Technical Report No. 5

BONDING ELASTOMER LAYERS TOGETHER BY A SURFACE OXIDATION REACTION

by

R. J. Chang, A. N. Gent, C. C. Hsu and K. C. Sehgal

Institute of Polymer Science
The University of Akron
Akron, Ohio 44325

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NR 092-555

October 31, 1979

Unclassified

Submitted for publication in Journal of Applied Polymer Science

Self-adhesion, bonding, elastomers, polybutadiene, cis-polyisoprene, natural rubber, antioxidants, oxidation, kinetics of bonding

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Bonding Elastomer Layers Together by a Surface Oxidation Reaction

Summary

An experimental study has been carried out of the self-adhesion of thin layers of three crosslinked elastomers: cis-polyisoprene (natural rubber) and two polybutadienes. For the polybutadiene materials, the strength of self-adhesion was found to be strongly dependent upon the time of exposure of the two surfaces to air before they were brought into contact. The strength rose dramatically during the first hour or so of exposure so that in some instances the layers could not later be forcibly separated. When the surfaces were exposed to air for longer periods before joining them, the strength of self-adhesion fell to low levels again. This remarkable enhancement in self-adhesion, caused by prior exposure to air for short periods, did not take place on exposure to nitrogen or for samples of cis-polyisoprene, and it was reduced or delayed in samples containing antioxidant. It is therefore attributed to surface oxidation reactions that can lead to the formation of interfacial covalent bonds with polybutadiene but not with polyisoprene. Some possible reaction schemes are discussed.
Introduction

Little previous work has been published on the self-adhesion of crosslinked elastomers. The adhesion of a crosslinked elastomer to an inert rigid substrate is known to be dependent upon the speed and temperature of the test, due to viscoelastic effects (1,2), but at low rates of peeling and at high temperatures, when the contribution of viscoelastic energy losses is small, the work of separation appears to reach a limiting low value of the order of \( 1 \text{ J/m}^2 \) of interfacial area (2). Adhesion of crosslinked elastomers to each other appears to follow the same general pattern and the work of separation is of the same order as for adhesion to inert rigid substrates (3).

Recently, however, extremely strong adhesion has been observed between some crosslinked elastomer layers when they have been exposed to air for short periods at room temperature before being pressed together (4). In some cases the layers could not later be forcibly separated. Details of these observations and a tentative explanation of the phenomenon in terms of oxidative crosslinking reactions at the surfaces are given in the remainder of this paper. The effect could have direct application both to the practical problem of adhering elastomer layers together and also, if the proposed explanation is correct, as a means of studying surface oxidation reactions.
**Experimental**

Thin layers of each crosslinked elastomer were prepared by a molding process using mix formulations given in the Appendix. The molding times and temperatures used for bringing about cross-linking are also given there. Each material was molded between a layer of thin cotton cloth, thickness 0.26 mm, which adhered to one surface and a layer of Mylar film, Type 300A (E. I. du Pont de Nemours and Co.) which adhered to the other surface. The elastomer layer itself was about 0.3 mm thick.

After preparation in this way, the samples were stored at room temperature for about 24 h. The Mylar film was then removed from each of two sheets, exposing the elastomer surfaces to air. After a given period of time two elastomer surfaces were brought into contact and pressed together under a light force for a further period, generally 24 h, at room temperature. The strength of self-adhesion was then measured by peeling the adhering strips apart, as shown in Figure 1. During the peel experiment, the cloth backing served to prevent any significant extension of the peeled sections.

All peel tests were carried out at room temperature and at a rate of clamp separation of 83 μm/s. These conditions are not sufficiently gentle to eliminate viscoelastic contributions.
Figure 1. Test method
to the work of separation altogether. Only a qualitative interpretation of the experimental results is therefore attempted.

