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Development of high damping acrylic rubber/sliding graft copolymer composites†

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Aiming at fabricating high damping rubber composites, the acrylic rubber ACM was incorporated with sliding graft copolymer (SGC) materials. SGC is a novel supramolecular material with sliding crosslink junctions, and it acts as a high damping phase in ACM/SGC composites. Fourier transform infrared spectroscopy reveals the presence of two types of hydrogen bonds in ACM/SGC composites. Micro-structure analysis shows a clear sea-island phase structure. SGC particles disperse fairly uniformly in the ACM matrix. A wide interphase region exists between these two phases, indicating the good blend compatibility between ACM and SGC. The damping performance of ACM/SGC composites under dynamic shear strain and frequency condition significantly improved with the increase in SGC content. Specifically, the loss factor ($\tan \delta$) value of ACM/SGC (100/40) composite increased by 120% compared with that of neat ACM, according to the RPA results. The significantly improved damping property can be ascribed to the interfacial hydrogen bonds and the pulley effect of SGC molecules.

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1 Introduction

Damping material can absorb vibrations and noise by converting mechanical energy into heat, which is highly desirable for many applications, such as seismic isolation of buildings and transportation.¹ In addition, for seismic isolation application, damping properties in the frequency range of 0.5–7 Hz for large amplitudes (up to 200%) is most important. Polymer materials, such as rubber, are the most commonly used damping materials because of their excellent vibration absorption capability, which can be attributed to the unique viscoelastic properties of polymers.² However, the damping performance of polymer materials is mainly affected by its glass transition behavior. In general, the effective damping (loss factor, $\tan \delta > 0.3$) temperature range of a homo-polymer is approximately 20–30 °C near its glass transition temperature (T_g), but this range is unsuitable for most practical applications.^{3,4} Moreover, the $\tan \delta$ value of pristine polymer materials is usually insufficient.

Many methods, such as polymer blends,^{5–7} copolymers,⁸ and interpenetrating polymer networks (IPNs),^{1,9} have been used for damping modification to improve the damping capabilities of

polymer materials. Zhao *et al.* investigated the morphology and damping performance of natural rubber (NR)/nitrile rubber (NBR)/epoxidized natural rubber (ENR) ternary rubber blends. The effective damping temperature range of this ternary blends became broader than that of pristine NR, and damping performance of rubber blends at around room temperature range was effectively improved.¹⁰ Chern *et al.* examined the damping performance of IPNs of polyurethane-modified epoxy and polyurethane and found that the damping ability was enhanced by the formation of IPN structure.⁹ However, these modifications usually lead to a decrease in $\tan \delta$ or a reduction in mechanical properties. As new damping composites, polymer/small organic molecular hybrids have been widely explored.^{11–18} Wu prepared high damping polymer composites by introducing hindered phenol AO-80 to binary hybrids of chlorinated polyethylene and acrylic rubber (ACM).¹⁹ Zhao *et al.* prepared high damping rubber composites by adding hindered phenol AO-80 to a NBR matrix.²⁰ Consequently, the strength and damping performance of rubber composites were enhanced. However, these hybrid materials suffer from the instability of damping properties. This is because these organic phenol molecules tend to diffuse to the surface of rubber composites if the affinity between rubber molecules and phenol molecules is weak.²¹ Moreover, the damping property of rubber materials in dynamic shear strain and frequency ranges has been rarely investigated in these studies, although this property is essential for some applications, such as seismic isolation of buildings.

As a novel type of polyrotaxane (PR) derivative, sliding graft copolymer (SGC) has been widely investigated because of its specific properties.²² SGC molecules contain a polyethylene

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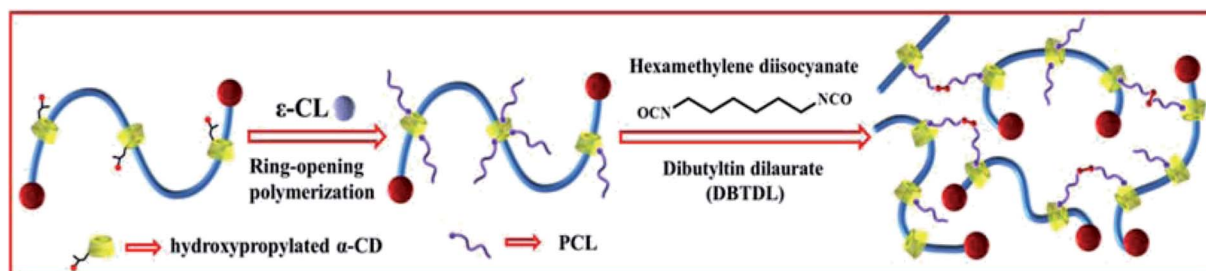


Fig. 1 Schematic of the preparation of SGC.

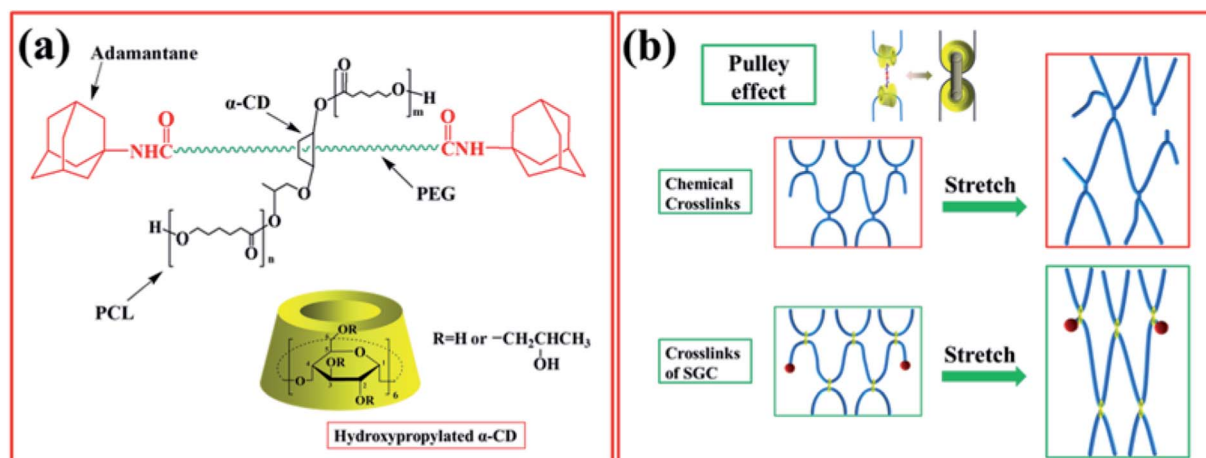


Fig. 2 (a) Chemical structure of SGC. (b) Schematic of the pulley effect.

