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Viscous dissipative energy density as a general adhesion characteristic of adhesive tape†

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The strength of adhesive tapes is generally evaluated using a peel or probe-tack test. However, these tests strongly depend on sample geometry considering the tape thickness of the backing film and pressure sensitive adhesive (PSA) layer. In this study, we propose the viscous dissipative energy density as a new parameter for evaluation, which is equivalent to the adhesion energy divided by the maximum elongation of the PSA layer during separation. Rubber-based adhesive tapes of different styrene contents were prepared, and their adhesion performance was evaluated based on the proposed parameter. Peel and probe-tack tests using tapes of various thicknesses were conducted with *in situ* observation using a high-speed camera. In the peel test, the adhesion energy can be obtained by converting the peel force using the tape width and peel angle; in the probe-tack test, it can be obtained by integrating the force divided by the contact area and elongation of the PSA layer. The adhesion energy divided by the maximum elongation from the probe-tack test showed an identical linear relationship with the strain rate of the PSA layer on a logarithmic scale, independent of the tape thickness. This relationship obtained from the peel test was slightly higher with more scattering. This is because the elongation of the PSA layer was distributed along the curved backing film, whereas it was almost uniform under the probe-tack test. Therefore, we confirmed that the viscous dissipative energy density can be used effectively as a material parameter, particularly when obtained from the probe-tack test.

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Introduction

The strength of adhesive tapes is usually evaluated using a peel test owing to its simplicity. The term “strength” is generally defined as a material property, which is independent of the geometry of the material. However, the peel force strongly depends on the peel angle^{1–3} and thicknesses of the backing film and pressure sensitive adhesive (PSA) layer.^{3,4} The influence of testing conditions has been considered in stress analysis, although this is rather complicated because the elongated PSA layer is distributed along the shape of the backing film [Fig. 1(a)].^{5–8} The bending deformation of the backing film generates a compressive stress ahead of the peel zone and the tensile stress sharply increases, followed by a plateau stress in the large elongation of fibrillation, as described in Fig. 1(b).⁹

The probe-tack test is easier to interpret than the peel test as the elongation of the PSA layer is almost uniform owing to the small deformation of the backing film during the separation

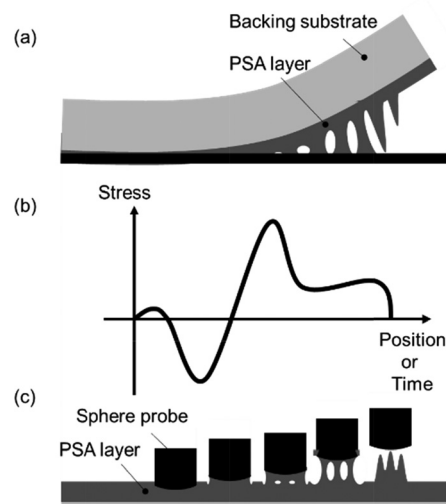


Fig. 1 Schematics of the relationship between (a) peel tests and (c) probe-tack tests in terms of (b) their stress profiles.

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process. The maximum normal stress after several seconds of contact time is defined as the “tack,” which is obtained during the initial elongation of the PSA layer. Subsequently, the normal stress sharply decreases with the expansion of the cavities, and then increases gradually or plateaus with fibrillation toward detachment [Fig. 1(c)].^{10–14} Although “tack” also depends on the testing conditions considering the probe shape,^{15,16} contact time,¹⁷ and thickness of the PSA layer,¹⁸ it is characterized by the expansion of the cavity near the PSA layer–probe surface interface. The maximum stress increases for a large contact area with a thin PSA layer and can be predicted using the cavity growth criterion as a function of the contact radius divided by the PSA thickness.^{19–22}

The above-described release process of the probe-tack test [Fig. 1(b) and (c)] corresponds to the stress distribution in the peel test [Fig. 1(a) and (b)]. Cavity growth is also observed during the peel test, and the maximum stress corresponds to the origin of cavity formation.³ This implies that integrating the normal stress over the entire peel zone provides the external work required for the separation of the PSA layer, which is equivalent to the measured peel force. In other words, the stress transition in the probe-tack test is measured simultaneously as the resultant force in the peel test. Pandey *et al.*²³ compared the peel with probe-tack tests using tapes with various styrenic block copolymers and confirmed the same correlation between these tests. This indicates that the local debonding mechanisms under the peel and tack tests are equivalent, and the adhesion energy can be defined as a material property.

The strain rate dependence of the adhesion energy owing to the viscoelasticity of the PSA layer must also be considered. The adhesion energy predominantly originates from the viscous dissipative energy consumed during the elongation of the PSA layer and follows the power law of the strain rate.²⁴ Based on these perspectives, we proposed a generalized characteristic of the adhesive tape—a ratio of the adhesion energy and the maximum elongation of the PSA layer.³ Adhesive tapes of various thicknesses were tested using a peel test with various peel angles and peel velocities, and the ratio of the adhesion energy and the maximum elongation was found to have an identical linear relationship with the strain rate on a logarithmic scale under identical confinement conditions. This relationship should also hold for the probe-tack test and may be uniquely determined if the debonding process is exactly the same as that in the peel test.

In this study, we investigated whether the same relationship between adhesion energy divided by the maximum elongation of the PSA layer and strain rate can be obtained from peel and probe-tack tests. Two types of adhesives with different viscoelastic properties were prepared and validated. Probe-tack and peel tests were conducted at three different velocities while observing the separation process using a high-speed camera to investigate their equivalence.