The work $W_a$ required to peel apart a unit area of the interface was calculated from the peel force $F$ per unit width of the test specimen (1,2),

$$W_a = 2F.$$

**Experimental results**

(i) The bonding phenomenon

As shown in Figures 2 - 4, the work required to separate two crosslinked strips of polybutadiene was found to depend strongly upon the period of exposure to air before they were brought into contact. Initially rather low, the strength of self-adhesion rose dramatically as the period of exposure to air increased, so that after exposure for 1-2 h before the strips were pressed together, they could not subsequently be separated without tearing the elastomer layers. For exposure periods greater than about 2 h the subsequent adhesion became weaker, and it fell rapidly as the exposure time was increased still further. Eventually, after exposure periods of about 10 days, the self-adhesion of the elastomer layers was again quite small, comparable to the initial value.
Figure 2. Work $W_a$ of separation vs time of exposure to air or to nitrogen before bringing strips of Cis-4 polybutadiene into contact. Peroxide recipe (0.1 per cent dicumyl peroxide).
Figure 3. Work $W_a$ of separation vs time of exposure to air or to nitrogen before bringing strips of Diene 35 NFA into contact. Peroxide recipe (0.05 per cent dicumyl peroxide).
Figure 4. Work $W_a$ of separation vs time of exposure to air or nitrogen before bringing strip of Diene 35 NFA into contact. EV sulphur recipe.
If the protective Mylar films were removed from the elastomer layers in an atmosphere of nitrogen (in a glove box) and the two layers were then pressed together, the resulting strength of self-adhesion was found to be low and constant, independent of the period which elapsed between removing the Mylar films and joining the elastomer layers together, Figures 2 and 3. Thus, the striking increases, and later decreases, in self-adhesion shown in Figures 2 and 3 appear to be a result of reaction with atmospheric oxygen.

Similar effects were noted for peroxide cures, Figures 2 and 3, and for sulphur cures, Figure 4, so that the bonding reaction does not appear to be specific to the nature of the crosslink in the elastomer layers. In other experiments, carried out in the dark and in a glove bag containing dry air, the development of self-adhesion took place equally rapidly so that the absence of light, humidity, and any ozone that might be present in the laboratory atmosphere does not impede the bonding reaction in any way. It is thus associated solely with exposure of polybutadiene layers to air, before bringing them into contact.
When a more powerful antioxidant, phenyl-β-naphthylamine, was employed in place of Agerite Resin D in the mix formulation, the enhancement of self-adhesion required longer times of exposure to air to develop, and did not reach the same high values, Figure 4. This result again suggests that an oxidation reaction is responsible for the remarkable degree of adhesion developed between crosslinked sheets of polybutadiene after exposure to air for relatively short periods at room temperature.

Further experiments were carried out to ascertain whether the actual process of peeling away the Mylar films induced an unusually rapid reaction of the elastomer with oxygen. The Mylar films were removed slowly in a nitrogen atmosphere and then the elastomer strips were left undisturbed for about three hours. After this time, air was admitted to the glove bag. The subsequent adhesion of the elastomer strips was found to follow the same time dependence as before, upon the period of exposure to air before joining. Thus, transitory effects of the peeling process itself do not seem to be responsible for the observed interfacial bonding.

Experiments were also carried out with other protective films in place of Mylar: aluminum foil (Alcoa Wrap No. 5186, Aluminum Company of America) and Teflon-coated aluminum foil.
Penfoil, Penetone Corporation). After removal of these foils from the elastomer surface, the subsequent strength of adhesion between two similar elastomer layers was found to vary with the period of exposure to air in substantially the same way as with Mylar. However, the strengths obtained for samples prepared with the Teflon-coated foil were appreciably lower, about one-third, of those obtained with Mylar and aluminum foil. This may be due to some contamination of the elastomer surface by Teflon, or possibly to a less-smooth surface finish. Nevertheless, all three foils yielded qualitatively similar results, indicating that the development of interfacial bonds as a result of air exposure is not due to a specific effect of Mylar film on the elastomer.