glycol (PEG) backbone, onto which numerous α -cyclodextrins (α -CD) rings are threaded, and the end of the main chains is capped with bulky adamantane groups. The ring components of SGC are grafted by polycaprolactone (PCL) chains.^{23–29} The grafting reaction is achieved through the ring-open polymerization of ϵ -caprolactone, initiated by hydroxyl groups ($-\text{OH}$) of the hydroxypropylated α -CDs. SGC network is obtained by cross-linking the terminals of PCL chains with hexamethylene diisocyanate (HMDI).^{30–33} The preparation schematic of SGC is illustrated in Fig. 1. One of the key features of SGC materials is that the crosslink junctions of these polymer chains are topologically interlocked. Unlike traditional chemical or physical cross-links, these figure-of-eight cross-links are not fixed and can pass along the backbone chains freely, similar to a pulley, to equalize the tension of threading polymer chains. Thus, this feature is also called pulley effect [Fig. 2(b)].³⁴ The pulley effect feature of SGC materials has been confirmed through some characterization methods, such as X-ray scattering, small-angle quasi-elastic light scattering, and small-angle neutron scattering.^{34,35} SGC materials exhibit some unique properties, such as small stress relaxation,^{36,37} low compression set, and large $\tan \delta$ in a wide frequency range because of the pulley effect.³⁸ This novel SGC material has been applied to scratch-resistant coatings,³⁹ but few studies have focused on its application in damping materials.

Previously, our group introduced SGC to NR matrix as a high damping phase to prepare damping materials.³⁸ However, ENR

was used as a compatibilizer because of the poor affinity between SGC and NR molecules, thus complicating the manufacturing process. Thus, in our present study, we introduced SGC to the ACM rubber matrix to prepare high damping rubber composites. The obtained results suggest that SGC exhibits a good dispersion in ACM rubber and can form hydrogen bonds with the rubber matrix. The ACM/SGC composites have a significantly improved damping performance under dynamic shear strain and frequency condition. In this contribution, the microstructures and damping performance were systematically investigated and the relationship between properties with the unique structures has been revealed.

2 Experimental

2.1 Materials

Acrylic rubber (ACM, AR-801, TOPHE Industry Co., Ltd., Japan), used as a matrix in this study, was obtained through the polymerization of ethyl acrylate and composed of active chlorine as cure site groups. Its chemical structure and vulcanization mechanism are illustrated in Fig. 3. A sliding graft copolymer (SGC) with a molecular weight of 6 00 000 (Tianjin Weirui Supramolecular Materials Technology Co., Ltd., China) was also utilized. The received SGC material contains crosslink agent (HMDI). Other compounding ingredients (Sinopharm Chemical Reagent Co., Ltd., China) were used without further purification.



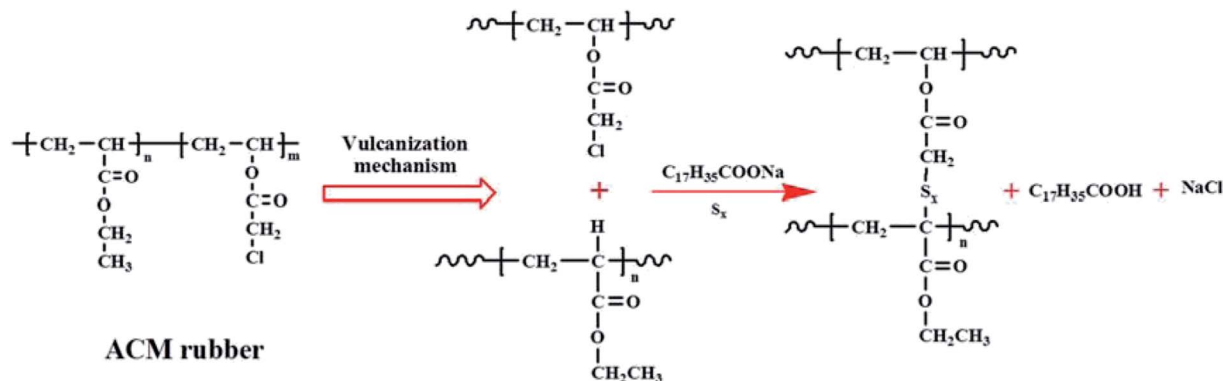


Fig. 3 Chemical structure and vulcanization mechanism of acrylic rubber.

Table 1 Compounding formulation of ACM/SGC composites

Ingredients	Content (phr ^a)
ACM	100
SGC	0/10/20/30/40
Stearic acid	1
Sodium stearate	4
Potassium stearate	1
Sulfur	1

^a Parts per hundred of rubber by mass.

2.2 Preparation of ACM/SGC composites

The formulation used to prepare the ACM/SGC composites is shown in Table 1.

The as-received SGC material was crosslinked in an oven at 170 °C for 3 h. During this process, the terminals of the PCL side chains were cross-linked by hexamethylene diisocyanate.

The ACM rubber was plasticated on a two-roll mill at room temperature and compounded with other ingredients except sulfur. The crosslinked SGC was mixed into the ACM blends on a two-roll mill at 120 °C, and then cooled to room temperature. Sulfur was mixed with the ACM/SGC blends at room temperature, and the blends were vulcanized by compression molding at 180 °C and 15 MPa. The ACM/SGC composites are denoted as ACM/SGC (100/*n*) in the subsequent discussion, where *n* indicates phr of SGC.

2.3 Characterization methods

A transmission electron microscope (TEM, The Tecnai G220 S-TWIN, FEI, Ltd., USA) was employed to observe the micro-morphology of ACM/SGC composites. All of the specimens were sectioned to ultra-thin thickness with a glass knife on an ultramicrotome (Leica EM UC7, Germany). An atomic force microscope (AFM Multi Mode 8 Bruker, Germany) was employed to analyze the surface morphology of the composites in Peak Force Quantitative Nano Mechanics (QNM) mode, and the samples were polished using a cryo-ultramicrotome (Leica EM UC7, Germany).

