

Secondary Relaxation Behavior of Some Diene Polymers

It is now well-known that the chain microstructure of a polymer influences its secondary relaxation behavior. A case in point is the diene polymers. It has been found that below the glass transition temperature there are two distinct secondary loss peaks for the *trans* isomer of polybutadiene but not for the *cis* isomer.¹ Two hypothetical molecular mechanisms were proposed by Morgan, Nielsen, and Buchdahl² for this observation. The purpose of this work is to compare the dynamic mechanical loss curves of three polymers containing the diene groups in an effort to further scrutinize this interesting behavior.

Samples of polybutadiene (PB) were supplied by the Phillips Petroleum Company. The *cis*-PB contains 93% *cis*, 4% *trans* and 3% vinyl, while the *trans*-PB contains 73% *trans*, 17% *cis*, and 10% vinyl structures. They were crosslinked by 0.16% and 0.32% of dicumyl peroxide, respectively. A sample of polybutadiene (PB), $[(CH_2)_3-CH=CH-]_n$, was obtained from the Goodyear Rubber and Tire Company with 82% *trans* content and cured with 0.5% dicumyl peroxide. A sample of polyoctenamer (PO), $[(CH_2)_6-CH=CH-]_n$, also provided by Goodyear, had about 40% *trans* diene groups and was similarly cured. Dynamic mechanical experiments were carried out on a vibrating-reed acoustic spectrometer previously described.³ Samples were cut to appropriate reed dimensions. One end of the sample was then lightly clamped in the spectrometer, while a metal plate was bolted to the free end. This plate served to drive the reed in forced oscillation and was also used to measure deflection. Following a rapid quench into liquid nitrogen and reclamping, the entire assembly was enclosed in a chamber and submerged in liquid nitrogen. If desired, the chamber, which contained some liquid nitrogen, was pumped to reduce the reed temperature toward the triple point, 63°K. Modulus and loss measurements were taken during warm-up at 0.5-1°K/min.

Figure 1 shows the loss data for the two PBs. The large upturns at 170°K for *cis*-PB and 210°K for *trans*-PB are attributable to their respective primary glass (β) transition. Below these temperatures the *trans* isomer has a γ relaxation at 150°K (104 Hz) and a very small and broad δ relaxation around 88°K (121 Hz). Previously, Gupta¹ found loss peaks at 160° and 110°K for *trans*-PB at a measurement frequency about 2000 Hz, and Morgan et al.² located a similar relaxations at 132°K and 85-103°K (7.5 Hz). The loss location of our data for *trans*-PB falls between that of these two authors as may be seen in Figure 2. This Arrhenius plot summarizes the glassy transitions of the polymers considered here and is particularly useful in that virtually all data refer to maxima in $\tan \delta_G$ versus T curves. (The horizontal bars indicate uncertainty in the broad δ -peak position.)

The activation energies of about 9 and 6 kcal/mole for the γ and δ peaks of *trans*-PB correspond to those for other polymers in this region of the Arrhenius plots. Our data for *cis*-PB, as those of the other two authors, show no resolved loss peak but a large shoulder on the side of the T_g peak, this

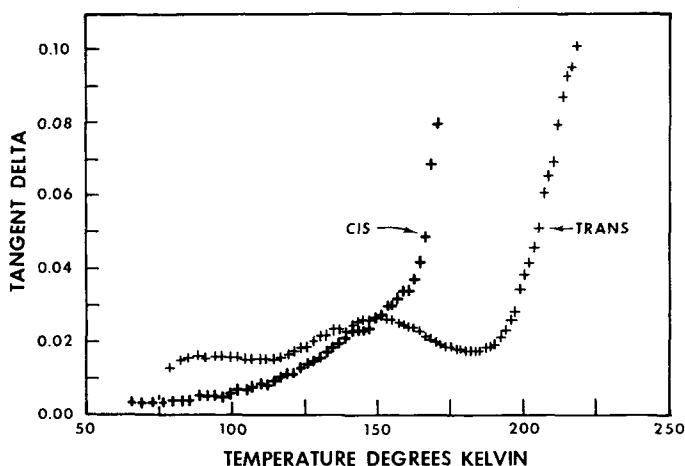


Fig. 1. Loss tangent as function of temperature for *trans* polybutadiene and *cis* polybutadiene.

