes. Measurements swept through a frequency range from 10^{-2} Hz up to 10^{5} Hz at a temperature range varying from 25 °C to 90 °C at 5 °C steps, under an rms excitation voltage of 1 V.

3.3.4 Dynamic mechanical analysis

The measurements were performed using a TA Q800 Instrument operated in tensile mode. All the scans were done from -100 °C to 130 °C. The test conditions were as follows: rate of heating equal to 3 °C/min, resonant frequency equal to 1 Hz and strain amplitude equal to 0.1%. The tensile storage modulus (E'), loss modulus (E'') and damping factor (tan δ) were evaluated.

3.4 Results and discussion

3.4.1 Morphology

SAXS analysis was performed on every sample before and after pressing in order to check the type (cylindrical, spherical, lamellar...) and the stability of the morphology as well as the orientation of both PS domains and clay nanoparticles. The diffraction patterns, before and after pressing, were similar, indicating that the compression molding did not induce any significant change in the initial morphologies. In addition, SAXS patterns were recorded at different temperatures up to 140 °C in order to check for changes in the morphology and state of order that might be encountered during thermal annealing or dielectric spectroscopy scans, as those ones were carried out as a function of temperature. The relative plots indicating peak positions were integrated as well from the 2D patterns. In the following sections, the different samples are analyzed and classified depending on PS block and clay nanoparticles orientations in three different categories: totally isotropic, totally aligned and partially aligned morphologies.

3.4.1.1 Totally isotropic morphologies

Isotropic morphologies with totally random orientation of PS nanodomains and clay nanoparticles were obtained in all the samples prepared by solvent casting independently from the polystyrene block content (13, 20 or 30 wt%). Typical diffraction patterns corresponding to SEBS-13-solution and SEBS-20-solution materials are reported in Figure A

II-1 of the supporting file (annex II). The patterns feature concentric rings, in the 3 directions of measurement indicating isotropic orientation of both PS domains and nanoclay. In the case of the pure copolymer, the concentric rings are related to the scattering from PS domains while in the nanocomposite, this scattering is overlapped with more intense scattering from clay nanoparticles. Similar features were observed in nanocomposites prepared from SEBS-30-solution (SAXS data available in Figure A I-4 of annex I relative to (E Helal et al., 2015)). TEM micrographs corresponding to SEBS-30-20A-solution nanocomposite are reported in Figure A II-5 of the supporting file (Annex II). They confirm the isotropic distribution of clay tactoids and PS cylinders in samples prepared by solvent casting method.

Figure 3-2 presents SAXS radial plots of scattering intensity as a function of scattering vector q corresponding to neat materials and nanocomposites prepared by solvent casting from SEBS-20 and SEBS-13 matrices. Results regarding samples prepared from SEBS-30 were already published in a previous study (E Helal et al., 2015). The profiles were integrated for 5 different temperatures ranging from 25 °C to 140 °C. In Figure 3-2(a) and Figure 3-2(b) corresponding to respectively neat SEBS-20-solution and its nanocomposite, the relative qpositions of Bragg peaks follow the sequence $1:\sqrt{3}:\sqrt{7}$ characteristic of a hexagonally packed cylindrical structure. It can be seen that the intensity of the second peak increased slightly with increasing temperature, as indicated by the arrows, which infers a positive effect of thermal annealing in reaching equilibrium morphologies and consequently improving longrange order (Hadjichristidis et al., 2011). In Figure 3-2(c), corresponding to the neat SEBS-13-solution, the same features were observed. However, in the case of the nanocomposite (Figure 3-2(d)), the first peak was detected at the same position but the second peak was very weak or completely absent. In addition, no obvious improvement was observed with increasing temperature. These results indicate that the state of order is altered in the presence of clay nanoparticles and does not improve with annealing. In particular, this behavior might indicate improved dispersion and degree of intercalation that, in addition to random nanoparticles distribution, help to freeze the block copolymer structure even when subjected to thermal annealing and to reduce the mobility of polymer chains located at the interphase. A higher clay interlayer spacing, d, and consequently intercalation degree has been already

reported for nanocomposites prepared by solvent casting from SEBS-13 (d=3.59 nm) compared to nanocomposites prepared from SEBS-30 (d=3.09 nm), both containing Cloisite 15A nanoparticles (Carastan et al., 2008). This observation will be correlated later with dielectric spectroscopy results.



Figure 3-2: Scattering intensity as function of scattering vector q at different temperatures of: (a) SEBS-20-solution and (b) SEB-20-20A-solution nanocomposite, (c) SEBS-13-solution and (d) SEBS-13-20A-solution nanocomposite

3.4.1.2 Totally aligned morphologies

Diffraction patterns corresponding to samples prepared by sheet die extrusion from SEBS-20, SEBS-30 and SEBS-30-MA are not shown in this manuscript since they exhibit totally aligned morphologies in the extrusion direction. Typical patterns were previously reported by co-authors in reference (Carastan et al., 2014) for SEBS-30 materials. TEM micrographs of SEBS-30-MA-20A-extrusion nanocomposites exhibiting aligned and mostly exfoliated clay layers are reported in Figure A II-6 of the supporting file (annex II). Aligned morphologies were not obtained in the case of SEBS-13-extrusion as this material has limited tendency to align due to the low fraction of PS block (Leice G Amurin et al., 2016). The case of SEBS-13-extrusion will be treated in the following section within partially aligned morphologies.

3.4.1.3 Partially aligned morphologies

In addition to totally random morphologies and totally aligned morphologies, some intermediate configurations were successfully achieved depending on the content of styrene and the process used. More specifically, materials prepared from SEBS-20 by film blowing extrusion exhibit two distinct morphologies depending on the blowing ratio: R1=1 or R3=3. Figure 3-3 shows the diffraction patterns ((a)-(b)) and the TEM images ((c)-(e))corresponding to SEBS-20-film blowing R1 and its nanocomposite SEBS-20-20A-film blowing R1. In particular, the sections examined by TEM were cut perpendicular to the flow direction (i.e. parallel to XZ plane indicated in Figure 3-1). Diffraction patterns corresponding to neat SEBS-20-R1, presented in Figure 3-3(a), show evidence of hexagonal packing of PS cylinders in the Y direction while in the X and Z directions, two sets of Bragg peaks can be observed indicating the alignment of the majority of PS cylinders parallel to Y axis (the initial direction of extrusion as indicated in Figure 3-1). Relative to nanocomposites, diffraction patterns corresponding to SEBS-20-20A-R1 (Figure 3-3(b)) show strong diffuse spots related to the scattering of nanoclay along the X and Y directions, which indicates dominant alignment of nanoclay parallel to the XY plane (the initial direction of the flow). The presence of maxima in the diffuse spots is characteristic of intercalated nanoclay

structure, which was confirmed by X-ray diffraction results reported by co-authors for similar samples (Leice G Amurin et al., 2016; Carastan et al., 2014). TEM images confirm preferential alignment of PS domains and nanoclay in the main direction of the flow (when PS domains appear as small well-defined circles as indicated by the full-line white rectangle in Figure 3-3(e)). Nevertheless, they reveal the appearance of additional directions of orientation for both PS cylinders and nanoclay, in less important fractions though. In some regions, a fraction of PS cylinders rotated across the thickness (as indicated by the white dotted rectangles in Figure 3-3(e)). However, the angle of orientation is in most of the cases lower than 90°, meaning that they do not succeed in orienting perfectly along the thickness. Besides, less strong scattering is also observed in the Z direction of the nanocomposite diffraction patterns (Figure 3-3(b)), which indicates orientation of some clay tactoids in the YZ plane (across the thickness of the sample). This observation is consistent with the TEM image of Figure 3-3(c) showing that some small clay nanoparticles (indicated by the arrows) are aligned perpendicular to the main orientation of bigger tactoids. This effect could be promoted by the presence of a normal force (parallel to Z axis) induced by the tubular die. However, the orientation of a small fraction of clay particles in directions other than the main flow direction was reported even for samples prepared by sheet die extrusion (Carastan et al., 2014). Radial plots of scattering intensity as a function of scattering vector q of the neat material and the nanocomposite are presented in Figure A II-2 of the supporting file (annex II). The $1:\sqrt{3}:\sqrt{7}$ sequence characteristic of hexagonal structure is observed in both the neat and the nanocomposite confirming the dominance of hexagonal cylindrical structure.



Figure 3-3: (a)-(b) 2D SAXS patterns of: (a) SEBS-20-film blowing R1 and (b) SEBS-20-20A-film blowing R1, (c)-(e) TEM images of SEBS 20-20A- film blowing R1 at 3 different magnifications showing imperfect alignment of PS cylinders and clay tactoids: (d) zoom on the section defined by black square in (c), (e) the white square indicates regions featuring perfect alignment of PS cylinders in the extrusion direction while dotted white rectangles indicate other directions of alignment (PS domains were stained with RuO4 (dark phase))

Figure 3-4 shows SAXS diffraction patterns and TEM images of neat SEBS-20-film blowing R3 and its nanocomposite SEBS-20-20A- film blowing R3. The diffraction patterns, presented in Figure 3-4(a)-(b), reveal the existence of a less perfect hexagonal packing of PS cylinders along the Y axis, compared to SEBS-20- film blowing R1. Besides, Bragg peaks featured in the X and Z directions are wide and almost form concentric rings indicating deviation from the dominant alignment of cylinders along the Y direction (as observed in SEBS-20-R1) and formation of a network of isotropic PS cylinders parallel to the XY plane due to the high blowing ratio and consequently important lateral forces. This observation is further supported by the TEM images presented in Figure 3-4(c)-(d), showing that more PS

domains are oriented in isotropic directions different from the initial flow direction. In Figure 3-4(b), corresponding to the diffraction pattern of SEBS-20-20A-film blowing R3 nanocomposite, similar features could be observed in the X and Y directions with more intense and narrow scattering indicating more perfect alignment of nanoclay parallel to the XY plane. However, by contrast to SEBS-20-20A-film blowing R1, no specific scattering related to the orientation of clay tactoids in the Z direction is detected. This fact is most likely due to the high lateral forces induced by the R3 blowing ratio that are able to overcome the normal forces applied by the tubular die. Only lateral movements of nanoclay should be possible; thus alignment in the initial direction of the flow is maintained. The radial plots corresponding to SEBS-20-film blowing R3 and its nanocomposite SEBS-20-20A-film blowing R3, presented in Figure A II-3 of the supporting file (annex II), exhibit again the sequence $1:\sqrt{3}:\sqrt{7}$ characteristic of hexagonal structure. However, it is worth noting that the intensities of the peaks decreased, compared to R1 samples, which might indicate a lack of order due to the increased stretching induced by the high blowing ratio. Moreover, the effect of increasing temperatures in improving the intensities of the peaks is less obvious in these samples, which might indicate a permanent alteration in the block copolymer ordered state due to the important applied deformations. All these observations regarding lack of long range order and alteration of initial alignment in the extrusion direction are in a good agreement with previous results reported by co-authors in references (Leice G Amurin et al., 2016; Carastan et al., 2013), stating that samples where initially aligned PS cylinders are subjected to high strain values in the transversal direction exhibit misalignment due to the rotation of PS domains that tend to align parallel to the deforming force direction. However, this rotation is incomplete due to the competition between the alignment of PS domains and copolymer molecules which don't have enough time to relax. This stretching results as well in breakage of cylinders in shorter domains and reduced long range order. More details regarding these phenomena could be consulted in the cited references (Leice G Amurin et al., 2016; Carastan et al., 2013).



Figure 3-4: (a)-(b) 2D SAXS patterns of: (a) SEBS-20-film blowing R3 and (b) SEBS-20-20A- film blowing R3 nanocomposite, (c)-(d) TEM images of SEBS-20-film blowing R3 at different magnifications (PS domains (darker phase) were stained with RuO4)

As it was mentioned before, the materials prepared from SEBS-13 matrix by sheet die extrusion do not exhibit totally aligned morphologies (Leice G Amurin et al., 2016). Therefore, their case will be discussed in this section. SAXS diffraction patterns and radial plots corresponding to these samples were integrated, in a similar trend to the previous samples, and reported in Figure A II-4 of the supporting file (annex II).

In the diffraction patterns (Figure A II-4(a)-(b) of annex II), only the Z direction is reported for the case of the pure material, as the 3 directions were similar. In particular, concentric rings related to the scattering from the PS phase were observed in the 3 directions, for both the neat copolymer and the nanocomposite. This fact indicates that the majority of PS domains were not able to orient by the applied shear forces during extrusion and remain randomly distributed due to the low polystyrene fraction. However, the scattering from the nanoclay features direction dependence (Figure A II-4(b) of annex II). In fact, 2 strong spots aligned parallel to Z direction corresponding to Bragg peaks associated with scattering from nanoclay were observed in the XZ and YZ planes. These signals indicate the alignment of nanoclays parallel to the extrusion direction. Since this alignment may potentially induce simultaneous orientation of a fraction of PS cylinders, a partially isotropic PS/partially oriented PS configuration will be rather considered for the case of SEBS-13-20A-extrusion nanocomposite. Radial plots (Figure A II-4(c)-(d) of annex II) reveal similar characteristics compared to the samples prepared by solvent casting in terms of dominance of hexagonally packed cylindrical morphology and evolution of order degree with increasing temperature and presence of nanoparticles.

In Table 3-3, a summary of all the studied samples, a brief description of their morphologies, i.e. the orientation of clay tactoids and PS cylinders, as well as the corresponding illustrations of the morphologies are presented. The nomenclature of the samples is also updated to recall the concluded morphologies. In particular, the samples that are neither totally aligned nor random will be referred to as "partially oriented" to indicate the presence of more than one preferential orientation of PS cylinders or clay tactoids or both of them. The direction of the applied electric field that was used for dielectric spectroscopy measurements, discussed in the following section, is specified in one of the illustrations as well and remains valid for all the samples.

It is worth noting that since the achieved orientations of PS cylinders and clay particles are dependent on the processing technique, the obtained morphologies are metastable and may change if the samples are subjected to additional processing steps. Therefore, a verification of the morphology is needed if further steps are required. For instance, some studied samples in this manuscript were hot pressed to obtain films of a precise thickness. To avoid potential modification of the morphology, the pressing time and temperature were limited as possible. The stability of the morphology after this step was checked by SAXS.

	Nomenclature	Morphology description	Morphology sketch
sion	SEBS-20-oriented SEBS-30-oriented SEBS-30-MA-oriented	Uniaxially oriented PS cylinders in the flow direction along Y axis	Clay
Extru	SEBS-20-20A-oriented SEBS-30-20A-oriented SEBS-30-MA-20A- oriented	Uniaxially oriented PS cylinders and nanoclay in the flow direction along Y axis	PEB PS
asting	SEBS-13-isotropic SEBS-20-isotropic SEBS-30-isotropic	Isotropic PS cylinders	
Solvent c	SEBS-13-20A-isotropic SEBS-20-20A-isotropic SEBS-30-20A-isotropic	Isotropic PS cylinders and nanoclay	field
	SEBS-20-partially oriented R1	Mainly oriented PS cylinders in the flow direction along Y axis Minor fraction oriented across the thickness along Z axis	
blowing	SEBS-20-20A- partially oriented R1	Mainly oriented PS cylinders and nanoclay in the flow direction along Y axis Few PS cylinders and nanoclay oriented across the thickness parallel to YZ plane	
Film	SEBS-20- partially oriented R3	Isotropic orientation of cylinders in the flow direction parallel to XY plane	
	SEBS-20-20A- partially oriented R3	Oriented nanoclay in the flow direction along Y axis Isotropic orientation of cylinders in the flow direction parallel to XY plane	
ısion	SEBS-13- partially oriented	Partially isotropic PS cylinders Partially oriented PS cylinders in the flow direction along Y axis	
Extru	SEBS-13-20A- partially oriented	Partially isotropic PS cylinders Partially oriented PS cylinders in the flow direction along Y axis Oriented nanoclay in the flow direction along Y axis	

Table 3-3: Nomenclature, morphology description and morphology sketch of the nanocomposites

3.4.2 Dielectric properties

The complex dielectric permittivity of an insulating material measures its response to an applied electric field either through polarization mechanisms or charge carrier fluctuations (Kremer & Schönhals, 2012). The corresponding equation is described in the supporting file (equation (A II.1) in annex II).

In this manuscript, the dielectric response of the studied materials has been characterized as functions of frequency and temperature. To better understand the origin and the dynamics related to the relaxation modes observed in each material as well as to evaluate the contribution from charge fluctuations, depending on the orientation of PS cylinders and nanoclay, dielectric permittivity spectra of all nanocomposites were fitted according to equation (A II.2) (Kremer & Schönhals, 2012), described in annex II, which comprises a power law term to describe the contribution of charge fluctuations (A.K. Jonscher, 1983; Andrzej K Jonscher, 1996) and a sum of Havriliak-Negami (HN) functions to take into account the observed dipolar dielectric relaxation processes (Havriliak & Negami, 1967; Hernández et al., 2010; A.K. Jonscher, 1983; Vo et al., 2011). There are as many terms in the sum of HN functions as there are observed relaxations. Commercially available software was used to obtain the curve-resolved spectra. Both real and imaginary parts of the dielectric permittivity were considered for the fitting, computed based on nonlinear procedures. However, only curve resolved spectra of the imaginary part are reported in the manuscript. More details regarding the fitting procedure are available in the supporting file (annex II).

It is worth noting that the fitting of the experimental data to equation (A II.2) was applied to the dielectric spectra of all nanocomposites in the temperature range from 40 °C to 90 °C, where no specific molecular relaxation processes are supposed to occur in the neat material as reported by previous studies (H. Chen et al., 2011; Kofod et al., 2011). In fact, the segmental relaxation associated with the bulk glass transition of the rubber phase is too fast and could not be observed within this temperature window while the relaxation associated with the glass transition of PS phase is too slow and could not be observed either.

Figure 3-5 shows the imaginary part of the dielectric permittivity of neat SEBS-20 (Figure 3-5(a)) and its nanocomposites (Figure 3-5(b)-(e)), prepared by the three processing techniques mentioned previously, at different temperatures ranging from 30 °C to 90 °C. Only SEBS-20-partially oriented R3 was presented in Figure 3-5(a) as the other control samples, without clay, prepared by sheet die extrusion or solvent casting feature similar dielectric behavior. Moreover, for the sake of simplicity, the real part of the dielectric permittivity is not reported in this manuscript since relaxation phenomena are observable in both real and imaginary parts and the analysis of the imaginary part is more straightforward. Typical spectra of real part of the dielectric permittivity could be consulted in Figure A II-8 of the supporting file (annex II).

The real part of the dielectric permittivity of the neat SEBS is almost frequency independent and nearly equal to 2.35. At increasing temperatures, both the real and the imaginary parts of the dielectric permittivity of the neat copolymer remained equal to or slightly decreased below the value exhibited at room temperature (as it could be seen in the insert of Figure 3-5(a) and Figure A II-8 (annex II)). This decrease is probably related to the decreased polymer density at increasing temperatures, which is linearly affecting the relative permittivity (Blythe & Bloor, 2005). In addition, the high frequency relaxation peak related to the glass transition of the rubber phase shifts towards higher frequencies with the temperature increase (not observed in the studied range) causing a decrease of the dielectric losses in the 10^3 to 10^5 Hz frequency window.

In all the nanocomposites, both ε ' and ε '' increased simultaneously compared to neat SEBS (Figure 3-5 of the manuscript, Figure A II-7 and Figure A II-8 of annex II), up to orders of magnitude, particularly at low frequencies and high temperatures, indicating an important contribution originating from the inclusion of nanoclay to the low frequency dispersion (A.K. Jonscher, 1983; Andrzej K Jonscher, 1996). In particular, a pronounced increase is observed in the isotropic samples featuring random clay tactoids distribution compared to the samples where clay particles are aligned perpendicular to the electric field (Figure 3-5(e) compared to Figure 3-5(b)-(d)). This behavior is most likely attributed to an anisotropic

nature of clay conductivity which might be considerably lower across the nanoplatelets thickness than along their surface and the interlayer spacing, covered by a diffuse ionic layer (David et al., 2013; Tokarský et al., 2016). Another possible contribution to this remarkable increase of permittivity could be attributed to the presence of a fraction of polar organic solvent that was trapped during the solvent casting process used for the preparation of the isotropic samples.

It is worth noting that there is only little information regarding the experimental values of dielectric permittivity and electrical conductivity of nanoclay which strongly depends on the type of surface modification and the density of adsorbed cations (Uddin, 2008). The dielectric properties of an organically modified clay tactoid, similar to the one used in this study, were estimated through modeling by David et al. (David et al., 2013). In particular, the pure direct current conductivity of clay was estimated to be equal to 10^{-9} S/m.

The fitting of the dielectric permittivity to equation (A II.2) confirmed that all the nanocomposites exhibit, in addition to the charge fluctuation term, two relaxation modes that were not observed in neat materials. Similar behavior was reported in the literature for clay nanocomposites in general (David & Fréchette, 2013; David et al., 2013; Tomer et al., 2011) and clay/rubber nanocomposites specifically (Hernández et al., 2010; Vo et al., 2011). The charge fluctuation term is few orders of magnitude higher in the isotropic nanocomposite. In particular, considering the case of pure electronic conductivity in the fitting process (n=0 in equation (A II.2) of annex II), the values of conductivity σ_0 are in the range 10⁻⁹ to 10⁻¹¹ S/m for the isotropic nanocomposite while they are in the range 10^{-12} to 10^{-14} S/m in the case of oriented and partially oriented nanocomposites. In terms of relaxations, the first peak is located at low frequencies and is often overshadowed by the low frequency dispersion. It is attributed to Maxwell-Wagner-Sillars (MWS) polarization and is at the origin of the step-like increase observed in the real part of dielectric permittivity at the lowest frequencies. The second relaxation process was observed at intermediate frequencies and is shifted to higher frequencies at increasing temperatures. It is at the origin of the second step-like increase observed in real permittivity at intermediate frequencies. This additional relaxation could be

probably due to a MWS process from water absorbed at nanoparticles surface for instance (David & Fréchette, 2013; Kremer & Schönhals, 2012) or to local segmental relaxation of rubber chains with reduced mobility located at polymer/clay interfaces, which is slower than the main segmental relaxation responsible for the bulk T_g of the elastomer (L. Chen et al., 2009; Y. Gao et al., 2014; Ghanbari et al., 2013; Hernández et al., 2010; Hernández et al., 2012; Holt et al., 2014; Klonos et al., 2015; Yu Lin et al., 2015; Qu et al., 2011; Robertson & Rackaitis, 2011; Tsagaropoulos & Eisenberg, 1995; Tsagaropoulos & Eisenburg, 1995; Vo et al., 2011). In the case of SEBS, the dynamic T_g of the PEB rubbery block is in the vicinity of -40 °C, depending on the polystyrene block fraction (Leice G Amurin et al., 2016; Saleem et al., 2014). An example of this fit is presented in Figure 3-5(f). It corresponds to the dielectric loss spectrum of SEBS-20-20A-partially oriented R3 nanocomposite at 75 °C (Different colors are used to denote the different contributions).

Moisture absorption in nanocomposites was reported several times in literature, especially for epoxy-based nanocomposites containing polar groups and/or nanoparticles modified with functional groups of relatively low hydrophobicity (David & Fréchette, 2013; Glaskova & Aniskevich, 2009; I Preda et al., 2012). The absorbed water could form an interfacial conductive layer between the nanoparticles and the matrix material, which gives rise to an interfacial loss process (Kremer & Schönhals, 2012). In this study, the hypothesis of moisture absorption is discarded as SEBS is an apolar copolymer exhibiting very low water uptake and Cloisite 20A, the organically-modified clay used, features high hydrophobic character due to its dimethyl ditallow ammonium based modifier (Darie et al., 2014). It has also been reported that 24 hours of immersion in water was not found to have a significant impact on the dielectric response of HDPE containing 10wt% of organically-modified clay (David, Zazoum, Fréchette, & Rogti, 2015).



Figure 3-5: Imaginary part of the dielectric permittivity as a function of temperature of: (a) neat SEBS-20-partially oriented R3, (b) SEBS-20-20A-oriented, (c) SEBS-20-20A- partially oriented R1, (d) SEBS-20-20A-partially oriented R3, (e) SEBS-20-20A-isotropic and (f) an example of the fitting corresponding to SEBS-20-20A-partially oriented R3 at 75 °C (Different colors are used to denote the different contributions)

3.4.2.1 Effect of orientation on dielectric behavior

In order to confirm the origin of the second relaxation process, i.e. glass transition of the interfacial rubbery chains with reduced mobility, and to evaluate the effect of PS cylinders and nanoclay orientations on it, the temperature dependence of the relaxation times relative to this process was analyzed. At each temperature, the relaxation time corresponding to the peak maximum frequency was determined from the HN fit according to equation (A II.3) (Kremer & Schönhals, 2012), described in the supporting file (annex II).

Figure 3-6 presents the relaxation times relative to the relaxation process observed at intermediate frequencies as a function of inverse temperature for all the nanocomposites prepared from SEBS-20. The temperature dependence of the relaxation times is well described by the empirical Vogel-Fulcher-Tammann (VFT) equation (equation (A II.4) described in the supporting file (annex II)), which is usually used to fit the segmental relaxation process related to the bulk glass transition (Kremer & Schönhals, 2012). This behavior supports the hypothesis of molecular relaxation at the origin of the second peak observed at intermediate frequencies. The parameters of the VFT fit of each nanocomposite were computed and listed in Table 3-4. Besides, a dielectric "interfacial" glass transition temperature T_{gi} was associated to this relaxation mode in analogy with the glass transition temperature assigned to the segmental relaxation of the bulk rubber phase. This temperature was estimated by extrapolating the VFT fit to an extremely long relaxation time, conventionally chosen equal to 100 seconds (Kremer & Schönhals, 2012; Vo et al., 2011). The calculated temperatures are listed in Table 3-4 as well. In particular, the obtained values of the VFT parameters τ_0 , B and T₀ related to this new relaxation mechanism governing the mobility of the interfacial chains are in the same range of those reported previously by several groups for different rubber based nanocomposites where a similar phenomenon was observed such as Vo et al. (Vo et al., 2011) for their SBR based nanocomposites, Lin et al. (Yu Lin et al., 2015) for their Poly(vinylacetate)/silica nanocomposites and Wu et al. (Wu, Tang, Guo, Zhang, & Jia, 2013) for their Butadiene-Styrene-Vinyl pyridine rubber/Graphene Oxide nanocomposites.

In addition, the kinetic fragility index F, which characterizes how rapidly the dynamics of a material slow down as it is cooled toward the glass transition temperature, was estimated according to equation (A II.5) (Angell, 1991; Böhmer, Ngai, Angell, & Plazek, 1993) described in the supporting file (annex II). Considering that the behavior of the relaxation time as function of temperature is described by the VFT equation, the derivative could be calculated for each sample. The calculated values of F are reported in Table 3-4 as well.

It could be seen from Figure 3-6 that the sample with isotropic orientation features the highest relaxation times and consequently the slowest dynamics while the oriented dynamics. nanocomposite exhibits faster Interestingly, both partially oriented nanocomposites R1 and R3 exhibit even more reduced relaxation times especially toward the highest studied temperatures. This speed up of chain dynamics was reported in literature for rubber-based nanocomposites at increasing clay and graphene oxide nanoparticles loadings and it was attributed to suppression of cooperativity of polymer chains confined in the intercalated structures, when the confinement volume becomes comparable to that of the cooperative rearranging regions (Hernández et al., 2010; Schönhals, Goering, Schick, Frick, & Zorn, 2004; Wu et al., 2013). Moreover, at decreasing temperatures, the partially oriented samples feature increased dependence on temperature and converge to the behavior of the isotropic nanocomposite. This special behavior might be an indication of strong intermolecular coupling, i.e. the relaxation times diminish more rapidly with increasing temperatures (Bohmer, Angell, Richert, & Blumen, 1994; I Preda et al., 2013; Vilgis, 1993). The analysis of the calculated interfacial glass transition temperature T_{gi} reveals that the new relaxation process is located, depending on the nanocomposite configuration, in the temperature range between 26 and 35 °C; much higher than the bulk glass transition of PEB phase which will be studied by DMA analysis in a following section. Furthermore, this T_{gi} seems to depend as well on the orientation of both nanoclay and PS cylinders, as it was expected from the observed difference of the dependence of relaxation times on temperature. Overall, the oriented samples prepared by extrusion exhibit the lowest interfacial glass transition temperature, which might indicate that this controlled configuration helps decreasing the induced molecular chains confinement, and consequently the interaction

strength in the interphase region. In fact, the isotropic and the partially oriented samples exhibit characteristic T_{gi} up to 9 degrees higher, which is in good agreement with their morphology featuring more complex patterns and disorder. Indeed, the calculated values of fragility index confirm that partially oriented and isotropic samples feature more fragile behavior and consequently higher intermolecular coupling and interaction strength at the interface.



Figure 3-6: Temperature dependence of the relaxation times corresponding to SEBS-20 nanocomposites prepared by different processes: symbols correspond to relaxations times retrieved from the HN function and solid lines correspond to VFT fitting

Table 3-4: VFT fitting parameters corresponding to SEBS-20 based nanocomposites

Samples	τ ₀ (s)	T ₀ (K)	B (K)	T _{gi} (K)	F
SEBS-20-20A-partially oriented R1	4.24E-09	271	869	308	87.8
SEBS-20-20A-partially oriented R3	7.33E-07	280	483	306	95.8
SEBS-20-20A-oriented	1.05E-08	229	1620	299	43.7
SEBS-20-20A-isotropic	6.00E-04	285	232	305	80.1

While T_{gi} is indicative of interaction strength, the dielectric strength of the related relaxation mechanism partially corresponds to the number of interacting dipoles involved in the relaxation process, according to the Debye-Fröhlich-Kirkwood theory (Fröhlich, 1949; Kirkwood, 1939; Kremer & Schönhals, 2012). Therefore, a weaker signal is expected to correspond to less interacting polymer compared to a stronger signal. However, it is difficult to compare quantitatively the signal strength among different samples due to some inaccuracies such as in sample thickness measurements. Thus, this parameter could be considered only qualitatively (Vo et al., 2011). Illustrative values of dielectric strength corresponding to each sample are reported in Table 3-5, at two temperatures: 60 °C and 80 °C. It could be seen that the isotropic sample featured the highest dielectric strength values followed by the partially oriented samples in a second level while the oriented sample featured the lowest values. Hence, the amount of polymer chains with restricted mobility, located at the interphase, is larger in these samples featuring complex morphologies, which is consistent with the conclusions made earlier from T_{gi} results.