Characterization

Adhesion energy

The external work required for separating the probe from the PSA layer in a probe-tack test depends on the area affected by

the probe displacement (x) and measured load (F). When the initial contact area of the probe on the PSA surface is A and the maximum elongation of the PSA layer is a_f , the external work per unit area (W_{tack}) can be expressed as

$$W_{\text{tack}} = \frac{\int_0^{a_f} F dl}{A} \quad (1)$$

The deformation of the backing film is typically suppressed within the elastic range by sandwiching the adhesive tape or using a rigid glass plate. However, the deflection of the backing is not negligible compared with the elongation of the PSA layer. This implies that the probe displacement x is the sum of the deflection of the backing and the elongation of the PSA layer. Therefore, the energy consumed by only the PSA layer ($G_{\text{c,tack}}$) can be extracted by subtracting the elastic strain energy of the PET backing (U_e) from the external work of the probe (W_{tack}).

$$G_{\text{c,tack}} = W_{\text{tack}} - U_e \quad (2)$$

here, it should be noted that $G_{\text{c,tack}}$ and U_e are values normalized by the initial contact area. The relationship between the displacement of the backing and the reaction force can be preliminarily determined by sufficiently pressing the adhesive tape using a probe from the backside, as described in our previous study.²² The elongation of the PSA layer during the probe-tack test was then extracted by subtracting the displacement of the backing from the probe displacement. The area under the force divided by the initial contact area and elongation of the PSA layer represents $G_{\text{c,tack}}$.

In a peel test, when the width of the tape is b and the peel velocity is V , a peeled area bVt is generated in t seconds. To fix the peel angle (θ), sliding the substrate plate at the same velocity V is necessary. Therefore, the net displacement of the peel arm is $Vt(1 - \cos \theta)$. The external work per peeled area (W_{peel}) provided at time t may be expressed using the peel force P .

$$W_{\text{peel}} = \frac{P(1 - \cos \theta)}{b} \quad (3)$$

W_{peel} is consumed by the deformation of the PSA layer and backing film.⁸ The energy consumed in the PSA layer consists of the surface energy of the newly generated surfaces and the viscoelastic dissipation energy owing to elongation during the peeling process. The dissipation energy of the PET backing results from plastic deformation *via* bending deformation (U_p). Therefore, the energy consumed by the PSA layer ($G_{\text{c,peel}}$) in the peeling process should be obtained by subtracting the plastic dissipation energy of the PET backing from the external work per peeled area at time t .

$$G_{\text{c,peel}} = W_{\text{peel}} - U_p \quad (4)$$

The plastic dissipative energy during the bending deformation of the backing film can be related to the bilinear approximation of the stress–strain curve and quantitatively estimated from the elastic modulus (E), yield strain (e_y), and slope after yielding (αE).⁷ The strain energy of the peeling arm does not



have to be considered because the geometry of the peeling arm is maintained during the stable peeling process.

Average strain rate of the PSA layer

Assuming a uniformly elongated entire PSA layer, the strain rate of the probe-tack test is calculated as follows, using the release velocity (V) and initial thickness of the PSA layer (a_0):

$$\dot{\epsilon}_{\text{tack}} = \frac{V}{a_0} \quad (5)$$

In the peel test, the elongation of the PSA layer varies with the bending of the backing; thus, the strain rate is not constant in the peel zone. Villey *et al.*²⁵ calculated the average strain rate of the PSA layer using the time derivative of the logarithmic strain in the peel zone. By averaging the integration of the strain rate over the curved backing, the average strain rate in the peel ($\dot{\epsilon}_{\text{peel}}$) zone can be expressed as

$$\dot{\epsilon}_{\text{peel}} = \frac{V}{l_s} \ln \frac{a_f}{a_0} \quad (6)$$

where V is the peel velocity, l_s is the length of the peel zone along the curved backing film, a_f is the maximum elongation of the PSA layer, and a_0 is the initial thickness of the PSA layer.

Strain rate dependence of the adhesion energy

The adhesion energy is considered to be generated by the viscous dissipative energy (U_v) through the elongation of the entire PSA layer.²⁶ We next defined the viscous dissipative energy density u_v , which is the viscous dissipative energy divided by the elongated volume of PSA.³

$$u_v = \frac{U_v}{L} = \frac{\int_0^{a_f} \sigma A dl}{\int_0^{a_f} A dl} = \frac{G_c - G_{c_0}}{a_f} \quad (7)$$

here, G_{c_0} is the minimum value of the adhesion energy at an extremely low strain rate with a negligibly small elongation of the PSA layer.²⁷ We assumed that the cross section is constant, considering the initial contact area. This means that the actual geometries of the formed cavity and fibrillation are not considered, and the PSA layer is considered to be elongated uniformly over the deformed volume. When the strain rate is sufficiently high, the critical adhesion energy is negligible, and the generalized adhesion strength can be simplified to G_c/a_f .

Experiments

Sample preparation and characterization

The PSAs used in this study were composed of a styrene-based triblock polymer Quintac[®] SL-188 and SL-189 (ZEON Corporation, Japan). SL-189-based PSA was used extensively in our previous study for the peel test;³ therefore, it was compared with SL-188-based PSA, which was designed to have a higher styrene content as a hard segment (Table 1). Differences in viscoelastic behavior were evaluated using dynamic mechanical analysis (DMA). An ARES G-2 rheometer (TA Instruments, USA) with two parallel plates (diameter = 8 mm) was used in the shear mode for each sample (thickness = 2 mm). The oscillation

Table 1 General properties of PSA samples

PSA sample	SL-188-based	SL-189-based
Styrene content [wt%]	30	18
Total molecular weight [$\times 10^3$]	120	170
Hardness [Shore A]	66	45

frequency and temperature ramp rate were 6.28 rad s⁻¹ and 4 °C min⁻¹, respectively.

The adhesive tapes with SL-188 or SL-189-based PSA were prepared by coating their polymer compound dissolved in a solvent (toluene) onto a 50- or 25- μm thick PET backing. The thickness of the PSA layer was adjusted by passing it through an applicator with the gap set to 375 or 125 μm . Overall, PSA tapes with three different thicknesses were prepared and named as ‘‘Applicator gap/PET thickness’’ (Table 2). The thicknesses of the PSA layers were not simply the applicator gap minus the backing thickness owing to volatilization, but they were obtained based on their weight and density.