Differential scanning calorimetry (DSC) measurements were performed using a DSC 204F1 calorimeter (Mettler-Toledo). The

specimens were heated from −80 °C to 160 °C at a heating rate of 10 °C min^{−1} in a nitrogen atmosphere. Fourier transform infrared spectroscopy (FTIR) measurements were performed on a Bruker Tensor 27 FTIR spectrometer. The FTIR spectra of ACM/SGC composites in the wavenumber range from 400 cm^{−1} to 4000 cm^{−1} were acquired from film specimens with a thickness of about 2 mm in an attenuated total reflection (ATR) mode. Temperature-dependent FTIR measurements were performed using an FTIR spectrometer connected with heating equipment in a transmission mode. Samples were heated from 30 °C to 170 °C at a rate of 10 °C min^{−1}.

The damping performance of the ACM/SGC composites was determined using a rubber process analyzer (RPA 2000, Alpha Technologies, USA). Dynamic shear strain sweep was conducted by varying the possible shear strain ranges from 0.1% to 200% at a constant frequency of 1 Hz at 30 °C. Dynamic frequency sweep was performed from 0.5 Hz to 7 Hz at a constant shear strain of 100% at 30 °C. Dynamic mechanical measurements were conducted on a dynamic mechanical thermal analyzer (DMTA, VA3000, 01dB Co., Ltd., France). Temperature sweep was performed at a frequency of 10 Hz, a tensile strain of 0.3%, and a heating rate of 3 °C min^{−1}.

3 Results and discussion

3.1 Intermolecular interactions and microstructure characterizations of ACM/SGC composites

Hydrogen bonding is a type of electrostatic attraction between an electron-deficient hydrogen atom and a highly electronegative atom. This interaction has been proved to be capable to improve the damping performance of rubber composites.^{17,20} FTIR is an effective tool to investigate hydrogen bond interactions among functional groups. Fig. 4 shows the FTIR spectra of ACM/SGC composites.

Fig. 4(a) shows the infrared spectra of the ACM/SGC composites and pristine SGC in the stretching vibration region of carbonyl group (C=O). In the FTIR curve of SGC, there are two distinct broad peaks: the peak around 1700–1750 cm^{−1} and the peak around 1657 cm^{−1}. And the broad peak around 1700–1750 cm^{−1} contains one main peak and two shoulder peaks. In order to investigate these broad peaks in detail, peak-



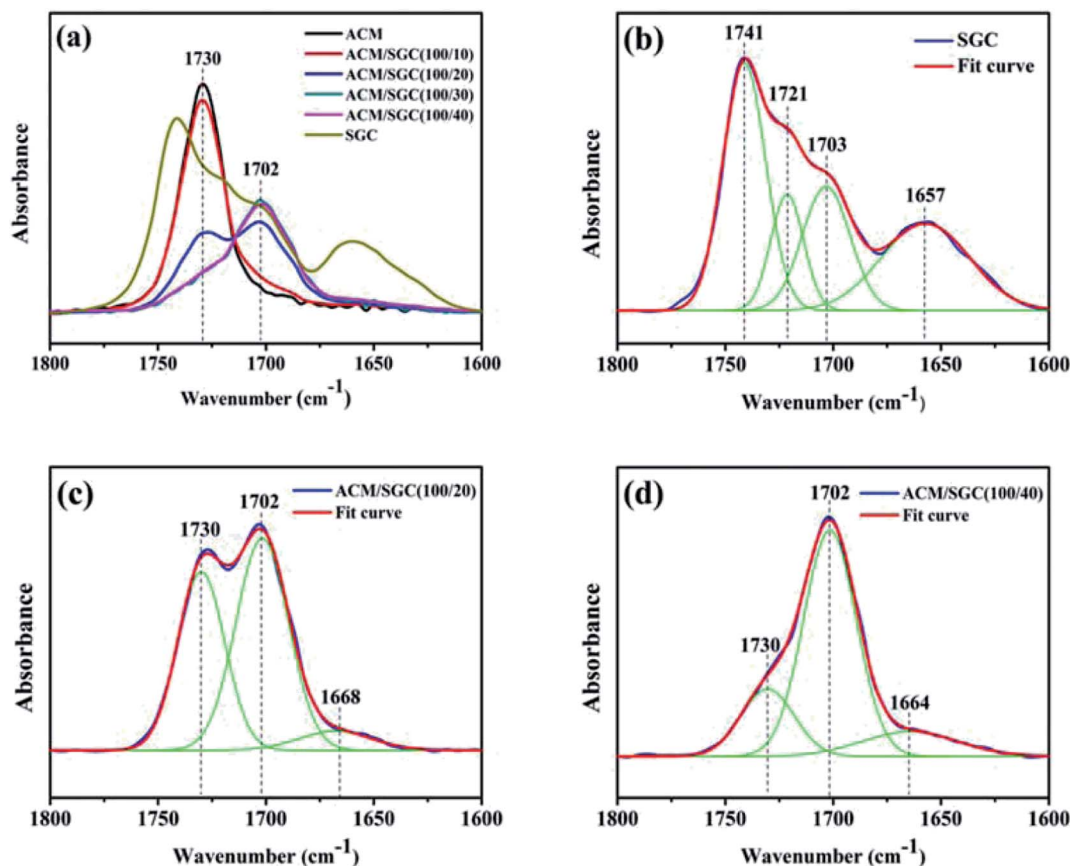


Fig. 4 FTIR spectra of ACM/SGC composites. (a) Infrared spectra of ACM/SGC composites in the carbonyl stretching vibration region. (b–d) Peak-fitting results of infrared spectra of pristine SGC, ACM/SGC (100/20) and ACM/SGC (100/40) composite in the carbonyl stretching vibration region.