Samples	Δε at 60 °C	Δε at 80 °C
SEBS-20-20A-partially oriented R1	0.369	0.379
SEBS-20-20A-partially oriented R3	0.383	0.345
SEBS-20-20A-oriented	0.295	0.273
SEBS-20-20A-isotropic	0.540	0.528

Table 3-5. I	Dielectric	strength c	of SEBS-20	based nanocom	posites at 6	0°C ar	nd 80 °0	С
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3.4.2.2 Effect of styrene content and diameter of PS cylinders

In order to elucidate the effect of nanoclay interaction with each phase of the block copolymer on the polymer chain dynamics at the interphase region, comparisons were made between isotropic samples and oriented samples prepared from the different block copolymer matrices containing respectively 0.13, 0.2 and 0.3 polystyrene fractions. The dielectric spectra of the samples prepared from SEBS-30 are available in reference (E Helal et al., 2015) while the results related to SEBS-13 based nanocomposites are reported in

Figure A II-7 of the supporting file (annex II). Examples of the fitting according to equation (A II.2) of the ESI file, for each type of nanocomposite are presented in Figure 3-7 (Different colors are used to denote the different contributions). In particular, nanocomposites prepared from SEBS-13 are well fitted by a power law term and two HN functions (similar to previous samples prepared from SEBS-20), such as the example reported in Figure 3-7(a), corresponding to SEBS-13-20A-isotropic. SEBS-30-20A nanocomposites, in comparison, are rather fitted by three HN functions in addition to a power law term (Figure 3-7(b)). The third relaxation peak has less dielectric strength than the two other peaks, i.e. the MWS peak and the interfacial glass transition peak, and is most likely attributed to PS/PEB interphase region, which starts to be more important with increasing PS content.



Figure 3-7: Examples of fitting according to equation (A II.2) in annex II of respectively:
(a) SEBS-13-20A-isotropic at 65 °C (ɛ" presented in log scale due to high contribution from charge fluctuation) and (b) SEBS-30-20A-oriented at 70 °C (Different colors are used to denote the different contributions)

Relaxation times corresponding to the interfacial glass transition relaxation peak are plotted as a function of temperature in Figure 3-8. The results corresponding to nanocomposites prepared from SEBS-20 discussed earlier (Figure 3-6) are reported again for the sake of comparison. The VFT equations corresponding to each sample are plotted as well, while the VFT parameters are summarized in Table 3-6. In particular, SEBS-13 nanocomposites, featuring the lowest content of styrene, show similar values and dependence on orientation of their interfacial T_{gi}, compared to samples prepared from SEBS-20, while SEBS-30 nanocomposites, containing the highest fraction of styrene, show the lowest values and sensitivity to orientation featuring T_{gi} values in the range of 6 to 9 °C. The fragility index values confirm this observation as well. This fact might indicate that although clay nanoparticles exhibit more affinity to the aromatic rings in PS blocks (Carastan et al., 2014), they affect more the mobility of chains in the rubbery PEB phase, and consequently are located more in contact with this phase. This latter observation is consistent with TEM images reported in Figure 3-3(d)-(e), showing the presence of intercalated nanoclay that either cross PS cylinders or are located completely outside PS cylinders if they are relatively big. Another possible reason could be related to the cylinder diameter, which is reduced with decreasing PS block content, providing more space for the contact between the clay nanoparticles and the PEB blocks. In fact, the diameter of PS cylinders in SEBS-13 is estimated to be equal to 9.7 nm according to calculations based on SAXS results (not reported in the manuscript), while in SEBS-20 and SEBS-30, the estimated diameter is respectively equal to 12.5 nm, according to the TEM image reported in Figure 3-3(e), and 13.5 nm according to a TEM image reported in reference (E Helal et al., 2015). As a consequence, in samples prepared from SEBS-30, the contact volume between clay nanoparticles and rubber chains might be reduced by the increased diameter of glassy PS rods, in a similar way in both oriented and isotropic samples, resulting in less sensitivity to orientation.

From another point of view, it could be seen that the relaxation times corresponding to SEBS-13 and SEBS-30 nanocomposites are both smaller than those corresponding to SEBS-20 nanocomposites. This behavior could be partially related to the bulk glass transition temperature of PEB phase, which might depend on its overall fraction in the block copolymer. In fact, the T_g of a specific block is usually different from the glass transition temperature of the corresponding homopolymer, and it depends on the interphase region between the blocks. In general, only when the compatibility between the blocks of a block copolymer is weak, a sharp interface is observed due to strong segregation and the T_g of each block is equal to the T_g of the corresponding homopolymers (Adhikari et al., 2003). However, in the case of SEBS, the pendant groups of the hard PS phase might be mixed to

the soft PEB phase forming an interphase region, as predicted from the small relaxation peak observed in SEBS-30 (Figure 3-7(b)). It will be shown later, by DMA measurements, that the Tg of each block increases with increasing fraction of that block in SEBS block copolymer.

In general, it is expected that the block copolymer matrices featuring higher bulk T_g of their PEB phase exhibit higher interfacial glass transition temperature and slower dynamics at a given temperature above the T_{gi} , if other factors such as the orientation and the interaction of nanoparticles with the block copolymer phases are maintained unchanged. This behavior is partially achieved. In fact, SEBS-30 based nanocomposites exhibit the lowest bulk T_g of the PEB block as what will be shown later by DMA, and simultaneously the fastest interfacial dynamics and the lowest T_{gi} compared to materials prepared from SEBS-13 and SEBS-20. However, one should keep in mind that the motion of polymer chains at the interphase region does not depend only on the corresponding bulk T_g of the rubber phase, but it depends also on other parameters including the interaction of nanoparticles with each block and the orientation as concluded earlier.



Figure 3-8: Temperature dependence of the relaxation times corresponding to SEBS-13 and SEBS-30 nanocomposites prepared by different processes: symbols correspond to relaxation times retrieved from the HN function and solid lines correspond to VFT fitting, plots corresponding to SEBS-20 nanocomposites are presented again for comparison

Samples	τ ₀ (s)	T ₀ (K)	B (K)	T _{gi} (K)	F
SEBS-13-20A- partially oriented	6.80E-09	244	1280	299	55.4
SEBS-13-20A-isotropic	7.16E-06	284	366	307	94.8
SEBS-30-20A-oriented	5.70E-10	220	1530	279	53.3
SEBS-30-20A-isotropic	2.59E-07	250	642	282	76.8

Table 3-6: VFT fitting parameters corresponding to SEBS-13 and SEBS-30 based nanocomposites

3.4.2.3 Effect of nanoparticles location and intercalation degree

To further understand the effect of the nanoparticles contact with one block or another on the observed interfacial peak, samples where nanoparticles are completely located in PEB phase are studied and compared to the previous results. The affinity of nanoparticles to PEB phase was increased by the attachment of maleic anhydride (MA) to it. MA exhibits great compatibility with the organic groups attached on the nanoparticles surface. Furthermore, this increased affinity results in more pronounced intercalation of the elastomer chains in clay galleries, disrupting the regular stacked layer structure of the organoclays and giving rise to mostly exfoliated structure, as reported in our previous study (E Helal et al., 2015) (TEM micrographs of SEBS-30-MA-20A nanocomposites exhibiting exfoliated structure are available in Figure A II-6 of the supporting file (annex II)). Hence, a higher fraction of elastomer chains is expected to be involved in this interphase region compared to the samples prepared without MA. The comparison of relaxation times between SEBS-30-20A and SEBS-30-MA-20A samples, plotted in Figure 3-9, shows that the presence of MA attached to the rubber phase affected remarkably the dynamics of the interphase region. In particular, relaxation times increased in the sample containing the MA graft, which indicates slower dynamics. This behavior is expected due to the selective location of clay in the rubber phase, which increases the amount of elastomer chains attached to the interface. In addition, the degree of clay exfoliation is improved in the presence of MA, which results in a larger number of interfaces and less efficient restriction of chains motion compared to intercalated

structures. Hence, the cooperativity of interfacial chains is higher, resulting in increased relaxation times (Wu et al., 2013). These hypotheses are in agreement with the increased estimated T_{gi} equal to 20 °C for SEBS-30-MA-20A, when compared to 6 °C for the sample prepared without MA. These values were extrapolated from the VFT fitting parameters reported in Table 3-7. The comparison of the dielectric strength values of the two nanocomposites reported in Table 3-8 supports the same conclusion as well. In fact, the dielectric strength of the sample containing MA is considerably higher than that of the nanocomposite without MA.

Table 3-7: VFT fitting parameters corresponding to SEBS-30-MA-20A-oriented nanocomposite

Samples	τ ₀ (s)	T ₀ (K)	B (K)	T _{gi} (K)	F
SEBS-30-MA-20A-oriented	7.91E-10	237	1430	293	58.7

Table 3-8: Dielectric strength corresponding to SEBS-30-20A vs. SEBS-30-MA-20A nanocomposites prepared by extrusion

Samples	Δε at 60°C	Δε at 80°C
SEBS-30-20A-oriented	0.252	0.233
SEBS-30-MA-20A-oriented	0.865	0.830



Figure 3-9: Temperature dependence of the relaxation times corresponding to SEBS-30-MA-20A-oriented nanocomposite: symbols correspond to relaxations times retrieved from the HN function and solid lines correspond to VFT fitting (plot corresponding to SEBS-30-20A-oriented nanocomposite is presented again for comparison)

Overall, BDS data revealed the appearance of a new relaxation mode attributed to rubber chains with restricted mobility located at polymer/nanoparticle interfaces, in all the nanocomposites prepared from SEBS-13, SEBS-20, SEBS-30 and SEBS-MA-30 grades. A T_{gi} associated with this process is estimated to occur in the range 6 to 35 °C depending on the nanocomposite morphology. To confirm the molecular origin of this relaxation process, DMA was performed. Results will be discussed in the following section.

3.4.3 Dynamic mechanical analysis

Dielectric spectroscopy data discussed in this paper were mainly performed in the temperature range from 30 to 90 °C and analyzed as a function of frequency. In order to extend the study of relaxation phenomena to lower temperatures, to confirm the origin of the new relaxation peak observed in BDS and to evaluate the dependence of bulk glass transition temperatures of both PS and PEB blocks, respectively T_{gPEB} and T_{gPS}, on styrene fraction and

clay addition, dynamic mechanical analysis was carried out in a large temperature range from -100 °C to 130 °C. In fact, in this range, both T_{gPEB} and T_{gPS} are expected to appear as peaks in DMA tan δ graphs. Besides, any potential peak that will appear in addition will be certainly due to a molecular relaxation, by contrast to BDS spectra, where relaxation phenomena observed in hybrid materials could be due to either a molecular origin or separation of charges at inner dielectric boundary layers (MWS polarization).

Figure 3-10 shows tan δ graphs corresponding to selected samples prepared from SEBS-30 (graphs corresponding to samples from SEBS-13 and SEBS-20 are not reported). It is worth noting that the DMA measurements were conducted in the transverse direction for all the studied samples (perpendicular to the main direction of PS cylinders alignment). In general, all curves show evidence of two peaks which correspond to T_{gPEB} (at low temperatures) and T_{gPS} (at high temperature). The values of T_{gPEB} and T_{gPS} corresponding to each material are reported in Table 3-9, except T_{gPS} corresponding to SEBS-13 samples where no clear peaks were observed up to 130 °C. In a general trend, the glass transition temperature of each block, either PS or PEB, and the height of its corresponding peak increases with increasing fraction of that block in SEBS. This behavior is due to the fact that more chains of the considered block are involved in the glass transition phenomena leading to a larger damping. Besides, the T_g temperature and the intensity of the peak depend on the orientation of PS cylinders. Indeed, TgPS of SEBS-20- partially oriented R1 material prepared by film blowing is 6 degrees higher than SEBS-20-oriented, 106 °C vs.100 °C. Simultaneously, the intensity of T_{gPS} is reduced and that of T_{gPEB} is considerably higher. This behavior might be due to the fact that these samples are not completely aligned in the extrusion direction which leads to an increasing fraction of PEB chains in the transverse direction, which is the direction of the test, as well as more hindrance of PS chains mobility.

In the nanocomposites, the addition of clay results, in some cases, in the reduction of T_g peak height and the broadening of the peak, such as the peaks corresponding to T_{gPS} in both SEBS-30-20A-oriented and SEBS-30-MA-20A-oriented nanocomposites, as shown in Figure 3-10. These behaviors respectively indicate lower number of chains participating in the bulk glass transition and wider distribution of chains mobility due to the restriction of motion imposed by clay nanoparticles. In addition, the glass transition temperatures of both blocks are either maintained or shifted to higher or lower temperatures. Shifts of bulk T_g to both higher and lower temperatures have been reported in the literature. The experimental results reported in the literature are in general not conclusive due to the complex nature of various polymers and nanocomposites that were investigated (Yu Lin et al., 2015; Rittigstein & Torkelson, 2006). Different reasons were suggested to explain the phenomena depending on the cases. In particular, the increase was mainly attributed to restriction of chain mobility at the interface (Yu Lin et al., 2015), while the decrease was attributed to different reasons such as a specific form of mechanical coupling between the filler, the bound layer and the bulk rubber (Arrighi, McEwen, Qian, & Prieto, 2003; Mélé, Marceau, Brown, de Puydt, & Albérola, 2002), an increase of mobility (Hao, Böhning, & Schönhals, 2007) or an increase of free volume and decrease of molecular packing density (Hao, Böhning, Goering, & Schönhals, 2007).



Figure 3-10: tan δ of different SEBS-30 materials (prepared by sheet die extrusion) from -100 °C to 130 °C at 1 Hz

Sample	T _{gPEB} (°C)	T _{gPS} (°C)
SEBS-13-partially oriented	-23	-
SEBS-13-20A-partially oriented	-27	-
SEBS-20-oriented	-26	100
SEBS-20-20A-oriented	-26	100
SEBS-20- partially oriented R1	-26	106
SEBS-20-20A- partially oriented R1	-20	110
SEBS-30-oriented	-35	107
SEBS-30-20A-oriented	-34	105
SEBS-30-MA-20A- oriented	-33	104

Table 3-9: Tgs of both PS and PEB blocks in different samples of SEBS based nanocomposites

In addition to the bulk glass transitions of PS and PEB blocks, nanocomposites prepared from SEBS-30 seem to exhibit an additional low intensity peak occurring at intermediate temperatures between T_{gPEB} and T_{gPS}, which might correspond to the glass transition of interfacial rubber chains with reduced mobility (as indicated by the arrow in Figure 3-10). However, this peak is not well resolved and as a consequence, resort to curve fitting was necessary. Although, there is no theoretical expression to describe dynamic mechanical relaxations as a function of temperature, the use of some empirical equations which reproduce satisfactorily the asymmetry of tan δ peaks has been reported in the literature. One such is the exponentially modified Gaussian (EMG) which was used by Tsagarapoulos et al. (Tsagaropoulos & Eisenberg, 1995) and Cowie et al. (Cowie, Arrighi, Cameron, McEwan, & McEwen, 2001) to characterize Tg peaks and particularly peaks related to Tg of interfacial layers observed in several homopolymers containing relatively high fractions of fillers such as silica nanoparticles and cellulose tricarbanilate. Another model based on the threeparameter asymmetric double sigmoid (ADS) equation was successfully used by Arrighi et al. (Arrighi et al., 2003) to characterize Tg of the interfacial layer observed in styrene butadiene rubber (SBR) containing silica nanoparticles.
In this study, we adopted a model based on ADS equation, similarly to reference (Arrighi et al., 2003), to describe both bulk T_g peaks and the new interfacial T_g peak. More details about this equation and its characteristic parameters are available in reference (Arrighi et al., 2003) and annex II (equation (A II.6)). The general model is composed of three ADS terms to describe the three peaks. An additional ADS term with a large asymmetry in the high temperature side was also considered to take into account the increasing baseline towards high temperatures. For consistency, the same parameters were taken for the baseline term in the neat SEBS-30-oriented and the nanocomposites. Examples of the fitting of both neat SEBS-30-oriented and SEBS-30-20A-oriented nanocomposites showing the resolved T_g peaks are reported in Figure 3-11.



Figure 3-11: Fitted tan delta curves showing bulk Tgs of PS and PEB blocks as well as Tgi of the interfacial layer (an additional ADS term is considered for the increasing baseline):(a) SEBS-30-oriented and (b) SEBS-30-20A-oriented

The peak corresponding to the glass transition of the interfacial layer, T_{gi} , in both nanocomposites, resulting from the curve-fitting process, is characterized by low intensity and a broad shape indicating a large distribution of chains mobility in the interfacial layer. Temperatures corresponding to T_{gi} peaks maxima are reported in Table 3-10. Taking the difference between these values and T_{gPEB} values reported in Table 3-9, T_{gi} of the interfacial rubbery layer is estimated to occur 72 degrees higher than the bulk in the case of SEBS-30-20A-oriented and 86 degrees higher than the bulk in the case of SEBS-30-MA-20A-oriented.

This result is consistent with many findings in the literature expecting the glass transition of tethered chains confined at the interface to occur 70 to 110 degrees higher than the bulk Tg (Tsagaropoulos & Eisenberg, 1995). It is worth noting that these values of T_{gi} determined from DMA data at 1 Hz are higher than those estimated earlier by BDS through the extrapolation of the VFT equation at very low relaxation times (100s): 38 °C vs. 6 °C for SEBS-30-20A-oriented and 53 °C vs. 20 °C for SEBS-MA-30-20A-oriented. However, the trend is maintained. Hence, to check the similarity of the results given by both BDS and DMA techniques, dielectric permittivity of SEBS-30-oriented and SEBS-30-20A-oriented materials were mapped in an extended temperature range [-100, 150 °C] and dielectric loss spectra were plotted as function of temperature at 1 Hz (Figure A II-9 and Figure A II-10 in annex II). At this frequency, the peak corresponding to T_{gPEB} is observed at -45 °C, which is 10 degrees higher than DMA and the peak corresponding to T_{gi} is observed at 40 °C, which is 2 degrees higher than the value obtained by DMA. This difference between the results given by the two methods is observed even when equivalent formalisms are used: electric modulus vs. mechanical modulus. Thus, it is rather due to experimental conditions such as different heating rates (Talja & Roos, 2001).

The fraction of bound polymer chains participating in the interfacial glass transition, F_{Bound} , could be determined by dividing the area under T_{gi} peak by the sum of the areas of all T_g peaks. Values of integrated area under each peak and estimated bound fractions are reported in Table 3-10 as well. The bound fraction in SEBS-30-20A-oriented is around 7.5% while in SEBS-30-MA-20A-oriented, it is around 12%. This is expected as the higher degree of exfoliation in the latter nanocomposite results in more interfacial area. Taking into account data about the average particle size of clay nanoparticles in SEBS-30 nanocomposites, which were published previously by coauthors (Carastan et al., 2014), the thickness of interfacial layer around each particle could be estimated in a similar fashion to the work of Arrighi et al. (Arrighi et al., 2003). More details regarding this procedure could be consulted in annex II (equations (A II.7) to equation (A II.10)).

Sample	T _{gi} (°C)	Apeb	A _{PS}	AInt	F _{Bound}	T(nm)
SEBS-30-20A-oriented	38	9.475	10.15	1.59	0.075	10.5
SEBS-30-MA-20A-oriented	53	9.42	10.04	2.58	0.12	7.6

Table 3-10: Interfacial Tgi, bound fraction and interfacial layer thickness in SEBS-30 based nanocomposites

The average tactoid thickness, average lateral dimensions and average number of clay layers per tactoid were determined based on the TEM quantitative analysis of clay dimensions in SEBS-30-20A and SEBS-30-MA-20A nanocomposites reported by Carastan et al. (Carastan et al., 2014). These values are listed in Table 3-11.

Table 3-11: Average dimensions of clay tactoids based on TEM quantitative analysis published by Carastan et al. (Carastan et al., 2014)

	Average thickness (nm)	Average lateral size (nm)	Average number of layers	
SEBS-30-20A-oriented	7.3	243.3	3	
SEBS-30-MA-20A- oriented	3	116.3	2	

To calculate the thickness of the interfacial layer in both nanocomposites, the clay tactoid was considered as a solid particle surrounded on both lateral sides with constrained rubber chains. Since the thickness of one tactoid is much smaller than its lateral dimension, the fraction of chains that could be constrained along the thickness was neglected. Then, the calculated interfacial layer thickness, t, was adjusted to take into account that some chains are rather confined in the interlayer spacing, considering that the thickness of 1 clay layer is equal to 0.94 nm (Carastan et al., 2014; Vermogen et al., 2005).These approximations are illustrated in Figure 3-12. With these approximations the calculated values of interfacial layer thickness are respectively equal to 7.6 nm for SEBS-30-MA-20A nanocomposite and 10.5 nm for SEBS-30-20A. They are in agreement with findings from the literature which reported estimations of thickness of the interfacial layer. Whether evaluated experimentally (Arrighi et al., 2003; Yu Lin et al., 2015) or theoretically (Y. Gao et al., 2014; Ghanbari et

al., 2013; Heid, 2015), the values reported in the literature did not exceed few nanometers ranging from 2 to 10 nm in most of the reported studies.



Figure 3-12: Scheme of the interfacial layer covering the lateral side of clay particles: (a) First approximation: clay tactoid as a solid particle, (b) part of the interfacial layer is between clay galleries

Hence, the fitting model based on ADS equation describes satisfactorily the bulk glass transitions of both PEB and PS blocks as well as the Tgi of interfacial chains with restricted mobility. It leads to an approximation of the interfacial layer thickness well in agreement with values reported in the literature. However, it is worth noting that an important hypothesis was assumed in the calculation of the interfacial layer thickness, which is the absence of an immobilized layer corresponding to polymer chains that are tightly bounded and don't participate in neither bulk Tg nor interfacial Tgi (Tsagaropoulos & Eisenberg, 1995). In fact, the total number of chains participating in glass transition phenomena in a nanocomposite is generally lower than the number of chains participating in the glass transition of the neat material. Thus, by comparing the total area under Tg peaks in respectively the nanocomposite and the pure material, the fraction of immobilized layer could be determined and excluded when calculating the thickness of the interfacial layer (Arrighi et al., 2003). The thickness of the immobilized layer was estimated in the literature to be in the range 0.5 to 2 nm (Arrighi et al., 2003). In our case, comparing the tan delta curves of the neat SEBS-30 and its corresponding nanocomposites, we could see that the total area is higher in the case of nanocomposites, which gives the impression that more chains are involved in the glass transition phenomena compared to the neat copolymer. This behavior is puzzling and for this reason comparison with neat and estimation of the immobilized fraction were not possible. As a consequence, the estimated thicknesses of the interfacial layer might be smaller in reality. Another behavior that requires more

investigation is the fact that the T_{gi} peak related to interfacial rubbery chains was not observed in SEBS-13 and SEBS-20 nanocomposites by DMA, by contrast to BDS. Only shifts of bulk T_g temperatures were obvious by this technique. However, for nanocomposites prepared from SEBS-30 and SEBS-30-MA, the new relaxation process was confirmed by both DMA and BDS. Furthermore, T_{gi} temperatures estimated by both techniques were in agreement.

3.5 Concluding remarks

In this study, nanocomposites of SEBS block copolymer with cylindrical morphology and organically modified clay nanoparticles have been successfully prepared with different configurations of PS cylinders and clay nanoparticles. In particular, isotropic vs. oriented vs. partially oriented morphologies have been successfully prepared by different processing techniques and confirmed by SAXS and TEM.

In all the nanocomposites, a glassy interphase region was formed and gave rise to a new relaxation mode corresponding to the interfacial rubbery chains with restricted mobility. A dielectric "interfacial" glass transition T_{gi} , associated with this process, is estimated to occur at temperatures ranging from 6 up to 35 °C, depending on the orientation of both PS domains and clay nanoparticles, the fraction of polystyrene block as well as the location and the degree of exfoliation of clay nanoparticles, which was tuned through the use of maleic anhydride graft. The thickness of the interfacial layer was estimated through fitting of DMA data and was found to be in the range 7.5-10.6 nm for nanocomposites prepared from SEBS-30 and SEBS-30-MA.

Overall, the relaxation mode related to this interphase region was sensitive to the following key parameters:

- Distribution of interfaces: oriented vs. isotropic vs. partially oriented; which affects the area and the volume of the "interphase region" as well as the mobility of polymer chains. In particular, samples featuring random or partially oriented morphologies were found to exhibit

higher interaction, dielectric strength and T_{gi} , up to 9 degrees higher, compared to their oriented counterparts. Besides, partially oriented samples prepared by film blowing extrusion featured the most "fragile" behavior and the fastest dynamics at temperatures above estimated T_{gi} , compared to oriented and isotropic samples. This behavior could be attributed to suppression of cooperativity of interfacial chains motion due to strong confinement in this configuration.

- Ratio of PS/PEB, which affects the T_g of the bulk PEB block, the diameter of PS cylinders as well as the amount and the mobility of PEB chains located in the interphase region. In particular, the T_{gi} was found to decrease and to be less sensitive to orientation effects, with increasing styrene content, most likely due to reduced amount of rubber chains involved in the interphase region and increasing diameter of PS cylinders resulting in a similar trend of confinement in both oriented and non-oriented samples.

- Location of clay nanoparticles either in PS or PEB blocks, which was modulated by the use of maleic anhydride attached to the rubber phase. In particular, in the presence of MA, clay particles were located exclusively within the elastomer block, resulting in increased amount of rubber chains involved in the interphase region and consequently higher dielectric strength associated to the relaxation process. Moreover, the degree of exfoliation of clay tactoids is improved, which induced slower dynamics compared to intercalated structures where strongly confined rubber chains exhibit reduced cooperativity and consequently reduced relaxation times.

Finally, the charge fluctuation contribution to the dielectric loss was dependent as well on the morphology. In particular, the conduction loss was reduced by 2 to 4 orders of magnitude in the samples with controlled architecture, i.e. oriented and partially oriented, as clay nanoparticles were quasi-aligned perpendicular to the electric field in these configurations.

Associated content

Supporting electronic information (ESI) file (reported in annex II): 2D and 1D SAXS patterns, TEM micrographs and dielectric spectroscopy data related to the studied nanocomposites are available. The procedure of the estimation of the interfacial layer thickness is explained as well.

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CHAPTER 4

ARTICLE 3: THERMOPLASTIC ELASTOMER NANOCOMPOSITES WITH CONTROLLED NANOPARTICLES DISPERSION FOR HV INSULATION SYSTEMS: CORRELATION BETWEEN RHEOLOGICAL, THERMAL, ELECTRICAL AND DIELECTRIC PROPERTIES

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Abstract

Thermoplastic elastomer nanocomposites based on respectively polystyrene-*b*-poly(ethylene*co*-butylene)-*b*-polystyrene (SEBS) and polystyrene-b-poly(ethylene-co-butylene)-bpolystyrene grafted maleic anhydride (SEBS-MA) block copolymers and containing functionalized zinc oxide (ZnO) nanoparticles have been investigated as candidate materials for high voltage (HV) insulation systems. The dispersion of the organically modified ZnO nanoparticles has been successfully tuned through the MA graft and the block copolymer nanostructure. In particular, nanocomposites with signs of rheological percolation, indicating the formation of a network between individually dispersed nanoparticles and polymer chains, have been obtained at ZnO content as low as 5 wt% (0.9 vol%). This behavior resulted in an enhancement of the thermal conductivity and better control of the electrical conductivity while maintaining breakdown strength and dielectric losses in the same range of the unfilled insulating matrices. Furthermore, the resistance to surface erosion by partial discharges was significantly improved: in the presence of 5 wt% of individually dispersed ZnO nanoparticles, the eroded volume was reduced 10 times.

Keywords: thermoplastic elastomer, Zinc Oxide nanoparticles, nanodielectric, rheological percolation, HV insulation...

4.1 Introduction

Due to the immiscibility of their blocks that are covalently bonded, block copolymers can self-organize into spherical, cylindrical, lamellar or more complex ordered nanodomains, depending on several parameters such as the block copolymer composition, the chemical interaction between the blocks and the molecular weight (Mai & Eisenberg, 2012). Owing to this nanostructure, several block copolymers, including polystyrene-b-poly(ethylene-cobutylene)-b-polystyrene (SEBS) and polystyrene-b-poly(ethylene-co-butylene)-b-polystyrene grafted maleic anhydride (SEBS-MA) thermoplastic elastomers, have recently attracted considerable attention as template matrices offering the possibility of nanoparticles dispersion control, in the aim of producing nanocomposites and master batches with tailored dispersion and interphase region. The expected high performance materials have a great potential in different applications including dielectric applications (Sarkar & Alexandridis, 2015). However, to the best of our knowledge, only few studies reported specific dielectric characterization of thermoplastic elastomeric block copolymer based nanocomposites (E Helal et al., 2015; Helal, Demarquette, David, & Fréchette, 2015; Kofod et al., 2011; Mc Carthy et al., 2009; Mi et al., 2014; Saleem et al., 2014), taking into consideration the possibility of tuned nanoparticles dispersion, in the aim of producing high performance nanodielectrics (Emna Helal et al., 2015; Sarkar & Alexandridis, 2015).

In addition to the nanoarchitecture, thermoplastic elastomeric block copolymers can play a functional role in dielectric applications such as dielectric elastomer actuators and HV insulation materials. In fact, these materials exhibit good mechanical properties combining both elastomer and thermoplastic properties (Balsamo et al., 2006; Holden et al., 2004), good electromechanical coupling (B. Kim et al., 2011) as well as good resistance to moisture absorption (Liu et al., 2011; Z. Ma et al., 2010). Currently, the use of thermoplastic elastomeric block copolymers, in nanocomposites generally and nanodielectrics specifically, is mainly limited to their role as compatibilizers to improve the dispersion of nanoparticles

such as nanoclay, metal oxides and more recently carbon nanotubes (Liang & Tjong, 2006) in polyolefin matrices or as water treeing retardant agents in HV cables (Liu et al., 2011; Z. Ma et al., 2010). Recent studies have started investigating these materials in a more functional aspect. For instance, blends of polyolefins and thermoplastic elastomers have been evaluated as candidates for HV cable insulation and specifically for recyclable high voltage direct current (HVDC) cables in replacement to cross-linked Polyethylene (XLPE) cables (Han & Sengupta, 2014; Helal, Demarquette, David, & Fréchette, 2016; Helal, Demarquette, David, & Fréchette, 2014; I. Hosier et al., 2007; Y. Zhou et al., 2015). Preliminary results revealed that these blends feature improved mechanical flexibility, thermal properties, volume resistivity and resistance to surface erosion (Helal et al., 2016; I. Hosier et al., 2007; Y. Zhou et al., 2015). However, they suffer from space charge accumulation, which is a bottle-neck especially in HVDC transmission (Hayase et al., 2006; Mizutani, 2000). This problem might be addressed by the incorporation of specific nanoparticles able to increase the density of charge traps and consequently decrease charge carriers mobility, in a similar trend to what has been reported for homopolymer nanocomposites (Fleming et al., 2008; Ju et al., 2014; Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016; Tian et al., 2011, 2012; Tian et al., 2015). But, better understanding of the nanoparticles impact on the thermoplastic elastomer phase is required first. Therefore, exhaustive dielectric, thermal and electrical characterization of thermoplastic elastomer nanocomposites might be a crucial step to evaluate their potential as nanodielectrics and as master batches to be blended with industrial polyolefins for application in HV and HVDC insulation systems.

An important property governing the performance of HV insulation systems is space charge distribution. In fact, accumulation of space charges in HV insulation materials subjected to high electric fields might lead to inhomogeneous distribution of the electric field and localized large electrical stresses, especially in DC transmission. To avoid the deterioration of the insulation in the regions suffering from elevated electrical stresses, large thicknesses of the insulating material are often required to reduce the magnitude of the peak stresses and to increase the breakdown voltage, which leads to expensive insulation systems and difficulty to dissipate heat. To reduce space charge accumulation, one approach is the use of non-linear

resistive fillers. At low loadings, these fillers were reported to increasing the charge trap density and consequently decrease charge carriers mobility and prohibit space charge accumulation in the bulk material (Fleming et al., 2008; Ju et al., 2014; Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016; Tian et al., 2011, 2012; Tian et al., 2015). At higher loadings, an improvement of the non-linear feature of electrical conductivity is reported and leads to efficient reduction and homogenization of electrical stresses within the insulation (Hong, Schadler, Siegel, & Mårtensson, 2003; Hong et al., 2006; Varlow et al., 2007). As the introduction of fillers may result in higher dielectric losses, an improvement of thermal conductivity is also desirable, through the choice of appropriate fillers, in order to increase the rate at which heat is dissipated in the system (Varlow et al., 2007).