The 50- μm thick PET film was trimmed into a dumbbell shape to conduct tensile tests using a Compact Tabletop Tester EZ-LX HS (Shimadzu Corporation). The crosshead speeds were set to 0.05, 0.5, and 5 mm s⁻¹, identical to those in the peel test. The material properties of the PET backing required for the prediction of U_p were obtained in our previous study, as listed in Table 3.³

Probe-tack test

The probe-tack test using a polymethyl methacrylate (PMMA) spherical probe (diameter = 9.5 mm) was conducted using a Compact Tabletop Tester EZ-LX HS (Shimadzu Corporation, Japan) at room temperature (22 °C). The adhesive tape was sandwiched between aluminum plates with a 12 mm diameter hole at the center [Fig. 2(a)] to suppress the deflection of the PET backing. The contact time of a probe is generally a few seconds. However, in this experiment, the probe was held for 20 min at the position when it reached a compressive force of 0.1 N to realize the same contact time as the peel test. The probe approached the PSA surface at a velocity of 0.1 mm s⁻¹, and then, it was pulled away from the PSA surface at release velocities of 0.03, 0.3, and 3 mm s⁻¹ after 20 min of holding under a fixed displacement. The probe was cleaned with alcohol before each test, and static electricity was removed using a blower-type static eliminator BF-X2MB (SHISHIDO ELECTROSTATIC, Ltd, Japan). The contact area from the back of the tape was observed during the test using a high-speed camera HAS-L1 (DITECT, Co., Ltd, Japan) [Fig. 2(b)].

Table 2 Thickness of tapes

Tape name	375/50	375/25	125/50
Applicator gap [μm]	375	375	125
PET backing [μm]	50	25	50
PSA [μm]			
SL-188	108	83.1	34.3
SL-189	103	87.9	32.0



Table 3 Material properties of the PET backing

Material	Velocity [mm s ⁻¹]	<i>E</i> [MPa]	ϵ_y [%]	α
PET backing	0.05	3.58×10^3	4.80	0.0111
	0.5	3.61×10^3	5.11	0.00994
	5	3.42×10^3	5.22	0.0115

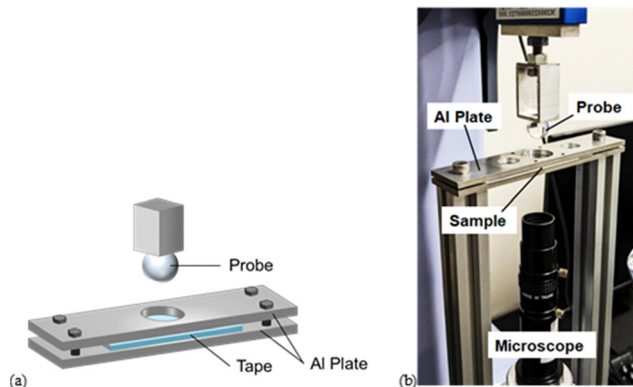


Fig. 2 (a) Sample preparation and (b) experimental setup of the probe-tack test.

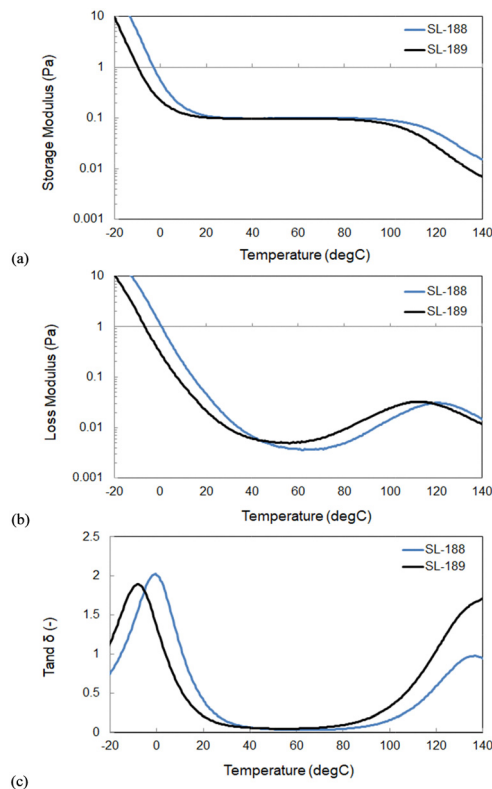
Peel test

The peel forces of the three samples (Table 1) were measured at room temperature (22 °C) using a VPA-2 versatile peel analyzer (Kyowa Interface Science Co., Ltd, Japan). The tapes were carefully attached to a PMMA plate, whose surface was thoroughly cleaned with alcohol to minimize the influence of contamination, and left for 20 min before the test. The PMMA plate was renewed for each test to ensure that the surface conditions remained consistent. The surface conditions exhibited a similarity with the probe employed in the probe-tack test, as examined by the force curve obtained by AFM (atomic force microscopy), as presented in the ESI.† The peel angle was set to 90°, and the peel velocities were 0.05, 0.5, and 5 mm s⁻¹. Each peel test was observed from the side and back of the PMMA plate using the same high-speed camera. Three images from the side were captured at arbitrary observation moments to measure l_s and a_f using the image-processing software ImageJ. The average strain rate of the PSA layer under all peel test conditions was calculated by substituting these values in eqn (6).

Results

Viscoelastic properties of PSA materials

The DMA results are shown in Fig. 3(a)–(c). Although a higher storage modulus was expected from the SL-188 sample because of the higher styrene content (Table 1), it showed the same plateau value as SL-189 (0.11 MPa) around room temperature. The storage modulus was higher below 10 °C and above 80 °C, and the loss modulus was higher below 40 °C, but lower above 40 °C. These differences correspond to a shift in the glass

Fig. 3 (a) Storage modulus, (b) loss modulus, and (c) $\tan \delta$ of the PSA sample measured by dynamic mechanical analysis (DMA).

transition temperature [Fig. 3(c)]. Therefore, the increased styrene content does not significantly change the overall viscoelastic behavior but contributes to the shift of the glass transition temperature to higher values.

Influence of the tape thickness on the probe-tack test

The results of the probe-tack test obtained from the 375/50, 375/25, and 125/50 tapes are shown in Fig. 4(a)–(c), respectively, with SL-188 and SL-189 on the left and right, respectively. The vertical axis represents the nominal stress (equal to the load divided by the initial contact area), and the horizontal axis represents the elongation of the PSA layer (equal to the displacement of the probe subtracted by the displacement of the PET backing). The origin of the elongation of the PSA layer was set to a point at which the load shifted from negative to positive during the release process.