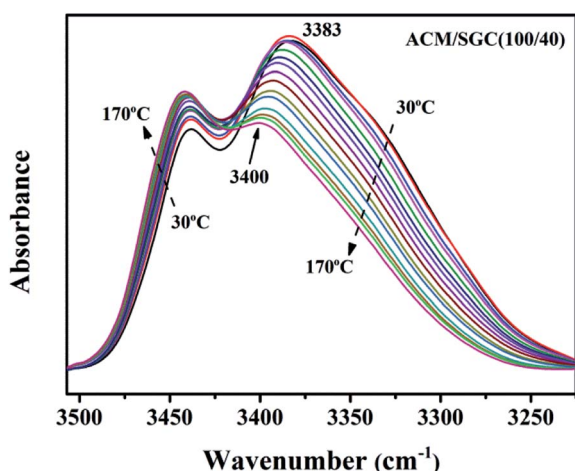


Fig. 5 The *in situ* diffuse reflectance FTIR spectra of ACM/SGC (100/40) composites under different temperatures (from 30 °C to 170 °C with a temperature interval of 10 °C).

fitting analysis was conducted. Fig. 4(b) shows the peak-fitting results of infrared spectrum of SGC. The peak at around 1741 cm⁻¹ corresponds to the stretching vibration of free C=O groups on the PCL side chain of SGC molecules, and the peak around 1721 cm⁻¹ is attributed to the stretching vibration of

carbamate carbonyl⁴⁰ (the crosslink junctions of SGC contain carbamate groups). The peak at around 1657 cm⁻¹ corresponds to the stretching vibration of amide carbonyl^{41,42} (which exists on the terminals of PEG main chains). The peak at around 1703 cm⁻¹ is attributed to the hydrogen-bonded C=O groups. The hydrogen bonds are formed between the C=O groups on the PCL and the hydroxyl groups (OH) in SGC molecules. Compared with the free C=O groups, the peak position of hydrogen-bonded C=O groups shifts from 1741 cm⁻¹ to 1703 cm⁻¹, which is called “red shift”. Because some OH groups on α -CDs were not grafted by PCL chains, they can still participate in the formation of hydrogen bonds. Hence, the OH groups participating in the formation of hydrogen bonds could come from the α -CDs or the terminal of PCL chains.

As shown in Fig. 4(a), there is one single peak at around 1730 cm⁻¹ in the FTIR curve of ACM, which corresponds to the stretching vibration of free C=O groups on the ACM side chains. After the addition of SGC, there emerged a shoulder peak at around 1702 cm⁻¹ in the curves of ACM/SGC composites. As there are no other peaks, such as those at 1741 cm⁻¹ and 1721 cm⁻¹, emerging in the fitted curves of ACM/SGC spectra, the peak at 1702 cm⁻¹ should not come from SGC itself. In fact, the peak at 1702 cm⁻¹ corresponds to the hydrogen-bonded C=O groups.⁴³ Hydrogen bonds are formed between the OH groups of SGC and the C=O groups of ACM, leading to the shift



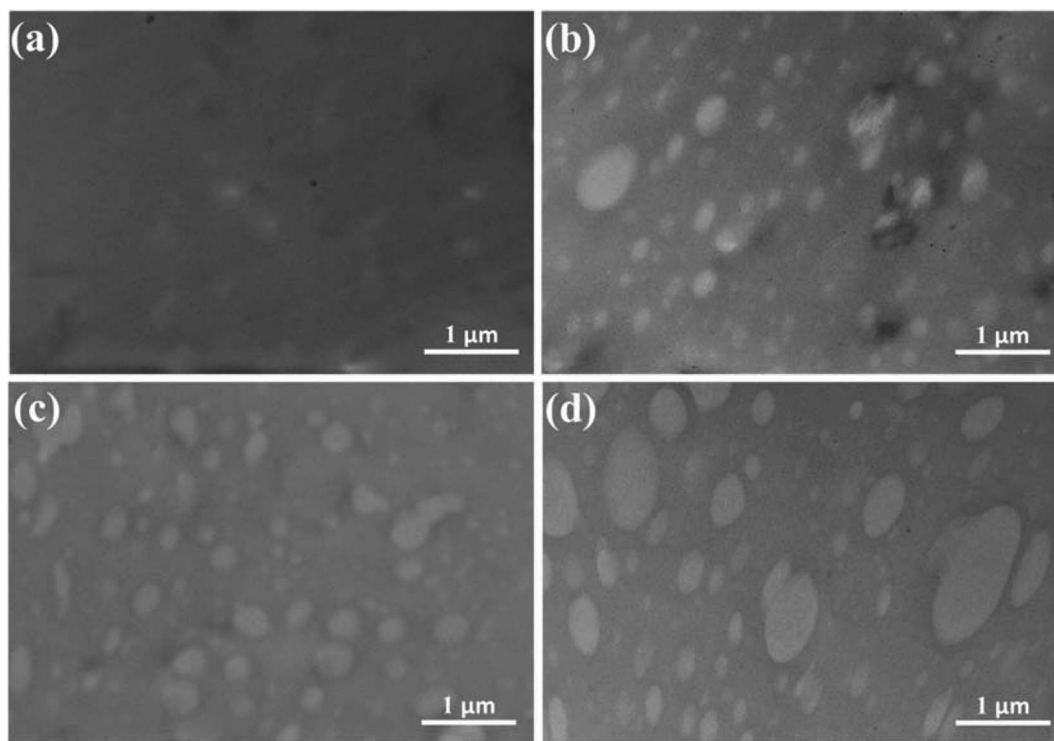


Fig. 6 TEM images of ACM/SGC composites. (a) ACM; (b) ACM/SGC (100/10); (c) ACM/SGC (100/20); (d) ACM/SGC (100/40).

in peak position from 1730 cm^{-1} to 1702 cm^{-1} . Fig. 4(c) and (d) illustrate the peak-fitting results of the infrared spectrum of ACM/SGC (100/20) and ACM/SGC (100/40), respectively. As SGC content increases, the peak intensity of the hydrogen-bonded C=O (1702 cm^{-1}) increases, while that of the free C=O (1730 cm^{-1}) decreases. This phenomenon reveals that the number of hydrogen bonds increases as SGC content increases.

In summary, two types of hydrogen bonds are found in ACM/SGC composites: (1) hydrogen bonds between the OH groups of SGC and the C=O groups of PCL chains and (2) hydrogen bonds between the OH groups of SGC and the C=O groups of ACM molecules.