One type of nanoparticles that is of particular interest when dealing with HV insulation systems is Zinc Oxide (ZnO) nanoparticles. In fact, ZnO is a large band gap semiconductor characterized by nonlinear electrical conductivity (Hong et al., 2006; Varlow et al., 2007), relatively high thermal conductivity (Huang et al., 2011) and ability to absorb UV light released by partial discharges (S. Chen et al., 2010). It was studied in nanocomposites for insulation applications, mainly polyethylene based systems (S. Chen et al., 2010; Fleming et al., 2008; Hong et al., 2003; Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016; Tian et al., 2011, 2012; Tian et al., 2015). Its incorporation led to several beneficial properties including the ability to significantly decrease space charge accumulation (Fleming et al., 2008; Tian et al., 2011; Tian et al., 2015) as well as to improve resistance to electrical treeing (Tian et al., 2012), which are stringent requirements to be fulfilled by any potential candidate for HV insulation in order to ensure longer lifetime of the insulation.

In this study, the morphology and rheological properties of nanocomposites of SEBS and SEBS grafted maleic anhydride (SEBS-MA) containing ZnO nanoparticles have been investigated. In particular, the dispersion of ZnO nanoparticles was tuned to obtain samples featuring agglomerations vs. samples featuring nanoscale dispersion and controlled interactions between polymer chains and nanoparticles at the interface. For that, functionalized nanoparticles exhibiting good affinity with maleic anhydride (MA) were used

in order to improve the dispersion and to initiate the process of selective localization in the phase exhibiting the highest affinity, which is the elastomer block. This tuned dispersion is expected to affect strongly the quality of the interphase region and consequently the engineering properties (David & Fréchette, 2013; Michel Fréchette et al., 2014; Lau, Vaughan, Chen, Hosier, & Holt, 2013; Sarkar & Alexandridis, 2015). Therefore, a correlation between the obtained morphologies and respectively the rheological, electrical, thermal and dielectric properties was established in a trial to evaluate the potential of thermoplastic elastomer nanocomposites for use in HV insulation systems and to predict their optimal performance.

4.2 Experimental section

4.2.1 Materials

SEBS powder grade G1652, of density 0.915 g/cm³ and SEBS-MA pellets grade FG1901 were purchased from Kraton. Both block copolymers contain 30 wt% of polystyrene (PS) endblocks dispersed in the form of nanodomains within a poly(ethylene-co-butylene) (PEB) midblock matrix. The maleic anhydride (MA) content is equal to 1-2 wt% in SEBS-MA and is grafted on the PEB elastomer block as illustrated in Figure 4-1. Zinc oxide nanoparticles were supplied in the form of 40 wt% colloidal suspension in ethanol from Sigma Aldrich. The particles are organically modified with an alkyl ammonium salt and have a particle size below 130 nm in general and an average particle size around 35 nm. All the materials were used as received.



Figure 4-1: Schematic description of SEBS block copolymer with MA group grafted to elastomer block ("Polystyrene-block-poly(ethylene-ran-butylene)-blockpolystyrene-graft-maleic anhydride,")

4.2.2 Processing

The nanocomposites were prepared by solution blending using toluene as a solvent. The suspension of nanoparticles was first mixed with toluene and disagglomerated using Sonic Ruptor 400 ultrasonicator for 8 minutes in continuous mode at 70% of the maximum power. The necessary amount of SEBS or SEBS-MA was subsequently added and dissolved in the mixture by means of magnetic stirring overnight. The solution was simultaneously heated at 60 °C. Subsequently, the obtained mixture was poured into a Petri dish and kept under fume hood until complete evaporation of solvent. The nanocomposites were finally dried for at least 48 hours at 60 °C under vacuum. The investigated nanoparticles loadings in the final samples were equal to: 1 wt%, 5 wt%, 10 wt% and 20 wt%. These concentrations were checked by means of thermal gravimetric analysis. All the samples were preheated for 10 minutes and pressed for 5 minutes at 200 °C under 10 MPa to obtain films with specific thicknesses suitable for further characterization. The samples prepared from SEBS and SEBS-MA will be referred to respectively as SEBS-X and SEBS-MA-X, where X is the weight fraction of ZnO nanoparticles in the nanocomposite. For example, SEBS-MA-5 corresponds to the nanocomposite made from SEBS-MA as a matrix and containing 5 wt% ZnO.

4.2.3 Characterization

Cross-sections of different nanocomposites were observed to characterize the dispersion of nanoparticles either by Scanning Electron Microscopy (SEM) or Transmission Electron Microscopy (TEM). Hitachi S-3600N microscope was used for SEM micrographs. The samples were cryogenically fractured using a Leica Microtome and a Tungsten knife operated at -100 °C. A gold layer of thickness 20 nm was sputtered on the cross-section of the samples before imaging. A Jeol JEM-2100F microscope was used for TEM micrographs. Ultrathin sections of the samples were obtained using an ultramicrotome operated at -100 °C and placed on copper grids.

The morphology of the block copolymer was characterized by Atomic Force Microscopy (AFM). The images were obtained using a Veeco atomic force microscope operated in tapping mode under ambient conditions at a scanning rate equal to 1 Hz. Samples for AFM imaging were prepared by spin coating of nanocomposites suspensions directly onto a glass substrate, allowing free evaporation of the solvent in open air. The obtained thin films were then subjected to the same thermal treatment applied to the free-standing films during vacuum drying and compression molding steps: 48 hours at 65 °C and 15 minutes at 200 °C.

The interactions at the nanoparticle-polymer interfaces were studied by means of Fourier Transformed Infra-Red (FTIR) spectroscopy. The spectra were recorded in the transmission mode in the wavelengths range from 500 to 4000 cm⁻¹ with spectral resolution of 4 cm⁻¹, using a Nicolet 6700 Spectrometer.

Rheological measurements were performed at 200 °C to characterize the evolution of dynamic moduli of the nanocomposites in the molten state. A strain-controlled rheometer MCR 501 Anton Paar was used. Small amplitude oscillatory shear (SAOS) tests were performed in the linear viscoelastic regime under a strain equal to 0.5% in the frequency range from 10⁻² to 300 rad.s⁻¹. 25 mm-diameter parallel plate geometry was used and 1 mm sample gap was used.

The complex dielectric permittivity of the different samples was measured at room temperature using a Novocontrol broadband spectrometer. Specimens of 20 mm in diameter were placed between parallel plated brass electrodes to form a plane-plane capacitor. Measurements swept through a frequency range from 10^{-2} Hz up to 10^{5} Hz under an excitation voltage of 3V. Besides, temperature ranges between -100 °C and 120 °C were considered for selected samples.

The AC short-term breakdown strength of the samples was measured in a dielectric oil environment using ball-type electrodes of 4-mm diameter. A ramp of 1 kV/s and a frequency of 60 Hz were applied until breakdown occurred. Samples of average thickness around 200 μ m were used for this test. But, since the thickness of the samples was not strictly uniform, all the measurements were corrected to equal the breakdown strength of a 200 μ m thick film using the power law relationship (equation (4.1)) between the breakdown field and the film thickness in order to obtain comparable results (Takala et al., 2010). The data were then retrieved considering a two-parameter Weibull distribution, by means of commercial software.

$$E_2 = E_1 \left(\frac{d_2}{d_1}\right)^{-0.4} \tag{4.1}$$

Where:

- d1: the real thickness of the sample;
- E1: the measured dielectric strength at d1;
- d₂=200 µm: the thickness selected for normalization;
- E₂: the estimated dielectric strength at the desired thickness d₂.

To evaluate the resistance of the fabricated materials to surface erosion, samples of average thickness equal to 500 µm were exposed to partial discharges for 30 hours, using a point-toplane geometry set-up operated in open air, as represented in Figure 4-2. A sinusoidal voltage of approximately 7 kV_{RMS} and a frequency of 300 Hz were applied to the high-voltage rod electrode. The electrode tip is made from nickel-plated carbon steel and has a 1-mm radius. The samples were fixed to the ground plane electrode using Kapton tape and separated from the tip by an air gap of 200 μ m. Then, the eroded volume was evaluated by the use of a mechanical profilometer. Before mapping the samples with the profilometer, the samples were carefully cleaned in an ultrasonic bath, to remove the debris caused by partial discharges.



Figure 4-2: Sketch of the set up for resistance to surface erosion test

The thermal conductivity of the samples was evaluated using a DTC-25 guarded heat flow meter (TA instruments) in accordance with ASTM E1530 standard. Samples of 2 inches diameter and average thickness of 500 μ m were subsequently placed between the upper heating plate and the lower cooling plate of the device. A pressure of 15 psi was applied on the upper plate to ensure good contact between the sample and the plates. The upper plate was heated to 45 °C while the lower plate was cooled to 15 °C, creating a Δ T over the sample and leading to a medium sample temperature of 25 °C. Samples have been allowed for thermal stabilization, which occurs usually within 2 to 3 hours, in the measuring chamber. A heat flux transducer attached to the lower plate measures the heat flow Q through the sample, from which the thermal conductivity can be deduced using equation (4.2).

$$\lambda = \frac{Q_{/A}}{\Delta T_{/t}} \tag{4.2}$$

Where:

A: surface of the sample;

t: thickness of the sample;

Q: heat flow;

 ΔT : temperature difference.

Finally, DC electrical conductivity of the nanocomposites was measured under ambient conditions, using a two-probe method. 40 mm diameter samples of thickness around 500 μ m were used. Different electric fields ranging from 17 kV/cm to 170 kV/cm were applied using a High Voltage DC source. The charge and discharge currents were measured using a Keithley 6487 Picoammeter. The data were collected for 600 s to reach quasi-steady state currents. The characteristic I-V curve corresponding to each nanocomposite was subsequently plotted.

4.3 **Results and discussion**

4.3.1 Dispersion and morphology

The nanoparticles dispersion in both sets of nanocomposites, respectively SEBS-ZnO and SEBS-MA-ZnO, was first characterized by SEM. The SEM micrographs reported in Figure 4-3 reveal that agglomerations are present in the case of SEBS-ZnO nanocomposites (indicated by the arrows in Figure 4-3(a)) while they are absent in the case of SEBS-MA-ZnO nanocomposites. To investigate further the dispersion in the latter case, TEM was performed on SEBS-MA-5 nanocomposite. The images reported in Figure 4-4 reveal homogeneous and quasi-individual dispersion of nanoparticles. Their size distribution was investigated by particle size analysis reported in annex III of the supporting file (Figure A III-1 and Figure A III-2). In particular, the size of more than 60% of the dispersed ZnO nanoparticles and their small agglomerations ranges between 20 nm and 60 nm, which is equal to or larger than the size of the SEBS block copolymer nanodomains estimated to be in the range 15-30 nm for both blocks according to reference (Leice G Amurin et al., 2016) and AFM images reported in Figure 4-7. AFM results will be discussed further later in the manuscript. Hence, it is difficult to locate the nanoparticles exclusively in one block, either PS or PEB. However, it is expected that the nanoparticles exhibit higher affinity to the PEB block due to favorable interaction between the surface of ZnO nanoparticles and the MA groups grafted onto the PEB block. To confirm this potential interaction, FTIR was performed. The spectra are reported in Figure 4-5.



Figure 4-3: SEM micrographs of: (a) SEBS-5 (arrows indicate the agglomerations) and (b) SEBS-MA-5



Figure 4-4: TEM micrographs of SEBS-MA-5 nanocomposite

Figure 4-5(a) shows the FTIR spectra of SEBS-MA based materials in the wavenumbers range from 1500 to 2000 cm⁻¹. In particular, the spectrum of SEBS-MA shows absorption bands at respectively 1776, 1740 and 1713 cm⁻¹ indicating the presence of carbonyl-containing groups, mainly maleic anhydride and potentially maleic acid (Barra, Crespo, Bertolino, Soldi, & Pires, 1999; "Characteristic IR Absorption Frequencies of Organic Functional Groups," ; Xiong et al., 2013; X. Zhou, Yu, Lin, & Chen, 2013). In the spectra of the nanocomposites, the absorption bands at 1713 and 1776 cm⁻¹ are still detected but the peak

intensity at 1713 cm⁻¹ seems to be considerably reduced. This observation is rather confirmed if we consider a ratio between this absorption band and a reference band (not affected by the potential interaction between ZnO nanoparticles and MA) such as the one observed at 1583 cm⁻¹ and assigned to the vibrations of the benzene rings of PS block (G. Chen, Liu, Chen, & Qi, 2001; Gupta, Wang, Hanssen, Hsia, & Datla, 1995). In fact, the ratio decreases especially at large ZnO content. For instance, it drops down from approximately 2.3 for the neat SEBS-MA to 1 for SEBS-MA-20 nanocomposite. Besides, the absorption band at 1713 cm⁻¹ is slightly shifted to lower wavenumbers at increasing ZnO content. Simultaneously, the intensity of the absorption band at 1740 cm⁻¹, which corresponds actually to the stretching vibration of ester carbonyls ("Characteristic IR Absorption Frequencies of Organic Functional Groups,"), seems to be increasing with increasing ZnO concentration, if we compare with the same reference, i.e. the absorption band at 1583 cm⁻¹. It reaches 2 for SEBS-MA-20 nanocomposite compared to 0.62 for the neat SEBS-MA. This behavior supports the hypothesis of formation of ester bonds upon the reaction between MA and hydroxyl groups on the surface of ZnO nanoparticles. Figure 4-5(b) shows the FTIR spectra of SEBS-MA based materials in the wavenumbers range from 3000 to 3800 cm⁻¹. As indicated by the arrow, a broad absorption band appeared between 3200 and 3600 cm⁻¹ in the spectra of all the nanocomposites. It most likely corresponds to stretching vibration of hydrogen-bonded hydroxyl groups. It is broad due to overlapping of several O-H stretching modes (Kuo & Chang, 2001). This behavior indicates the formation of hydrogen bonds as well after the reaction between MA and hydroxyl groups on the nanoparticles surfaces (Ndiaye et al., 2011; Sari, Moradian, Bastani, & Stribeck, 2012; Xiong et al., 2013; X. Zhou et al., 2013). In summary, in the nanocomposites studied in the present work, ZnO nanoparticles are organically modified with an alkyl ammonium salt which is one of the functional groups that are commonly used for the preparation of pre-intercalated clay layers. Although the used nanoparticles are spherical, this modification helps to improve the compatibility with the organic polymer matrix by decreasing the hydrophilic character of the nanoparticles. Besides, maleic anhydride graft attached to the PEB block can be considered as a second effective functional group which is able to bond with hydroxyl groups that partially cover the surface of the nanoparticles, forming ester and hydrogen bonds. The resulting complexes act as in-situ

formed cross-linked structures that improve further the dispersion, as seen in TEM micrographs (Figure 4-4), and affect drastically the rheological properties which will be discussed later in the manuscript.



Figure 4-5: FTIR spectra of SEBS-MA-ZnO nanocomposites: (a) in the absorption range 1500-2000 cm⁻¹ and (b) in the absorption range $3000-3800 \text{ cm}^{-1}$

A scheme of the possible reaction mechanism is presented in Figure 4-6 (Ndiaye et al., 2011; Sari et al., 2012). The system studied in this manuscript is very similar to many clay containing-maleated polymer nanocomposites that feature an excellent exfoliation of clay due to simultaneous surface treatment by organic groups, which result in pre-intercalated clay layers, and strong interaction between MA and hydroxyl groups at the edge of clay layers improving the peeling mechanism and consequently the exfoliation of clay into individual layers (Iwasa, Ueda, Shibayama, & Fukatani, 2002; L.A. Utracki, 2004).



Figure 4-6: Reaction scheme between maleic anhydride and the surface of the nanoparticles (R: alkyl ammonium salt)

As a consequence of the good interaction with MA, ZnO nanoparticles tend to be individually dispersed and exhibit higher affinity to PEB phase to which the MA group is grafted. However, as the size of the nanoparticles is comparable to or greater than the size of the PEB domains, it is thermodynamically challenging to locate them inclusively in the rubber phase (Emna Helal et al., 2015; Sarkar & Alexandridis, 2015). Only the nanoparticles with the smallest size will potentially locate in the rubber phase. In order to reveal the obtained SEBS morphology and the possibility of selective location of nanoparticles, some TEM grids were treated with ruthenium tetroxide RuO₄ to enhance contrast between the different phases. However, the staining was not successful as ZnO reacted with RuO₄ (TEM micrographs of the treated samples can be consulted in Figure A III-3 of the supporting file (annex III)). Thus, AFM was carried out instead to overcome this problem. In general, lamellar morphology is expected when SEBS block copolymer is prepared from solvent blending using toluene

without annealing at elevated temperatures (E Helal et al., 2015). However, the compression molding at 200 °C and the addition of nanoparticles can distort the block copolymer morphology (Cano, Gutierrez, & Tercjak, 2013). Figure 4-7(b) shows AFM images corresponding to SEBS-MA-1 nanocomposite subjected to the same thermal treatment as the compression molded samples. A distorted "fingerprint" morphology is observed, where PS domains (bright phase) are mostly in the form of short cylinders (Ganguly & Bhowmick, 2008; Seppala, 2012). This morphology is observed in the neat SEBS as well (Figure 4-7(a)), but with better long-range order. The nanoparticles appear as bright spots. The smallest ones are more in contact with the PEB block (dark phase) while the biggest ones do not selectively locate. This observation agrees well with TEM and FTIR results.



Figure 4-7: AFM images of: (a) Neat SEBS-MA, (b) SEBS-MA-1 nanocomposite, (c) Zoom on the region indicated by the square in (b) (showing the same magnification as (a))

4.3.2 Rheological properties

SAOS results corresponding to respectively SEBS-ZnO and SEBS-MA-ZnO nanocomposites containing different nanoparticles concentrations are reported in Figure 4-8. The measurements were done in the linear regime in the frequency range from 10⁻² to 300 rad.s⁻¹. Besides, the low frequency range was extended down to 10⁻³ rad.s⁻¹ for selected samples to evaluate further the non-terminal behavior. Plots corresponding to dynamic storage and loss

moduli, G' and G'', are reported in Figure 4-8(a)-(b) while those corresponding to the complex viscosity modulus $|\eta^*|$ are reported in Figure 4-8(c)-(d).

By comparison with the linear viscoelastic response of unfilled SEBS and SEBS-MA, a substantial effect of ZnO nanoparticles was observed in both sets of nanocomposites. With increasing ZnO content, both storage modulus, G', and loss modulus, G'', of the nanocomposites were progressively enhanced and became less dependent on the angular frequency throughout the test range. The reduced dependence on frequency indicates that the nanocomposites gradually exhibit solid-like behavior which is attributed to the confinement of polymer chains and the formation of ZnO networks, due to good adhesion between the nanoparticles and the polymer matrix (Hong et al., 2006). This behavior was more prominent in SEBS-MA-ZnO nanocomposites compared to SEBS-ZnO nanocomposites. In fact, for SEBS-ZnO nanocomposites, a slight increase of both storage and loss moduli, was exhibited in the nanocomposites compared to the pure material due to the confinement of the polymer chains in the presence of the nanoparticles. In SEBS-MA-ZnO nanocomposites, the same behavior is exhibited but the degree of increase is higher than one order of magnitude, compared to the neat, for both moduli. This behavior is most likely due to the improved dispersion and affinity to PEB block which were more efficient in confining polymer chains. Moreover, a plateau in G'- ω curve in log-scale was approached for all the nanocomposites prepared from maleated SEBS, even at low ZnO content equal to 1 wt%. This plateau was more evident when extending the frequency range to 10^{-3} rad.s⁻¹, as seen in Figure 4-8(a). The low frequency slope, calculated in the range 10⁻¹ rad.s⁻¹ to 10⁻² rad.s⁻¹, decreased from 0.17 for neat SEBS-MA, which corresponds to hexagonally-packed cylindrical morphology that was probably improved during the SAOS test (Carastan et al., 2008), down to 0.03 for SEBS-MA-5 nanocomposite. Since the rheological measurements in the low frequency regime reflect the long range motion of polymer chains, this particular non-terminal behavior means that the long-range motion is hindered due to the formation of a network between the ZnO nanoparticles and the polymer chains, i.e. rheological percolation, resulting in incomplete relaxation of the chains(C. Gao et al., 2014; Q. Zhang & Archer, 2002). Moreover, the values of complex viscosity, $|\eta^*|$, increased in the whole frequency range, as shown in Figure 4-8(c), along with a more pronounced shear-thinning behavior at higher ZnO content. Comparing SEBS-MA-ZnO nanocomposites containing respectively 10 and 20wt% nanoparticles, same values of G', G'' and $|\eta^*|$ can be observed. Hence, it seems that the rheological percolation occurs between 5wt% (0.9 vol%) and 10wt% (1.8 vol%). This range is below the predicted theoretical rheological percolation for spherical nanoparticles (Q. Zhang & Archer, 2002). This fact seems to support the hypothesis of good interaction between the nanoparticles and the rubber phase of the block copolymer which may result in lowering the threshold for rheological percolation.





Figure 4-8: SAOS measurements of SEBS-ZnO vs. SEBS-MA-ZnO nanocomposites: (a)-(b) storage modulus G' (●) and loss modulus G'' (■) and (c)-(d) complex viscosity modulus |η*|

In order to investigate if the non-terminal behavior observed in SEBS-MA-ZnO nanocomposites is indeed due to the rheological percolation and not due to a change of SEBS block copolymer morphology during the SAOS test to body-centered cubic spherical morphology, which is also characterized by a zero-slope in G'- ω plot as a non-terminal

behavior (Carastan et al., 2008; Kossuth, Morse, & Bates, 1999), a time sweep test were performed. The time sweep test describes the evolution of viscoelastic moduli as function of time under a constant temperature, frequency and strain. In this test, a change of morphology will be manifested by a sudden change of the storage modulus (Carastan et al., 2008). In literature, changes of morphology in block copolymer nanocomposites have been reported, especially in the case where nanoparticles are selectively located in one phase and consequently increase its volume fraction (Cano et al., 2013; Carastan et al., 2008). Figure 4-9 shows the evolution of the storage moduli of unfilled SEBS-MA and SEBS-MA-5 nanocomposite for 8 hours at a fixed frequency equal to 0.05 rad.s⁻¹. G' of both samples remained constant over time which confirms that the morphology of the block copolymer didn't change. Hence, the plateau observed in G'- ω plots of the nanocomposites, is rather due to rheological percolation.



Figure 4-9: Time sweep test at 200 °C, 0.05 rad.s⁻¹ and 0.5% strain

4.3.3 Low-field dielectric characterization

4.3.3.1 Complex dielectric permittivity

Figure 4-10 reports respectively the values of real and imaginary parts of the complex dielectric permittivity, ε' and ε'' respectively, of both SEBS-MA-ZnO and SEBS-ZnO nanocomposites sets, measured at room temperature. The comparison of dielectric response of both types of nanocomposites shows that the dielectric behavior is strongly affected by the presence of MA and the quality of ZnO dispersion. For the unfilled matrices, it can be seen that SEBS-MA has slightly higher real and imaginary permittivities compared to SEBS. This is most likely due to the presence of MA polar groups. Regarding the SEBS-MA nanocomposites, it can be seen from Figure 4-10(a) and Figure 4-10(b) that up to 10wt% loading, there is no significant increase of either the imaginary or the real part of the complex permittivity. This has been frequently observed in the case of well dispersed ceramic filled polymer nanocomposites (Praeger, Andritsch, Swingler, & Vaughan, 2014; I. Tsekmes, Kochetov, Morshuis, & Smit, 2014), particularly when functional groups creates covalent bonding between particles and matrix (Bouchaib Zazoum et al., 2016). The most popular explanation for this behavior is the interlayer theory suggesting an interlayer zone between the nanofiller and the matrix having different dielectric properties than both the matrix and the particles (for example a lower permittivity) (Heid, David, & Fréchette, 2016; I. A. Tsekmes, 2016). However, when hydrophilic filler, such as a metallic oxide, is used, the occurrence of a water layer also surrounding the particle is frequently observed which invariably leads to a strong relaxation peak usually within the 10^{-2} to 10^{5} Hz frequency window (see discussion in the paragraph below). This was not observed in the case of SEBS-MA nanocomposites for loadings below 20 wt%.

The poor dispersion and the presence of agglomerations lead to much higher dielectric losses (and consequently permittivity) in the case of the SEBS-ZnO nanocomposites as it can be seen in Figure 4-10(c) and Figure 4-10(d). Due to the distribution of the agglomerate size, very broad relaxation mechanisms were observed from 5wt% loading. At a high

nanoparticles loading equal to 20wt%, a strong increase of dielectric losses was observed for both nanocomposites. In the case of the SEBS-MA-20, a clear and rather narrow interfacial relaxation peak appeared in the vicinity of 10^2 - 10^3 Hz suggesting a somewhat narrow distribution of filler size. On the other hand, SEBS-20 shows larger increase of its dielectric losses over the whole frequency range showing a very broad relaxation mechanism possibly resulting from the overlapping of several interfacial relaxation processes related to different agglomerate size and properties. Step-like increases related to these relaxation peaks were simultaneously observed in real permittivities of these samples, resulting in increased values of real permittivity, especially at low frequencies. The observed relaxation peaks might have different origins including the presence of a water layer absorbed at the interface between ZnO nanoparticles and polymer chains. In fact, water sorbed from the ambient atmosphere constitutes a conductive layer leading to an interfacial relaxation observed at intermediate frequencies (Kremer & Schönhals, 2012). This phenomenon was reported in the literature (Couderc, David, Frechette, & Medidoub, 2013; David & Fréchette, 2013; Glaskova & Aniskevich, 2009; I. Hosier, Praeger, Vaughan, & Swingler, 2015; Kofod et al., 2011; Lau et al., 2013), for polymers filled with several types of hydrophilic nanofillers, among them metal oxides, which were demonstrated to effectively absorb humidity. The geometry and the fraction of this conductive layer depend strongly on the quality of the interface and consequently the quality of the dispersion. Thus, in materials suffering from the presence of agglomerations, different interfaces lead to different thicknesses of water layers that might coexist resulting in more than one relaxation peak, observed at distinguished frequencies and featuring different dielectric strengths (Kremer & Schönhals, 2012; Lau et al., 2013). However, in specific cases where nanoparticles are functionalized with an adequate surface modification, reduced or negligible effect of humidity was reported (David & Fréchette, 2013; David et al., 2015; Lau et al., 2013). Moisture absorption can also affect dipolar relaxation dynamics by enhancing the molecular mobility of its related dipoles and consequently shifting the corresponding peak to higher frequencies (Kofod et al., 2011).



Figure 4-10: Complex dielectric permittivity at room temperature of SEBS-MA-ZnO vs. SEBS-ZnO nanocomposites: (a) Real part of permittivity of SEBS-MA-ZnO nanocomposites, (b) Imaginary part of permittivity of SEBS-MA-ZnO nanocomposites, (c) Real part of permittivity of SEBS-ZnO nanocomposites and (d) Imaginary part of permittivity of SEBS-MA-ZnO nanocomposites

To follow the evolution of the different observed relaxations, isothermal dielectric measurements were performed on unfilled materials and selected nanocomposites at temperatures ranging from 25 °C up to 120 °C. Dielectric loss spectra of unfilled matrices are reported in Figure 4-11(a) and Figure 4-11(b) while those of nanocomposites containing 5 wt% ZnO are reported in Figure 4-11(c) and Figure 4-11(d) (Data corresponding to nanocomposites containing 1 wt% ZnO are reported in Figure A III-4 of the supporting file (annex III)). First, it can be seen that values of ε '' of both neat materials increased at low frequencies and elevated temperatures due to increased charge carrier fluctuation leading to both direct conduction and most likely electrode polarization. Besides, no specific relaxation

process was detected even in the case of SEBS-MA, by contrast to some studies reporting relaxation related to dipolar fluctuations of MA polar groups (Böhning et al., 2005). As a consequence, unfilled SEBS and SEBS-MA don't undergo any relaxation processes in the temperature range studied (Kofod et al., 2011).

From Figure 4-11(c) and Figure 4-11(d), it can be seen that at elevated temperatures and low frequencies below 1 Hz, the dielectric loss of SEBS-MA-5 exhibits strong frequency dependence, especially at temperatures higher than 60 °C, and exceeds largely that of SEBS-5 nanocomposite, which features only a broad relaxation peak in the intermediate frequency range shifting to higher frequencies with increasing temperatures. The large increase of ε " of SEBS-MA-5 nanocomposite at low frequencies indicates an important contribution from conductivity. The sample becomes even slightly conductive at temperatures higher than 60 °C. An additional contribution from electrode polarization might also be present and overlap potential relaxation phenomena. The slope of the ε "-frequency curve in log-scale at elevated temperatures is between -0.9 and -0.86 for SEBS-MA-5 nanocomposite, which supports the assumption of presence of non-ohmic conduction (Kremer & Schönhals, 2012). This effect was also observed in the rest of SEBS-MA-ZnO nanocomposites and seems to reach its maximum at 5wt% ZnO. However, it was not observed in SEBS-ZnO nanocomposites even at 20 wt% ZnO.

The improved dispersion, homogeneous distribution and reduced interparticle distance achieved in SEBS-MA-ZnO nanocomposites are most likely at the origin of this interesting behavior. Moreover, the formation of complexes upon the interaction of MA with functional groups on the surface of the nanoparticles as well as the established network between the nanoparticles and polymer chains, as confirmed by rheological properties, might modify the nature of the interfaces and charge carriers activity, which ultimately leads to a much lower percolation threshold, under favorable heating conditions.



Figure 4-11: Imaginary part of the dielectric permittivity as functions of frequency and temperature of: (a) Neat SEBS-MA, (b) Neat SEBS, (c) SEBS-MA-5 nanocomposite and (d) SEBS-5 nanocomposite

To evaluate the increase of AC conductivity, the real part of the complex electrical conductivity, σ ', might be estimated using equation (4.3). It includes the contribution of direct conduction as well as contribution from dielectric losses associated with relaxation processes and electrode polarization.

$$\sigma' = \omega \varepsilon_0 \varepsilon'' \tag{4.3}$$

Where:

- σ ': real part of complex conductivity;
- ε₀: vacuum permittivity;
- ε '': imaginary part of complex permittivity at angular frequency ω .

In Table 4-1, values of σ' , at respectively 25 °C and 100 °C, are reported for different nanocomposites. These values have been calculated at a relatively low frequency equal to 10^{-1} Hz in order to compare them with values of dc conductivity that will be discussed later in the manuscript. It is worth mentioning that, in the case of the studied samples, an important increase of electrical conductivity is not expected, since the intrinsic electrical conductivity of semi-conductive ZnO nanoparticles is relatively low and doesn't increase considerably with temperature. It can be seen that in the case of neat materials and SEBS-ZnO nanocomposites, the increase of conductivity with increasing temperature from 25 °C to 100 °C doesn't exceed 1 order of magnitude generally. In the case of SEBS-MA-ZnO nanocomposites, the conductivity increases by 3-4 orders of magnitude. This effect seems to saturate at 5 wt%, which is in line with the rheological properties.