In both the SL-188 and SL-189 PSAs, the maximum nominal stress and slope increased with increasing release velocity. The maximum nominal stress was the lowest in the 375/50 tape and the highest in the 125/50 tape. Conversely, the maximum elongation was the longest in the 375/50 tape and the shortest in the 125/50 tape. It follows that the thinner PSA layer resulted in a higher stress with a shorter elongation. The maximum nominal stress of SL-188-based tape was higher than that of SL-189-based tape for all thicknesses. Overall, the nominal stress through large deformation was not significantly affected



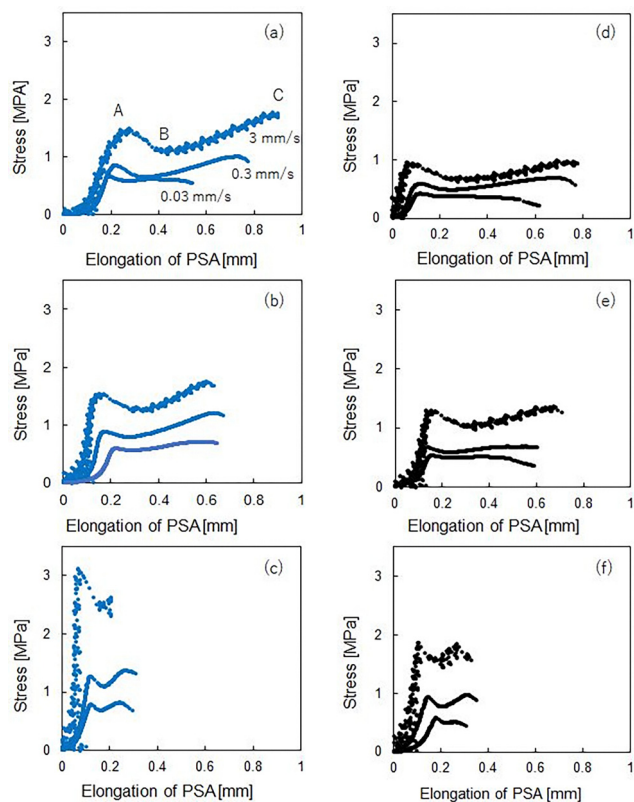


Fig. 4 Measured normal stress of the probe-tack test and the elongation of the PSA layer using all the prepared tapes at different separation velocities. SL-188-based: (a) 375/50, (b) 375/25, and (c) 125/50; SL-189-based: (d) 375/50, (e) 375/25, and (f) 125/50.

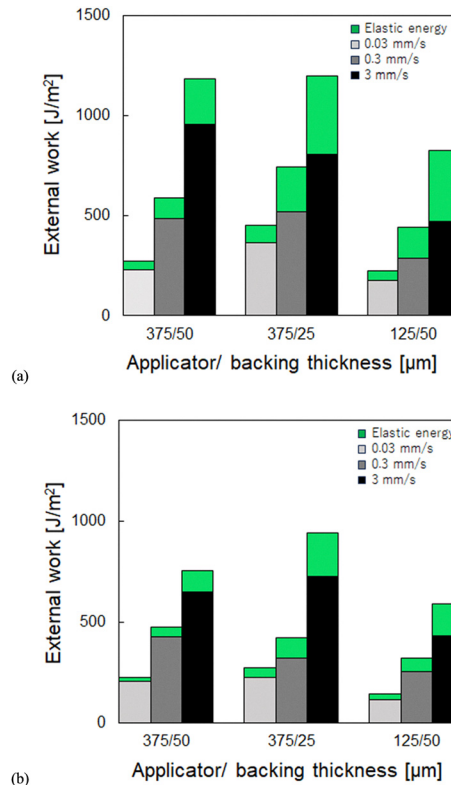


Fig. 5 Adhesion energy, which is the external work subtracted by the elastic strain energy (represented in green), calculated from the probe-tack test using (a) SL-188-based and (b) SL-189-based tapes. The colors represent the separation velocities of 0.03 mm s^{-1} .

by the elongation but was sensitive to the strain rate, which is a characteristic of typical extensional viscosity.²⁶

The adhesion energies (G_{c_tack}) of the backing for the SL-188- and SL-189-based tapes were calculated *via* the area under the plots shown in Fig. 5(a) and (b), respectively. The release velocities of 0.03 , 0.3 , and 3 mm s^{-1} are shown in light gray, gray, and black, respectively. The external work (W_{tack}) was also calculated as the area under the nominal stress and probe displacement curves, and their difference represents the elastic strain energy (U_e) of the PET backing, which is shown in green. A higher adhesion energy was generated by the higher release velocity, regardless of the tape thickness and PSA materials. Compared to the 375/50 tape, the thinner backing and PSA layer of the 375/25 and 125/50 tapes, respectively, generated more elastic strain energy.

Fig. 6(a)–(c) show the observed images of the contact area through the backing PET under a digital microscope for the 375/50, 375/25, and 125/50 tapes of the SL-188-based PSA, respectively. The images were captured at the loads of the local maxima, local minima, and immediately before detachment, represented by points A, B, and C in Fig. 4(a), respectively. The cavities appeared near the local maxima, followed by a sharp decrease toward the local minima owing to the instant expansion of these cavities. A finger-like shape was also formed around the outer edge of the contact area. The load increased

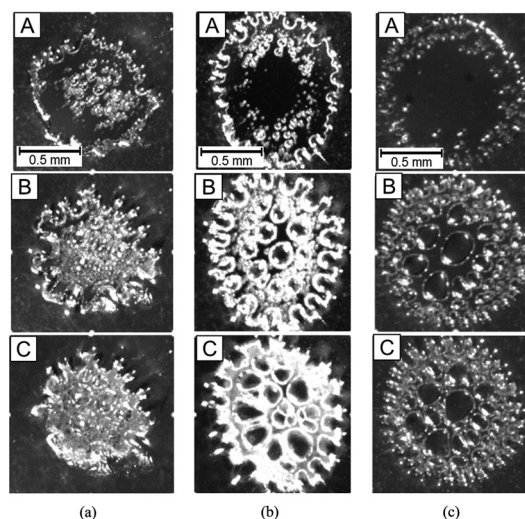


Fig. 6 Images of the backside of SL-188-based tapes observed through a digital microscope: (a) 375/50, (b) 375/25, and (c) 125/50 at the local maxima (point A), local minima (point B), and just before detachment (point C).