The fact that the adhesion developed at the interface between the elastomer layers is extremely strong, so that the layers cannot be pulled apart in some instances, suggests that covalent chemical bonds are formed between them. When the adhering layers were immersed for several hours in reagents which can sever hydrogen bonds, for example, formic acid, acetic acid, and formamide, the layers remained firmly adhered together, suggesting that primary chemical bonds are, indeed responsible for the strong adhesion.
In striking contrast to the strong self-adhesion developed in polybutadiene materials, no comparable effect was observed with natural or synthetic \emph{cis}-polyisoprene. Indeed, the self-adhesion of crosslinked layers of natural rubber was found to be decreased somewhat by prior exposure to air. Now, it is well-known that oxidative reactions lead to further crosslinking and hardening of polybutadiene vulcanizates whereas, in contrast, they generally lead to softening of polyisoprene vulcanizates as a result of molecular scission (5–8). Thus, the observed adhesion phenomenon is wholly consistent with the development of interfacial bonding by means of oxidative processes in those polymers for which molecular interlinking is the principal result of oxidation. A possible reaction scheme is discussed later.

Samples of polybutadiene, obtained at different times, were found to respond quite differently to air exposure. It is thought that the antioxidants added to the material by the manufacturer may have been changed over a period of years. Even after extraction with hot acetone, however, the samples did not develop self-adhesion as rapidly, or to the same degree, as earlier samples. Either the added antioxidants are difficult to remove by this method or the new material is itself more resistant to
oxidation. This aspect of oxidative interlinking needs further examination. It suggests that the process is strongly affected by minor changes in structure or composition.
(ii) **Kinetics of bonding**

Initially, the strength of self-adhesion appeared to increase in direct proportion to the time $t$ of prior exposure to air, Figures 2-4. The early stages of development of self-adhesion could therefore be described by two parameters; the intercept of these linear relationships at $t = 0$, representing the self-adhesion in the absence of any exposure to air, and the slope of the lines, representing the rate of increase of self-adhesion with time of prior exposure. These values are listed in Table 1.

The initial work of separation varied between about 10 and 100 J/m$^2$ for the various materials examined. (The same values were obtained in a nitrogen atmosphere.) They appeared to be inversely correlated with the degree of crosslinking so that the stiffer, more highly crosslinked materials exhibited generally lower values for the work $W$ of separation at $t = 0$. This feature may be due in part to elastic stresses set up during contact, tending to separate the surfaces; in part to the effect of molecular network strand length on the adhesive strength under threshold conditions (2); and in part to variations in mechanical hysteresis under the present experimental conditions. Although the range of initial self-adhesion values was relatively large, and the subject clearly merits detailed study, it is not considered further in this investigation of oxidative effects.
<table>
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<tr>
<th>Recipe (See Appendix)</th>
<th>$W_a$ at $t = 0$ (J/m$^2$)</th>
<th>$\frac{\partial W_a}{\partial t}$ at $t = 0$ (J/m$^2$/s)</th>
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<td><strong>Diene 35 NFA Polybutadiene</strong></td>
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<tr>
<td>Peroxide</td>
<td>8</td>
<td>0.3</td>
</tr>
<tr>
<td>EV sulphur, 1% Agerite Resin D</td>
<td>28</td>
<td>0.6</td>
</tr>
<tr>
<td>EV sulphur, 1% PBNA</td>
<td>18</td>
<td>0.06</td>
</tr>
<tr>
<td>Sulphur, no PBNA</td>
<td>35</td>
<td>0.5</td>
</tr>
<tr>
<td>Sulphur, 1% PBNA</td>
<td>90</td>
<td>0</td>
</tr>
<tr>
<td><strong>Cis-4 Polybutadiene</strong></td>
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</tr>
<tr>
<td>Peroxide</td>
<td>30</td>
<td>0.2</td>
</tr>
<tr>
<td>EV Sulphur, 1% Agerite Resin D</td>
<td>55</td>
<td>1.1</td>
</tr>
<tr>
<td><strong>Natural Rubber</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Peroxide</td>
<td>27</td>
<td>-0.001</td>
</tr>
<tr>
<td><strong>Natsyn 200 Polysoprene</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Peroxide</td>
<td>15</td>
<td>0.001</td>
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</table>
As shown in Table 1, the rates of increase in work of separation with the period of prior exposure to air were found to be surprisingly large in many cases. Easily-measurable effects were observed after exposure times of only a few seconds at ambient temperature. Although the oxidation of unsaturated elastomers is known to be a facile reaction (5-8), the present materials do not undergo major changes in appearance or in bulk physical properties during several months exposure to air at ambient temperature. The observed large effects of short periods of exposure to air on their self-adhesion are therefore quite unexpected.