	σ' at 25 °C (S.cm ⁻¹)	σ' at 100 °C (S.cm ⁻¹)
SEBS	3.0 x 10 ⁻¹⁸	3.6 x 10 ⁻¹⁷
SEBS-1	3.6 x 10 ⁻¹⁸	1.2 x 10 ⁻¹⁶
SEBS-5	2.3 x 10 ⁻¹⁷	4.4 x 10 ⁻¹⁷
SEBS-20	3.2 x 10 ⁻¹⁶	4.5 x 10 ⁻¹⁶
SEBS-MA	2.3 x 10 ⁻¹⁷	8.1 x 10 ⁻¹⁷
SEBS-MA-1	3.6 x 10 ⁻¹⁸	1.9 x 10 ⁻¹⁵
SEBS-MA-5	7.8 x 10 ⁻¹⁸	6.4 x 10 ⁻¹⁴
SEBS-MA-20	3.0 x 10 ⁻¹⁷	1.7 x 10 ⁻¹⁴

Table 4-1: Values of real part of complex conductivity at 10⁻¹ Hz

As observed in room temperature spectra reported in Figure 4-10, additional relaxation peaks in the intermediate frequencies range are detected in the dielectric loss spectra of some nanocomposites, especially at elevated ZnO loadings. In order to investigate these relaxation processes, isothermal measurements at higher temperatures were performed on both SEBS-MA-20 and SEBS-20 nanocomposites. Their corresponding spectra of ε " are reported in Figure 4-12(a) and Figure 4-12(b). In particular, Figure 4-12(a) shows that SEBS-MA-20 exhibits strong conduction at low frequencies eventually associated with a MWS peak, similarly to what have been observed earlier in the spectrum of SEBS-MA-5 (Figure

4-11(c)). In addition, a relaxation peak is observed initially around 10^3 Hz and shifted towards higher frequencies at elevated temperatures (indicated by an arrow). In SEBS-20, two peaks were observed initially at room temperature, at two different frequencies as indicated by the arrows in Figure 4-12(b). At increasing temperatures, both peaks were shifted to higher frequencies as well and seem to merge in one peak with higher asymmetry at the low-frequency side.

These new relaxations observed in the materials containing high loadings of nanoparticles might be related to the presence of a conductive water layer adsorbed at the interface between the polymer and the metal oxide nanoparticles as mentioned earlier. Although it was reported in several studies that relaxation processes affected by moisture absorption exhibit a distinguished behavior consisting on the displacement of the peak maximum to lower frequencies starting from a certain temperature (Böhning et al., 2005; Couderc et al., 2013; David & Fréchette, 2013; Lau et al., 2013), a regular trend of the displacement of the peak towards higher frequencies was also reported in other studies (Ciuprina et al., 2010; David & Fréchette, 2013). In our case, the latter behavior is observed. Besides, it seems that the temperature dependence of the peak maximum frequency is stronger in SEBS-ZnO compared to SEBS-MA-ZnO.

In order to confirm if the additional relaxations are corresponding to a conductive water layer adsorbed at the interface between the nanoparticles and polymer chains, nanocomposites containing 5 wt% and 20 wt% ZnO from both sets, were dried for 3 days at 65°C under vacuum before rescanning their dielectric spectra. In fact, if humidity absorption is at the origin of these relaxations, corresponding peaks are expected to shift to lower frequencies or completely disappear after drying (Böhning et al., 2005; Couderc et al., 2013; David & Fréchette, 2013; Kofod et al., 2011; Lau et al., 2013). In the case of SEBS-MA-ZnO nanocomposites, no change of the dielectric spectra was observed after drying, for both studied ZnO contents: 5 wt% and 20 wt% ZnO (Figure A III-5 and Figure A III-6 in annex III). However, in the case of SEBS-ZnO nanocomposites, some changes were observed after

drying (Figure A III-7 in annex III). In particular, peak maximum frequencies of the observed relaxations were not significantly affected but the dielectric losses increased mainly at low frequencies and a new relaxation process is observed at elevated temperatures. This behavior is in agreement with findings from the literature reporting that the effect of moisture is reduced on treated nanoparticles and that in the dry state, the dielectric losses might increase due the presence of coupling agents (David & Fréchette, 2013; David et al., 2015; Lau et al., 2013). As a consequence, it was not possible to confirm a correlation between the relaxation processes observed at intermediate frequencies and the presence of water layer surrounding the nanoparticles, especially in the case of SEBS-MA-ZnO nanocomposites. Probably, additional drying is necessary to observe significant changes but overall it can be concluded that moisture absorption is not a major issue in the studied SEBS-ZnO and especially SEBS-MA-ZnO samples. If the effect of moisture absorption is negligible, the observed relaxations might be caused by the co-existence of different types of interfaces: polymer-particle and particle-particle, especially at high nanoparticles loadings and in the presence of agglomerations, resulting in broad and multiple relaxations (Figure 4-12(b)). Another possible origin for the observed relaxations, especially at high loadings of nanoparticles, is the existence of an interphase region where polymer chains might feature restricted mobility (Michel Fréchette et al., 2014; Yu Lin et al., 2015; Tsagaropoulos & Eisenburg, 1995). In this case, further investigations are required to confirm these phenomena.



Figure 4-12: Imaginary part of electric modulus as functions of frequency and temperature of: (a) SEBS-MA-20wt% ZnO nanocomposite and (b) SEBS-20wt% ZnO nanocomposite

In order to evaluate the dielectric losses of the nanocomposites compared to the pure materials and between themselves at operational conditions, values of the dissipation factor tan δ equal to the ratio ϵ''/ϵ' , at a fixed frequency equal to 66 Hz (close to power frequencies) and different temperatures: 25, 40 and 60 °C were reported in Table 4-2 for the 5, 10 and 20 wt% nanocomposites (1wt% nanocomposites did not show any significant increase of the dissipation factor). At a first glance, it can be seen that at this frequency, nanocomposites prepared from SEBS-MA don't exhibit higher losses compared to nanocomposites prepared from SEBS despite the fact that they show stronger conductivity at low frequencies and elevated temperatures and that maleated SEBS features higher losses compared to unmaleated SEBS. Besides, both sets of nanocomposites, containing 1 wt% and 5wt% nanoparticles, have dielectric losses similar to (less than one order of magnitude increase) or lower than unfilled matrices even at 60 °C. At higher ZnO contents equal to 10wt% and 20wt%, losses increase up to 2 orders of magnitude compared to unfilled materials. However, SEBS-MA-ZnO feature lower losses compared to SEBS-ZnO. For instance, the maximum dissipation factor for SEBS-20 nanocomposite in the studied temperature range is equal to 4.57 x 10⁻² compared to 3.78 x 10⁻³ for SEBS-MA-20. Overall, by controlling the dispersion of ZnO in SEBS-MA, it is possible to incorporate up to 20wt% ZnO while limiting the increase of the loss factor to less than 1 order of magnitude. At lower concentrations up to 5wt% ZnO, no additional dielectric losses are induced.

ZnO	25 °C		40 °C		60 °C	
(wt%)	SEBS	SEBS-MA	SEBS	SEBS-MA	SEBS	SEBS-MA
0	6.33 x 10 ⁻⁰⁴	1.5 x 10 ⁻⁰³	1.54 x 10 ⁻⁰⁴	9.94 x 10 ⁻⁰⁴	1.99 x 10 ⁻⁰⁴	8.62 x 10 ⁻⁰⁴
5	3.34 x 10 ⁻⁰³	1.04 x 10 ⁻⁰³	1.28 x 10 ⁻⁰³	7.6 x 10 ⁻⁰⁴	1.34 x 10 ⁻⁰³	2.03 x 10 ⁻⁰³
10	1.18 x 10 ⁻⁰²	1.37 x 10 ⁻⁰³	5.39 x 10 ⁻⁰³	1.40 x 10 ⁻⁰³	2.45 x 10 ⁻⁰³	5.02 x 10 ⁻⁰³
20	4.57 x 10 ⁻⁰²	3.78 x 10 ⁻⁰³	3.23 x 10 ⁻⁰²	2.56 x 10 ⁻⁰³	3.13 x 10 ⁻⁰²	3.51 x 10 ⁻⁰³

Table 4-2: Dissipation factor (tan $\delta = \epsilon^{\prime\prime}/\epsilon^{\prime}$) at 66 Hz for different nanocomposites
4.3.4 High-field dielectric characterization

In this section, several dielectric characterizations, performed to evaluate the dielectric performance at high electric fields, will be discussed.

4.3.4.1 Short-term breakdown strength

Figure 4-13(a) and Figure 4-13(b) present Weibull plots of breakdown strength corresponding respectively to SEBS-ZnO and SEBS-MA-ZnO sets of nanocomposites. In SEBS-ZnO set, the values of breakdown strength decrease progressively, when increased loadings of ZnO nanoparticles are incorporated, due to the presence of nanoparticles agglomerations which act as defect centers distorting the electric field around them. This behavior has been repeatedly reported in literature for ZnO contents equal to 5wt% or higher (S. Chen et al., 2010; Cheng, Guo, Wang, & Zhang, 2013; Hong et al., 2003; Tian et al., 2012). In particular, an initial drop from 107 kV/mm down to approximately 86 kV/mm was observed in nanocomposites containing 5 wt% up to 10 wt% ZnO, both exhibiting similar performance. Then, a second drop down to 73 kV/mm was observed when ZnO content is increased up to 20 wt%. This second large decline is an indication that a network of agglomerated ZnO nanoparticles. This observation will be discussed later in correlation with dc electrical conductivity and thermal conductivity results showing consistent increase at 20wt% ZnO loading.

In SEBS-MA-ZnO nanocomposites set, a different behavior was observed. First of all, the breakdown strength of the maleated SEBS decreased to 93 kV/mm compared to 107kV/mm for non maleated SEBS. This drop is most likely due to lower stiffness in SEBS-MA (Kollosche & Kofod, 2010). Regarding the nanocomposites, it seems that the incorporation of nanoparticles has only marginal effect. In fact, the breakdown strength remained equal or slightly higher than the unfilled SEBS-MA even at ZnO loading as high as 20wt%. This behavior can be the sum of two opposite mechanisms, both resulting from nanoscale dispersion and homogeneous distribution of the nanoparticles, which compensate each other:

- Homogeneous distribution of particles and their nanosize results in large interfacial area and consequently large amounts of charge traps and efficient scattering of electrons, inducing an initial increase of the breakdown strength (Fleming et al., 2008; Smith et al., 2008; Tian et al., 2011, 2012).
- Reduced interparticle distance due to improved dispersion, especially at high nanoparticles loadings, leads to easier tunneling. Moreover, mismatch of permittivity between the nanoparticles and the polymer matrix leads to local electric field distortion around them (Hayase et al., 2006). Both factors might induce a decrease of the breakdown strength that neutralizes the initial increase.

Comparing both sets of nanocomposites, values of breakdown strength of nanocomposites prepared from SEBS-MA are higher than those prepared from SEBS due to the role of tuned dispersion in maintaining the breakdown strength in the same range of the SEBS-MA matrix. But, overall the dielectric strength of all the nanocomposites is lower than that of pure non maleated SEBS. The reduction varies from 30% in the worst case to 10% in the best case, which corresponds to SEBS-MA-5 nanocomposite. Also, it is worth noting that the shape factor β is relatively low in both sets of samples, even the unfilled ones, due to probably the complex morphology of the matrix, i.e. two blocks of different mechanical and dielectric strengths.



Figure 4-13: Breakdown strength of: (a) SEBS-ZnO and (b) SEBS-MA-ZnO nanocomposites

4.3.4.1 Resistance to surface erosion by partial discharges

Resistance to surface erosion by partial discharges is an important criterion to evaluate the aging of polymer insulation in operating conditions. In fact, when subjected to high electric fields, the insulation surface can be degraded and its effective thickness reduced. In this section, we evaluate the resistance to surface erosion of the studied nanocomposites, by estimating the eroded volume induced in each sample after being subjected to a specific electric field. The eroded volumes, calculated using mechanical profilometer data, corresponding to the samples exposed to a voltage equal to 7 kV_{rms} for 30 hours are presented in Figure 4-14. In general, a decrease of the sample eroded volume can be associated with an increase in its resistance to electrical discharges. Comparing unfilled materials, the resistance to surface erosion of SEBS-MA is 10% lower than SEBS. Considering the nanocomposites, both SEBS-ZnO and SEBS-MA-ZnO sets exhibit excellent resistance to surface erosion by partial discharges compared to unfilled matrices, even at low ZnO content equal to 1wt%. This behavior has been commonly reported for nanocomposites and microcomposites containing fillers featuring good resistance to partial discharges (Brockschmidt, Pohlmann, Kempen, & Gröppel, 2011; S. Chen et al., 2010; Heid et al., 2015; Huang et al., 2014), specifically ZnO which is known for its effective shielding of ultraviolet radiation, emitted during the process of corona aging and playing an important role in the degradation of the polymer insulation (S. Chen et al., 2010). However, it is obvious from the reported results that SEBS-MA-ZnO nanocomposites exhibit better performance compared to SEBS-ZnO nanocomposites. For instance, the eroded volume was reduced by almost 60% upon addition of 1wt% nanoparticles to SEBS-MA compared to only 18% when the same amount of nanoparticles is incorporated in SEBS. This behavior is of particular interest. In fact, it demonstrates that the resistance to erosion by partial discharges cannot be explained only by the intrinsic shielding property of ZnO, as in this case one would expect similar performance for similar ZnO content (David & Fréchette, 2013). Thus, the superior performance of SEBS-MA-ZnO is most likely related to the improved dispersion and interaction in SEBS-MA-ZnO nanocomposites which results in large interphase layer with a resistance higher than that of the unfilled matrix and consequently more homogeneous eroded surface and less erosion

depth (Mappings of the eroded area corresponding to different samples are available in Figure A III-8 in annex III). However, in SEBS-ZnO nanocomposites, the inhomogeneous distribution and the presence of agglomerations may cause local electric field enhancements, which lead to locally increased electrical and thermal stresses of the nanocomposite, resulting in intense erosion of the surrounding organic matrix. Results in agreement with these findings were reported in the literature. For instance, comparison of epoxy microcomposites and nanocomposites filled with Alumina showed higher resistance to erosion in the case of nanocomposites (Z. Li, Okamoto, Ohki, & Tanaka, 2011). Also, it was found that nanocomposites filled with high contents of agglomerated polyhedral oligomeric silsesquioxane (POSS) molecules exhibit lower resistance than nanocomposites filled with lower POSS loadings but featuring better dispersion (Heid et al., 2015; Heid, Fréchette, et al., 2016).

At 10 wt% and 20 wt% ZnO loadings, the eroded volume in respectively SEBS-MA-10 and SEBS-MA-20 was almost negligible and it was difficult to estimate it properly from the mapping data. Hence, it seems that the beneficial effect of ZnO nanoparticles saturates at 5 wt% loading, which corresponds to rheological percolation and implicitly to the maximum interphase volume, as estimated from SAOS measurements earlier. At this loading, the eroded volume is reduced by 90%.



Figure 4-14: Resistance to surface erosion by partial discharges

4.3.4.1 DC electrical conductivity: dependence on nanoparticles networks

Non-linear electrical conductivity is an important property in polymer insulation systems in order to reduce space charge accumulation especially under DC electric field. In nanocomposites, the electrical conductivity behavior is often strongly dependent on the distribution of nanoparticles and the possibility of creating conductive pathways, in addition to the intrinsic conductivity of the nanoparticles.

Figure 4-15 presents plots of current density vs. electric field for both sets of nanocomposites: SEBS-ZnO and SEBS-MA-ZnO (Figure 4-15(a)) as well as the evolution of DC electrical conductivity as function of ZnO loading and applied electric field (Figure 4-15(b)). Since SEBS-MA-ZnO nanocomposites exhibit improved dispersion, homogenous nanoparticles distribution and consequently less contact between the nanoparticles, one would expect that these materials will show lower conductivity and will require higher electric fields to exhibit a non-linear I-V relationship compared to SEBS-ZnO nanocomposites (Hong et al., 2006). The data reveal that all the materials feature an improved non-linear behavior in the studied electric field range, especially at electric fields

higher than 50kV/cm (corresponding to a change of the slope), except the unfilled SEBS-MA and SEBS-MA-5 nanocomposite (Figure 4-15(a)). However, different dependences on ZnO content and electric field have been observed for each set: SEBS-ZnO vs. SEBS-MA-ZnO. In SEBS-ZnO nanocomposites, a non-linear behavior is observed starting from 5wt% ZnO content. Besides, a similar trend of current density increase as function of nanoparticles loadings is observed, independently from the applied electric field, which indicates that the resulting increase of conductivity observed in these samples is rather depending on the fraction of agglomerated nanoparticles networks that were progressively formed in the material at increasing loadings due to imperfect dispersion and inhomogeneous distribution.

Regarding SEBS-MA-ZnO materials, it can be observed that the calculated slope for unfilled SEBS-MA is higher than 1 (all the slopes are calculated starting from 50 kV/cm), which corresponds to ohmic conduction. This behavior might be due to the presence of MA polar groups. At 5wt% ZnO loading, no non-linear electrical behavior was observed up to 170 kV/cm. The slope of the I-V plot is even lower than that of unfilled SEBS-MA, which might be explained by the presence of deep trapping sites that limit the current transport through the material (S. Chen et al., 2010; Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016). Similar behavior was reported in the literature for nanocomposites containing low loadings of ZnO up to 5wt%. A significant decrease of conductivity was even observed in some cases (Fleming et al., 2008; Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016). Besides, the increase of the current density of SEBS-MA-ZnO materials, at increasing ZnO wt%, seems to be more dependent on the electric field compared to SEBS-ZnO nanocomposites. In particular, at low electric field (17 kV/cm), no increase of current density was observed up to 10 wt% ZnO. At higher electric fields such as 119 kV/cm and 170 kV/cm, the current density starts increasing at 10 wt% ZnO. The improvement induced by increasing ZnO content from 5 wt% to 10wt% was significantly higher in SEBS-MA-ZnO nanocomposite compared to SEBS-ZnO nanocomposites. This behavior is most likely due to reduced interparticle distance enabling efficient electron tunneling activated at high electric fields (Hong et al., 2003). At increased ZnO loading from 10 wt% up to 20wt% ZnO, a jump of the current density of approximately two orders of magnitude was observed at low electric fields in SEBS-MA-ZnO nanocomposites compared to 1 order of magnitude for SEBS-ZnO nanocomposite. However, at high electric fields, a similar trend of increase is observed which is expected due to the high concentration of nanoparticles. In a summary, the electrical conduction in SEBS-MA-ZnO nanocomposites is thought to be due to tunneling between ZnO nanoparticles. The slope of the current-voltage curve in log scale, which reflects the non-linearity, exhibits a maximum at 10 wt% ZnO (Figure 4-15(a)). This behavior might be correlated with the rheological percolation estimated between 5wt% and 10wt% from SAOS measurements discussed earlier. In SEBS-ZnO nanocomposites, the conduction is rather governed by the formation of paths of touching particles. At increasing ZnO concentrations, conduction paths extend through the entire sample, which is in line with increasing slope of non-linearity (Hong et al., 2003, 2006; Zohrevand, Ajji, & Mighri, 2014).

DC electrical conductivity values reported in Figure 4-15(b) illustrate also the different behaviors exhibited by SEBS-ZnO and SEBS-MA-ZnO nanocomposites as functions of ZnO content and electric field, as discussed earlier. Besides, the values corresponding to neat SEBS-MA and SEBS-MA-5 nanocomposite are close to AC conductivity values reported in Table 4-1, despite the fact that they are measured at higher electric fields and under DC field. However, for the rest of the samples exhibiting stronger non-linearity, the measured values of DC conductivity are higher.



Figure 4-15: (a) current density vs. DC electric field plots of SEBS-MA-ZnO vs. SEBS-ZnO nanocomposites , (b) DC conductivity of SEBS-MA-ZnO vs. SEBS-ZnO nanocomposites as functions of ZnO loading (wt%) and electric field

4.3.5 Thermal conductivity

Relatively high thermal conductivity is also a desirable property in insulating materials in order to facilitate heat dissipation and to avoid the deterioration of the insulation. In general, the medium to transport thermal energy in solid insulations is phonons, i.e. lattice vibrations (W. Cui et al., 2011; Heid et al., 2015; Heid, Fréchette, et al., 2016; Huang, Iizuka, Jiang, Ohki, & Tanaka, 2012). Hence, one would expect that in nanocomposites, the formation of organized networks of thermally conductive particles is necessary to achieve an improvement of the matrix thermal conductivity. In this section, we will evaluate the potential of controlled spatial distribution offered by the established network between ZnO nanoparticles and polymer chains in enhancing the thermal conductivity of the studied nanocomposites.

Thermal conductivity values of both SEBS-ZnO and SEBS-MA-ZnO sets of nanocomposites are reported in Figure 4-16. The accuracy of the method used is within 5%. A distinct behavior has been observed in each set. In particular, in SEBS-ZnO nanocomposites, the thermal conductivity remained constant up to 10wt% ZnO content, and then increased by 20% at 20 wt% ZnO. This abrupt increase indicates that thermal networks might start to form in a large scale only at this relatively high loading due to the presence of agglomerations in SEBS-ZnO nanocomposites and the spherical shape of the nanoparticles. This observation is consistent with the observed decline of breakdown in SEBS-20 nanocomposite. At ZnO contents lower than 20wt%, phonon scattering at nanoparticle-polymer and nanoparticle-nanoparticle interfaces is dominant in SEBS-ZnO nanocomposites and impedes improvement of thermal conductivity (Huang et al., 2012).

In SEBS-MA-ZnO samples, a higher conductivity of neat SEBS-MA, compared to SEBS, can be observed at first. This is most-likely related to a molecular rearrangement of the repetitive units of the copolymer in the presence of MA graft resulting in a denser and more organized thermal network (Murillo & López, 2015). It was demonstrated in several studies that controlling the chemistry of amorphous polymers at molecular level might reduce phonon scattering and consequently enhance the thermal transport (Z. Guo et al., 2014; T.

Zhang & Luo, 2016). Upon the incorporation of 5wt% ZnO nanoparticles, an increase of thermal conductivity by 10% compared to unfilled SEBS-MA and by more than 20% compared to unfilled SEBS is achieved. At higher contents of ZnO equal to 10wt% and 20wt%, the thermal conductivity didn't increase further and probably a slight decrease is even induced. This behavior infers that in SEBS-MA-ZnO nanocomposites, the formation of thermally conductive pathways starts at lower loadings of nanoparticles, compared to SEBS-ZnO, most likely due to rheological percolation that was confirmed by SAOS. This is in agreement with the conclusions of recent studies suggesting that creating ordered structures at the nanoscale within an amorphous polymer matrix would improve the thermal conductivity of the polymer (Heid, Fréchette, et al., 2016). Simultaneously, phonon scattering has more chances to occur especially at elevated nanoparticles loadings, as a result of good dispersion and the presence of large number of interfaces. However, it is worth noting that the improved compatibility and adhesion at nanoparticle-polymer interfaces reduces significantly the impact of scattering (W. Cui et al., 2011; Heid et al., 2015; Heid, Fréchette, et al., 2016; Huang et al., 2012).



Figure 4-16: Thermal conductivity of SEBS-ZnO vs. SEBS-MA-ZnO nanocomposites

4.4 Concluding remarks

In this study, the dispersion of ZnO nanoparticles within SEBS block copolymer was tailored through the use of MA graft attached to the rubber block. Although the nanoparticles were of spherical shape, rheological percolation was observed at low ZnO content between 5 wt% and 10 wt%. This network formed between the nanoparticles and polymer chains was accompanied by an improvement of thermal conductivity and excellent resistance to surface erosion, at lower ZnO contents not exceeding 5 wt%, compared to non-maleated SEBS-ZnO samples where the dispersion was not tailored. Simultaneously, a better control of the electrical conductivity as function of the electric field and nanoparticles loading was enabled, which might reduce efficiently the space charge accumulation either by increasing trapping sites density or improving the non-linearity feature of the conductivity. Besides, the dielectric losses and the breakdown strength of these samples with tuned dispersion were in similar range compared to unfilled SEBS and SEBS-MA at power frequencies and operational temperatures. Furthermore, moisture absorption was found to have a negligible effect on the dielectric response of these nanocomposites. This combination of properties makes SEBS-MA-ZnO nanocomposites a promising candidate for HV insulation systems. These features were enabled at relatively low contents of nanoparticles by using a block copolymer matrix as a nanostructured template and MA as a functional graft, leading to rheological percolation. The correlation between rheological properties and electrical, thermal and dielectric properties brings the attention to viscoelastic properties as an efficient tool to evaluate the control of dispersion and predict the dielectric performance. Interestingly, this model might be applied to other types of nanoparticles with desirable properties such as magnesium oxide (MgO) for instance, which is reported to reduce significantly space charge accumulation. Application wise, it might be interesting to investigate the resulting thermoplastic elastomer nanocomposites as master batches to blend with polyolefins commonly used in HV insulation systems such as polyethylene and polypropylene, in order to improve their performance.

Associated content

Supporting electronic information (ESI) file: Additional TEM micrographs, dielectric spectroscopy data and mappings of eroded area related to the studied nanocomposites are available online.

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CHAPTER 5

ARTICLE 4: POLYETHYLENE/THERMOPLASTIC ELASTOMER/ZINC OXIDE NANOCOMPOSITES FOR HIGH VOLTAGE INSULATION APPLICATIONS: DIELECTRIC, MECHANICAL AND RHEOLOGICAL BEHAVIOR

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Abstract

Blends of polyethylene (PE) with nanocomposites of polystyrene-b-poly(ethylene-cobutylene)-b-polystyrene grafted maleic anhydride (SEBS-MA) thermoplastic elastomer filled with Zinc Oxide (ZnO) nanoparticles have been studied as potential candidates for applications in HV insulation systems including HVDC cables. In particular, the dielectric and mechanical properties of PE/SEBS-MA/ZnO blend nanocomposites have been evaluated and compared to those of PE/ZnO homopolymer nanocomposites prepared as a reference. PE/ZnO materials were characterized by homogeneous distribution of nanoparticles and presence of agglomerations attributed to insufficient compatibility between the metal oxide nanoparticles and the polyolefin matrix. However, nanoscale dispersion was achieved in SEBS-MA/ZnO and PE/SEBS-MA/ZnO nanocomposites due to improved compatibility between the nanoparticles and SEBS-MA. Besides, in PE/SEBS-MA/ZnO blend nanocomposites, ZnO nanoparticles remained exclusively confined in SEBS-MA or at the interface between PE and SEBS-MA. In terms of dielectric properties, the unfilled blend PE/SEBS-MA featured reduced breakdown strength and resistance to surface erosion by partial discharges in comparison with neat PE. However, upon addition of ZnO, PE/SEBS-MA/ZnO presented higher performance when compared to PE/ZnO nanocomposites. At 1

wt% ZnO loading, the resistance to surface erosion of PE/SEBS-MA/ZnO increased by 45% higher than neat PE/SEBS-MA, 38% higher than unfilled PE and 30% higher than PE/ZnO nanocomposite containing the same ZnO loading. Besides, PE/SEBS-MA/ZnO blend nanocomposites exhibited dielectric losses lower than PE/ZnO nanocomposites at power frequencies and temperatures up to 80 °C. The breakdown strength of both sets of nanocomposites decreased compared to unfilled materials, at large loadings of nanoparticles. However, smaller reduction was observed in the case of PE/SEBS-MA/ZnO nanocomposites due to improved nanoparticles dispersion. Finally, PE/SEBS-MA/ZnO nanocomposites featured enhanced mechanical flexibility when compared to PE/ZnO nanocomposites.

Keywords: thermoplastic elastomer, polymer blend nanocomposite, nanodielectrics, selective nanoparticles dispersion, HV insulation systems...

5.1 Introduction

Thermoplastic materials such as Polyethylene (PE) and Polypropylene (PP) are common materials used within high voltage insulation systems. Among them, low density polyethylene (LDPE) has been extensively used in cable insulation due to its excellent dielectric properties and attractive processability. However, LDPE has relatively poor thermal and mechanical stability at high temperatures which could be improved by crosslinking, for instance. Therefore, currently LDPE is mainly used in its cross-linked form (XLPE), which is unfortunately unrecyclable (Arora & Mosch, 2011) and vulnerable to space charge accumulation due to the presence of by-products from the crosslinking process (Fu et al., 2007).

High density polyethylene (HDPE) and especially PP feature better thermal and mechanical stability due to their higher melting points and might constitute potential candidates for recyclable high voltage (HV) insulation materials. But, they suffer from increased brittleness at low temperatures (Arora & Mosch, 2011). To adapt better the properties of HDPE and PP to the purpose of recyclable HV insulation, blending was investigated as one of the solutions. In particular, blends of HDPE with LDPE and/or thermoplastic elastomers were considered

(Helal et al., 2016; Helal et al., 2014; I. Hosier et al., 2007; I. L. Hosier et al., 2010; B Zazoum, David, & Ngô, 2013). Recently, blends of PP and thermoplastic elastomers have been investigated as well (Du et al., 2016; Hamzah, Jaafar, Jamil, & Kamarol, 2014; Y. Zhou et al., 2015). The blends have shown mainly an interesting improvement of the mechanical flexibility at the cost of a reduction in breakdown strength and resistance to surface erosion by partial discharges as well as increased space charge accumulation (Helal et al., 2016; Helal et al., 2014; B Zazoum et al., 2013; Y. Zhou et al., 2015). Addition of nanoparticles to these blends might be a powerful tool to compensate these drawbacks, in a similar trend to what has been reported in the case of PE based nanocomposites. In fact, it has been well established that the addition of nanoparticles, including metallic oxide nanoparticles such zinc oxide (ZnO) and magnesium oxide (MgO), to homopolymer induces a reduction in space charge accumulation as well as an increase of breakdown strength and resistance to surface erosion (David & Fréchette, 2013; Fabiani et al., 2016; Fleming et al., 2008; Michel F Fréchette et al., 2010; M. Guo, Fréchette, David, & Demarquette, 2016; Hayase et al., 2006; Amir M Pourrahimi et al., 2016; Amir Masoud Pourrahimi et al., 2016; Smith et al., 2008; Tian et al., 2011, 2012; Tian et al., 2015). In the case of HDPE and PP based blends, few studies have been reported regarding the effect of nanoparticles (Du et al., 2016; Hamzah et al., 2014; Helal et al., 2016; B Zazoum et al., 2013). Among them, the recent study published by Du et al. (Du et al., 2016), reporting that the addition of ZnO nanoparticles to PP/Polyolefin elastomer (POE) blends results in suppression of space charge accumulation while maintaining an excellent flexibility ensured by the POE phase. However, there is still work to be done in this area, especially in the case of HDPE. Therefore, in this study, a blend of HDPE with polystyrene-b-poly(ethylene-co-butylene)-b-polystyrene grafted maleic anhydride (SEBS-MA) thermoplastic elastomer containing ZnO nanoparticles will be investigated. In fact, SEBS and SEBS-MA are often added in minor fractions to polymer matrices either as compatibilizers to improve the dispersion of nanoparticles including nanoclays, metal oxides and carbon nanotubes (Liang & Tjong, 2006), when a polyolefin matrix is used or as water treeing retardant agent in polymer insulating materials (Liu et al., 2011; Z. Ma et al., 2010). Besides, recently few studies have started exploring blends of polyolefins with SEBS or SEBS-MA, where the thermoplastic elastomer is acting as a structural phase of the blend

affecting its morphology and the spatial distribution of nanoparticles in it (Emna Helal et al., 2015; Helal et al., 2016; Sarkar & Alexandridis, 2015).