again toward complete separation with further expansion of the cavities to fill the gaps between them. These separation processes were clearer for the 375/25 [Fig. 6(b)] and 125/50 tapes



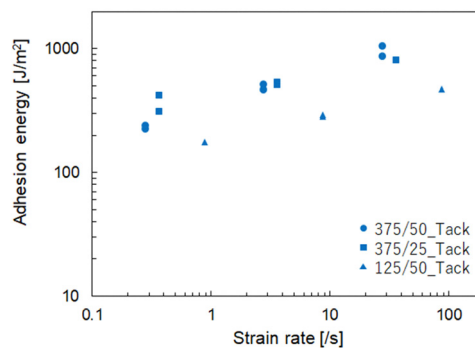
[Fig. 6(c)]. The 375/50 tape produced smaller cavities with a more irregular contact area pattern [Fig. 6(a)].

Fig. 7(a)–(c) show the observed images for the same set of tapes used in SL-189-based PSA. The 375/50 tape of the SL-189-based PSA exhibited only a few large cavities. The way outer edge of the contact area was also less clear than that of the 375/50 tape for the LS-188-based PSA. The images of the 375/25 and 125/50 tapes are similar to those shown in Fig. 6(b) and (c) for the SL-188-based PSA. Although the observed images of the 375/50 tape differ, cavity growth was observed from the local maxima to the minima of the nominal stress (indicated by points A and B, respectively) in all the tapes.

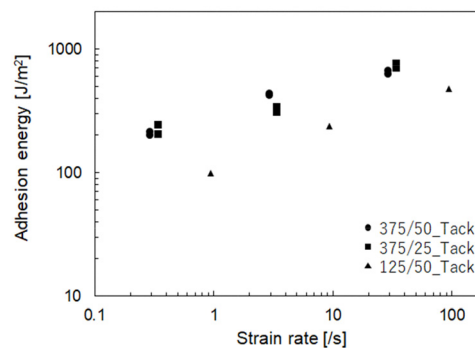
It was ascertained for both tapes that the contact area did not decrease until detachment. The nominal stress increased sharply to the cavity generation indicated by point A, decreased upon cavity growth, and then plateaued or slightly increased until detachment through fibrillation. Therefore, most of the adhesion energy was generated during the fibrillation process.

Viscous dissipative energy density for tapes with different styrene contents

Fig. 8(a) and (b) show the relationship between the adhesion energy (G_{c_tack}) and the strain rate obtained from eqn (5) for LS-188-based and SL-189-based tapes, respectively: The circles, squares, and triangles represent the results for the 375/50, 375/25, and 125/50 tapes, respectively. For all tapes, a linear relationship was observed between the adhesion energy and strain rate on a logarithmic scale. Furthermore, the 375/50 and 375/25 tapes exhibited similar relationships, regardless of the backing thickness. Conversely, the 125/50 tape showed a lower



(a)



(b)

Fig. 8 Relationship between the adhesion energy and strain rate of the PSA layer obtained from (a) SL-188-based tape and (b) SL-189-based tape with the marker shapes representing the thickness of the tapes; 375/50 tapes (circles), 375/25 tapes (squares), and 125/50 tapes (triangles).

adhesion energy. These results are similar to those for the SL-189-based tapes [Fig. 8(b)].

The adhesion energy divided by the initial PSA thickness (G_c/a_0) and the maximum elongation of the PSA layer (G_c/a_f) are shown in Fig. 9(a) and (b), respectively. Although little difference was observed between the different tapes, there was considerable variation in G_c/a_0 for the SL-188-based tape at a low strain rate of $d\varepsilon/dt = 0.3 \text{ [s}^{-1}\text{]}$. In addition, there was almost no difference for G_c/a_0 between the SL-188 and SL-189 tapes at a high strain rate of $d\varepsilon/dt = 100 \text{ [s}^{-1}\text{]}$. By contrast, G_c/a_f was almost consistent at all strain rates. Furthermore, the trendline of the SL-188-based tapes was higher than that of the SL-189-based tapes, but the slope of the linear relationship on a logarithmic scale was almost similar to that of the power law ($G_c/a_f \sim (d\varepsilon/dt)^n$ with $n = 0.23$).

This trend is analogous to the DMA results showing a shift to a higher glass transition temperature and corresponds to the nominal stress shown in Fig. 4, in which the SL-188-based tape showed slightly higher values over all the strain rates. Therefore, the adhesion energy divided by the maximum elongation of the PSA layer represents the adhesion performance more appropriately than that divided by the initial PSA thickness. This result indicates that the elongation of the PSA layer is not proportional to the initial thickness because the deformation along the interface with the probe is constrained by maintaining the initial contact area until detachment.

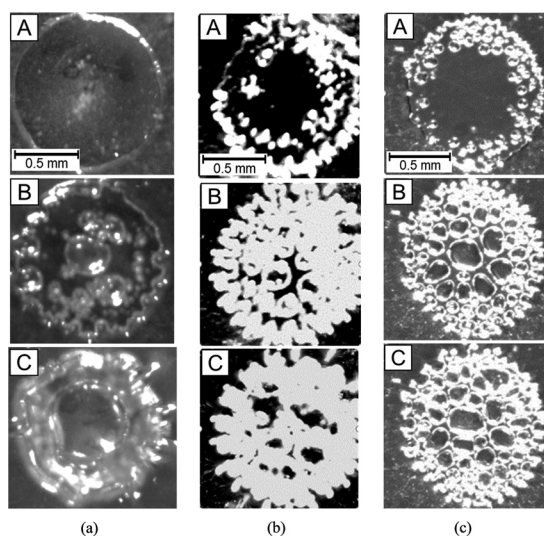


Fig. 7 Images of the backside of SL-189-based tapes observed through a digital microscope: (a) 375/50, (b) 375/25, and (c) 125/50 at the local maxima (point A), local minima (point B), and just before detachment (point C).