It is noteworthy that similar effects are shown by both peroxide and sulphur recipes of both types of polybutadiene. Thus neither a particular microstructure of the polybutadiene nor a particular type of crosslink are required for the development of interfacial bonds after prior exposure to air. There are some indications, however, that the process is more rapid with an EV
sulphur recipe than with a peroxide recipe, and when the anti-
oxidant phenyl-β-naphthylamine is employed the crosslinking
process takes much more slowly, if at all.

As indicated in Table 1, samples of cis 1,4-polyisoprene
did not exhibit any comparable tendency to form interfacial
bonds. Natural rubber samples tended to show a lower degree
of self-adhesion after exposure to air whereas synthetic cis-
polyisoprene samples tended to show somewhat greater self-adhesion
after exposure periods of a few hours. In both cases, however,
the effects were much smaller than for the polybutadiene materials.

Some experiments were carried out to determine the rate at
which interfacial bonding took place. As shown in Figure 5, the
maximum degree of self-adhesion was attained relatively rapidly,
within about 30 minutes after the elastomer layers were brought
into contact, and it did not change significantly during several
hours thereafter. As the bonding process is attributed to an
oxidative reaction in the surface regions of the elastomer layers,
any oxygen that has dissolved in these regions and is available
to continue the reaction when the layers are brought into contact
will be used up eventually and the bonding process will come to
an end. Apparently the oxidative reaction only continues
Figure 5. Work $W_a$ of separation vs time of contact after exposure to air for 30 minutes and 45 minutes. Cis-4 polybutadiene, crosslinked with 0.1 per cent dicumyl peroxide.
for about 30 minutes or so after the layers are joined together. This is comparable to the previous time of exposure to air. It indicates that the chemical reaction leading to interfacial bonding is a rather rapid one, even at ambient temperatures.

(iii) Contact angle measurements

In order to determine whether changes in the elastomer surface on exposure to air could be detected by other means, some measurements were made of the contact angles for small droplets of water placed upon the elastomer surface at various times after removing the Mylar cover sheet. The results are given in Table 2. On exposure to air, significant decreases in the contact angle, by 4 to 7 degrees, took place within the first hour, followed by further small decreases during the next day. No comparable change was observed when the surfaces were exposed to a nitrogen atmosphere, indicating that the effect is again due solely to oxidation.

It is noteworthy, first, that this effect follows roughly the same time dependence as the development of self-adhesion discussed previously, and, secondly, that it is readily observed in natural rubber (although somewhat smaller in magnitude), even though this material did not develop any self-adhesion after exposure to air.
Table 2: Changes in water contact angle $\theta$ with time $t$ of exposure to air or to nitrogen

<table>
<thead>
<tr>
<th>$t$ (h)</th>
<th>$\theta_{\text{air}}$ (degrees)</th>
<th>$\theta_{\text{nitrogen}}$ (degrees)</th>
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<tr>
<td>0</td>
<td>88.9 ± 1.4</td>
<td>88.9 ± 1.4</td>
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<tr>
<td>1</td>
<td>82.5 ± 1.5</td>
<td>88.9 ± 1.4</td>
</tr>
<tr>
<td>3</td>
<td>80.6 ± 3.3</td>
<td>88.2 ± 2.0</td>
</tr>
<tr>
<td>24</td>
<td>80.8 ± 2.6</td>
<td>87.8 ± 1.8</td>
</tr>
<tr>
<td>120</td>
<td>80.8 ± 2.6</td>
<td></td>
</tr>
</tbody>
</table>