In this work, SEBS-MA/ZnO nanocomposites with excellent nanoparticles dispersion were initially fabricated by solvent blending following a method reported in a previous study (Emna Helal et al., 2015) then blended with HDPE using a melt compounding process (Helal et al., 2016). The morphology of the as-prepared PE/SEBS-MA/ZnO blend nanocomposites was studied through microscopy and characterization of the linear viscoelastic properties. Besides, measurements of complex dielectric permittivity, AC short-term breakdown strength and resistance to surface erosion by partial discharges were performed to evaluate the blend nanocomposites dielectric behavior, in comparison with reference PE/ZnO nanocomposites prepared by melt compounding. The mechanical properties of the nanocomposites have been also investigated, viz. the tensile strength and the elongation at break.

5.2 Experimental

5.2.1 Materials

HDPE was purchased from Chevron Phillips in pellets form (grade HXM50100). Its density is equal to 0.948 g/cm³ and its melt flow index is equal to 10 g/10 min at 190 °C and under a load of 21.6 Kg. The grade FG1901 of SEBS block copolymer, which is grafted with maleic anhydride (MA), was supplied from Kraton in pellets form as well. Its density is equal to 0.915 g/cm³. It is composed of two polystyrene (PS) endblocks in the form of nanometric cylinders dispersed within the poly(ethylene-*co*-butylene) (PEB) elastomer midblock. The PS block fraction is equal to 30wt% while the MA content is equal to 1-2 wt% and is attached to the PEB block. The ZnO nanoparticles were purchased from Sigma Aldrich in the form of 40wt% colloidal suspension in ethanol. They are modified with an alkyl ammonium salt and feature an average particle size around 35 nm. All the materials were used as received.

5.2.2 Processing

In a first step, SEBS-MA/ZnO nanocomposites were prepared by solvent casting method. In a second step, SEBS-MA/ZnO nanocomposites were blended with polyethylene by melt compounding to achieve PE/SEBS-MA/ZnO blend nanocomposites. Moreover, PE/ZnO nanocomposites were prepared by melt mixing using the same processing conditions, for the sake of comparison. Before further characterization, all the samples were hot pressed at 160 °C to obtain thin films. The detailed fabrication procedure is published in previous studies (Emna Helal et al., 2015; Helal et al., 2016). The list and composition of the obtained films are summarized in Table 5-1. Note that mass fractions of PE and SEBS-MA in the obtained blends are equal.

	PE (wt%)	SEBS-MA (wt%)	ZnO (wt%)
PE	100	0	0
PE/1	99	0	1
PE/5	95	0	5
PE/10	90	0	10
PE/SEBS-MA	50	50	0
PE/SEBS-MA/1	49.5	49.5	1
PE/SEBS-MA/5	47.5	47.5	5
PE/SEBS-MA/10	45	45	10

Table 5-1: Nomenclature and composition of PE/ZnO and PE/SEBS-MA/ZnO nanocomposites

5.2.3 Characterization

Cross-sections of different nanocomposites were observed to characterize the dispersion of nanoparticles either by Scanning Electron Microscopy (SEM) or High Resolution Scanning Electron Microscopy (HRSEM). Hitachi S-3600N microscope was used for SEM images

while Hitachi SU-8230 microscope was used for HRSEM images. A Leica microtome equipped with a tungsten knife was used to fracture the samples in liquid nitrogen at -100 °C. In order to characterize the morphology of the prepared blends, some samples were immersed overnight in a large volume of toluene and stirred gently at room temperature, after fracturing, to extract the SEBS-MA component. Prior to observation, the cross-sections were coated with a 20 nm thick layer of gold to prevent charge accumulation.

Rheological measurements were performed at 160 °C to characterize the linear viscoelastic properties of the nanocomposites in the molten state. In particular, small amplitude oscillatory shear (SAOS) tests were performed by means of Anton Paar MCR 501 strain-controlled rheometer equipped with 25 mm-diameter parallel plate geometry. The strain value was fixed to 0.5% while the angular frequency range swept from 0.01 to 300 rad.s⁻¹.

The complex dielectric permittivity of the materials was characterized at room temperature by means of a Novocontrol broadband spectrometer. In particular, 20-mm-diameter disks were prepared from the samples and placed between parallel plated brass electrodes to form a plane-plane capacitor. Then, measurements were performed in the frequency range from 10^{-2} Hz up to 10^{5} Hz. Besides, a temperature range between 25 °C and 100 °C was considered for selected samples.

AC short-term breakdown strength tests were performed at ambient temperature using a Bauer DTA100 equipped with a pair of 4 mm-diameter ball-type electrodes. A constant AC voltage ramp of 2 kV.s⁻¹ was progressively applied at a frequency equal to 60 Hz, until breakdown occurred. During the measurement, the samples were fixed between the two electrodes and immersed in Luminol TRi insulating fluid. 12 specimens of each material, of average thickness around 250 μ m, were considered for a statistical analysis by means of two-parameter Weibull distribution. However, the samples thickness was not strictly uniform. Thus, to reduce the errors related to thickness, all the measured breakdown strength fields were normalized to the average thickness of 250 μ m using a power law equation reported in

reference (Takala et al., 2010). The normalized breakdown strengths were used for the statistical analysis.

To evaluate the resistance of the fabricated materials to surface erosion, samples of average thickness equal to 500 μ m were subjected to prolonged AC stress under electrical discharge conditions, using a point-to-plane geometry set-up operated in open air. The samples were fixed to the ground plane electrode using Kapton tape and separated from the tip of the rod electrode by an air gap of 200 μ m. The radius of the rod electrode is equal to 1 mm. During the test, a sinusoidal AC voltage (7 kV_{RMS}, 300 Hz) was applied to the rod electrode while the plane electrode was connected to ground. The erosion time was fixed to 40 hours which is equivalent to 200 hours at 60 Hz. Then, the eroded volume of each sample was evaluated by means of a mechanical profilometer (Dektak 150).

More details regarding the procedures used for dielectric spectroscopy measurements, AC short-breakdown tests and resistance to surface erosion tests are available in previous studies (Emna Helal et al., 2015; Helal et al., 2016).

The mechanical properties of all the nanocomposites were measured using STM Alliance machine equipped with 1 kN load cell and operated in the tensile mode according to ASTM D638-14 (*Standard Test Method for Tensile Properties of Plastics*, 2014). The stretching rate was selected equal to 10mm/min. Dog-bone shaped specimens were used with a gauge length of 9.53 mm, a width of narrow section equal to 3.15 mm and a thickness of 3.15 mm. 5 specimens were tested for each material. All the specimen were prepared by injection molding using a melt temperature equal to 200 °C, a mold temperature equal to 60 °C and an injection pressure equal to 50 MPa.

170

5.3 Results and discussion

5.3.1 Scanning electron microscopy (SEM)

The dispersion of ZnO nanoparticles in the fabricated nanocomposites was first characterized by SEM at low magnifications. Samples not subjected to solvent extraction were considered at this step. Micrographs corresponding to untreated PE/ZnO nanocomposites containing different concentrations of nanoparticles are presented in Figure 5-1. It can be seen that at increasing content of nanoparticles, microscale agglomerations of increasing size are observed (as indicated by the arrows in Figure 5-1(c) and Figure 5-1(d)), which indicates poor dispersion of nanoparticles in polyethylene matrix. In micrographs corresponding to untreated PE/SEBS-MA/ZnO reported previously in (Helal et al., 2016), no agglomerations were observed. Besides, chemical analysis by energy dispersive X-ray spectroscopy (EDX), reported in the same study (Helal et al., 2016), revealed homogeneous distribution of Zinc element all over the sample.



Figure 5-1: SEM micrographs of PE/ZnO nanocomposites: (a) unfilled PE, (b) PE/1, (c) PE/5 and (d) PE/10

To study the morphology of PE/SEBS-MA/ZnO blend nanocomposites, samples treated by solvent extraction to dissolve SEBS-MA were considered. Micrographs corresponding to these samples are reported in Figure 5-2. Figure 5-2(a), corresponding to the unfilled blend, indicates the formation of a co-continuous morphology, characterized by a sponge shape. Interestingly, the incorporation of ZnO nanoparticles in the blend results in a considerable change of morphology (Figure 5-2(b)-(d)), even at nanoparticles concentration as low as 1wt%. More specifically, the co-continuous morphology of the unfilled blend disappeared giving place to more droplet dispersion type morphology where SEBS-MA phase is in the form of elongated droplets. This switch of morphology is the result of the increase of SEBS-MA viscosity during the melt processing, upon the addition of ZnO nanoparticles, which leads to breaking the threads. The viscosity increase hypothesis will be discussed in a following section of the manuscript, in correlation with SAOS measurements performed on PE/SEBS-MA/ZnO blends and reported in Figure 5-4 as well as SAOS measurements

corresponding to SEBS-MA/ZnO master batches and reported previously in (Helal, David, Fréchette, & Demarquette, under review; Emna Helal et al., 2015).

Apart from the switch of morphology, SEM micrographs reported in Figure 5-2 doesn't indicate any presence of agglomerations. This observation confirms that the dispersion of nanoparticles was considerably improved. However, the location of nanoparticles cannot be identified at this step. To determine their location, two methods were adopted: an analytical method based on the wetting coefficient theory as well as an experimental method based on HRSEM.



Figure 5-2: SEM micrographs of samples treated by solvent extraction: (a) unfilled PE/SEBS-MA, (b) PE/SEBS-MA/1, (c) PE/SEBS-MA/5 and (d) PE/SEBS-MA/10

PE/SEBS-MA/5 nanocomposite was considered for HRSEM. Two specimens were investigated: one treated with toluene to dissolve the SEBS-MA phase and one untreated where both phases are present. The HRSEM micrographs are reported in Figure 5-3. In particular, micrographs corresponding to the untreated specimen and the treated specimen are reported respectively in Figure 5-3(a) and Figure 5-3(b). Besides, Figure 5-3(c) and Figure 5-3(d) correspond to two zooms on respectively a PE region and an SEBS-MA region, selected from Figure 5-3(b). From the SEM image corresponding to the untreated sample (Figure 5-3(a)), it is difficult to accurately distinguish the two phases of the blend, i.e. PE and SEBS-MA, due to insufficient contrast between the two polymers (which could be due to the fact that the examined section was not thin enough). However, a nanoscale dispersion and inhomogeneous distribution of the nanoparticles is observed (regions surrounded by yellow dotted lines in Figure 5-3(a) correspond to regions containing nanoparticles). The examination of the sample treated with toluene at low magnification shows the presence of islands which correspond normally to the extracted SEBS-MA domains. A zoom inside one of these domains (region surrounded by red dotted line in Figure 5-3(b)) shows multiple traces of nanoparticles (Figure 5-3(d)). However, no nanoparticles were observed outside the SEBS-MA region (region surrounded by blue continuous line in Figure 5-3(b) and presented in Figure 5-3(c)). Hence, it can be concluded that the nanoparticles are enclosed in SEBS-MA phase and at the interface PE/SEBS-MA, due to the good affinity between the MA graft and the functional groups on the surface of ZnO nanoparticles.



Figure 5-3: HRSEM micrographs of PE/SEBS-MA/5 nanocomposite: (a) SEBS-MA not extracted, (b) SEBS-MA extracted, (c) Zoom inside PE region surrounded by blue continuous line in (b): poor ZnO phase, (d) Zoom inside SEBS-MA region surrounded by red dotted line in (b): poor ZnO phase, (d) Zoom inside SEBS-MA region surrounded by red dotted line in (b): rich ZnO phase (Colors are available in the online version)

5.3.2 Thermodynamic prediction of ZnO nanoparticles localization: Wetting coefficient

In order to support the conclusions made from HRSEM analyses regarding the localization of ZnO nanoparticles, a thermodynamic prediction was performed to estimate the ability of the nanoparticles to be wetted by the two components of the blend: PE and SEBS-MA. In fact, at thermodynamic equilibrium, nanoparticles in a polymer blend tend to locate in a position that minimizes the interfacial energy. This equilibrium position might be predicted by the

evaluation of the wetting coefficient W_a which is estimated according to equation (5.1), adapted from Young's equation (Tadros, 2015; Trifkovic, Hedegaard, Sheikhzadeh, Huang, & Macosko, 2015; X. Zhao et al., 2013). In the case of the studied system, W_a is equivalent to the contact angle of the nanoparticle at the interface between PE and SEBS-MA (Vandebril, Vermant, & Moldenaers, 2010).

$$W_a = \frac{\gamma_{ZnO/PE} - \gamma_{ZnO/SEBS-MA}}{\gamma_{PE/SEBS-MA}}$$
(5.1)

Where:

Wa: wetting coefficient;

 $\gamma_{1/2}$: interfacial tension between components 1 and 2.

 W_a values above 1 or below -1 mean that ZnO nanoparticles would preferentially locate in SEBS-MA or in PE, respectively. If the value of W_a is between -1 and 1, the difference between ZnO/PE and ZnO/SEBS-MA interfacial tensions is less than PE/SEBS-MA interfacial tension which induces the nanoparticles to localize at the interface in order to minimize the interfacial energy.

The interfacial tension between two components could be determined experimentally or theoretically. The experimental method might be difficult and not accurate especially in the case of nanoparticle-polymer interfaces (Trifkovic et al., 2015). Therefore, to avoid this complication, the values of the interfacial tension used in this manuscript were derived according to Owens-Wendt mean equation (equation (5.2)), knowing the surface tension of each component (Owens & Wendt, 1969). The values of surface tension corresponding to PE, SEBS-MA and ZnO nanoparticles at the processing temperature 160 °C have been extrapolated from values reported in literature (Demarquette, Da Silva, Brandi, & Gouvêa, 2000; Moreira & Demarquette, 2001; "Solid surface energy data (SFE) for common polymers," ; Švab, Musil, Šmit, & Makarovič, 2007; Torchinsky & Rosenman, 2009; Trifkovic et al., 2015; Wilkinson, Clemens, & Harding, 2004) and summarized in Table 5-2.

$$\gamma_{1/2} = \gamma_1 + \gamma_2 - 2(\sqrt{\gamma_1^d \gamma_2^d} + \sqrt{\gamma_1^p \gamma_2^p})$$
(5.2)

Where:

 $\gamma_{1/2}$: interfacial tension between component 1 and 2;

 γ_1 : surface tension of component 1;

 γ^{p} : polar component;

 γ^{d} : dispersive component.

Table 5-2: Total surface tension and Dispersive and Polar components for polymers and ZnO nanoparticles estimated at 160 °C

	-dγ/dT (mJ.m ⁻² .K ⁻¹)	γ (mJ.m ⁻²)	γ^{d} (mJ.m ⁻²)	γ ^p (mJ.m ⁻²)
PE (Demarquette et al., 2000; Moreira & Demarquette, 2001; "Solid surface energy data (SFE) for common polymers," ; Trifkovic et al., 2015)	0.057	27.72	27.72	0
SEBS-MA (Švab et al., 2007; Wilkinson et al., 2004)	0.045	30	21.6	8.4
ZnO (Torchinsky & Rosenman, 2009)	-	34	25	9

The calculated values of interfacial tension between each two components and the estimated wetting coefficient are reported in Table 5-3. The estimated W_a indicates that the nanoparticles should be mainly localized in SEBS-MA phase and probably at the interface PE/SEBS-MA. This result is expected due to the interaction between the nanoparticles surface and the MA graft. It is also in line with the results of microscopy reported earlier. As a consequence, the theoretical method represents a straightforward tool to predict the localization of nanoparticles. However, it is worth mentioning that this theoretical prediction has several limitations. The major one is the difficulty to get accurate values of surface tension from the literature in the case of SEBS-MA block copolymer and ZnO nanoparticles,

which takes into account the exact physical properties and applied chemical modification, for instance. Also, in this study, the value of the temperature coefficient, $d\delta/dT$, of SEBS-MA was assumed equal to that of SEBS. Besides, the values of surface tension calculated for ZnO nanoparticles in reference (Torchinsky & Rosenman, 2009) and used in this study, correspond to nanoparticles treated to decrease their surface tension and improve their compatibility with polymers. Unfortunately, their chemical modification differs from that applied to the nanoparticles used in this work. But, these values remain reliable since they are also close to those reported for several organically modified clays (Trifkovic et al., 2015) functionalized in a similar fashion to the nanoparticles used in this study, which are modified by an alkyl ammonium salt. In addition to errors related to surface tension values, in the general case, nanoparticles localization in a polymer blend is not only influenced by thermodynamic considerations, but also by kinetic effects. In fact, interplay between nanoparticle compatibility with each polymer phase, processing sequence and rheology is often involved (Trifkovic et al., 2015).

	Estimated values
$\gamma_{ZnO/PE} (mJ.m^{-2})$	9.07
γzno/sebs-ma (mJ.m ⁻²)	0.13
γpe/sebs-ma (mJ.m ⁻²)	8.78
Wa	1.01

Table 5-3: Estimated interfacial tension and wetting coefficient values

5.3.3 Rheological properties

The morphology of PE/SEBS-MA blends and the improved quality of dispersion in PE/SEBS-MA/ZnO nanocomposites were studied by microscopy, as discussed earlier. However, this technique only gives insights at a local micro and nanoscale. A useful technique that might give information about the dispersion and morphology at a larger macroscale is rheology. In fact, by performing small amplitude oscillatory shear (SAOS)

tests for example, the variation of the viscoelastic properties of the polymer matrix in the molten state as function of the nanoparticles dispersion could be probed. Generally, many homopolymers such as polyethylene exhibit a terminal behavior distinguished by a high slope and drop of the elastic modulus, at low frequencies. However, in the presence of nanostructures such as nanoparticles or nanodomains of a block copolymer for example, this slope may decrease and the elastic modulus might increase due to confinement of polymer chains between the nanostructures. This effect is more obvious with increasing amount of nanoparticles.

In Figure 5-4, plots of storage modulus G' and complex viscosity modulus $|\eta^*|$ as function of angular frequency ω are reported, respectively in Figure 5-4(a) and Figure 5-4(b), for both PE/ZnO and PE/SEBS-MA/ZnO nanocomposites. The viscoelastic properties of unfilled SEBS-MA and SEBS-MA/ZnO master batch, used to prepare PE/SEBS-MA/5 nanocomposite and referred to as MB5, are also reported for the sake of comparison. In the case of PE/ZnO nanocomposites, it can be seen that G' and $|\eta^*|$ of PE remain roughly the same after the addition of the nanoparticles. This behavior is valid for ZnO concentrations up to 10wt% ZnO (only results corresponding to 5 wt% ZnO are reported in Figure 5-4) and indicates that the interaction between ZnO nanoparticles and PE chains is relatively weak. In the case of PE/SEBS-MA/ZnO materials, a different behavior is observed. First of all, it can be seen that G' of unfilled PE/SEBS-MA blend increased compared to neat PE especially at low frequencies. This behavior is partially attributed to the characteristic non-terminal behavior of SEBS-MA as a block copolymer (Carastan et al., 2008) and also to the deformation of SEBS-MA domains in the blend (Graebling, Muller, & Palierne, 1993). Considering PE/SEBS-MA/ZnO nanocomposites, a consistent increase of G' and $|n^*|$ compared to unfilled PE/SEBS-MA is exhibited in the whole frequency range and especially at low frequencies. Simultaneously, the slope of G' vs. ω plot decreases at low frequencies. The same behavior is also featured by MB5 compared to unfilled SEBS-MA, with a higher improvement rate. Moreover, a plateau in G' vs. ω plot is exhibited at low frequency indicating good nanoparticles dispersion and establishment of a network between ZnO nanoparticles and SEBS-MA block copolymer chains attributed to the interaction between

MA graft and functional groups on the surface of ZnO nanoparticles. More details regarding this viscoelastic behavior of SEBS-MA/ZnO nanocomposites can be found in prior studies (Helal et al., under review; Emna Helal et al., 2015). The resulting efficient confinement of the macromolecular chains by the nanoparticles is at the origin of the increased dynamic viscosity of SEBS-MA and subsequently the blend in the whole ω range studied. This increase justifies the switch of morphology of PE/SEBS-MA/ZnO nanocomposites compared to unfilled PE/SEBS-MA reported in Figure 5-2, if we consider that the shear viscosity during the melt processing is well approximated by the dynamic viscosity, in respect with Cox-Merz approximation, which is valid for different polymer systems at relatively low shear rates (Dealy & Wang, 2013). In fact, although the value of the shear rate during the processing was not estimated quantitatively in this study, due to the different parameters that are involved in its estimation (Bousmina, Ait-Kadi, & Faisant, 1999), it is not expected to exceed few hundreds of reciprocal seconds (Bousmina et al., 1999; Kohlgrüber, 2008), which is in the range of the studied angular frequencies of the SAOS tests.



Figure 5-4: SAOS measurements of PE/ZnO vs. PE/SEBS-MA/ZnO nanocomposites: (a) storage modulus G' and (b) complex viscosity modulus $|\eta^*|$ as function of angular frequency ω

5.3.4 Complex dielectric permittivity

Complex dielectric permittivity of the studied nanocomposites has been characterized as functions of frequency and temperature. Data corresponding to room temperature measurements are reported in Figure 5-5. In particular, plots of imaginary permittivity ε '' as function of frequency, corresponding to PE/ZnO and PE/SEBS-MA/ZnO nanocomposites are reported respectively in Figure 5-5(a) and Figure 5-5(b). Besides, plots of ε '' as function of frequency corresponding to SEBS-MA/ZnO nanocomposites, used as master batches to prepare respectively PE/SEBS-MA/5 and PE/SEBS-MA/10 nanocomposites, are reported in Figure 5-5(c) for the sake of comparison. The master batches are referred to as MB5 and MB10 respectively. More details regarding the dielectric spectroscopy results of SEBS-MA/ZnO nanocomposites are reported in (Helal et al., under review).

All PE/ZnO nanocomposites show large relaxation peaks in the vicinity of 10¹-10² Hz, which correspond most likely to Maxwell-Wagner-Sillars (MWS) interfacial process. At increasing ZnO concentration, the peak is shifted to lower frequencies. This behavior is in part consistent with the MWS theory suggesting that, if the conductivity of the matrix is small and can be neglected, which is the case for both PE and SEBS-MA, an increase in the volume fraction of the fillers induces a slight increase of the relaxation time and consequently a decrease of the peak maximum frequency (David & Fréchette, 2013; Kremer & Schönhals, 2012; Tsangaris, Kouloumbi, & Kyvelidis, 1996). However, this effect is too small to justify the observed experimental results. Instead, the formation of bigger agglomerates with lower effective conductivity, at increasing ZnO concentrations, is the likely explanation for this behavior. PE/SEBS-MA/ZnO nanocomposites exhibit an interfacial relaxation process as well, but less intense and at higher frequencies in the vicinity of 10²-10³ Hz. Similar to the behavior observed in PE/ZnO nanocomposites, the relaxation peak is shifted to lower frequencies at increasing ZnO content.

The difference in frequencies windows corresponding to the relaxation phenomena observed respectively in PE/ZnO and PE/SEBS-MA/ZnO might be correlated with the location of

nanoparticles in either PE or SEBS-MA, since the MA graft attached to SEBS interacts with the surface of ZnO nanoparticles, modifying the properties of the interface (Helal et al., under review). In fact, MB5 and MB10 nanocomposites used as master batches exhibit relaxation peaks at around 10³ Hz. Besides, the peak featured in PE/SEBS-MA/ZnO nanocomposites is more similar to SEBS-MA/ZnO nanocomposites in terms of dielectric strength and peak position indicating that ZnO nanoparticles are potentially located in SEBS-MA phase. This observation supports the conclusions made earlier from the microscopy and thermodynamic study, which suggest that the nanoparticles stay confined inside SEBS-MA phase and probably at SEBS-MA/PE interfaces.

As far as the insulation application is concerned, PE/SEBS-MA/ZnO nanocomposites feature the lowest losses at room temperature, as compared to both PE/ZnO nanocomposites and SEBS-MA/ZnO master batches as shown in Figure 5-5(c), especially at power frequencies. Moreover, the real permittivities of the nanocomposites remain low and close to those of unfilled PE and PE/SEBS-MA, all values in the range 2.2-2.4 (not reported in this manuscript).



Figure 5-5: Plots of imaginary permittivity ε'' as function of frequency measured at room temperature corresponding to: (a) PE/ZnO nanocomposites, (b) PE/SEBS-MA/ZnO nanocomposites and (c) comparison with SEBS-MA/ZnO nanocomposites

To investigate further the observed interfacial relaxation process and the effect of absorbed humidity on it, isothermal frequency sweeps of the complex dielectric permittivities of both PE/5 and PE/SEBS-MA/5 nanocomposites have been performed in the temperature range from 25 °C to 100 °C, before drying and immediately after drying them for 5 days in a vacuum oven at 65 °C. The results are reported in Figure 5-6. The comparison of the spectra recorded before (Figure 5-6(a)-(b)) and after drying (Figure 5-6(c)-(d)) reveals that there is a little effect of drying, which indicates that the quantity of absorbed humidity is negligible. This behavior is expected and has been already reported several times in the literature, when nanoparticles with adequate surface modification are used (David et al., 2015; Fleming et al., 2008; Lau et al., 2013).

Apart from the drying aspect, a drastic change in the dielectric behavior was observed between PE/ZnO and PE/SEBS/ZnO nanocomposites. For PE/5 nanocomposite, the relaxation peak was found to shift towards higher frequencies at increasing temperatures, which is the usual trend for nanocomposites for which the conductivity of the inclusion increases with temperature. This behavior is more obvious in the case of the dried sample since the release of absorbed water as the temperature increases in the case of the undried samples slightly counterbalance the increase of the inclusions conductivity. The PE/SEBS-MA/5 nanocomposite (Figure 5-6(d)) exhibits a considerably reduced dielectric strength and broadness compared to PE/5 nanocomposite (Figure 5-6(c)) in the studied temperature range. This behavior is in line with the absence of agglomerations in PE/SEBS-MA/ZnO nanocomposites and the improved interaction with the nanoparticles which modifies the interphase properties. Similar results have been reported when improved compatibility between the nanoparticles and the polymer matrix is ensured (David & Fréchette, 2013; Smith et al., 2008; Bouchaib Zazoum, Eric David, & Anh Dung Ngô, 2014; Bouchaib Zazoum et al., 2016). However, it could be seen that at elevated temperatures starting from 50 °C, a dominant contribution from charge fluctuations to the dielectric loss spectra, is observed for the PE/SEBS-MA/5 nanocomposite, especially for frequencies below 10 Hz, while being completely absent in the case of PE/5 nanocomposite. At 90 °C, the dielectric loss of PE/SEBS-MA/5 nanocomposite becomes even slightly higher than PE/5 nanocomposite at power frequencies.

Electrode polarization might be responsible of part of this increase. However, a considerable improvement of the nanocomposites conductivity by few orders of magnitude is the likely explanation. In fact, this behavior has been also observed in the dielectric spectra of SEBS-MA/ZnO nanocomposites, as reported in (Helal et al., under review), and was attributed to the interaction of MA with functional groups on the surface of the nanoparticles that leads to an improved dispersion and the formation of a network between ZnO nanoparticles and the block copolymer chains. As a consequence, SEBS-MA/ZnO nanocomposites become slightly conductive at favorable heating conditions. Moreover, the increase of conductivity reaches a

threshold at about 5 wt% ZnO. This effect seems to be ultimately transferred to the PE/SEBS-MA blends leading to the observed behavior.



Figure 5-6: Isothermal frequency sweeps of the imaginary permittivity ε" of PE/5 and PE/SEBS-MA/5 nanocomposites at different temperatures between 25 °C and 100 °C, performed before and after drying: (a) undried PE/5, (b) undried PE/SEBS-MA/5, (c) PE/5 dried and (d) PE/SEBS-MA/5 dried for 5 days at 65 °C.

5.3.5 AC short-term breakdown strength

After normalization to the thickness of 250 μ m, the retrieved data corresponding to the AC short term breakdown tests were analyzed by means of a two-parameter Weibull distribution

to evaluate the breakdown strength of the studied nanocomposites. Commercial software was used for the analysis.

The cumulative Weibull distribution function of breakdown strength is given by equation (5.3). More details regarding the use of Weibull distribution for dielectric strength estimation could be found in IEEE standard 930 ("IEEE Guide for the Statistical Analysis of Electrical Insulation Breakdown Data," 2005).

$$P = 1 - \exp\left[-\frac{E}{E_0}\right]^{\beta}$$
(5.3)

Where:

P: is the cumulative probability of failure at an electric field less or equal to E;

E₀: is the scale parameter corresponding to the breakdown strength for which the cumulative failure probability is equal to 63.2%;

 β : is the shape parameter which measures the scatter of the data. The larger β is the narrower is the range for breakdown strength.

Weibull plots as well as scale and shape parameters values corresponding to PE/ZnO and PE/SEBS-MA/ZnO nanocomposites are reported respectively in Figure 5-7(a) and Figure 5-7(b). In particular, the value of the breakdown strength of neat PE is equal to 162 kV/mm. Upon addition of nanoparticles up to 5 wt%, the breakdown strength decreased by about 10%. At 10 wt % ZnO loading, a considerable decrease of breakdown by 34% was induced. This behavior might be explained by the presence of agglomerations, especially at large concentrations of nanoparticles, which result in local voids and defects where the electric field is enhanced. This negative effect is dominating the potential positive effect consisting of increasing charge traps density and more efficient electron scattering (Cheng et al., 2013; Tian et al., 2012). The unfilled PE/SEBS-MA blend features a reduction of breakdown strength compared to neat PE by 10% as well, which is in line with its reduced mechanical strength (to be discussed later) and the presence of PE/SEBS-MA interfaces. Upon the addition of ZnO nanoparticles to the blend up to 5wt%, the breakdown strength of the blend was maintained or slightly reduced by 5%. This behavior might be correlated with the
improved compatibility and dispersion of nanoparticles inside SEBS-MA phase, which ensure a good balance between electron scattering and local electric field enhancements (Cheng et al., 2013; Smith et al., 2008; Tian et al., 2011, 2012). At this level of ZnO loadings, dielectric breakdown strength values of PE/ZnO and PE/SEBS-MA/ZnO nanocomposites are comparable and the initial drop induced by the thermoplastic elastomer phase was compensated by the improved role of ZnO dispersion. At 10wt% ZnO, the dielectric strength of PE/SEBS-MA/10 nanocomposite exhibits a drop by 15% compared to the unfilled blend but remains higher than PE/10 nanocomposite: 123 vs.107 kV/mm. Overall in both sets of nanocomposites, it seems that 5wt% ZnO is an optimum loading at which the breakdown of the nanocomposites reaches its maximum.