Hydrogen bonds will gradually weaken as the temperature increases, or even dissociate at high temperature. In order to further prove the existence of hydrogen bonds in ACM/SGC composites, FTIR measurements at different temperatures were conducted. Fig. 5 shows the FTIR spectra of ACM/SGC (100/40) composites in the hydroxyl stretching vibration region at different temperatures. There are two distinct peaks observed at around $3200\text{--}3500\text{ cm}^{-1}$. The peak at around 3383 cm^{-1} can be attributed to the hydrogen-bonded OH groups, and the peak at around 3440 cm^{-1} can correspond to the free OH groups. As the temperature increases, the peak intensity of hydrogen-bonded OH groups decreases gradually, and the peak position shifts from 3383 cm^{-1} to 3400 cm^{-1} , which is called the “blue shift”. Moreover, the peak intensity of free OH groups increases gradually, and the peak position shows no distinct change. These results can be attributed to the dissociation of hydrogen bonds. As the temperature increases, hydrogen-bonded OH groups dissociate and more free OH

groups emerge. FTIR measurements under different temperatures prove the existence of hydrogen bonds successfully.

Fig. 6 shows the TEM images of ACM/SGC composites. Fig. 6(a) is the TEM image of pristine ACM rubber. Only a few dark particles are observed in the picture, which correspond to the compounding ingredients in the vulcanization system. In the other pictures, ACM/SGC composites present a distinct sea-island phase structure. The bright particles represent SGC phase and the dark phase corresponds to ACM rubber matrix. The SGC phase is uniformly dispersed in the ACM rubber matrix, and the average particle size of SGC is less than $1\text{ }\mu\text{m}$. This dual-phase morphology indicates that ACM and SGC are not fully miscible. Furthermore, the particle size of SGC is very small, which indicates the good blend compatibility between these two phases. However, the particle size of some SGC particles in ACM/SGC (100/40) composite (Fig. 6(d)) is larger than that of the others. This phenomenon can be attributed to the aggregation of SGC particles with the increase in SGC content. In addition, FTIR results confirmed the existence of hydrogen bond interactions between ACM and SGC, which would facilitate the homogeneous dispersion of SGC in ACM matrix. However, the interphase interaction region cannot be clearly distinguished in these TEM images. The PCL side chains of SGC can slide freely along the main chains because of the “pulley effect” and diffuse into the rubber matrix to form entanglement with ACM molecular chains. Therefore, a wide interphase interaction region is expected to form between the ACM and SGC phases. This structure can be further verified by the AFM results.

Fig. 7 illustrates a series of AFM Young's modulus maps of the polished surface of ACM/SGC composites. The color scale



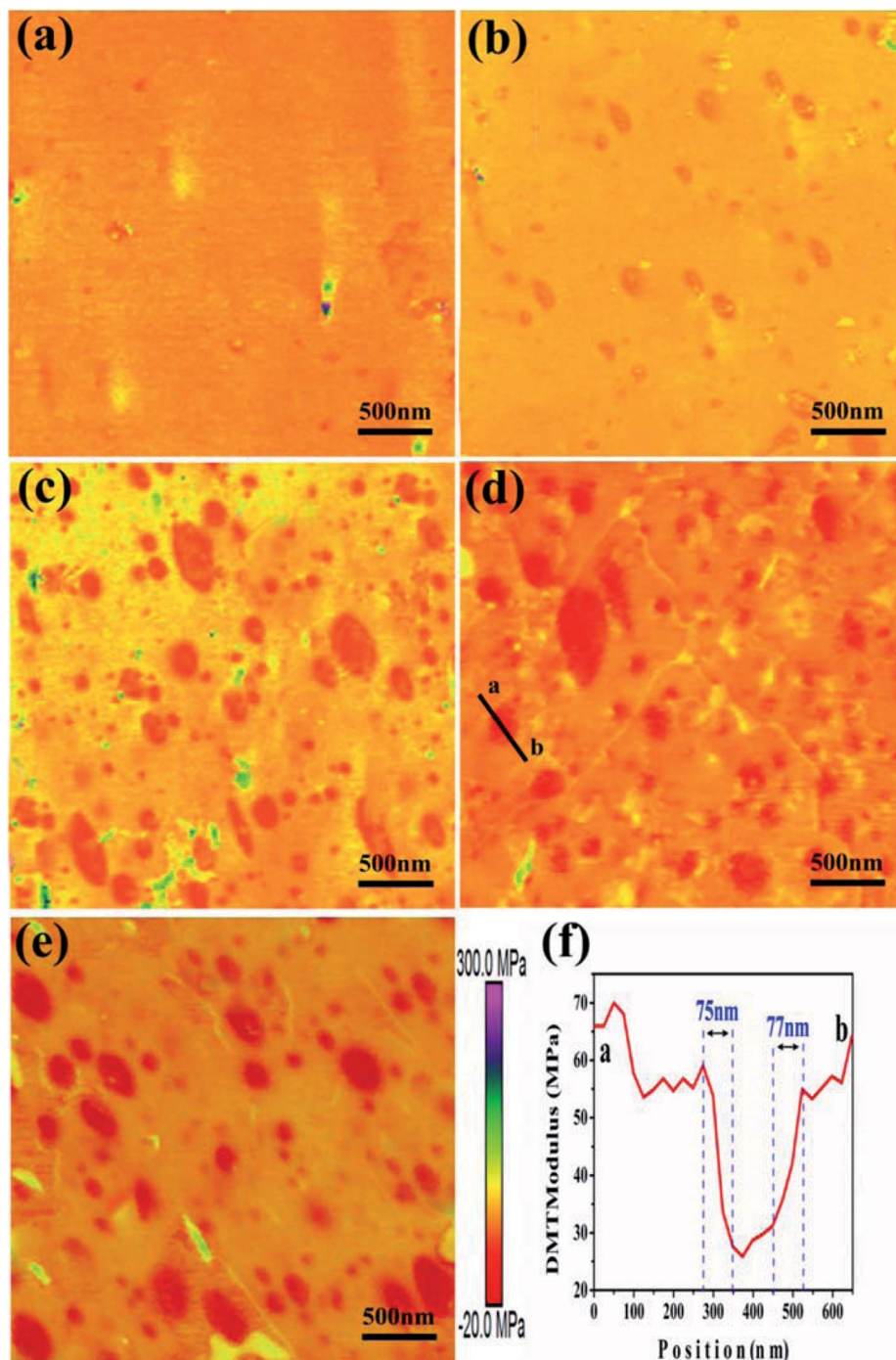


Fig. 7 AFM-Young's modulus maps of ACM/SGC composites. (a) ACM; (b) ACM/SGC (100/10); (c) ACM/SGC (100/20); (d) ACM/SGC (100/30); and (e) ACM/SGC (100/40). (f) Cross-section along the black line a–b in (d).