**Diene 35 NPA Polybutadiene (Peroxide Recipe)**

**Cis-4 Polybutadiene (Peroxide Recipe)**

**Natural Rubber (Peroxide Recipe)**

<table>
<thead>
<tr>
<th>$t$ (h)</th>
<th>$\theta_{\text{air}}$ (degrees)</th>
<th>$\theta_{\text{nitrogen}}$ (degrees)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>91.0 ± 2.0</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>83.8 ± 2.1</td>
<td></td>
</tr>
<tr>
<td>24</td>
<td>82.0 ± 2.5</td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>89.2 ± 1.5</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>85.2 ± 2.7</td>
<td></td>
</tr>
<tr>
<td>24</td>
<td>84.2 ± 1.6</td>
<td></td>
</tr>
</tbody>
</table>
Thus, an oxidative reaction appears to take place readily in the surface regions of all the elastomers, as revealed by corresponding changes in the contact angle, but it only results in strong adhesion for the polybutadiene materials. This suggests that interfacial bonding is due to a particular feature of the oxidation of polybutadiene materials, not shown by polyisoprene. One particular mechanism is outlined below.

Proposed interfacial bonding mechanism

Oxidation of polyolefins is reported to involve two main propagation steps (5-8),

\[ \text{R} + \text{O}_2 \rightarrow \text{RO}_2^* \]

\[ \text{RO}_2^* + \text{R}'\text{H} \rightarrow \text{RO}_2\text{H} + \text{R}'^* \]

In the first, a polymer radical reacts with oxygen to form the peroxy radical which, in the second step, abstracts H from a nearby group to form a hydroperoxide and a second radical. Moreover, the hydroperoxide itself decomposes slowly, generating further radical species, so that the process is autocatalytic.

Now, this general reaction scheme does not account for hardening during oxidation. Another reaction must therefore be invoked: the addition of polymer radicals to other polymer molecules to form intermolecular bonds.

\[ \text{R} + \text{R}' \rightarrow \text{RR}'^* \]
This reaction is known to occur in polybutadiene by addition to the C-C double bond but not to a significant degree in polyisoprene where the radicals appear to be less reactive. It can thus account for the interfacial bonding observed with polybutadiene but not with polyisoprene. Moreover, it will become of proportionately greater importance as the concentration of oxygen becomes lower, i.e., as oxidation continues in the interfacial region after the elastomer layers have been brought into contact. We therefore infer that the observed adhesion is due to the attack of polymer free radicals, generated during oxidation, upon molecules lying on the other side of the interface, to bring about covalent interlinking.
Conclusions

The following general conclusions are obtained:

(i) Exposure to air at ambient temperature causes a surprisingly rapid oxidation of the surface of polybutadiene and polyisoprene layers. This reaction causes a marked decrease in the contact angle of water within one hour.

(ii) When layers of polybutadiene are brought into contact during this oxidation reaction, interfacial bonds are formed and the layers become firmly adhered together. A significant increase in adhesion can be detected after exposure to air for only one minute or less.

(iii) When the surface oxidation is substantially complete (after several hours in the samples studied here) then the surfaces no longer adhere strongly together when they are brought into contact.

(iv) Samples of polyisoprene also undergo surface oxidation but do not adhere strongly together.

(v) The bonding reaction in polybutadiene is attributed to macro-radicals generated during oxidation which add to C-C double bonds in molecules lying across the interface and form covalent interfacial bonds.

(vi) The bonding reaction is highly sensitive to the presence of certain antioxidants. It was much slower and less pronounced when phenyl-β-naphthylamine was incorporated, for example.
Acknowledgements

This work was supported by a research grant from the Office of Naval Research (Contract N00014-76-C-0408). Helpful suggestions and advice were received from Dr. J. N. Henderson of the Goodyear Tire and Rubber Company.

References


Appendix

The following materials, mix recipes, and reaction conditions, were employed for preparing test sheets.

(i) Elastomers

The elastomers used were:

(a) Polybutadiene, having cis-1,4, trans-1,4, and 1,2 contents of about 36%, 54% and 10%, respectively (Diene 35 NFA, Firestone Tire and Rubber Company).