Figure 5-7: Weibull probability plots of: (a) PE/ZnO nanocomposites and (b) PE/SEBS-MA/ZnO nanocomposites

5.3.6 Resistance to surface erosion by partial discharges

The resistance to surface erosion by partial discharges is an important property to evaluate the long-term dielectric performance of insulating materials under operational conditions, especially high electrical stresses. In general, a reduction in the eroded volume caused by corona discharges is equivalent to an improved resistance. In Figure 5-8, eroded volumes corresponding to PE/ZnO and PE/SEBS-MA/ZnO nanocomposites, subjected to a voltage equal to 7 kV_{RMS} for 40 hours at 300 Hz, are reported. The relative evolution of the eroded volume corresponding to each nanocomposite is calculated compared to the unfilled reference, either unfilled PE or unfilled PE/SEBS-MA.

Comparing unfilled PE and PE/SEBS-MA, it could be seen that the blend exhibits again lower performance, with an increase of the eroded volume by approximatively 13% compared to neat PE. Upon the addition of ZnO nanoparticles, both sets of nanocomposites exhibit improved resistance. In a general trend, the eroded volumes decrease with increasing nanoparticles content. This is expected due to higher resistance of inorganic species to partial discharges compared to organic compounds (Brockschmidt et al., 2011; S. Chen et al., 2010; Heid et al., 2015). Moreover, Zinc oxide is well known by its shielding effect to UV light emitted during the process of partial discharges (S. Chen et al., 2010).

Comparing both sets of nanocomposites, it can be observed that PE/SEBS-MA/ZnO nanocomposites feature the best resistance, despite the low performance of the unfilled blend. For example, the eroded volume of PE/SEBS-MA/1 is 45% less than unfilled PE/SEBS-MA, 38% less than unfilled PE and 30% less than PE/1 nanocomposite. At higher loading equal to 10wt%, the eroded volume of PE/SEBS-MA/10 is 82% less than PE/SEBS-MA, 79% less than PE and 24% less than PE/10. Since the comparison is made considering the same amount of nanoparticles, the observed difference between both sets of nanocomposites is most likely related to the dispersion and distribution of the nanoparticles. In fact, the improved nanoscale dispersion of ZnO nanoparticles in SEBS-MA results in reduced distortion of the electric field and highly improved resistance inside the SEBS-MA phase. Resistance to surface erosion measurements performed on SEBS-MA/ZnO samples and reported in (Helal et al., under review) support this conclusion. However, in PE/ZnO nanocomposites, the presence of agglomerations might lead to local field enhancements and severe erosion of the surrounding polymer (Heid et al., 2015; Heid, Fréchette, et al., 2016). In PE/SEBS-MA/ZnO blend nanocomposites, SEBS-MA/ZnO domains are distributed in form of droplets, as seen in Figure 5-2(b)-(d). These droplets featuring high resistance to erosion improve the performance of PE matrix and lead to an overall performance between that of PE/ZnO nanocomposites and that of SEBS-MA/ZnO nanocomposites used as master batch.

Hence, an improved control of the blend morphology in order to achieve co-continuity or more homogeneous distribution of SEBS-MA/ZnO islands might be beneficial to achieve higher performance. The shape of SEBS-MA/ZnO domains might be of importance as well. For instance, elongated domains could increase further the efficiency of SEBS-MA/ZnO phase in the blend.



Figure 5-8: Resistance to surface erosion by partial discharges for PE/ZnO vs. PE/SEBS-MA/ZnO nanocomposites

5.3.7 Mechanical properties

Mechanical properties of both PE/ZnO and PE/SEBS-MA/ZnO nanocomposites have been characterized in tensile mode. Typical stress-strain curves corresponding to different samples are reported in Figure 5-9 while estimated values of elastic modulus E, elongation at break EB, ultimate tensile strength UTS and elongation at ultimate tensile strength EUTS are reported respectively in Figure 5-10(a)-(d). The reported stress-strain curves show that PE/ZnO and PE/SEBS-MA/ZnO feature different behaviors in terms of stiffness and flexibility. In particular, unfilled PE and its corresponding PE/ZnO nanocomposites exhibit

high rigidity. However, they fracture shortly after necking at relatively low strains. These results are in agreement with other studies reporting similar values of elongation for HDPE based materials (Bhattacharyya, Chen, & Zhu, 2014; H. Ku, Wang, Pattarachaiyakoop, & Trada, 2011; Kumar et al., 2016; J.-H. Lee, Jung, Hong, Rhee, & Advani, 2005). It is worth noting that the absence of a large cold drawing region might be correlated with the used strain rate. In particular, reducing the strain rate may result in larger cold drawing regions and consequently larger values of elongation (Cai & Song, 2015). Unfilled PE/SEBS-MA blend exhibits a different behavior characterized by a reduced elastic modulus and an improved plasticity compared to PE, leading to a considerable improvement of toughness overall. Finally, PE/SEBS-MA/ZnO nanocomposites exhibit an intermediate behavior in terms of stiffness and elongation.



Figure 5-9: Typical stress-strain curves of selected PE/ZnO vs. PE/SEBS-MA/ZnO materials

Considering the values of elastic modulus E of the studied materials reported in Figure 5-10(a), it could be seen that indeed blending PE with SEBS-MA reduces largely E from 385 MPa to about 65 MPa. This behavior is expected due to the low stiffness of SEBS-MA. In PE/ZnO nanocomposites, the addition of increasing ZnO content up to 5wt% leads to an

initial small increase of E up to 400 MPa. However, at 10wt% ZnO, the modulus drops to about 370 MPa. The increase of stiffness with the addition of metal oxide nanoparticles is expected due to their high rigidity. However, at elevated concentrations, the formed agglomerations impede the load stress transfer from the polymer matrix to the nanoparticles leading to reduced strength (Pang et al., 2015). In PE/SEBS-MA based nanocomposites, the presence of ZnO nanoparticles results in considerable improvement of stiffness compared to the unfilled blend, up 95% at 5 wt% ZnO loading. This effect could be explained by the good dispersion of ZnO in SEBS-MA and the good adhesion at the interface which increase efficiently the stiffness of SEBS-MA phase and consequently the blend. This behavior compensates partially for the initial drop of stiffness caused by the thermoplastic elastomer. However, the stiffness of PE/SEBS-MA/ZnO nanocomposites remains in general considerably lower than PE/ZnO materials. UTS values corresponding to PE/SEBS-MA/ZnO are also lower than PE/ZnO materials, as shown in Figure 5-10(c). In both sets, the presence of ZnO nanoparticles induces small enhancements, mainly at 1wt% and 5wt% ZnO loadings. Overall, tensile strength values of PE/SEBS-MA/ZnO are in the range 20-25 MPa while those of PE/ZnO are in the range 40-45 MPa.

In terms of flexibility, EB and EUTS, shown respectively in Figure 5-10(b) and Figure 5-10(d), are largely improved in unfilled PE/SEBS-MA. Upon the addition of ZnO nanoparticles to PE/SEBS-MA, the flexibility is reduced, which is expected due to the rigid nature of the nanoparticles, but remains considerably high compared to PE/ZnO materials. In particular, PE/SEBS-MA/1 nanocomposite features a good balance of strength and ductility, ensuring an overall enhancement of toughness compared to PE.



Figure 5-10: Mechanical properties of PE/ZnO vs. PE/SEBS-MA/ZnO nanocomposites: (a) Elastic Young's modulus E, (b) Elongation at break EB, (c) Ultimate tensile strength UTS and (d) Elongation at ultimate tensile strength EUTS (error bars stand for standard deviation)

5.4 Concluding remarks

In this study, the dielectric and mechanical properties of PE/SEBS-MA/ZnO nanocomposites have been evaluated, as potential candidates for HV insulation systems, in comparison with conventional PE/ZnO nanocomposites prepared by melt compounding. A correlation with the morphology and nanoparticles dispersion has been established as well. The main conclusions are the following:

- The use of SEBS-MA/ZnO master batch and a short mixing time with PE allows maintaining a good dispersion of the nanoparticles and their selective localization

inside the SEBS-MA phase and potentially at the interfaces PE/SEBS-MA. The increase of SEBS-MA viscosity upon addition of ZnO nanoparticles results in a switch of morphology from co-continuity in the case of unfilled PE/SEBS-MA blend to elongated droplets, in the case of PE/SEBS-MA/ZnO nanocomposites.

- The improved dispersion of nanoparticles in PE/SEBS-MA/ZnO compared to PE/ZnO results in better control of dielectric loss increase and breakdown strength decrease especially at elevated ZnO concentrations. Besides, the selective dispersion of ZnO in SEBS-MA droplets results in more efficient resistance to surface erosion by partial discharges inside the droplets and ultimately in the whole blend, compared to PE/ZnO nanocomposites where agglomerations are formed. This effect is more obvious at low concentrations of ZnO.
- Due to the presence of the elastomer PEB block, the breaking elongation of unfilled PE/SEBS-MA is improved compared to PE while the tensile strength is reduced. The presence of ZnO nanoparticles results in a considerable decrease of the elongation while improving slightly the tensile strength. But overall, PE/SEBS-MA-ZnO nanocomposites show improved mechanical flexibility compared to PE/ZnO nanocomposites and higher toughness at low ZnO loading equal to 1wt%.

Due to these improved electrical properties and mechanical flexibility, PE/SEBS-MA/ZnO nanocomposites have good potential for applications in HV insulation systems, specifically in HVDC cables as a replacement to XLPE. Therefore, the investigation of different fractions of PE and SEBS-MA in the blend as well as ZnO contents lower than 1 wt%, are recommended to optimize further the initial properties of the unfilled blend and to ensure a good trade-off between mechanical and dielectric/electrical properties in PE/SEBS-MA/ZnO blend nanocomposites.

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CHAPTER 6

ARTICLE 5: TUNING THE MECHANICAL AND DIELECTRIC PROPERTIES OF CLAY-CONTAINING THERMOPLASTIC ELASTOMER NANOCOMPOSITES

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Abstract

In this study, the mechanical strength and the AC short term breakdown strength of polystyrene-*b*-poly(ethylene-*co*-butylene)-*b*-polystyrene (SEBS) thermoplastic elastomer clay-containing nanocomposites have been investigated as function of their morphologies. The SEBS/clay nanocomposites with tailored morphologies were prepared previously by different processing techniques. They featured different orientations of clay platelets as well as polystyrene (PS) block nanodomains, namely: isotropic, oriented and partially oriented morphologies. In unfilled SEBS matrices, the mechanical strength was mainly tuned by the orientation of PS block nanodomains. A good correlation between the dielectric breakdown strength and the mechanical stiffness was observed overall: the higher the mechanical strength was, the higher the breakdown strength was. In the nanocomposites, the orientation of clay platelets as well as the degree of order and the characteristic sizes of the block

copolymer domains were seen to affect strongly the breakdown strength behavior in addition to the mechanical strength. In particular, the partially oriented morphology achieved by film blowing extrusion exhibited the maximum increase of the breakdown strength by 25% with optimized mechanical stiffness evaluated between that of the oriented morphology as a lower limit and that of the isotropic morphology as an upper limit.

Keywords: thermoplastic elastomer nanocomposite, tailored morphology, mechanical stiffness, dielectric breakdown strength

6.1 Introduction

Block copolymers are a special class of materials composed of immiscible polymer blocks that are usually self-organized in well-ordered nanodomains. The morphologies adopted by these nanodomains depend on several parameters such as the affinity between the blocks, their molecular weights and their relative proportions (Helfand, 1975; Sarkar & Alexandridis, 2015). Besides, these morphologies can be easily tuned by different external forces that can be applied either during melt compounding or solvent casting fabrication processes (Leice G Amurin et al., 2016; Daniel, Hamley, & Mortensen, 2000; Liedel, Pester, Ruppel, Urban, & Böker, 2012; Mansky et al., 1998; Ruppel et al., 2013).

Due to these features, block copolymers have been investigated recently as templated multiphase matrices to tailor nanoparticles dispersion in polymer nanocomposites (Balazs et al., 2006; Bockstaller et al., 2005; Kao et al., 2013; J. Y. Lee, Park, Yang, Cho, & Kim, 2003; Park et al., 2003; Sarkar & Alexandridis, 2015). Currently, this approach is considered of high importance to develop techniques for designing an advanced generation of nanocomposites with prescribed morphologies and engineering properties.

In this context, some recent studies investigated different morphologies of polystyrene-*b*-poly(ethylene-*co*-butylene)-*b*-polystyrene (SEBS) triblock copolymers and their claycontaining nanocomposites (Carastan et al., 2013; Carastan et al., 2014; De Sousa Jr et al., 2014; Helal et al., 2017; E Helal et al., 2015). SEBS belongs to the class of styrenic thermoplastic elastomers and features in addition to its ordered nanostructure, excellent combination of mechanical properties (Balsamo et al., 2006; Holden et al., 2004), good resistance to water treeing (Liu et al., 2011; Z. Ma et al., 2010), good electromechanical coupling (B. Kim et al., 2011) and high compatibility with polyolefins (Agari et al., 1993). As a block copolymer, SEBS is composed of two polystyrene (PS) blocks and one poly(ethylene-*co*-butylene) (PEB) elastomer midblock. It was widely used as a compatibilizer (Liang & Tjong, 2006) and as a dielectric elastomer actuator (Kofod et al., 2011; Mc Carthy et al., 2009; Mi et al., 2014; Saleem et al., 2014; Stoyanov et al., 2011).

The studies performed on SEBS and SEBS/clay nanocomposites demonstrated that different orientations of PS ordered nanodomains and clay nanoplatelets can be achieved by appropriate processing techniques (Leice G Amurin et al., 2016; Carastan et al., 2013; Carastan et al., 2014; De Sousa Jr et al., 2014; Helal et al., 2017; E Helal et al., 2015). For instance, isotropic lamellar or cylindrical morphologies were obtained by solvent casting process when adequate thermal annealing procedures were applied and appropriate fractions of PS blocks were present (E Helal et al., 2015). Furthermore, oriented morphologies where PS nanodomains and clay nanoplatelets are simultaneously aligned in the same direction were obtained by extensional and shear forces applied during a sheet die extrusion process (Leice G Amurin et al., 2016; Carastan et al., 2013; Carastan et al., 2014). In addition, more complex morphologies where PS domains and clay platelets have more than one preferential direction of alignment were achieved by film blowing extrusion. More details regarding these morphologies are available in the cited publications (De Sousa Jr et al., 2014; Helal et al., 2017). The abovementioned changes in spatial distribution of PS domains and nanoclays were correlated with changes in the mobility of elastomer chains located in the interfacial region. An interfacial glass transition (Tgi), several tens of degrees higher than the glass transition of the bulk PEB block, was attributed to these interfacial chains. In particular, the lowest Tgi temperatures were attributed to nanocomposites with aligned morphologies indicating less mobility restriction in this configuration compared to the other morphologies (Helal et al., 2017).

198

The control of morphology ultimately resulted in changes in effective dielectric properties as well. This behavior is partially expected due to the observed changes in the interfacial region which usually governs the properties of nanocomposites. In particular, increase of the breakdown strength was observed when SEBS-30 matrix (30 wt% PS domains) was filled with clay content equal to 5 wt%. Moreover, the comparison between isotropic and oriented morphologies of this nanocomposite revealed that the maximum increase of breakdown strength was equal to 45% and corresponded to the nanocomposite featuring oriented morphology, with both clay platelets and PS domains simultaneously aligned perpendicularly to the applied electric field (E Helal et al., 2015). This effect is in agreement with findings from the literature reporting that the addition of nanoclay to polyolefin matrices, increases the dielectric breakdown strength and their alignment perpendicular to the applied electric field improves further the results by increasing the tortuosity of the paths taken by the charge carriers (David et al., 2013; Liao et al., 2013b; Liao et al., 2014; Tomer et al., 2011). However, the dielectric strength of oriented unfilled SEBS matrix was reduced, compared to isotropic unfilled SEBS, which unfortunately neutralizes the increase observed by the alignment of clay.

This behavior is most likely related to the reduction of mechanical stiffness upon the alignment of PS nanodomains perpendicular to the electric field direction. In support to this hypothesis, few studies reported that the breakdown strength of SEBS was reduced with decreasing mechanical stiffness (Kollosche & Kofod, 2010; Kollosche et al., 2009). In fact, the electrodes attached to the specimen's surface during the breakdown test exert a compressive force that might result in appreciable deformations at an applied voltage below the intrinsic breakdown voltage, especially in soft elastomeric materials, leading to reduced dielectric strength and failure. This phenomenon is well-known as electromechanical breakdown. Some empirical equations were already established to describe the correlation between mechanical stiffness and electromechanical breakdown strength in rubbery materials (Blythe & Bloor, 2005).

In the same research orientation, this work aims at evaluating and validating the correlation between the breakdown strength and the mechanical strength of clay-containing SEBS nanocomposites in dependence with their designed morphologies, mainly the spatial distribution of PS block nanodomains and clay nanoplatelets. To this end, two sets of SEBS block copolymers with different ratios of PS blocks: 13 wt% and 20 wt% as well as their nanocomposites were investigated. The morphologies of these nanocomposites were tuned by means of different processing techniques.

6.2 **Processing and characterization**

6.2.1 Materials

Two grades of symmetric SEBS triblock copolymers donated by Kraton were used: G1643and G1645. These grades contain respectively 13 wt% and 20 wt% of PS blocks and are referred to in the manuscript respectively as SEBS-13 and SEBS-20. PS block nanodomains are ordered in the form of hexagonally packed cylinders within the poly(ethylene-*co*-butylene) (PEB) elastomer block. Montmorillonite clay grade Cloisite 20A purchased from Southern Clay (Gonzales, USA) and modified with dimethyl di (hydrogenated tallow) quaternary ammonium salt, was used as nanofiller for the nanocomposites preparation. In the rest of the manuscript, the abbreviation 20A will be used to design clay Cloisite 20A. All the materials studied in this paper were used as received. More details regarding the physical properties of the polymers as well as the modification and size of clay particles were provided in the previous studies published by co-authors (Leice G Amurin et al., 2016; Carastan et al., 2013; Carastan et al., 2014; De Sousa Jr et al., 2015).

6.2.2 Processing

The materials investigated in this study were prepared by different processing techniques to achieve different orientations of PS nanodomains and clay platelets. Three selected techniques were used: solvent casting, sheet die extrusion and film blowing extrusion at two air inflation ratios R1 and R3. In all the nanocomposites, the content of nanoclay used was 5 wt%. An illustrative scheme of the last two techniques based on extrusion is presented in Figure 6-1. Besides, a coordinate system that will be used to refer to test directions and related discussions is also presented. More details regarding the processing parameters corresponding to each technique and the procedure to obtain nanocomposite films are available in references (Leice G Amurin et al., 2016; Carastan et al., 2014; De Sousa Jr et al., 2014; Helal et al., 2017).



Figure 6-1: Cartoons illustrating: sheet-die extrusion (a), film blowing extrusion using a tubular die with two different ratios: R1 (b) and R3 (c) (the coordinate system is used as a reference for the following characterizations and discussions)

6.2.3 Summary of nanocomposite morphologies

The obtained morphologies of the studied materials depend mainly on the external forces applied during the selected fabrication technique and also on the fraction of the PS block. In a summary, nanocomposites with isotropic morphologies were obtained from both SEBS-13 and SEBS-20 grades when a solvent casting process was used. These materials are denoted isotropic. When a sheet die extrusion process was used, SEBS-20 based materials featuring PS cylinders and clay platelets simultaneously aligned parallel to the extrusion direction (Y direction according to Figure 6-1) were obtained. These materials are referred to as oriented.

The rest of the nanocomposites discussed in this manuscript feature more complex morphologies. In fact, the sheet die extrusion of SEBS-13 based materials induced the alignment of clay platelets parallel to (XY) plane and a fraction of PS nanodomains parallel to Y axis. However, another fraction of PS cylinders remained isotropic. The imperfect alignment of PS cylinders in SEBS-13 was attributed to the low fraction of PS block (Leice G Amurin et al., 2016). SEBS-20 based materials prepared by film blowing extrusion featured also complex morphologies depending on the air inflation ratio. At ratio R1, a major fraction of both PS domains and clay particles were aligned in the extrusion direction (Y axis) and a minor fraction was aligned across the thickness (parallel to Z axis). At ratio R3, clay platelets were oriented in the XY plane while PS domains were randomly distributed in the XY plane perpendicular to the thickness. This third class of materials is designated partially oriented.

An exhaustive characterization of the as-described morphologies by small angle X-ray scattering (SAXS) and transmission electron microscopy (TEM) was reported in previous publications (Leice G Amurin et al., 2016; De Sousa Jr et al., 2014; Helal et al., 2017). The nomenclature of these nanocomposites and short descriptions of their morphologies are provided in Table 6-1. More details and illustrative schemes can be found in (Helal et al., 2017) or Table 3-3 of this dissertation.

	Nomenclature	Morphology description	
Sheet-die Extrusion	SEBS-20-oriented SEBS-20-20A-oriented	PS cylinders uniaxially oriented in the main flow direction along Y axis Clay platelets oriented in the main flow direction parallel to XY plane	Z Field PER N N Diagonal
Solvent casting	SEBS-13-isotropic SEBS-13-20A-isotropic SEBS-20-isotropic SEBS-20-20A-isotropic	Isotropic PS cylinders Isotropic clay platelets	
owing extrusion	SEBS-20-partially oriented R1 SEBS-20-20A- partially oriented R1	Major fraction of PS cylinders oriented in the main flow direction along Y axis Major fraction of clay platelets oriented parallel to XY plane Minor fraction of PS cylinders and clay platelets oriented across the thickness parallel to XZ and YZ planes	
Film bl	SEBS-20- partially oriented R3 SEBS-20-20A- partially oriented R3	PS cylinders randomly distributed in the plane XY Clay platelets oriented parallel to XY plane	
Sheet-die Extrusion	SEBS-13- partially oriented SEBS-13-20A- partially oriented	Partially isotropic PS cylinders Partially oriented PS cylinders in the main flow direction along Y axis Clay platelets oriented parallel to XY plane	

Table 6-1: Nomenclature and morphology description of SEBS-20 and SEBS-13 based nanocomposites

6.3 Characterization

6.3.1 Mechanical properties

The mechanical properties of the block copolymers and their nanocomposites were measured using STM Alliance machine equipped with 1kN load cell and pneumatic grippers. The elastic Young's modulus and the elongation at break were evaluated in the tensile mode according to ASTM 412D (*Standard Test Methods for Vulcanized Rubber and Thermoplastic Elastomers*, 2016). The speed was selected equal to 50mm/min, lower than the speed recommended by the standard (500 mm/min) for more precision (Saleem et al., 2014). 5 specimens at least were tested for each material.

6.3.2 AC short-term breakdown strength

The AC short-term breakdown strength of the samples was measured using ball-type electrodes of diameter 4 mm in a dielectric oil environment. For each sample, 12 specimens were considered to calculate the dielectric strength using Weibull distribution. A voltage ramp of 2 kV/s at a frequency of 60 Hz was applied until breakdown occurred. All measurements were done at room temperature. Since the thickness of the samples was not strictly uniform, all the measurements were corrected to equal the breakdown strength of a 550 μ m thick film using the power law relationship between the breakdown field and the film thickness in order to obtain comparable results (Takala et al., 2010).

6.4 **Results and discussion**

6.4.1 Mechanical properties

The stress-strain curves of SEBS-20 and SEBS-13 sets of materials are respectively plotted in Figure 6-2 and Figure 6-3. The Young's modulus was evaluated by calculating the slope in the linear region up to 5% strain. The values are summarized in Figure 6-4. It is worth mentioning that the final objective is to evaluate the mechanical properties of the samples in the thickness direction (Z direction) in order to correlate them with the breakdown strength measurements, where the electric field was applied parallel to the thickness as indicated by the arrows in the illustrative cartoon in Table 6-1. Since it was technically difficult to evaluate the mechanical properties by compression, tensile measurements were adopted and the directions of the tests were varied, depending on the orientation of PS cylinders and clay platelets, in order to mimic the maximum the mechanical behavior across the thickness. Nevertheless, the results will be discussed carefully as some differences from the real behavior may exist.

For the samples prepared by solution, no specific recommendation for the test direction was needed as they are isotropic. For the extruded samples exhibiting specific orientation of PS cylinders and clay layers parallel to the Y axis, the tensile testing was performed in the transverse direction (parallel to X axis) which is equivalent to Z axis as far as the contribution of PS cylinders to the mechanical properties is concerned.

For the samples prepared by film blowing extrusion at a ratio R1, the tests were performed in the XY plane with an angle of 45° to take into account the orientation of a small fraction of PS cylinders that deviate from aligning parallel to the Y axis. This direction will be referred to as diagonal (Figure 6-1 and Table 6-1) and is believed to be equivalent to Z axis as far as the contribution of PS cylinders to the mechanical properties is concerned, since a minor fraction of PS cylinders were oriented in the thickness direction. For the samples prepared by film blowing extrusion at a ratio R3, the tensile tests were performed in the diagonal direction as well. Since the PS cylinders are randomly oriented in the XY plane, the mechanical properties in this direction are expected to be close to those of the isotropic sample. However, in the thickness direction, the mechanical properties should be equivalent to the case of oriented samples.

As noticed, all the specified test directions for the anisotropic samples take into account the real contribution of PS cylinders orientation to the mechanical properties but do not correspond to the real contribution of clay layers which are oriented parallel to the XY plan in which the tensile tests are performed and perpendicular to the Z axis (thickness direction).

Thus, the real mechanical properties of these samples in the thickness direction, especially in the elastic region, might be lower than the measured values due to the reduced mechanical strength of the "intrastack phase" as reported by Mishnaevsky et al. (Dai & Mishnaevsky Jr, 2013; Mishnaevsky Jr, 2012). However, since this imprecision is present in all the anisotropic samples, the evaluation of their relative performance compared to each other remains possible.

Figure 6-2 shows that all SEBS-20 based materials present comparable elongations at break and ultimate strengths (including those prepared by film blowing extrusion and not reported in the Figure). The same conclusion can be made for SEBS-13 set of materials as well (Figure 6-3). In terms of stiffness, all the nanocomposites showed in general improved stiffness compared to their pure materials, as it can be seen in Figure 6-4. Hence, the effect of nanoclays is more obvious on the elastic deformation rather than plastic deformation. This stiffening effect is partially attributed to the relatively high Young's modulus of clay platelets especially along their planes, as reported several times in the literature (Dai & Mishnaevsky Jr, 2013; Mishnaevsky Jr, 2012; Leszek A Utracki, 2010). Besides, the interfacial interactions have an important impact as well. In fact, the presence of clay platelets might prevent the molecules from orientation resulting in stiffness enhancement (Fang, Leng, & Gao, 2006; Helal et al., 2017; Vo et al., 2011).

Considering the set of materials prepared from SEBS-20 and the data reported in Figure 6-2 and Figure 6-4, it can be concluded that the oriented materials prepared by extrusion exhibit the lowest elastic moduli, respectively equal to 2.7MPa and 5.2MPa for the neat SEBS-20 and the SEBS-20-20A-oriented nanocomposite. This is expected since perpendicular to PS cylinders orientation, the mechanical properties are rather governed by the soft PEB phase featuring lower stiffness (Leice G Amurin et al., 2016; Wang et al., 2010). The materials prepared by film blowing extrusion at a ratio R1 exhibit improved elastic moduli, respectively equal to 3.3MPa and 5.7MPa for the neat SEBS-20-partially oriented R1 and its corresponding nanocomposite. This improvement is attributed to the orientation of a fraction of PS cylinders parallel to the thickness. However, it is worth noting that SEBS-20- partially

oriented R1 seems to be relatively inhomogeneous in the test direction. In fact, although the majority of the tested specimens exhibit behavior close to the average behavior presented in Figure 6-2 and Figure 6-4, few specimens exhibit higher stiffness with Young's modulus around 6.2 MPa (plots not presented in this manuscript).

The samples prepared by solution and by film blowing extrusion at a ratio R3 exhibit the highest improvement in stiffness. Furthermore, their elastic moduli are similar, for both cases: unfilled matrices and nanocomposites. This behavior confirms the hypothesis of isotropic morphology in the XY plane of samples prepared with a ratio R3 concluded from SAXS and TEM data discussed in a previous publication ((Helal et al., 2017). As a consequence, in the Z direction, the mechanical behavior of these materials (partially oriented R3) is rather close to that of the oriented samples measured in the transverse direction. This assumption will be considered later in the discussion of the breakdown strength results. In a summary, parallel to the thickness (Z direction), SEBS-20 based materials could be classified corresponding to their stiffness as following: oriented~ partially oriented R3< partially oriented R1< isotropic. This trend supports also the potential role of the interfacial interactions in the stiffening effect. In fact, it was reported in (Helal et al., 2017) that nanocomposites with isotropic and partially oriented morphologies exhibit more confinement in the interfacial region.

According to Figure 6-3 and Figure 6-4, the set of samples prepared from SEBS-13 exhibit generally higher elongation at break but lower stiffness compared to the samples prepared from SEBS-20. This behavior is expected due to the reduction of PS block fraction exhibiting higher mechanical strength compared to PEB block. Moreover, extruded SEBS-13 based samples, both the neat and the nanocomposite, exhibit lower stiffness compared to isotropic materials prepared by solution. This behavior supports the conclusion that SEBS-13 materials prepared by extrusion exhibit both oriented PS domains and isotropic PS domains, despite the fact that SAXS diffraction patterns corresponding to these materials reported in (Helal et al., 2017) indicate only isotropic PS cylinders.



Figure 6-2: Characteristic stress-strain plots of SEBS-20 based materials



Figure 6-3: Characteristic stress-strain plots of SEBS-13 based materials



Figure 6-4: Young's modulus of respectively SEBS-20 and SEBS-13 sets of materials

6.4.2 AC short-term Breakdown strength

The data corresponding to the AC short-term breakdown tests were retrieved using commercial software. Two-parameter Weibull distribution (equation (6.1)) was used for the estimation of the breakdown strength. More details regarding the statistical analysis of breakdown measurements could be found in IEEE 930 standard ("IEEE Guide for the Statistical Analysis of Electrical Insulation Breakdown Data," 2005).

$$P = 1 - \exp\left[-\frac{E}{E_0}\right]^{\beta} \tag{6.1}$$

Where:

P: is the cumulative probability of failure at an electric field less or equal to E;

E₀: is the scale parameter corresponding to the breakdown strength for which the cumulative failure probability is equal to 63.2%;

 β : is the shape parameter which measures the scatter of the data. The larger β is the narrower is the range for breakdown strength.

Weibull plots, scale factor and shape factor related to SEBS-20 and SEBS-13 sets of materials are reported respectively in

Figure 6-5 and Figure 6-6. An overall inspection of the results shows that all the nanocomposites exhibit higher dielectric strength compared to their unfilled parts and increased shape factors indicative of improved reliability. The highest increase of β is observed in samples prepared by solution as they are more homogeneous due to their isotropy.