bar (from red to purple) corresponds to an increase in the AFM modulus from low to high values. Fig. 7(a) presents the AFM modulus map of ACM rubber. Only a few high-modulus particles, such as the green and purple colored particles, are dispersed in the rubber matrix. These particles correspond to the compounding ingredients, such as sodium stearate, which are aggregated on the polished surface of the ACM rubber sample. These high-modulus particles can also be observed in the other AFM images. Fig. 7(b)–(e) show the AFM modulus

maps of ACM/SGC composites. Some red particles with the lowest modulus values, which correspond to SGC phase, are homogeneously dispersed in the ACM rubber matrix to form a sea-island phase structure. As the SGC contents increase, the particle size of SGC phase increases but remains less than 500 nm. The curve in Fig. 7(f) indicates the variation of the rubber modulus along the black line in Fig. 7(d). As is shown in Fig. 7(f), the modulus of the ACM rubber matrix and the SGC particle are around 55–60 and 30 MPa, respectively. The region



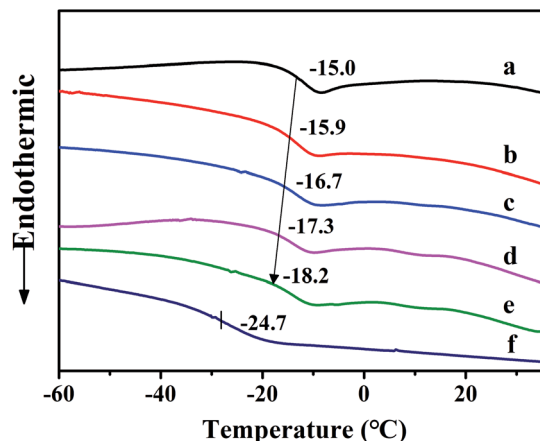


Fig. 8 DSC curves of ACM/SGC composites. (a) ACM; (b) ACM/SGC (100/10); (c) ACM/SGC (100/20); (d) ACM/SGC (100/30); (e) ACM/SGC (100/40); (f) SGC.

between these two phases is the interphase interaction region, whose width is almost half the diameter of the selected SGC particle. According to the TEM results, this finding can be attributed to the formation of the entanglement between PCL side chains of SGC molecules and ACM molecular chains. Therefore, a wide interphase interaction region exists between

ACM and SGC phases, indicating the good blend compatibility between these two phases.

Fig. 8 shows the DSC curves of ACM/SGC composites. Although the micro-morphology analysis reveals that the ACM/SGC composites form a phase separation structure, their T_g stacks together because of the good blend compatibility between these two phases. As a result, one distinct T_g is found in the DSC curves of ACM/SGC composites, and the glass transition region of these composites becomes much wider than that of pristine ACM. In Fig. 8, curve (a) represents the DSC curve of pristine ACM rubber, and T_g of ACM is about -15.0 °C. Curve (f) corresponds to the DSC curve of the cross-linked SGC material, and T_g of SGC is about -24.7 °C. This result suggests that the segments of SGC molecules possess higher mobility than those of ACM molecules. As a result, T_g of ACM/SGC composites shifts slightly to a lower temperature region as the SGC contents increase, compared with that of pristine ACM.

3.2 Damping properties of ACM/SGC composites

RPA measurements were conducted to investigate the damping performance of ACM/SGC composites at dynamic shear condition. Dynamic strain sweep was conducted by varying the possible shear strain ranges from 0.1% to 200% at a constant frequency of 1 Hz at 30 °C. Dynamic frequency sweep was carried out from 0.5 Hz to 7 Hz at a constant shear strain of

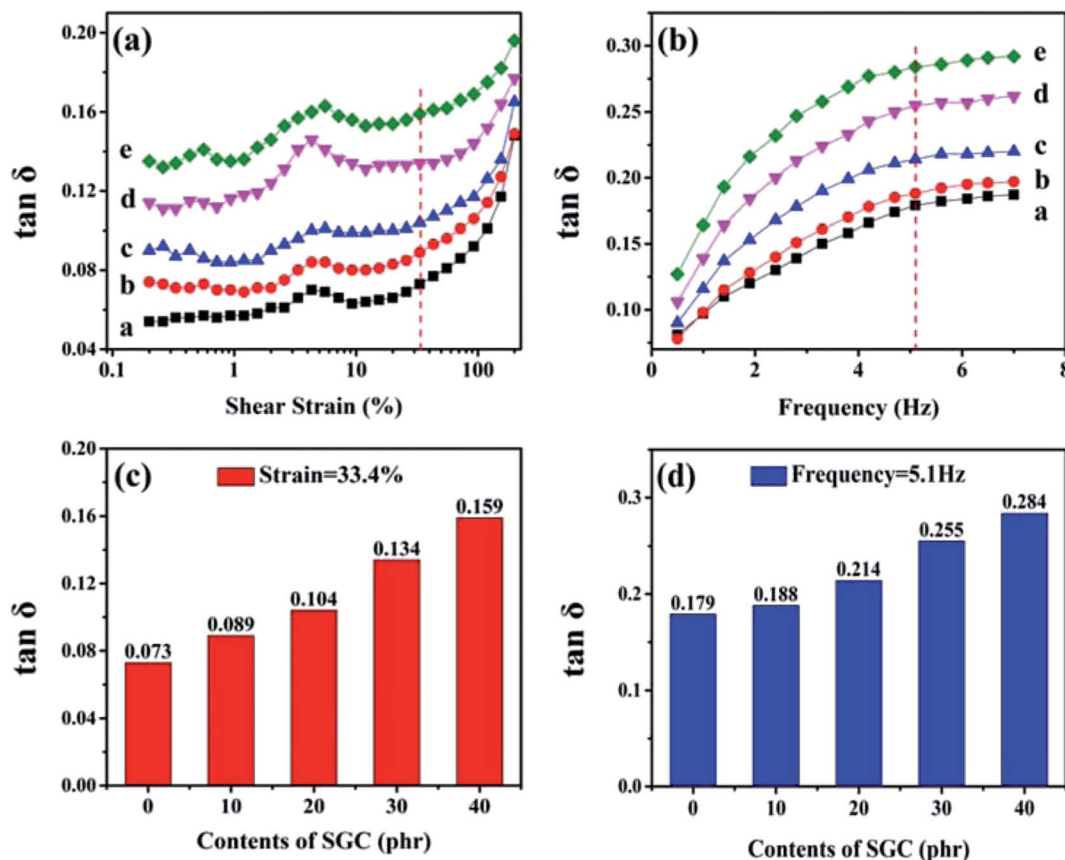


Fig. 9 RPA results of ACM/SGC composites: (a) variation of $\tan \delta$ under dynamic shear strain sweep; (b) variation of $\tan \delta$ under dynamic frequency sweep; (c) $\tan \delta$ value of rubber composites at the same shear strain (33.4%); (d) $\tan \delta$ value of rubber composites at the same frequency (5.1 Hz). (a) ACM; (b) ACM/SGC (100/10); (c) ACM/SGC (100/20); (d) ACM/SGC (100/30); (e) ACM/SGC (100/40).