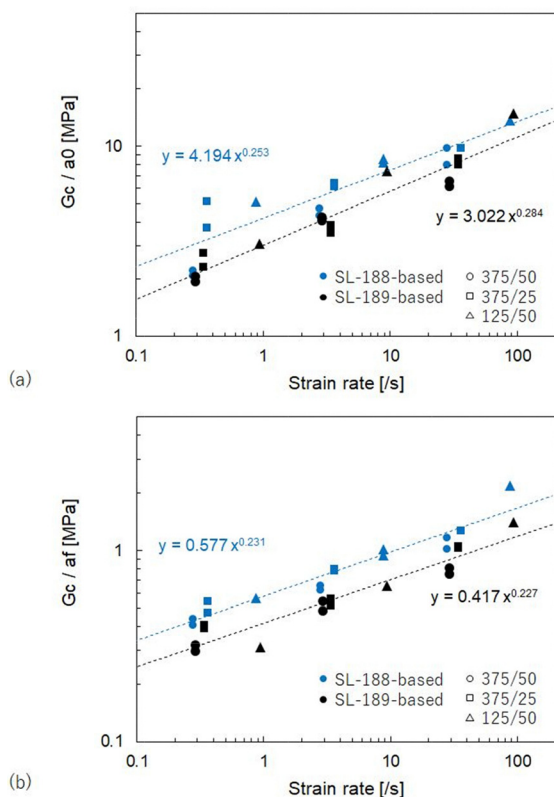


Fig. 9 The adhesion energy divided by (a) the initial PSA thickness and (b) the maximum elongation of the PSA layer, and their relationships with strain rate of the PSA layer obtained from SL-188-based tape (blue) and SL-189-based tape (black), with the marker shapes representing the thickness of the tapes; 375/50 tapes (circles), 375/25 tapes (squares), and 125/50 tapes (triangles).

Comparison with the peel test

Dependence of the tape thickness on the peel force

The external work obtained from the peel test for SL-188- and SL-189-based tapes is shown in Fig. 10(a) and (b), respectively. The peel forces were divided by the tape width to express the external work, using eqn (3) where $\theta = \pi/2$. The light gray, gray, and black colors represent the peel velocity of 0.05, 0.5, and 5 mm s⁻¹, respectively. The plastic dissipation energy of the backing PET caused by bending deformation, calculated using the values in Table 3, is expressed by the green portion of the bar graph. The remaining portion is the adhesion energy defined in eqn (4). A stable peel force could not be obtained from the 375/50 tape of the SL-188 sample at 5 mm s⁻¹ owing to stick-slip oscillation. Regardless of the thickness of the backing and PSA layers, the adhesion energy was higher for the PSA layer and increased with its strain rate, and the viscous dissipative energy during elongation became more significant as the strain rate increased. The plastic dissipative energy (U_p) of the backing also increased with increasing peel velocity. In both samples, the adhesion energy of the 375/50 tape was similar to that of the thinner backing of the 375/25 tape, but higher than that of the thinner PSA layer of the 125/50 tape.

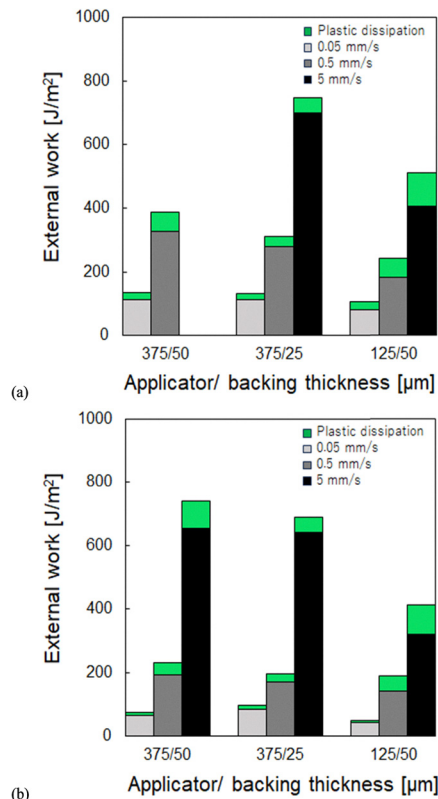


Fig. 10 Adhesion energy, defined as the external work subtracted by the plastic dissipation energy (represented in green), calculated from the peel test using (a) SL-188-based and (b) SL-189-based tapes. The light gray, gray, and black colors represent the separation velocities of 0.03, 0.3, and 3 mm s⁻¹, respectively.

Fig. 11 shows the captured images of the peeled areas of the SL-188-based tapes at a peel velocity of 0.05 mm s⁻¹. Images captured from the 375/50, 375/25, and 125/50 tapes are shown in Fig. 11(a)–(c), respectively. The upper and lower images were obtained from the side and back of the PMMA plate, respectively. Overall, the side view revealed that the curvature of the peeling shape strongly depends on the backing thickness; the 375/50 tape is similar to the 125/50 tape, but has a smaller bending deformation than the 375/25 tape. The peel zone,

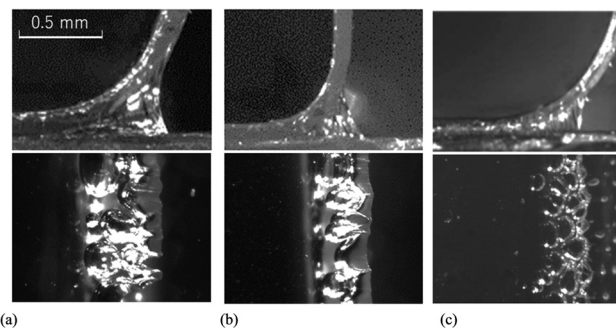


Fig. 11 Images of the SL-188-based tapes observed from the side (top) and through the backside (bottom) under a digital microscope: (a) 375/50, (b) 375/25, and (c) 125/50 tapes.



where the PSA layer was elongated, was more extensive for the thicker PSA layers of the 375/50 and 375/25 tapes. The back view revealed different debonding morphologies. The generation of cavities during peeling significantly depends on the thickness and viscoelastic properties of PSA; the thicker backing and thinner PSA layer of the 125/50 tape clearly showed cavities.