As far as unfilled materials are concerned, it could be seen that indeed the breakdown strength varied depending on the processing method and consequently on the morphology and mechanical properties of the block copolymer. In fact, oriented samples exhibit lower dielectric strength compared to the isotropic samples. Moreover, in the case of SEBS-20 set, SEBS-20- partially oriented R3 and SEBS-20-oriented exhibit comparable values around 56 kV/mm while SEBS-20-isotropic and SEBS-20-partially oriented R1 exhibit higher values respectively around 60kV/mm and 62 kV/mm. This observation partially matches the trend observed for the mechanical properties but a higher breakdown was expected for the isotropic sample which exhibits the highest elastic modulus. This behavior might be due to the fact that SEBS-20-isotropic features less perfect order compared to SEBS-20-bioriented R1. In fact, the decrease of order may result in reduced number of nanoscaled interfaces between the blocks and induce less efficient trapping of charge carriers (Samant et al., 2016). SAXS data reported in (Helal et al., 2017) support this hypothesis.

Surprisingly, SEBS-13 set of materials shows similar behavior compared to SEBS-20 materials with breakdown strength equal to 61kV/mm for the isotropic material prepared by solution and equal to 53kV/mm for the material prepared by extrusion. Although this behavior supports the hypothesis that the two samples prepared by different processing techniques do not exhibit the same PS orientation, it disagrees with the fact that these materials exhibit lower stiffness compared to SEBS-20 due to the reduced ratio of styrene and are consequently expected to exhibit lower dielectric strength. This unexpected high performance is probably due to the fact that the diameter of PS cylinders in SEBS-13, equal to 9.7 nm according to calculations based on SAXS results (not reported in this manuscript), is smaller than the diameter of PS cylinders in SEBS-20 estimated as 12.5 nm (Helal et al., 2017), which induces larger and more nanosized interfacial area where the charge carriers could be trapped and subsequently compensates for the deterioration of stiffness in SEBS-13. However, this hypothesis needs further investigation to be confirmed.

In terms of nanocomposites, the data reported in

Figure 6-5 and Figure 6-6 shows that the lowest improvement in breakdown strength, compared to the unfilled matrices, corresponds to isotropic nanocomposites prepared by solution. It is in the range of 6 to 7%. The nanocomposites featuring clay platelets aligned parallel to XY plane, prepared either by sheet-die extrusion or film blowing extrusion at ratio R3, exhibit higher improvement up to 10.6% due to alignment of nanoparticles perpendicular to the electric field. Unfortunately, this improvement does not compensate for the decline of the breakdown strength of the matrix caused by the simultaneous alignment of PS cylinders perpendicular to the thickness. A similar behavior was reported in our previous study (E Helal et al., 2015) concerning SEBS-30 based nanocomposites. However, a higher increase of breakdown strength, up to 45%, was achieved in SEBS-30-20A oriented nanocomposite compared to unfilled SEBS-30 oriented, indicating more efficient role of clay platelets as barriers to the flow of charge carriers. This higher performance might be related to improved dispersion, degree of orientation and interaction with polymer chains at increasing fraction of PS block. In fact, clay platelets exhibit better interaction with the relatively polar aromatic rings of PS block (Carastan et al., 2014). Also, the difference in the thickness between the

samples used to perform the breakdown tests (average thickness in the range 100-170 µm for SEBS-30 based nanocomposites compared to 550 µm for SEBS-20 and SEBS-13 based nanocomposites) might partially contribute to the superior performance of SEBS-30 nanocomposite. In fact, in solid insulations, the breakdown strength is often reported to increase with decreasing sample thickness. This film thickness effect has been attributed to potential changes in several parameters including the probability of defects, charge trapping and detrapping mechanism and material morphology (H. K. Kim & Shi, 2001; L. Zhao, Liu, Su, Pan, & Zhang, 2011). Hence, the reduced thickness of tested SEBS-30 materials might have resulted in more perfect packing and alignment of clay platelets and consequently higher improvement of breakdown strength compared to the unfilled material, as observed.

SEBS-20-20A-partially oriented R1 nanocomposite exhibits the highest increase in dielectric strength by 25% compared to its unfilled peer SEBS-20-R1 and 37% compared to conventionally extruded matrix SEBS-20-oriented. This increase is attributed to synergetic effects including the improved elastic moduli of the matrix induced by the alignment of some cylinders in the thickness direction, the high state of order of the block copolymer compared to isotropic and R3 samples and simultaneously the alignment of major fraction of clay tactoids perpendicular to the electric field (Helal et al., 2017).



Figure 6-5: Breakdown strength of SEBS-20 based nanocomposites



Figure 6-6: Breakdown strength of SEBS-13 based nanocomposites

6.4.3 Summary

The comparison of the trends followed by the tensile strength and breakdown strength results reveal an important correlation between these two properties in the case of unfilled SEBS-20 and SEBS-13 matrices. In fact, this correlation is a well-known behavior in soft elastomers and thermoplastic elastomers in general, due to the high possibility of deformation of soft materials under an applied electric field (Blythe & Bloor, 2005)

In the studied materials, since the mechanical strength of thermoplastic PS domains is greater than that of the rubbery PEB domains, the control of their orientation induced different mechanical behaviors of unfilled SEBS-20 and SEBS-13 matrices as function of their morphologies. Subsequently, the dielectric breakdown strength followed the same trend of the mechanical stiffness (both estimated in the same direction). In particular, when PS cylinders are aligned parallel to the direction of the applied electric field, the lowest mechanical strength and dielectric breakdown strength were simultaneously observed. In the nanocomposites, in addition to its dependence on the mechanical stiffness, the breakdown strength was mainly governed by the orientation of clay platelets compared to the applied electric field. In particular, the alignment of clay perpendicular to the applied electric field is the desired configuration. In this context, isotropic nanocomposites exhibited the lowest improvement of breakdown strength, compared to their unfilled pairs, despite their high mechanical stiffness. However, the partially oriented R1 nanocomposite exhibited the best combination of mechanical and dielectric breakdown strengths due to adjusted alignment of respectively clay platelets and PS cylinders. Other factors such as the state of order in the block copolymer and the quality of the interfaces affected also the breakdown strength.

6.5 Conclusion

In this study, it was demonstrated that the mechanical strength and ultimately the breakdown strength of SEBS/clay thermoplastic elastomer nanocomposites can be tuned through the design of their morphologies, using different processing techniques. The investigated

nanocomposites featured different morphologies that were classified as oriented, partially oriented or isotropic depending on the alignment of both clay platelets and PS cylinders. In unfilled SEBS matrices, the mechanical stiffness and the breakdown strength were strongly correlated and dependent on the orientation of PS nanodomains. But, in the nanocomposites, this correlation is less strong. Instead, the spatial distribution of clay platelets was the main factor governing the extent of the achieved improvement of breakdown strength compared to the unfilled matrix.

Overall, this control of morphology and functional properties is expected to increase the potential of these flexible nanocomposites in different dielectric applications including electrical insulation and dielectric elastomer actuators.

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CHAPTER 7

DISCUSSION

This thesis studied the design and processing of polymer nanocomposites with controlled morphologies, intended to serve as HV insulating materials. The parameters involved within the concept of controlled morphology are: state of dispersion (agglomerations vs. nanoscale dispersion), orientation and selective localization of the nanoparticles. To achieve this objective, this project was divided in three main parts as described in the introduction of this thesis. The work performed in these parts involved the investigation of two types of nanoparticles as well as a block copolymer and a blend of polyethylene with a block copolymer as template matrices. In the following, some of the challenges encountered in each part of the project are discussed. Besides, the achievement of the specific research objectives related to each part is evaluated.

7.1 Nanocomposites with tailored morphologies

The performance of insulating polymers used as HV insulating materials is affected by various physical properties including real and imaginary parts of dielectric permittivity, breakdown strength, resistance to partial discharges, thermal conductivity and mechanical strength. In polymer nanocomposites, it was observed that several engineering properties might be optimized by tailoring the nanocomposites morphologies. In this context, the analysis of the recent literature, discussed in chapter 1, has shown that specific approaches based on the use of multiphase polymer systems were developed to achieve tuned morphologies in nanocomposites. The multiphase materials, such as block copolymers and polymer blends, were used as template matrices and were able to induce different enhancements in the electrical, thermal and/or mechanical behavior depending on the targeted field. In particular, these approaches started gaining attraction in nanodielectrics. However, their application to the field of HV insulation is still limited.

In this thesis, these approaches were extended to polymer nanocomposites intended for use as HV insulating materials. Through chapter 2 to chapter 6, it was demonstrated that indeed these approaches have potential to improve the functional properties related to HV insulation.

The first challenge encountered at the early steps of this project was the choice of the block copolymer and the nanoparticles. In particular, SEBS was chosen due to its ordered nanostructure as a block copolymer, interesting mechanical properties as a thermoplastic elastomer and good resistance to water treeing. In terms of nanoparticles, clay platelets and ZnO nanospheres were selected. Both of them exhibit different shapes and chemical natures but have in common benefits in HV insulation and low costs.

The specific findings and challenges related to each part of the project are discussed in the next three sections.

7.2 SEBS/Clay nanocomposites

SEBS/clay nanocomposites with different morphologies were investigated, with stress on the effect of clay orientation, localization and degree of exfoliation on some key properties related to HV insulation; namely: dielectric losses, dielectric breakdown strength and mechanical strength. The control of localization and degree of exfoliation was ensured through the use of MA graft attached to the PEB block. In fact, the interaction between MA and hydroxyl groups at the edge of clay layers, which are already pre-intercalated due to the organic surface treatment, improves the peeling mechanism and consequently the exfoliation of clay into individual layers. Furthermore, this increased affinity favors the localization of clay platelets in the elastomer PEB block.

In terms of orientation, the control was achieved by specific processing techniques. In this context, oriented nanocomposites exhibiting alignment of both clay platelets and PS nanodomains along the thickness of nanocomposites films were obtained by sheet die extrusion process while isotropic nanocomposites were achieved by solvent casting process. The alignment of clay platelets improved efficiently the resistance to dielectric failure and

reduced the dielectric losses compared to isotropic morphologies. However, the alignment of PS domains reduced the mechanical stiffness of the matrix and subsequently its dielectric breakdown strength. To overcome this undesirable effect, more complex configurations, allowing taking advantage from clay alignment without sacrificing the dielectric and mechanical strength of the matrix, were needed. In this context, the morphologies, untitled "partially oriented" were prepared using a different processing technique: film blowing extrusion. In this process, a tubular die is used instead of a sheet die and the nanocomposite film is inflated with air at tunable blow-up ratios. As a result, radial forces are applied instead of only lateral forces (applied in the main extrusion direction when a sheet-die extrusion process is used). The applied blow-up ratio determines the ratio between the final and initial diameters of the fabricated tube and ultimately affects the orientation of both clay platelets and PS cylinders. In the as-obtained nanocomposite films, an important fraction of clay platelets were aligned along the thickness (parallel to the main extrusion direction) while PS domains were either isotropic in the plane perpendicular to the thickness or having a fraction of them oriented across the thickness as well. For this reason, these morphologies were referred to as "partially oriented". They succeeded in maintaining the mechanical strength of the matrix while improving the breakdown strength and limiting the dielectric losses, due to alignment of clay platelets.

To validate the correlation between mechanical and dielectric strength for these soft SEBS/clay nanocomposites exhibiting different morphologies, the mechanical properties were evaluated. In breakdown measurements, the electric field was applied perpendicular to the samples thickness. Therefore, evaluation of the mechanical stiffness across the thickness would be favored for more accuracy. However, it was technically difficult to evaluate the mechanical properties across the thickness by compression. Therefore, tensile mode was adopted and the directions of the tests were selected carefully in order to simulate to the best the morphology across the thickness, as described in chapter 6. The results confirmed the correlation between mechanical and dielectric strengths. This was another challenge encountered in this part of the project.

In order to better understand the role of the interface in such nanocomposites, an attempt was made to establish a relationship between the studied morphologies and their corresponding interfacial molecular dynamics. For this purpose, relaxation processes observed in broadband dielectric spectra (BDS) and dynamic mechanical analysis (DMA) spectra were analyzed. The studied systems were rather complex, with the two blocks of SEBS exhibiting distinct segmental relaxations in addition to the potential interfacial relaxations. In particular, a relaxation process attributed to rubber chains with restricted mobility located in the interface region was identified. Furthermore, the spatial distribution of nanoparticles affected this relaxation process. In a summary, it was found out that the isotropic morphologies, partially oriented morphologies and morphologies with clay selectively located in the rubber block, result in more restriction and stronger fragile behavior. It is worth noting that this relaxation process was observed in BDS spectra of all the nanocomposites but only for selected materials in DMA spectra. This behavior requires further investigation.

7.3 SEBS/ZnO nanocomposites

In the second part of this project, tailored dispersion of ZnO nanospheres in SEBS block copolymer was investigated. In particular, nanoscale dispersion and interaction between these isotropic nanoparticles with polymer chains were the tuned aspects. The control was again enabled through the use of MA groups grafted to PEB elastomer block and exhibiting good affinity with the nanoparticles surface. In fact, the organic treatment of the surface of ZnO nanoparticles helped decreasing their hydrophilic character and consequently improving their compatibility with the polymer matrix. Simultaneously, MA reacted with the remaining hydroxyl groups that partially cover the surface of the nanoparticles, forming ester and hydrogen bonds. The resulting complexes act as in-situ formed cross-linked structures that improve the dispersion and affect drastically the rheological properties. In particular, the storage modulus of SEBS/ZnO nanocomposites prepared with MA, was nearly constant at low angular frequencies (a plateau was formed). This interesting behavior indicates the formation of networks between the nanoparticles and polymer chains. It was observed at ZnO concentrations starting from 5 wt%.

The formation of networks was correlated with improvements in thermal conductivity and resistance to surface erosion at lower content of nanoparticles compared to SEBS/ZnO nanocomposites prepared without the MA graft. Besides, at low frequencies, the dielectric losses, rather dominated by electrical conduction phenomena in this range, were remarkably increased with temperature. This intriguing behavior was not observed in SEBS/ZnO nanocomposites prepared without MA. It is also believed to be related to the specific spatial distribution of nanoparticles and the established networks which render the nanocomposites slightly conductive. However, further investigations are required to confirm it. From a practical point of view, the efficiency of these materials as HV insulations might be reduced if elevated operational temperatures are required. However, at room temperature and intermediate temperatures up to 70 °C, which are more common, the dielectric losses of SEBS/ZnO prepared with MA are lower than those prepared without MA due to the improved dispersion.

Lastly, it is worth noting that although networks between polymer chains and ZnO nanoparticles were formed in the presence of MA, a total selective localization of the studied ZnO nanoparticles in the PEB block was unfortunately not possible because their dimensions were in the same range or slightly higher than the characteristic length scales of SEBS nanodomains, which imposes thermodynamic constraints. Hence, the use of ZnO nanoparticles of average diameter smaller than the dimensions of the PEB block could be interesting, since it facilitates the achievement of preferential localization and may improve further the observed property enhancements at even lower loadings of nanoparticles.

7.4 PE/SEBS/ZnO blend nanocomposites

The last part of this project aimed at producing materials with controlled morphology for practical use in HV insulation. In this context, blends of SEBS/ZnO nanocomposites with HDPE were investigated. The choice of ZnO-containing nanocomposites was justified by the confirmed role of ZnO nanoparticles in reducing space charge accumulation, which is a crucial property in HV cable insulations and especially HVDC cable insulations. The selective localization of nanoparticles in either PE or SEBS was the main parameter to

control at this step. It was confirmed by microscopy and by estimation of the wetting coefficient that ZnO nanoparticles were located in SEBS phase. This preferential localization was enabled by the high affinity of ZnO nanoparticles to the MA groups attached to the PEB block and by using short mixing time during melt compounding (to avoid migration of the nanoparticles to PE).

The as-obtained PE/SEBS/ZnO blend nanocomposites exhibited lower dielectric losses as well as improved resistance to surface erosion and mechanical flexibility compared to conventional PE/ZnO nanocomposites. In this context, tensile measurements confirmed that the elongation at break increased for PE/SEBS/ZnO nanocomposites due to the presence of SEBS thermoplastic elastomer while PE/ZnO nanocomposites broke shortly after necking. It is also worth mentioning that the measured values of elongation at break for both PE/SEBS/ZnO and PE/ZnO nanocomposites might be extended further if the tensile test speed is decreased.

Regarding the morphology of the blends, equal mass fractions of PE and SEBS were selected initially to achieve a co-continuous morphology. In the unfilled PE/SEBS blend, the achievement of a co-continuous morphology was confirmed by microscopy after solvent extraction of SEBS. However, the incorporation of ZnO in SEBS, in the presence of MA, induced an increase of viscosity and subsequently a switch of the blend morphology to droplets morphology where elongated domains of SEBS/ZnO constitute the dispersed phase. In the case of the studied blend nanocomposites, the co-continuous morphology is believed important to achieve a thermal conductivity path for instance and to increase further the efficiency of resistance to surface erosion. Moreover, it ensures the homogenous distribution of the nanoparticles in the matrix while maintaining the option of selective localization of the nanoparticles possible. Therefore, further investigation to optimize the morphology of PE/SEBS/ZnO blend nanocomposites would be interesting.

CONCLUSION

In this section, a summary of the main findings of this project is provided. Partial conclusions relative to each part of the project are presented in accordance with the order of their presentation in the thesis. At the end of the section, the general conclusion is stated.

SEBS/clay nanocomposites

In the first part of this project, SEBS/clay nanocomposites with different concentrations of clay particles up to 7.5 wt% and different fractions of PS block in SEBS respectively equal to: 13%, 20% and 30%, were investigated. Various morphologies were obtained by different processing techniques. In these morphologies, orientation and preferential localization of clay platelets were the main aspects of dispersion control. It was found that:

- The control of orientation of clay platelets and PS nanodomains in SEBS block copolymer was enabled by the use of different processing techniques and different grades of SEBS block copolymer, one of them having an MA graft attached to its rubber block. The used processing techniques were: solvent casting, sheet-die extrusion and film blowing extrusion. They led to respectively: isotropic, totally oriented and partially oriented morphologies, as discussed in chapter 3.
- In SEBS grade containing 30 wt% of PS block, 5 wt% clay platelets represented an optimum concentration in terms of dispersion and exfoliated clay fraction. This result was confirmed by TEM quantitative analysis. Moreover, this concentration coincided with the maximum improvement of the breakdown strength.
- The orientation of clay platelets induced a significant decrease of the dielectric losses especially at low frequencies and significant increase of the breakdown strength up to 45 % at 5 wt% clay. However, in the isotropic samples only a slight increase of the breakdown strength was achieved and significantly higher dielectric losses were exhibited.
- The orientation of PS nanodomains which occurs simultaneously with clay orientation, when a sheet die extrusion process is used, led to a decrease of the

breakdown strength of the polymer matrix, due to reduction of the mechanical strength. This reduction unfortunately counterbalances the increase observed from the orientation of clay.

- The partially oriented morphologies investigated in the purpose of limiting this negative effect, featured an optimized morphology were clay particles are mainly oriented along the thickness while a fraction of PS domains are randomly distributed along the thickness and another fraction oriented across the thickness. This morphology obtained by film blowing extrusion allowed avoiding the negative effect on breakdown strength and mechanical strength of the unfilled matrix encountered when aligning PS nanodomains while maintaining the advantages on breakdown strength and dielectric losses observed when clay platelets are aligned.
- When clay platelets are incorporated, rubber chains located at the interface exhibited restricted mobility. This behavior was confirmed by Broadband Dielectric Spectroscopy (BDS) and Dynamic Mechanical Analysis (DMA) measurements that both show the appearance of a relaxation peak related to the dynamics of this interfacial layer. The interfacial glass transition temperature was estimated to occur in the range 6 to 35 °C while the Tg of the bulk elastomer block is around -30°C. The thickness of the interfacial layer region was estimated, by fitting of DMA data, to be in the range 7-10 nm.
- The molecular dynamics of these interfacial chains were found to depend on the orientation of both PS nanodomains and clay nanoparticles, the fraction of PS block as well as the location and degree of exfoliation of clay platelets, which was tuned through the use of maleic anhydride graft. In particular, samples with lower degree of alignment, higher degree of clay exfoliation and preferential localization in rubber block, were found to exhibit higher interaction and higher interfacial glass transition temperatures. This behavior was attributed to strong confinement of interfacial glass transition temperature decreases and the effect of orientation was found to decrease.

The main conclusions of this part are: the orientation and localization of clay platelets in SEBS block copolymer can be controlled by means of the processing technique and selection
of the appropriate grafts or functional groups that have high affinity to functional groups attached to the surface of nanoparticles. This control of **orientation and selective localization** allows the tuning of the **dielectric losses**, **breakdown strength and mechanical strength**.

SEBS/ZnO nanocomposites

In the second part of this project, SEBS/ZnO nanocomposites were investigated. ZnO nanoparticles have a spherical shape and were modified with alkyl ammonium salt. Due to the isotropy of the nanoparticles, no emphasis was given to orientation. Instead, the spatial distribution of the nanoparticles and their interaction with polymer chains were the key parameters to tailor, in this part. This control was enabled through the use of MA grafted to rubber block. It was found that:

- The dispersion of ZnO nanoparticles in SEBS was improved in the presence of MA compared to the dispersion observed in SEBS without MA. This behavior is attributed to good interaction between MA and the surface of the nanoparticles.
- In the presence of MA, nanocomposites exhibited a distinguished rheological behavior characterized by a plateau of the storage modulus at low frequencies. This behavior observed at low ZnO content starting from 5 wt% indicates the formation of a network between ZnO nanoparticles and polymer chains, i.e. rheological percolation.
- This rheological percolation was accompanied by an improvement of thermal conductivity by 10% and excellent increase of resistance to surface erosion by partial discharges: the eroded volume was reduced by 90% at 5 wt% ZnO. Furthermore, the dielectric losses at low frequencies, which are dominated by electrical conduction phenomena in this range, increased remarkably as function of temperature. This behavior is also believed to be correlated with the established network.
- At power frequencies and operational temperatures, the dielectric losses of nanocomposites with controlled dispersion were lower than the nanocomposites not featuring controlled dispersion (prepared without MA). Besides, no noticeable

decrease of the dielectric breakdown strength compared to the unfilled matrix was observed up to 20wt% ZnO loading.

DC electrical conductivity measurements have shown that SEBS/ZnO nanocomposites exhibit in general a non-linear feature attributed to the presence of ZnO. In the nanocomposites with controlled dispersion, a higher control of this non-linearity as function of the electric field and ZnO content was possible. In addition, at 5wt% ZnO and in the presence of MA, SEBS/ZnO nanocomposite exhibited electrical conductivity values lower than those of unfilled SEBS. This interesting behavior might be caused by the formation of trapping sites that block charge carriers from propagating and accumulating in the bulk of the material.

The main conclusions of this part are: the **establishment of a network between ZnO nanoparticles and SEBS block copolymer** chains was enabled by the presence of MA groups attached to the elastomer block of SEBS. This morphology was accompanied by improvements of **thermal conductivity** and **resistance to surface erosion** as well as reduced dielectric losses at power frequencies and service temperatures. Besides, the breakdown strength was maintained in the same range as the unfilled matrix.

PE/SEBS/ZnO blend nanocomposites

In the last part of this project, blends of SEBS-MA/ZnO nanocomposites with HDPE were investigated. The choice of ZnO-containing nanocomposites was driven by the confirmed role of metal oxide nanoparticles in reducing space charge accumulation in polymer insulations. This property is a crucial requirement in the development of insulating materials suitable for high voltage direct current (HVDC) cables. In this immiscible blend of PE and SEBS, the selective localization of ZnO nanoparticles in either PE or SEBS was the main aspect of dispersion control that was highlighted. In particular, it was found out that:

• Due to the affinity of ZnO nanoparticles surface to MA graft, nanoparticles remained exclusively in SEBS phase and and potentially at the interfaces PE/SEBS but didn't migrate to PE. This behavior was predicted thermodynamically by the estimation of

the wetting coefficient and confirmed by high resolution scanning electron microscopy.

- The blends contained equal mass fractions of PE and SEBS polymers. The unfilled PE/SEBS blend featured a co-continuous morphology. However, in the presence of ZnO nanoparticles, the blend switched to elongated droplets morphology where the dispersed phase is composed of SEBS/ZnO domains. The morphology switch was attributed to the increase of SEBS viscosity upon the incorporation of ZnO nanoparticles in the presence of MA.
- PE/SEBS/ZnO blend nanocomposites featured improved nanoscale dispersion compared to PE/ZnO nanocomposites prepared by melt compounding which exhibited several agglomerations, especially at increasing ZnO content.
- At power frequencies and operational temperatures, the dielectric losses of PE/SEBS/ZnO blend nanocomposites are lower than those related to PE/ZnO reference nanocomposites. Besides, in both types of nanocomposites, an interfacial relaxation process was observed. In the blend nanocomposites, the broadness and the dielectric strength of the relaxation process were reduced compared to PE/ZnO nanocomposites, thanks again to the improved dispersion.
- The selective dispersion of ZnO in SEBS droplets induced more efficient resistance to surface erosion by partial discharges inside the droplets and ultimately in the whole blend, compared to PE/ZnO nanocomposites. This effect was more obvious at low concentrations of ZnO. In fact, elongated domains of SEBS/ZnO constitute an equivalent of micrometric particles featuring at the same time excellent resistance to surface erosion and good adhesion to PE matrix, due to the compatibility between the PEB rubber block and PE. In PE/ZnO nanocomposites, although the presence of the inorganic nanoparticles improved the resistance to surface erosion, the observed submicrometric and micrometric agglomerations exhibit poor adhesion to PE and act as week zones of high electrical stresses reducing the overall performance.
- The presence of the elastomer PEB block allowed PE/SEBS/ZnO nanocomposites to exhibit improved elongation at break compared to neat PE and PE/ZnO nanocomposites, and higher toughness when low ZnO loading (1wt%) was used.

The main conclusions of this part are: **blending** HDPE with SEBS thermoplastic elastomer and **selectively locating** ZnO nanoparticles in SEBS phase led to improvements in dielectric and mechanical performances, namely: **lower dielectric losses** at utility frequencies, **higher resistance to surface erosion** by partial discharges and **higher mechanical flexibility**. As a consequence of these improvements, PE/SEBS/ZnO blend nanocomposites have good potential for HV insulation applications.

General conclusion

In this project, nanocomposites with tailored nanoparticles spatial distribution have been studied as candidates for HV insulation. Within the concept of controlled dispersion, several aspects have been tuned, namely: state of dispersion, interaction between the nanoparticles and polymer chains, orientation and selective localization of the nanoparticles. The control was achieved using template matrices, i.e. block copolymers and polymer blends, and appropriate grafts to increase the affinity of nanoparticles to a specific block of the block copolymer. It was demonstrated in several occasions that the control of one or more of the abovementioned aspects is accompanied by an enhancement of one or more engineering properties that affect the performance of HV insulating materials. These properties include: dielectric breakdown strength, resistance to surface erosion by partial discharges, dielectric losses, thermal conductivity and mechanical properties. In an attempt to correlate the observed behaviors with nano and microstructure, the interfacial molecular dynamics were investigated by means of broadband dielectric spectroscopy and dynamic mechanical analysis. Interestingly, it was found out that the orientation and preferential localization of nanoparticles affect indeed the mobility of chains and the thickness of the interfacial region. To conclude, the findings of this work helped paving the way to a new generation of nanodielectrics and HV insulating materials featuring controllable properties and gave few insights to better understand the structure-property relationships behind.

RECOMMENDATIONS

In this project, it was demonstrated that well-defined nano/microstructures, achieved by controlling the spatial distribution of nanoparticles in multi-phase polymer matrices, have high potential for designing novel HV insulating materials with superior performance. To improve, consolidate, extend and/or up-scale the findings of this work, many additional studies might be worth investigation. In the following, few suggestions for future work are presented.

• SEBS/Clay nanocomposites

As far as SEBS/clay nanocomposites are concerned, the observed effect of orientation and preferential localization of clay platelets on the molecular dynamics of elastomer chains located at nanoparticle/polymer interface might be further investigated by carrying out modulated DSC measurements. In particular, the interfacial glass transition might be evaluated and correlated with the results of dielectric spectroscopy and dynamic mechanical analysis reported in chapter 3.

• SEBS/ZnO nanocomposites

The measurements of DC electrical conductivity carried out on SEBS/ZnO nanocomposites containing MA graft demonstrated that at 5 wt% ZnO, the electrical conductivity of the nanocomposite was even slightly lower than that of unfilled SEBS. This behavior indicates the formation of traps that block the space charges at an early stage from propagating and accumulating in the bulk and consequently reduce the overall conductivity. It is well in agreement with recent findings from the literature reporting a reduction in space charge accumulation and decrease of electrical conductivity of the insulating polymer matrix when low loadings of metal oxide nanoparticles (< 5 wt%) are incorporated. Thus, extending the study of the electrical conductivity of SEBS/ZnO nanocomposites to ZnO loadings below 5wt% is important to evaluate the possibility of reducing space charge accumulation in the bulk of the material by increasing the number of trapping sites and blocking the mobility of

charges carriers at an early stage. In fact, the controlled dispersion achieved in these nanocomposites have the potential to favor this behavior at concentrations of ZnO nanoparticles even lower than those reported in the literature, generally up to 5 wt%. Besides, achieving successful selective localization by using ZnO nanoparticles of average size lower than characteristic dimensions of SEBS block, might improve further this desired behavior.

Polyolefin/SEBS nanocomposite blends

In the last part of this project, blends of SEBS/ZnO nanocomposites with HDPE have been studied to evaluate their potential as new HV insulating materials recyclable and suitable for HVDC cables. Compared to conventional PE/ZnO nanocomposites, PE/SEBS/ZnO blend nanocomposites have shown their promise. Improving further the potential of these blends might be achieved by:

- Studying and determining the proportions of both PE and SEBS that lead to cocontinuity in PE/SEBS/ZnO blend nanocomposites, taking into account the increase of viscosity of SEBS after the addition of ZnO nanoparticles. In fact, the cocontinuous morphology might improve further several properties such as thermal conductivity and resistance to surface erosion, by enabling the formation of particles networks all over the material.
- PP is an interesting candidate for HVDC cable insulation that gained much attention recently. Application wise, the investigation of PP in blends with the studied SEBS nanocomposites is certainly beneficial to extend the potential of these materials in HV insulation and more specifically HVDC cables.

• Investigation of other nanoparticles

In the context of generalizing and up-scaling the approach of controlled dispersion for the development of new HV insulating materials, other nanoparticles featuring interesting properties might be considered. In particular, MgO as well as BN nanosheets and nanotubes constitute excellent candidates that have shown recently great improvements mainly in

thermal conductivity and suppression of space charge accumulation. The latter property is highly important in the development HVDC cable insulations.

Combinations of nanoparticles of different properties and geometries might also open opportunities to more complex nanocomposites designs and potential synergetic effects.

Besides, the synthesis of customized nanoparticles with a narrow size distribution and subjected to appropriate cleaning methods to remove residual impurities resulting from surface treatments would help controlling further the dielectric and electrical properties.

• Other measurements and modeling

To complement the reported characterizations and evaluate better the potential of the studied nanocomposites as candidates for HVDC cable insulations, space charge and DC short-term breakdown measurements could be of interest. Also, long-term measurements such as electrical treeing might give a better understanding of the influence of well-defined microstructures on the growth of electrical trees mean path.

Theoretical modeling and simulation of the electrical and thermal properties of the studied nanocomposites, such as the dielectric permittivity and the thermal conductivity, is also worth investigation. The modeling is expected to help elucidating the role of the interface in correlation with the controlled morphology of the nanocomposites. Ultimately, an easier prediction of structure-property relationships is pursued.