100% at 30 °C. Fig. 9(a) shows the variation of $\tan \delta$ with dynamic shear strain sweep. The $\tan \delta$ values of ACM/SGC composites increase significantly with the increase in SGC content. Specifically, Fig. 9(c) shows the $\tan \delta$ values of rubber composites at the same strain (33.4%). The $\tan \delta$ value of pristine ACM rubber is about 0.073 and that of ACM/SGC (100/40) is about 0.159. The $\tan \delta$ value of ACM/SGC (100/40) is about 1.2 times higher than that of pristine ACM.

Fig. 9(b) illustrates the variation of $\tan \delta$ with dynamic frequency sweep. Similarly, $\tan \delta$ values of ACM/SGC composites significantly increase with the increase in SGC content. For example, Fig. 9(d) shows the $\tan \delta$ values of rubber composites at the same frequency (5.1 Hz). Pristine ACM rubber and ACM/SGC (100/40) composite yield $\tan \delta$ of about 0.179 and 0.284, respectively; thus, the value of the latter is about 1.6 times as high as that of the former. Overall, these results indicate the significantly improved damping property of ACM/SGC composites.

For seismic isolation rubber bearings, $\tan \delta$ values higher than 0.1 (equivalent damping ratio of 10%) in the 0.2–5 Hz frequency range and 10–200% strain range are required, according to the ISO standard.⁴⁴ As can be concluded from the RPA results, after adding 20 phr SGC, the damping property of rubber composites under strain sweep meets the requirements. The damping property of all the ACM/SGC composites under frequency sweep meets the requirements.

As mentioned previously, the PCL side chains and crosslink junctions of SGC materials can slide freely along the backbone chains because of the pulley effect. During this sliding process, the internal friction among SGC molecular chains increases, resulting in improved energy dissipation and high $\tan \delta$ value. Hence, SGC acts as a high damping phase in rubber composites. Moreover, as proved in AFM results, there is a wide interfacial region between ACM and SGC phases because of the interfacial hydrogen bonds and molecular chain entanglement. During deformation, the interfacial hydrogen bonds will continuously rupture and re-associate, and the chain entanglement will increase the internal friction. As a result, these

behaviors consume a lot of energy. Hence, the damping property of the ACM/SGC composites is significantly improved.

The damping property of ACM/SGC composites with the variation of temperature under tensile deformation was tested by DMA measurements. The results are illustrated in Fig. 10. Only one $\tan \delta$ peak is observed in DMA curves for ACM/SGC composites, which indicates the good blend compatibility between ACM and SGC. The maximum $\tan \delta$ values of the composites slightly decrease compared with that of pristine ACM rubber, but is still higher than 2.0. Moreover, the $\tan \delta$ value of the ACM/SGC composites above room temperature is significantly improved. This improvement of damping performance above room temperature is exactly needed for seismic isolation application.

Furthermore, Table S1† lists the improvement in $\tan \delta$ of different rubber damping materials reported in other literature reports. It can be concluded that ACM/SGC composites prepared in this study possess significantly improved $\tan \delta$ compared with that of other materials listed in the table.

4 Conclusion

ACM/SGC composites with significantly improved damping performance were successfully prepared in the present study. Two types of hydrogen bonds are found in ACM/SGC composites. The ACM/SGC composites exhibit a clear sea-island phase structure. SGC phase is homogeneously dispersed in the rubber matrix and form a wide interphase interaction region with the ACM phase. Damping performance of ACM/SGC composites is significantly improved as the SGC contents increase. The $\tan \delta$ value of ACM/SGC (100/40) composite is about 1.2 times higher than that of pristine ACM according to the RPA results. The increase in damping property can be attributed to the pulley effect of SGC and the interfacial hydrogen bonds.

Conflicts of interest

The authors declare no competing financial interests.

Acknowledgements

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References

- 1 T. Wang, S. Chen, Q. Wang and X. Pei, *Mater. Des.*, 2010, **31**, 3810–3815.
- 2 Q. Liu, M. Li, Y. Gu, S. Wang, Y. Zhang, Q. Li, L. Gao and Z. Zhang, *Carbon*, 2015, **86**, 46–53.
- 3 C. Qin, W. Cai, J. Cai, D. Tang, J. Zhang and M. Qin, *Mater. Chem. Phys.*, 2004, **85**, 402–409.
- 4 X. Shi, Q. Li and A. Zheng, *Polym. Test.*, 2014, **35**, 87–91.

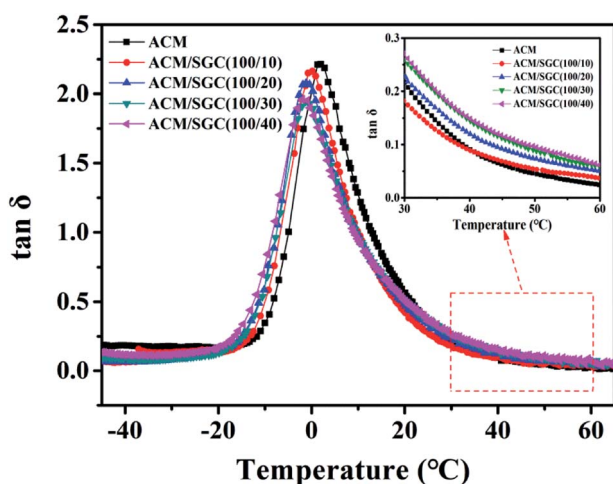


Fig. 10 Temperature dependence of $\tan \delta$ for ACM/SGC composites.