Fig. 12 shows the captured images of the peeled area of the SL-189-based tapes in the same manner as in Fig. 11. The general trend was the same as that for the SL-188 samples; however, the cavities were less evident in the 125/50 tape than in the SL-188-based tape. The maximum elongation was slightly lower.

Viscous dissipative energy density in the peel and probe-tack tests

The viscous dissipative energy density, which is the adhesion energy divided by the maximum elongation of the PSA layer, is plotted in terms of the strain rate in Fig. 13(a) for the SL-188-based tapes and Fig. 13(b) for the SL-189-based tapes. The maximum elongation (a_r) and length of the peel region along the backing (l_s) were measured from the captured images in Fig. 11 and 12 to calculate the strain rate using eqn (6). They are plotted as open markers, with those obtained from the probe-tack test as closed markers (Fig. 9). Circles, squares, and triangles represent the results for the 375/50, 375/25, and 125/50 tapes, respectively.

Although the relationship between the viscous dissipative energy density and the strain rate of the PSA layer was almost identical regardless of the thickness, the values of the peel test were slightly higher compared to those of the probe-tack test because of the shorter maximum elongation during this test. This difference may result from the crack propagation rate during detachment. In the probe-tack test, the crack propagates unstably upon detachment,²⁸ while the crack propagation is stable during the peel test. Our previous work clarified the “sticking effect” caused by the shearing deformation of the PSA layer along the interface with the adherend.²⁹ The initial contact area is maintained during elongation by the high shearing strain rate that is locally generated along its outer edge. The stable and slower crack propagation at the end of the

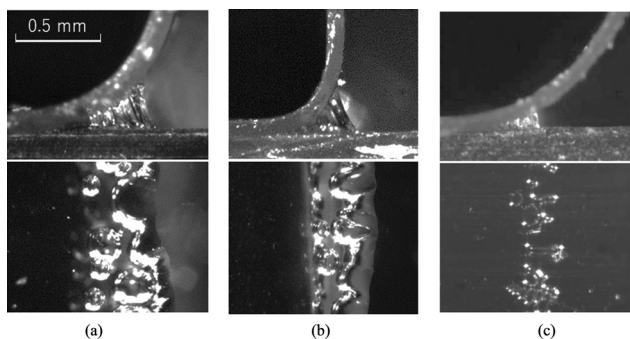


Fig. 12 Images of the SL-189-based tapes observed from the side (top) and through the backside (bottom) under a digital microscope: (a) 375/50 tape, (b) 375/25 tape, and (c) 125/50 tape.

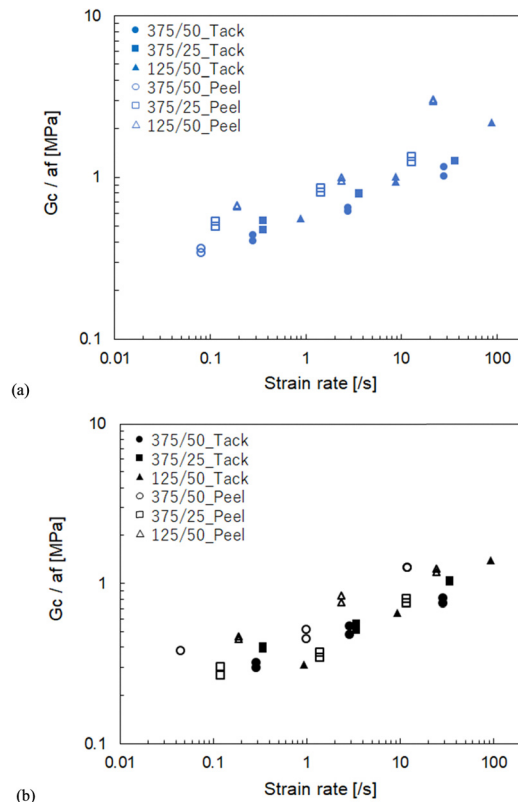


Fig. 13 Relationship between the generalized adhesion strength and strain rate of the PSA layer obtained from the (a) SL-188-based tape and (b) SL-189-based tape with open markers indicating the peel test and filled markers indicating the probe-tack test. The marker shapes represent the thickness of the tapes; 375/50 tapes (circles), 375/25 tapes (squares), and 125/50 tapes (triangles).

peel zone in the peel test has a lower shearing strain rate and thus results in a weaker sticking effect compared to the probe-tack test.

In addition, the results from the peel test were slightly more scattered than those of the probe-tack test. This is because the elongation of the PSA layer in the probe-tack test was almost uniform, while it varied in the peel region during the peel test. The elongation of the PSA layer in the peel test resulted from the nonuniform strain rate in the peel zone and the corresponding stress field. Therefore, the probe-tack test is more suitable for characterizing the adhesion strength of the tape as a material constant compared to the peel test. Moreover, the maximum elongation of the PSA layer in the probe-tack test can be easily obtained by subtracting the displacement of the backing at detachment from the probe position, while that of the peel test must be measured by *in situ* observation under a microscope.

Notably, the relationship between the viscous dissipative energy density and strain rate (Fig. 13) was obtained only from interfacial debonding. When the velocity was sufficiently low, the fracture mode shifted to a cohesive fracture inside the PSA layer.³⁰ Furthermore, the stick-slip phenomenon, which was observed at a peel velocity of 5 mm s^{-1} on the 375/50 tape of the SL-188-based PSA, has been reported to occur in the transition



region between interfacial debonding and glassy fracture owing to high-velocity peeling.^{31,32} The relationship between the viscous dissipative energy density and the strain rate obtained (Fig. 13) is valid only in the range of interfacial debonding with stable peeling.

Conclusions

In this study, adhesion tapes of SIS triblock copolymers with different styrene contents and various thicknesses were prepared. The probe-tack test was conducted at various release velocities. The adhesion energy, which is an integral of the stress–displacement relationships, was higher for adhesion tapes with higher styrene contents and thicker PSA layers. The adhesion energy was divided by the maximum elongation of the PSA layer, which is defined as the viscous dissipative energy density, and plotted in terms of the strain rate. A linear relationship on a logarithmic scale was obtained, independent of the tape thickness. These results suggest that the viscous dissipative energy density can be expressed solely by the power law of the strain rate. Adhesion tapes with higher styrene contents exhibited higher viscous dissipative energy density values, while keeping the same dependence on the strain rate.