ANNEX I

SUPPORTING ELECTRONIC INFORMATION FOR ARTICLE 1

Styrenic block copolymer-based nanocomposites: implications of nanostructuration and nanofiller tailored dispersion on the dielectric properties

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1. Location of nanoclays

The location of nanoparticles in PS or PEB phase of SEBS depends on the presence of maleic anhydride (MA) group. In fact, in the absence of MA, nanoclays have greater affinity to polar PS phase which is most likely due to interaction between aromatic ring of PS and the polar surface of clay layers (Carastan et al., 2014; Richard A. Vaia, Jandt, Kramer, & Giannelis, 1996). However, in the presence of MA which is grafted to PEB phase, nanoclays tend to locate more in this phase since MA has greater affinity to nanoclays compared to PS due to its high polarity. This hypothesis was confirmed for the case of extruded samples and published in (Carastan et al., 2014). In this section, the case of isotropic samples prepared by solution will be treated. In particular, an annealed sample containing 5wt% nanoclays will be presented in Figure A I-1 and an unannealed sample containing 5wt% nanoclays will be presented in Figure A I-2.



Figure A I-1: SEBS_5wt % nanoclays (Prepared by solution, annealed, stained for 30 minutes)



Figure A I-2: SEBS_MA_5wt % nanoclay (prepared by solution, unannealed, stained for 30 minutes): (a) lamellar structure, (b) localized cylindrical structure where nanoclays are located

Figure A I-1 shows that:

- In annealed samples, PS phase exhibits an hexagonal cylindrical packing.
- Both nanoclays (indicated by arrows) and PS cylinders (black and dark grey circles) are randomly dispersed and not aligned.
- The nanoclays are intercalated and located within PS cylinders.

Figure A I-2 shows that:

- In unannealed samples, both lamellar and hexagonal cylindrical structures co-exist.
- Lamellar morphology is dominant (figure A I-2(a)).
- Nanoclays are mostly located in the cylindrical morphology which may indicate that they facilitate the transition from lamellar to cylindrical morphology even without thermal treatment (figure A I-2(b)).
- Both nanoclays (indicated by arrows) and PS domains (black and dark grey) are randomly dispersed and not aligned.
- The nanoclays are exfoliated and located in PEB phase (grey) between PS domains (black or dark grey).

2. Quantitative analysis procedure

The quantitative analysis procedure adopted in this paper was published in references (Carastan et al., 2014; Carastan et al., 2010; Vermogen et al., 2005). In this section, the basic steps are summarized. For more details, the cited references could be consulted. The steps are as follows:

- For each nanocomposite, 1 image at 20K, 4 images at 50 K and 10 images at 100K magnifications are considered in order to have a good overview of clay tactoids size at different scales. Examples of typical images used for the analysis are presented in Figure A I-3.
- For a considered magnification, the thicknesses of a fixed number of tactoids (30 per image) chosen randomly in each image were measured by imageJ software.
- The number of layers in each tactoid is determined using the following equation:

$$N = \frac{(t_{tactoid} + d_{001} - t_{layer})}{d_{001}}$$

Where:

N: number of layers;

ttactoid: thickness of the tactoid;

t_{layer}: estimated thickness of a single layer taken equal to 0.94 nm according to (Vermogen et al., 2005);

d₀₀₁: interlayer distance measured taken equal to 3.73 nm for SEBS-MA-2.5% and 3.47 nm for SEBS-MA-7.5% nanocomposite. These values were measured by X-ray diffraction (XRD).

- Each tactoid is then classified in one of 6 classes ranging from individual layers to large tactoids containing more than 50 layers depending on its estimated number of layers. An example of this classification is provided in Table A I-1.
- The final proportions of each class were then calculated taking into account a ponderation factor of the covered area corresponding to each magnification. For example, an image taken at 100k has a coefficient of 1 while an image taken at 50k has a coefficient of 4. These final results were used to plot Figure 2-3 in the manuscript reported in Chapter 2.



Figure A I-3: Typical TEM images considered for quantitative analysis (related to SEBS_MA_2.5% extruded nanocomposite) taken at: (a) 20 K, (b) 50 K and (c) 100 K

Class	Number of layers per tactoid	Number of tactoids per class
1	1	-
2	2 to 3	-
3	3 to 5	93
4	Small tactoids (5 to 15)	27
5	Middle size tactoids (15-50)	-
6	Large tactoids (>50)	-
Total number of analyzed tactoids in the 4		120
images		

Table A I-1: Proportions of tactoids in each class calculated from TEM images of SEBS_MA_2.5% extruded nanocomposite taken at 50 K

3. SAXS analysis

To evaluate the alignment of nanoclay and PS domains, we considered two types of characterization:

- TEM: probes a small area but characterizes at the same time the state of exfoliation and the location of nanoclays.
- SAXS: probes a larger area than TEM and characterizes both the alignment of PS domain and nanoclay.

TEM images of extruded nanocomposites containing different loadings of nanoclay are reported in Figure A I-4(a)-(d) of the manuscript. They show evidence of nanoclay alignment except Figure 1(a) corresponding to the intercalated sample which is not as conclusive.

The SAXS analysis is reported in this annex as follows: Figure A I-4(a) presents the diffraction pattern of pure extruded SEBS while Figure A I-4(b) shows the diffraction pattern of extruded SEBS-7.5wt% clay nanocomposite and Figure A I-4(c) shows the diffraction pattern of extruded SEBS-MA-7.5wt% clay nanocomposite. Figure A I-4(d) shows the diffraction pattern of isotropic SEBS-5wt% clay nanocomposite prepared by solution for comparison. Similar results were also obtained with different clay concentrations and reported in (Carastan et al., 2014). It is worth mentioning that the incident X-Ray beam was oriented perpendicular to the extrusion direction (parallel to Y direction indicated in Figure A I-5).



Figure A I-4: Two dimensional SAXS patterns of (a) extruded pure SEBS,
(b) extruded SEBS-7.5wt% clay nanocomposite, (c) extruded SEBS-MA-7.5wt% clay nanocomposite and (d) SEBS-5wt% clay nanocomposite prepared by solution



Figure A I-5: X: direction of extrusion and Y: direction of the incident X-Ray beam

- The scattering patterns corresponding to the beam diffraction along the Y direction, shown in Figure A I-5(a), reveal a set of Bragg peaks corresponding to long order arrangement of PS cylinders in a two dimensional hexagonal packing and their alignment in a direction perpendicular to the Y direction (which is the direction of extrusion X as demonstrated in reference (Carastan et al., 2014)).
- In addition to the scattering from aligned PS domains, two strong diffuse spots corresponding to Bragg peaks associated with scattering from clay particles are observed in the SAXS patterns of extruded SEBS-7.5wt% and SEBS-MA-7.5wt% clay nanocomposites presented in Figure A I-4(b) and Figure A I-4(c). These signals indicate the alignment of nanoclay in the same direction of PS cylinders. The presence of maxima in the Figure A I-4(b) is characteristic of intercalated nanoclay while the absence of maxima in Figure A I-4(c) indicates scattering from exfoliated nanoclay.
- The scattering patterns in Figure A I-4(d) which correspond to SEBS-5wt% clay nanocomposite prepared by solution casting reveal concentric rings indicating that both PS domains and nanoclays are randomly dispersed in samples prepared by solution casting.

So far, these SAXS measurements clearly show the alignment of both PS domains and nanoclay in our extruded samples despite the fact that the TEM image presented in Figure

2-1(a) of the manuscript could have been misleading since it shows some randomly dispersed clay tactoids. In fact, one should keep in mind that TEM images only probe a small area whereas SAXS present a more global analysis of the sample.

ANNEX II

SUPPORTING ELECTRONIC INFORMATION FOR ARTICLE 2

Interfacial molecular dynamics of styrenic block copolymer-based nanocomposites with controlled spatial distribution

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1. 2D SAXS plots corresponding to SEBS-13 based nanocomposites prepared by solvent casting

Figure A II-1 shows diffraction patterns of SEBS-13 and SEBS-20 block copolymers and their corresponding nanocomposites SEBS-13-20A and SEBS-20-20A, prepared by solvent casting. Both samples exhibit isotropic rings, in the 3 directions (only the Z direction is presented as all the 3 directions were identical), characteristic of an isotropic morphology.



Figure A II-1. 2D SAXS diffraction patterns of: (a) SEBS-20-solution,
(b) SEBS-20-20A-solution nanocomposite, (c) SEBS-13-solution and
(d) SEBS-13-20A-solution nanocomposite (patterns corresponding to the Z direction, X and Y directions have similar features)

2. Scattering intensity as function of scattering vector q corresponding to SEBS-20film blowing R1 and SEBS-20- film blowing R3 and their nanocomposites, prepared by film blowing extrusion



Figure A II-2. Scattering intensity as function of scattering vector q at different temperatures of: (a) SEBS-20- film blowing R1 and (b) SEBS-20-20A- film blowing R1 nanocomposite



Figure A II-3. Scattering intensity as function of scattering vector q at different temperatures of: (a) SEBS-20-film blowing R3 and (b) SEBS-20-20A- film blowing R3 nanocomposite



3. SAXS data corresponding to SEBS-13-extrusion and its nanocomposite SEBS-13-20A-extrusion, prepared by sheet die extrusion

Figure A II-4. (a)-(b) 2D SAXS patterns of: (a) SEBS-13-extrusion (Z direction) and (b) SEBS-13-20A-extrusion nanocomposite, (c)-(d) Scattering intensity as function of scattering vector q at different temperatures of: (c) SEBS-13-extrusion and (d) SEB-13-20A-extrusion nanocomposite. Peak positions follow the $1:\sqrt{3}:\sqrt{7}$ characteristic of hexagonal morphology

4. TEM micrographs of SEBS-30-20A-solution nanocomposite

TEM images of SEBS-30-20A-solution nanocomposite without and with staining are reported in Figure A II-5 for the sake of comparison with samples exhibiting preferential alignment in the direction of extrusion such as SEBS-20-20A-R1 (TEM images in Figure 3-3(c)-(e) of the manuscript). Comparison between the 2 figures shows that, indeed in Figure A II-5, clay tactoids and PS cylinders are randomly distributed and isotropic within the matrix (In Figure A II-5(d), clay particles are indicated by the arrows) whereas most of the clay particles are oriented in one direction in Figure 3-3 of the manuscript.



Figure A II-5. TEM of SEBS-30-20A-isotropic, (a) and (b) unstained, (c) and (d) samples stained for 30 min with RuO₄ (dark phase corresponds to PS domains)

5. TEM micrographs of SEBS-30-MA-20A nanocomposites prepared by sheet die extrusion

TEM micrographs reported in Figure A II-6 show the presence of mostly exfoliated and aligned clay particles in nanocomposites prepared by sheet die extrusion using SEBS-30-MA as a matrix.



Figure A II-6. TEM micrographs of SEBS-30-MA-20A nanocomposites prepared by sheet die extrusion with different clay loadings: (a) 2.5wt% and (b) 7.5wt%

6. Complex dielectric permittivity and fitting equations

6.1.Complex dielectric permittivity

$$\hat{\varepsilon}(\omega) = \varepsilon'(\omega) - i\varepsilon''(\omega)$$
 (A II.1)

Where:

ω: angular frequency;

 $\hat{\varepsilon}$: complex dielectric permittivity;

 ε ': real part of the complex dielectric permittivity, which is related to the stored energy. It is also known as the dielectric constant;

 ε '': imaginary part of dielectric permittivity, which is related to the dissipated energy. It is also known as the dielectric loss.

6.2.Fitting equation

$$\hat{\varepsilon}(\omega) = b(i\omega)^{n-1} + \sum_{k=1}^{N} \left[\frac{\Delta \varepsilon_k}{(1+(i\omega\tau_k)^{\alpha_k})^{\beta_k}} + \varepsilon_{\infty k} \right]$$
(A II.2)
Power law term Havriliak-Negami (HN) term

Where:

 ω : angular frequency;

n: exponential factor, between 0 and 1 characterizing the nature of the charge hopping process (the case n=0 corresponds to pure electronic conduction with $b=\sigma_0/\epsilon_0$);

 τ_k : relaxation time related to relaxation process k;

 $\Delta \varepsilon_k$: difference between real permittivity values at very low and infinite frequencies. It is proportional to the area below the relaxation peak in ε " and known also as the dielectric strength;

 $\varepsilon_{\infty k}$: real permittivity at much higher frequencies than the relaxation frequency of the relaxation process k;

 α_k : width parameter characteristic of the slope at the low frequency side of the relaxation peak;

 β_k : asymmetry parameter. $-\alpha_k\beta_k$ gives the slope at the high frequency side of the relaxation peak;

N: number of total observed relaxations in the studied range of frequencies and temperatures.

6.3.Fitting procedure

Commercially available software was used to obtain the curve-resolved spectra. Both real and imaginary parts of the dielectric permittivity were considered for the fitting, computed based on nonlinear procedures. The mean square deviation (MSD) was provided automatically by the software at each fitting step, as criteria for the goodness of fit. It gives the mean value of quadratic deviations between observed and calculated ε '' values. The objective is to find a combination of fitting parameters ensuring a minimum MSD value. For all the fits performed in this study, a fit is considered stable and accepted if the MSD is relatively stable to 10⁻⁴ over 3 iterations. Besides, all the MSD values were lower than 0.05.

6.4. Relaxation time corresponding to the peak maximum frequency

$$\tau_{max} = \tau_{HN} * \left[\frac{\sin(\frac{\pi\alpha\beta}{2(\beta+1)})}{\sin(\frac{\pi\alpha}{2(\beta+1)})} \right]^{\frac{1}{\alpha}}$$
(A II.3)

Where:

 τ_{max} : relaxation time corresponding to the peak maximum frequency;

 τ_{HN} : relaxation time obtained from the HN fit (equation (A II.2));

 α : width parameter obtained from the HN fit (equation (A II.2));

 β : asymmetry parameter obtained from the HN fit (equation (A II.2)).

6.5.Vogel Fulcher Tammann (VFT) equation

$$\tau(T) = \tau_0 \exp(\frac{B}{T - T_0}) \tag{A II.4}$$

Where:

B: constant related to the apparent activation energy and cooperativity of underlying molecular motions;

 τ_0 : relaxation time at infinite temperature;

T₀: Vogel-Fulcher temperature, which is 30-70 K below the T_g in case of the main segmental relaxation α .

6.6.Fragility index

$$F = \frac{\partial \log_{10}(\tau)}{\partial (T_g/T)} | (T = T_g)$$
(A II.5)

Where:

F: fragility index;

 τ : the relaxation time expressed according to VFT equation (A II.4) (Considering the VFT equation for the relaxation time, the derivative could be calculated for each sample);

T_g: glass transition temperature.

6.7.Imaginary part of complex dielectric permittivity corresponding to SEBS-13 nanocomposites prepared by extrusion and solvent casting

Figure A II-7 shows dielectric loss spectra corresponding to pure SEBS-13 and SEBS-13-20A nanocomposites prepared by sheet die extrusion (clay oriented) and solvent casting (isotropic). In particular, remarkably higher losses are observed in the isotropic sample. Dielectric relaxation peaks could be observed in both samples and exhibit dependence on the morphology and orientation as reported in the manuscript.



Figure A II-7. Imaginary part of the dielectric permittivity as function of temperature of: (a) neat SEBS-13-partially oriented and (b) SEBS-13-20A-partially oriented (sheet die extrusion) and (c) SEBS-13-20A-isotropic (solvent casting)

6.8.Real part of complex dielectric permittivity corresponding to SEBS-13 nanocomposites prepared by sheet die extrusion and solvent casting

Figure A II-8 shows typical behaviors of the real part of dielectric permittivity as functions of frequency and temperature, of respectively neat SEBS, partially oriented nanocomposite and isotropic nanocomposite. In general, a relaxation peak in the dielectric loss spectrum leads to a step-like increase in the real part of permittivity since their frequency dependence is mathematically linked through the Kramers-Kronig relations.

The real permittivity of the neat exhibits no dependence on frequency and decreases slightly with increasing temperature due to the decrease of density. The real permittivity of the partially oriented nanocomposite exhibits slight dependence on frequency and increases with increasing temperatures especially at low frequencies, due to most likely Maxwell-Wagner-Sillars (MWS) interfacial polarization. The real permittivity of the isotropic sample exhibits stronger dependence on frequency and temperature. Step-like increases corresponding to relaxation phenomena are more obvious in this sample compared to the partially oriented sample.



Figure A II-8. Real part of the dielectric permittivity as functions of frequency and temperature: (a) neat SEBS-13- partially oriented and (b) SEBS-13-20A- partially oriented (sheet die extrusion) and (c) SEBS-13-20A-isotropic (solvent casting)

6.9. Dielectric permittivity at extended range of temperature

Dielectric measurements were carried out in an extended temperature range [-100°C, 150°C] for both SEBS-30-oriented and SEBS-30-20A-oriented in order to check the main glass transition of PEB phase and to correlate with DMA results. 3D maps of the imaginary part of the dielectric permittivity as functions of frequency and temperature are reported in Figure A II-9 while its evolution as a function of temperature and at a fixed frequency equal to 1 Hz are reported in Figure A II-10. The data show clearly the presence of a low temperature peak related to the glass transition of the PEB elastomer phase (α_{PEB}). At 1 Hz, the peak maximum occurs at -45°C. The peak attributed to the segmental motion of interfacial rubber chains (α_{Int}) is observed at intermediate temperatures, below the glass transition of PS phase. At 1

Hz, this peak maximum is around 40°C. Besides, an additional peak is observed towards higher temperatures (α '). This peak is more obvious at increased frequencies (Figure A II-10, at 1 kHz). Since this peak is observed only in the spectrum of the nanocomposite, it is believed to correspond to MWS interfacial polarization as discussed in the manuscript. At low frequencies and high temperatures, a remarkable increase of dielectric loss is observed in the spectra of both: the pure and the nanocomposite. This phenomenon is due to electrode polarization (α_{EP}) at the interface between the sample and the electrode.

It is worth noting that in this studied range of temperature, it was not possible to observe clearly the peak related to the segmental relaxation of PS block. However, the low-temperature wing of this peak might contribute to the observed spectra.



Figure A II-9. Dielectric spectroscopy maps as function of frequencies and temperatures of: (a) SEBS-30-oriented and (b) SEBS-30-20A-oriented



Figure A II-10. Imaginary part of the dielectric permittivity of SEBS-30-oriented vs. SEBS-30-20A-oriented as function of temperature at 1 Hz

7. Fitting of Dynamic Mechanical Analysis (DMA) results

7.1. Three parameter asymmetric double sigmoid (ADS) equation

$$P_{i}(T) = A(1 + \exp\left(\frac{T-\mu}{\sigma_{1}}\right))^{-1} \left[1 - \left(1 + \exp\left(\frac{T-\mu}{\sigma_{2}}\right)\right)^{-1}\right]$$
(A II.6)

Where:

P_i= peak I;

T: temperature;

A: height of the peak;

 μ , σ_1 , σ_2 : three mathematical parameters. When $\sigma_1=\sigma_2$, the function is symmetric about μ .

7.2. Estimation of the interfacial thickness

The fraction of bound polymer chains participating in the interfacial glass transition, F_{Bound} , could be determined by dividing the area under the interfacial Tgi peak by the sum of the areas of all Tg peaks, as it could be seen in equation (A II.7).

$$F_{Bound} = A_i / (A_{PS} + A_{PEB} + A_{Int})$$
(A II.7)

Where:

A_i: area under the peak i.

Subsequently, the total number of clay particles per nanocomposite volume unit, N, can be evaluated by equation (A II.8).

$$N = V_{Clay total} / V_{Clay particle}$$
(A II.8)

Where:

 $V_{Clay total}$: fraction of clay particles in a volume unit of the composite which is equal to 0.0214 (5wt% clay is equivalent to 2.14vol% in SEBS-30 considering that the density of Cloisite 20A is equal to 1.77 g/cm³);

V_{Clay particle}: the volume of one clay particle that could be determined knowing the average size of the tactoids.

Moreover, the fraction of polymer chains in a volume unit of the composite is given by equation (A II.9).

$$V_{Polymer} = 1 - V_{Clay total} = 1 - 0.0214$$
 (A II.9)

Where:

 $V_{Polymer}$: volume of polymer chains per volume unit (only the volume of clay particles is excluded).

The volume of chains with restricted mobility that participate in the interfacial glass T_{gi} is obtained either by multiplying the bound fraction obtained by the volume of polymer chains

in a volume unit or by multiplying the number of particles by the volume of the interfacial layer surrounding each particle according to equation (A II.10).

$$V_{Bound} = V_{Polymer} * F_{Bound} = (V_{clay particle+t} - V_{clay particle}) * N$$
(A II.10)

Where:

V_{bound}: volume of bound polymer chains in a volume unit;

V_{Clay particle+ t}: volume of one clay particle and interfacial layer surrounding it;

t: thickness of interfacial layer.

Finally, the thickness of the interfacial layer, t, was deduced from equation (A II.10).

ANNEX III

SUPPORTING ELECTRONIC INFORMATION FOR ARTICLE 3

Thermoplastic elastomer nanocomposites with controlled nanoparticles dispersion for HV insulation: correlation between rheological, thermal, electrical and dielectric properties

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1. Particle size analysis and size distribution of ZnO nanoparticles into SEBS-MA-5 nanocomposite

A TEM image of low magnification was selected to investigate the size distribution of ZnO nanoparticles/small agglomerations as dispersed in the nanocomposite. Initially, several processing steps were performed with ImageJ software to isolate the particles. Then, the particle size analysis was performed automatically as indicated in Figure A III-1. In particular, 504 particles were considered for the analysis and the particle size was determined based on Feret's diameter (given directly by the software). Based on the analysis, ZnO particles/agglomerations were classified in 12 classes of dimensions ranging from 0 to 220 nm as reported in Figure A III-2. This size distribution reveals that more than 60 % of ZnO particles/agglomerations, as dispersed in SEBS-MA, have sizes between 20 and 60 nm. This fact confirms the achievement of nanoscale dispersion.


Figure A III-1. Steps of particle size analysis: (a) TEM image considered for the analysis (related to SEBS-MA-5 nanocomposite, (b) cropped image, (c) particle domains isolated, (d) analyzed domains



Figure A III-2. Size distribution of ZnO nanoparticles/agglomerations in SEBS-MA-5 nanocomposite based on particle size analysis

2. TEM images corresponding to SEBS-MA-5 nanocomposite (sample stained with RuO₄)

The TEM images corresponding to SEBS-MA-5 stained sample, reported in Figure A III-3, indicate the dominance of cylindrical morphology. Unfortunately, the location of nanoparticles is not clear due to a potential undesirable interaction between RuO₄ with the zinc element. Many black spots, of size considerably larger than the average nanoparticles size, were observed throughout the treated sample in TEM micrographs. They correspond to the regions where ZnO reacted with the staining agent.



Figure A III-3. TEM images corresponding to SEBS-MA-5 nanocomposite after staining with RuO₄ (black spots correspond to regions where ZnO reacted with the staining agent)

3. Imaginary permittivity as functions of frequency and temperature of SEBS-1 and SEBS-MA-1 nanocomposites

Dielectric loss spectra corresponding to SEBS-1 and SEBS-MA-1 nanocomposites are reported respectively in Figure A III-4(a) and Figure A III-4(b). The measurements were performed at different temperatures ranging from 25 to 105 °C.



Figure A III-4. Imaginary permittivity, ε'', as functions of frequency and temperature corresponding to: (a) SEBS-1 and (b) SEBS-MA-1 nanocomposites

4. Effect of drying on dielectric response of SEBS-MA-5 nanocomposite

Dielectric loss spectra corresponding to SEBS-MA-5 nanocomposite before and after drying are reported respectively in Figure A III-5(a) and Figure A III-5(b). The measurements were performed at different temperatures ranging from 20 to 110 °C. Only a marginal effect could be observed.



Figure A III-5. ε'' of SEBS-MA-5 nanocomposite: (a) before drying and (b) after drying for 3 days at 65°C under vacuum

5. Effect of drying on the dielectric response of SEBS-MA-20 nanocomposite

Dielectric loss spectra corresponding to SEBS-MA-20 nanocomposite before and after drying are reported respectively in Figure A III-6(a) and Figure A III-6(b). The measurements were performed at different temperatures ranging from 20 to 110 °C. Only a small effect could be observed at low temperatures (Figure A III-6(c)).



Figure A III-6. ε '' of SEBS-MA-20 nanocomposite: (a) before drying, (b) after drying for 3 days at 65°C under vacuum and (c) comparison at selected temperatures

6. Effect of drying on the dielectric response of SEBS-20 nanocomposite

Dielectric loss spectra corresponding to SEBS-20 nanocomposite before and after drying are reported respectively in Figure A III-7(a) and Figure A III-7(b). The measurements were performed at different temperatures ranging from 20 to 110 °C. Several changes were obseved, mainly the increase of dielectric losses at low frequencies after drying (Figure A III-7(c)).



Figure A III-7. ε'' of SEBS-20 nanocomposite: (a) before drying, (b) after drying for 3 days at 65°C under vacuum and (c) comparison at selected temperatures

7. Mapping of eroded area

Mappings of the eroded area corresponding to neat materials and nanocomposites containing 1wt% ZnO are reported in Figure A III-8. To evaluate the performance of SEBS-MA-ZnO compared to SEBS-ZnO in terms of resistance to erosion, the erosion depth (estimated from the mapping) might be considered as a quantitative parameter. For instance, the maximum depth of erosion in neat SEBS is around 177 μ m (Figure A III-8(a)) while it is reduced to 128 μ m in SEBS-1 nanocomposite (Figure A III-8(c)). In unfilled SEBS-MA, the estimated erosion depth is equal to 145 μ m (Figure A III-8(b)). It was reduced to only 38 μ m in SEBS-MA-1 nanocomposite (Figure A III-8(d)).



Figure A III-8. Mapping of eroded area in different SEBS-ZnO vs. SEBS-MA-ZnO samples: (a) SEBS, (b) SEBS-MA, (c) SEBS-1 and (d) SEBS-MA-1

APPENDIX

VITA

EDUCATION

- Doctor of Philosophy École de Technologie Supérieure (ÉTS). Supervisor: Prof. Nicole R. Demarquette, Co-supervisors: Prof. Eric David and Prof: Michel Fréchette. From February 2013 to June 2017. Montréal, QC – Canada.
- Master of Science in Mechanical Engineering École Polytechnique de Tunisie.
 Supervisors: Prof. Moez Chafra and Prof. Zoubeida Ounaies. From September 2009 to December 2011. Tunis – Tunisie.
- Bachelor of Science in Engineering École Polytechnique de Tunisie. From September 2007 to September 2010. Tunis – Tunisie.

AWARDS AND SCHOLARSHIPS

- 2016: Sommet de Montréal sur l'innovation SMI 2016 Poster selected for oral presentation.
- 2016: American Society of Materials ASM 1st poster prize.
- 2016: École de Technologie Supérieure Bourse interne
- 2014&2015: École de Technologie Supérieure Conference travel awards
- 2015: Society of Plastics Engineers SPE Conference travel award
- 2010&2011: International Institute for Multifunctional Materials for Energy Conversion IIMEC- scholarships for two six months internships in USA universities within the context of collaboration between USA and Tunisian researchers.
- 2007-2010: Tunisian Ministry of Higher Education- monthly scholarship awarded to engineering students admitted at Tunisia Polytechnic School.

JOURNAL PUBLICATIONS

• E. Helal, E. David, M. Fréchette and N.R. Demarquette, Thermoplastic elastomer nanocomposites with controlled nanoparticles dispersion for HV insulation:

correlation between rheological, thermal, electrical and dielectric properties, *European Polymer Journal*, Volume 94, September 2017, pp. 68-86.

- E. Helal, L.G. Amurin, D.J. Carastan, R.R. de Sousa Jr., E. David, M. Fréchette and N.R. Demarquette, Interfacial molecular dynamics of styrenic block copolymer-based nanocomposites with controlled spatial distribution, *Polymer*, Volume 113, 24 March 2017, Pages 9–26.
- E. Helal, N.R. Demarquette, L.G. Amurin, E. David, D.J. Carastan and M. Fréchette, Styrenic block copolymer-based nanocomposites: Implications of nanostructuration and nanofiller tailored dispersion on the dielectric properties, *Polymer*, Volume 64, May 2015, pp. 139–152.
- E. Helal, C. Pottier, E. David, M. Fréchette and N.R. Demarquette, Polyethylene/Thermoplastic elastomer/Zinc Oxide nanocomposites for High Voltage insulation applications: dielectric, mechanical and rheological behavior, submitted to *Polymer* on March 2017 (under review).
- E. Helal, L.G. Amurin, D.J. Carastan, R.R. de Sousa Jr., E. David, M. Fréchette and N.R. Demarquette, Tuning the mechanical and dielectric properties of clay-containing thermoplastic elastomer nanocomposites, submitted to *Polymer Engineering and Science* on May 2017 (under review).
- E. Helal, Z. Ounaies and A.B. Meddeb, Processing and characterisation of two- and three-phase polymer-based nanocomposites for energy storage applications, International Journal of Microstructure and Materials Properties, Volume7, 2012, pp.417 – 427.

CONFERENCE PRESENTATIONS

- E. Helal, C. Pottier, C.V. Bouanga, E. David, M. Fréchette and N.R. Demarquette, Effect of blending with styrenic thermoplastic elastomer on dielectric performance of polyethylene/ZnO and maleated polyethylene/ZnO nanocomposites, Abstract submitted for Nano 2016, August 7-12th, 2016, Quebec, Canada
- E. Helal, N. R. Demarquette, E. David and M. Fréchette, Evaluation of dielectric behavior of polyethylene/thermoplastic elastomer blends containing zinc oxide (ZnO)

nanoparticles for high voltage insulation, IEEE Electrical Insulation Conference (EIC), pp. 592-595, June 19-22nd, 2016.

- E. Helal, N.R. Demarquette, E. David and M. Fréchette, Dielectric properties of thermoplastic elastomer/zinc oxide (ZnO) nanocomposites with controlled nanoparticles dispersion, Conference on Electrical Insulation and Dielectric Phenomena (CEIDP), October 18th-21st, 2015, Ann Arbor MI, USA, pp.447-450.
- E. Helal, N.R. Demarquette, E. David and M. Fréchette, Polystyrene-b-Poly (Ethylene-co-Butylene)-b-Polystyrene/Zinc Oxide Block Copolymer Nanocomposites: Rheological and Dielectric Properties, Annual Technical Conference of the Society of Plastics Engineers (ANTEC), March 23-25th, 2015, Orlando FL, USA.
- E. Helal, N.R. Demarquette, E. David and M. Fréchette, Polyethylene/styrenic block copolymer blends: Morphology and dielectric properties, Conference on Electrical Insulation and Dielectric Phenomena (CEIDP), October 19th-22nd, Des Moines IA, USA, 2014, pp. 767-770.
- E. Helal, N.R. Demarquette, E. David, D.J. Carastan, L.G. Amurin and M.F. Fréchette, Effect of Clay-Containing SEBS Nanocomposites Morphology on Their Thermal, Mechanical, Rheological and Dielectric Properties, Annual Technical Conference of the Society of Plastics Engineers (ANTEC), April 28-30th, 2014, Las Vegas NV, USA.

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