- 5 J. Wu, C. Li, Y. Wu, M. Leu and Y. Tsai, *Compos. Sci. Technol.*, 2010, **70**, 1258–1264.
- 6 L. Qu, G. Huang, J. Wu and Z. Tang, *J. Mater. Sci.*, 2007, **42**, 7256–7262.
- 7 C. Wu, C. Wei, W. Guo and C. Wu, *J. Appl. Polym. Sci.*, 2008, **109**, 2065–2070.
- 8 M. M. Mok, J. Kim and J. M. Torkelson, *J. Polym. Sci., Part B: Polym. Phys.*, 2008, **46**, 48–58.
- 9 Y. C. Chern, S. M. Tseng and K. H. Hsieh, *J. Appl. Polym. Sci.*, 1999, **74**, 328–335.
- 10 X. Zhao, K. Niu, Y. Xu, Z. Peng, L. Jia, D. Hui and L. Zhang, *Composites, Part B*, 2016, **107**, 106–112.
- 11 C. Wu, S. Akiyama, T. Mabuchi and K.-h. Nitta, *Polym. J.*, 2001, **33**, 792–798.
- 12 C. Wu, T.-A. Yamagishi, Y. Nakamoto, S. Ishida, K.-H. Nitta and S. Kubota, *J. Polym. Sci., Part B: Polym. Phys.*, 2000, **38**, 2285–2295.
- 13 P. Xiang, X. Zhao, D. Xiao, Y. Lu and L. Zhang, *J. Appl. Polym. Sci.*, 2008, **109**, 106–114.
- 14 D. Xiao, X. Zhao, Y. Feng, P. Xiang, L. Zhang and W. Wang, *J. Appl. Polym. Sci.*, 2010, **116**, 2143–2150.
- 15 X. Zhao, D. Xiao, S. Wu, Y. Feng, L. Zhang and W. Wang, *J. Appl. Polym. Sci.*, 2011, **120**, 906–913.
- 16 M. Song, X. Zhao, Y. Li, S. Hu, L. Zhang and S. Wu, *RSC Adv.*, 2014, **4**, 6719.
- 17 X. Zhao, G. Zhang, F. Lu, L. Zhang and S. Wu, *RSC Adv.*, 2016, **6**, 85994–86005.
- 18 D. Yang, X. Zhao, T. Chan, L. Zhang and S. Wu, *J. Mater. Sci.*, 2016, **51**, 5760–5774.
- 19 C. Wu, *J. Appl. Polym. Sci.*, 2000, **80**, 2468–2473.
- 20 X. Zhao, P. Xiang, M. Tian, H. Fong, R. Jin and L. Zhang, *Polymer*, 2007, **48**, 6056–6063.
- 21 Y. Gao, X. Wang, M. Liu, X. Xi, X. Zhang and D. Jia, *Mater. Des.*, 2014, **58**, 316–323.
- 22 J. Araki, *The 15th International Conference on Biomedical Engineering*. Springer, Cham, 2014, 43, pp. 20–23.
- 23 V. Nagarajan, A. K. Mohanty and M. Misra, *ACS Sustainable Chem. Eng.*, 2016, **4**, 2899–2916.
- 24 M. K. Thakur, R. K. Gupta and V. K. Thakur, *Carbohydr. Polym.*, 2014, **111**, 849–855.
- 25 V. K. Thakur, M. K. Thakur and R. K. Gupta, *Int. J. Biol. Macromol.*, 2013, **61**, 121–126.
- 26 V. K. Thakur, M. K. Thakur and R. K. Gupta, *Int. J. Biol. Macromol.*, 2013, **62**, 44–51.
- 27 V. K. Thakur, M. K. Thakur and R. K. Gupta, *Carbohydr. Polym.*, 2013, **97**, 18–25.
- 28 V. K. Thakur, M. K. Thakur and R. K. Gupta, *Carbohydr. Polym.*, 2013, **98**, 820–828.
- 29 V. K. Thakur, M. K. Thakur and R. K. Gupta, *Carbohydr. Polym.*, 2014, **104**, 87–93.
- 30 X. Li, H. Kang, J. Shen, L. Zhang, T. Nishi, K. Ito, C. Zhao and P. Coates, *Polymer*, 2014, **55**, 4313–4323.
- 31 D. Yang, F. Ge, M. Tian, N. Ning, L. Zhang, C. Zhao, K. Ito, T. Nishi, H. Wang and Y. Luan, *J. Mater. Chem. A*, 2015, **3**, 9468–9479.
- 32 X. Li, H. Kang, J. Shen, L. Zhang, T. Nishi and K. Ito, *Chin. J. Polym. Sci.*, 2015, **33**, 433–443.
- 33 J. Araki, T. Kataoka and K. Ito, *Soft Matter*, 2008, **4**, 245–249.
- 34 K. Ito, *Polym. J.*, 2007, **39**, 489–499.
- 35 J. Araki and K. Ito, *Soft Matter*, 2007, **3**, 1456.
- 36 N. Murata, A. Konda, K. Urayama, T. Takigawa, M. Kidowaki and K. Ito, *Macromolecules*, 2009, **42**, 8485–8491.
- 37 Y. Bitoh, N. Akuzawa, K. Urayama, T. Takigawa, M. Kidowaki and K. Ito, *Macromolecules*, 2011, **44**, 8661–8667.
- 38 W. Wang, D. Zhao, J. Yang, T. Nishi, K. Ito, X. Zhao and L. Zhang, *Sci. Rep.*, 2016, **6**, 22810.
- 39 Y. Noda, Y. Hayashi and K. Ito, *J. Appl. Polym. Sci.*, 2013, **131**, 9.
- 40 P. A. Edwards, G. Striemer and D. C. Webster, *Prog. Org. Coat.*, 2006, **57**, 128–139.
- 41 K. J. Howe, K. P. Ishida and M. M. Clark, *Desalination*, 2002, **147**, 251–255.
- 42 M. Zeng, Z. Fang and C. Xu, *J. Membr. Sci.*, 2004, **230**, 175–181.
- 43 J. Wang, M. K. Cheung and Y. Mi, *Polymer*, 2002, **43**, 1357–1364.
- 44 T. C./Sc. ISO 22762-1:2005 - Elastomeric seismic-protection isolators – Part 1: Test methods.