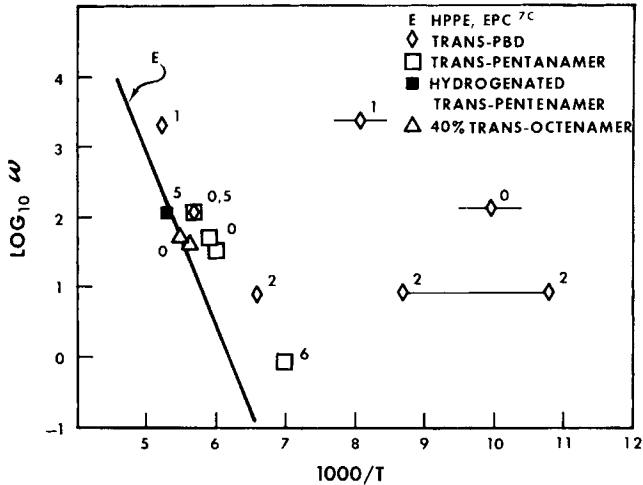


Fig. 2. Arrhenius plot of *trans* dienes and related polymers, maxima in $\tan \delta_G$ vs T : E = HPPE, EPC^{7c}; (\diamond) *trans*-PBD; (\square) *trans* pentenamer; (\blacksquare) hydrogenated *trans* pentenamer; (\triangle) 40% *trans* octenamer.

shoulder appearing roughly in the region of the γ peak of the *trans* polymer. It is conceivable that a small γ peak does exist for the *cis* polymer but is hidden by the closely adjacent T_g peak. This would need to be resolved by working at very low frequencies.

The polyethylene γ -peak position is indicated by a line in Figure 2. The line shown is the peak position found^{7c} in branched polyethylene (i.e., PE of limited crystallinity such as high-pressure PE and ethylene-propylene copolymer) which we believe to be representative of hydrogenated compounds discussed below.

Our loss data for polypentenamer are given in Figure 3. Below the primary glass transition of 170°K, there is but a single loss peak at 142°K (33 Hz, $\tan \delta = 0.044$). Another sample resulted in a substantially higher peak (145°K, 64 Hz, $\tan \delta = 0.067$) but of the same shape. The reason for this difference is unclear. Polypentenamer has also been measured by Sanui, MacKnight, and Lenz⁵ and by Gillham and Benci.⁶ An activation energy of 7.6 kcal/mole is computed (Fig. 2), but this may be a bit low because the peak temperature of the broad low-frequency peak⁶ is hard to establish. Again, our data fall between those of these two observers. One notes that the PP peak falls between the *trans*-PB and the PE peaks, rather closer to the *trans*-PB. This is as might be expected on the basis of the microstructure. When *trans*-PP is hydrogenated,⁵ an additional peak appears at ap-

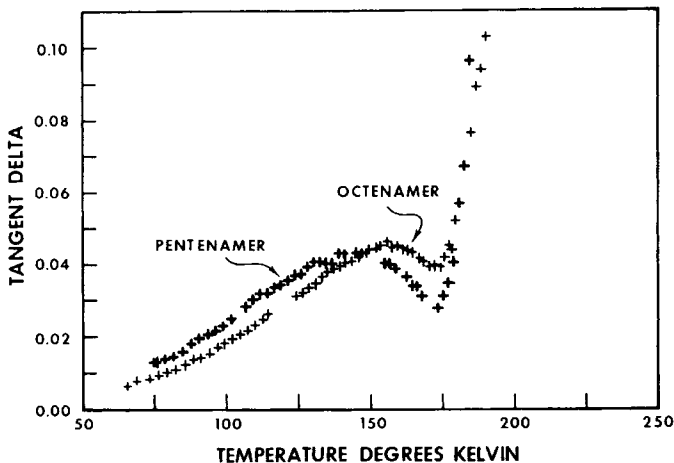


Fig. 3. Loss tangent as function of temperature for polyoctenamer and polypentenamer.

proximately the position of the irregular branch of PE. (Note that even fully hydrogenated material was only 85% crystalline.) The fact that we see two peaks in hydro-PP rather than a single shifted one suggests that the hydrogenation is relatively blocky. Furthermore, Gilliam and Benci⁶ have shown that unhydrogenated *cis*-PP and *trans*-PP both show a loss at low frequency as a broad peak of the same intensity. The situation thus differs from that in PB where only *trans*-PB shows a large peak.