Peel tests were also conducted using the same adhesion tapes, and their viscous dissipative energy densities were obtained. The results were slightly higher and more scattered than those obtained from the probe-tack test. The discrepancy from the probe-tack test results was caused by the slower crack propagation rate and the variation in the elongation of the PSA layer during the bending deformation of the backing film. These phenomena affect the strain rate dependence of the locally high shearing stress along the outer edge of the initial contact area (sticking effect) and the extensional viscosity, respectively. By contrast, the probe-tack test generated an almost uniform elongation of the PSA layer. Obtaining the maximum elongation of the PSA layer from the probe-tack test is easily achieved because *in situ* observations are not necessary. Therefore, the proposed viscous dissipative energy density is an effective parameter independent of testing conditions, particularly when obtained from the probe-tack test. As the effectiveness of this parameter was confirmed only for interfacial failure, we intend to investigate the transition of the fracture mode at sufficiently low or high testing velocities in future studies.

Conflicts of interest

There are no conflicts of interest to declare.

Data availability

Data used in this study will be made available upon request to the corresponding author.

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References

- 1 D. H. Kaelble, *Trans. Soc. Rheol.*, 1959, **3**, 161–180.
- 2 A. N. Gent and S. Y. Kaang, *J. Adhes.*, 1987, **24**, 173–181.
- 3 M. Yamada, K. Takahashi, N. Fujimura and T. Nakamura, *Eng. Fract. Mech.*, 2022, **271**, 108653.
- 4 K. Shitajima, N. Karyu, S. Fujii, Y. Nakamura and Y. Urahama, *J. Appl. Polym. Sci.*, 2015, **132**(27), DOI: [10.1002/app.42210](https://doi.org/10.1002/app.42210).
- 5 D. H. Kaelble, *Trans. Soc. Rheol.*, 1959, **3**, 161–180.
- 6 D. H. Kaelble, *Trans. Soc. Rheol.*, 1960, **4**, 45–73.
- 7 A. J. Kinloch, C. C. Lau and J. G. Williams, *Int. J. Fract.*, 1994, **66**, 45–70.
- 8 K.-S. Kim and N. Aravas, *Int. J. Solids Struct.*, 1988, **24**, 417–435.
- 9 D. H. Kaelble, *Trans. Soc. Rheol.*, 1965, **9**, 135–163.
- 10 H. Lakrout, P. Sergot and C. Creton, *J. Adhes.*, 1999, **69**, 307–359.
- 11 A. J. Crosby, K. R. Shull, H. Lakrout and C. Creton, *J. Appl. Phys.*, 2000, **88**, 2956–2966.
- 12 A. Zosel, *Int. J. Adhes. Adhes.*, 1998, **18**, 265–271.
- 13 K. Takahashi, R. Oda, K. Inaba and K. Kishimoto, *Soft. Matter.*, 2020, **16**, 6493–6500.
- 14 T. Yamaguchi, C. Creton and M. Doi, *Soft. Matter.*, 2018, **14**, 6206–6213.
- 15 Y. Y. Lin, C. Y. Hui and H. D. Conway, *J. Polym. Sci., Part B: Polym. Phys.*, 2000, **38**, 2769–2784.
- 16 K. Takahashi, F. Yanai, K. Inaba, K. Kishimoto, Y. Kozono and T. Sugizaki, *Langmuir*, 2021, **37**, 11457–11464.
- 17 C. Creton and L. Leibler, *J. Polym. Sci., Part B: Polym. Phys.*, 1996, **34**, 545–554.
- 18 P. Tordjeman, E. Papon and J.-J. Villenave, *J. Polym. Sci., Part B: Polym. Phys.*, 2000, **38**, 1201–1208.
- 19 R. E. Webber, K. R. Shull, A. Roos and C. Creton, *Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys.*, 2003, **68**, 021805.
- 20 K. R. Shull and C. Creton, *J. Polym. Sci., Part B: Polym. Phys.*, 2004, **42**, 4023–4043.
- 21 T. Yamaguchi, K. Koike and M. Doi, *Europhys. Lett.*, 2007, **77**, 64002.
- 22 K. Takahashi, Y. Yamagata, K. Inaba, K. Kishimoto, S. Tomioka and T. Sugizaki, *Langmuir*, 2016, **32**, 3525–3531.
- 23 V. Pandey, A. Fleury, R. Villey, C. Creton and M. Ciccotti, *Soft. Matter.*, 2020, **16**, 3267–3275.
- 24 J. Chopin, R. Villey, D. Yarusso, E. Barthel, C. Creton and M. Ciccotti, *Macromolecules*, 2018, **51**, 8605–8610.
- 25 R. Villey, P.-P. Cortet, C. Creton and M. Ciccotti, *Int. J. Fract.*, 2017, **204**, 175–190.
- 26 J. M. Piau, C. Verdier and L. Benyahia, *Rheol. Acta*, 1997, **36**, 449–461.



- 27 A. N. Gent and S.-M. Lai, *J. Polym. Sci., Part B: Polym. Phys.*, 1994, **32**, 1543–1555.
- 28 A. J. Crosby and K. R. Shull, *J. Polym. Sci., Part B: Polym. Phys.*, 1999, **37**, 3455–3472.
- 29 K. Takahashi, F. Yanai, K. Inaba, K. Kishimoto, Y. Kozono and T. Sugizaki, *Langmuir*, 2021, **37**(39), 11457–11464.
- 30 S. Sun, M. Li and A. Liu, *Int. J. Adhes. Adhes.*, 2013, **41**, 98–106.
- 31 P. P. Cortet, M. Ciccotti and L. Vanel, *J. Stat. Mech.*, 2007, P03005.
- 32 Y. Sugizaki and A. Suzuki, *Soft. Matter.*, 2019, **15**, 7080–7088.