(b) Polybutadiene containing about 92% cis-1,4 units (Cis-4 1203, Phillips Petroleum Company).

(c) Cis-1,4 polyisoprene (Natural rubber, Standard Malaysian Rubber 5).

(d) Cis-1,4 polyisoprene (Natsyn Type 200, Goodyear Tire and Rubber Company).

(ii) Peroxide recipes

Dicumyl peroxide was added to each elastomer in the following amounts by weight: 0.05 per cent with Diene 35 NFA, 0.1 per cent with Cis-4 polybutadiene, and 1.0 per cent with both types of cis-polyisoprene. Crosslinking was effected by heating for 2 h at 150°C for the polybutadiene mixes and 1.5 h at 150°C for the polyisoprene mixes.
(iii) **Sulphur recipes**

Two recipes based on sulphur as the crosslinking agent were used.

(a) **EV (efficient vulcanization) sulphur recipe**, parts by weight:

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<th>Parts by weight</th>
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<tr>
<td>Elastomer</td>
<td>100</td>
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<tr>
<td>Zinc oxide</td>
<td>5</td>
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<tr>
<td>Zinc 2-ethylhexanoate</td>
<td>2</td>
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<tr>
<td>2-morpholinothio-benzothiazole</td>
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<tr>
<td>Tetrabutylthiuram disulphide</td>
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<td>Sulphur</td>
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<td>Phenyl-α-naphthylamine or Agerite</td>
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</tr>
<tr>
<td>Resin D (R. T. Vanderbilt Company)</td>
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</tr>
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Crosslinking was effected by heating for 40-50 min at 140°C

(b) **Regular sulphur recipe**, parts by weight:

<table>
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<td>Elastomer</td>
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<td>Zinc oxide</td>
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<td>Stearic acid</td>
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<td>Benzothiazyl disulphide</td>
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<td>Tetramethylthiuram disulphide</td>
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Arnold Adicoff, Code 6058
Naval Weapons Center
Research Dept.
China Lake, CA  93555

B. J. Alley
AMSMI-RKC
US AMC
Redstone Arsenal, AL  35809

Dr. S. John Bennett
Thiokol Corp.
Brigham City, UT  84302

Robert A. Biggers
AFRPL/MKP
Edwards, CA  93523

John E. Branigan
AFRPL/RPRP
Edwards, CA  93523

G. W. Burdette
Code 753
Naval Weapons Center
China Lake, CA  93555

Dr. E. A. Burns
TRW Systems Bldg 0-1 Room 2020
One Space Park
Redondo Beach, CA  90278

C. D. Chandler
Hercules, Inc.
Radford Army Ammunition Plant
Radford, VA  24141

John E. Christian
Hercules/ABL
P.O. Box 210
Cumberland, MD  21502

Albert Z. Conner
Hercules, Inc.
Research Center
Wilmington, DE  19899

R. C. Corley
AFRPL/RCS
Edwards, CA  93523

D. B. Davis
Thiokol/Wasatch Division
P.O. Box 524
Brigham City, UT  84302

John H. DeRyke
Chemical Systems Div/United Technologies
P.O. Box 358
Sunnyvale, CA  94088

James Dietz
Thiokol Corp/Wasatch Division
P.O. Box 524
Brigham City, UT  84302

A. J. DiMilo
Propellant R&D
Aerojet General Corp.
P.O. Box 15847
Sacramento, CA  95813

R. J. DuBois
Hercules Inc/Bacchus Works
P.O. Box 98 - MS 8131
Magna, UT  84044

Hiram W. H. Dykes
AMSMI-RKC
US AMC
Redstone Arsenal, AL  35809

Dr. W. David English
TRW Systems Bldg 0-1, Room 2020
One Space Park
Redondo Beach, CA  90278

D. M. French
Code 6012
Naval Ordnance Station
Indian Head, MD  20640

W. A. Gogis, Fleet Support
Naval Ordnance Station
Indian Head, MD  20640