Polyoctenamer is an interesting analog of partly hydrogenated PP. Its loss curve is also shown in Figure 3. Here, two samples gave similar results. Note that only a single peak appears below T_g , which is shown in Figure 2 to fall just in the area of branched PE. No vestige of a *trans*-PB peak seems to remain. No activation energy can be determined since the frequency range is quite small. The loss peaks might be affected by the crosslinking, but we do not believe this is to be the case. It has been shown previously⁸ that crosslinking PE at levels 15 to 25 times higher in dicumyl peroxide has only a relatively small effect on peak shape and position.

In their work, Morgan, Nielsen, and Buchdahl² proposed two possible mechanisms for the β relaxations involving *trans*-PB and *trans*-polyisoprene. One mechanism (model A) involves the motion of C₃—C₄ in C₁—C₂=C₃—C₄—C₅—C₆, specifically a crankshaft-like rotation about the virtually colinear (vinyl) C₁—C₂ and C₅—C₆ bonds. The other mechanism proposed (model B) involves a libration of the C₂=C₃ unit in an all *trans* configuration of C₀—C₁—C₂=C₃—C₄—C₅ with hydrogen wagging but minimal motion of the C₁ and C₄ carbons themselves. Atoms C₀ and C₅ remain fixed. Models A and B are only possible for *trans* configuration around the double bond, whether PB, PP, or PO.

It is useful to interpret the additional data on PP and PO in the light of these models. Replacement of PB by PP changes the C₅—C₆ bond in model A from vinyl to allyl. In PO that bond becomes alkyl. In either case the chemical nature of the C₁—C₂ bond in model A remains unchanged. In model B the C₀—C₁ and C₄—C₅ bonds are biallyl (i.e., allyl to two double bonds) in PB, allyl/alkyl in PP and PO. We expect the vinyl bond to be weak (relatively free rotation), while the allyl bond should be rather similar to the alkyl bond. We have seen that the δ transition occurs in *trans*-PB^{0,1,2} (reference 0 refers to this work) but is absent for *cis*-PB,² *trans*-PP,⁰ and 40% *trans*-PO.⁰ The γ transition is found to be strong for *trans*-PB,^{0,1,2} probably weak for *cis*-PB,^{0,1,2} and present for *trans*-PP^{0,5,6} *cis*-PP,⁶ and 40% *trans*-PO.⁰ When we compare the proposed models to the data, we see that model A can account for the δ transition if we assume that two vinyl bonds are required for the rotor. Model B fails here, the change of an allyl to alkyl bond seems too slight to account for the loss of the transition when we go to PP. Neither model can account for the γ transition, at least in *cis*-PP.

The authors wish to thank Dr. G. Kraus of the Phillips Petroleum Company and Dr. K. W. Scott of the Goodyear Rubber and Tire Company for supplying the samples. The work at the University of California was supported by the Office of Naval Research. Reference to a company or product name does not imply approval or recommendation of that product by the U.S. Department of Agriculture to the exclusion of others that may be suitable.

References

1. R. P. Gupta, *J. Phys. Chem.*, **66**, 1 (1962).
2. R. J. Morgan, L. E. Nielsen, and R. Buchdahl, *J. Appl. Phys.*, **42**, 4653 (1971).
3. T. F. Schatzki and R. G. Meisenheimer, Shell Development Company Technical Report No. 141, 1963 (obtainable from TFS); J. Mathew, M.S.Ch.E. Thesis, University of California, Berkeley, 1976.
4. G. Kraus, C. W. Childers, and J. T. Gruver, *J. Appl. Polym. Sci.*, **11**, 1581 (1967).
5. K. Sanui, W. J. MacKnight, and R. W. Lenz, *Macromolecules*, **7**, 101 (1974).
6. J. K. Gillham and J. A. Benci, *J. Appl. Polym. Sci.*, **18**, 3775 (1974).
7. (a) T. F. Schatzki, *J. Polym. Sci.*, **57**, 496 (1962); (b) *Polym. Prepr. Am. Chem. Soc., Div. Polym. Chem.*, **6**, 464 (1965); (c) *J. Polym. Sci.*, **C14**, 139 (1966).
8. J. Mathew, M. Shen, and T. F. Schatzki, *J. Macromol. Sci. Phys.*, **B13**, 349 (1977).

JACOB MATHEW*
MITCHEL SHEN

Department of Chemical Engineering
University of California
Berkeley, California 94720

THOMAS F. SCHATZKI

Western Regional Research Center
U.S. Department of Agriculture
Berkeley, California 94710

Received September 19, 1977

* Present address: Proctor and Gamble Co., Cincinnati OH 45217.