ELASTOMER BONDING GUIDE



A Material Selection, Processing, and Troubleshooting Guide



TABLE OF CONTENTS

Introduction	3
The Elastomer and the Adhesive — Choosing the Right Combination	4
Elastomers	4
Compounding Effects	4
Sulfur Levels	4
Accelerators	5
Fillers	5
Waxes and Oils	5
Phthalate Ester Plasticizers	5
Anti-ozonants	5 7
Non-diene Elastomers	D
Elastonier Bienus	ə 7
Fasy to Uso	(
Elastomer Compatibility	(
Specialty Elastomers	1
Additional Considerations	1
Part Design	7
Molding Method	7
Prebaking	7
Specific Formulation for Specific Needs	7
Further Reading	7
Overview of the Bonding Process	8
Substrate Prenaration	Q
Removal of Oily Contaminants	g
Removal of Insoluble Materials	9
Maintaining Prepared Substrates	9
Adhesive Application	10
Dinning	. 10
Snraving	10
Brushing	. 10
Roll Coating	. 10
Tumbling	. 10
Molding and Finishing Operations	11
Molding Methods	. 11
Finishing Operations	. 11
Troublesheeting	19
Rubber (R) Failure	· 14
Rubber-Cement (RC) Failure	. 12 19
Cement-Metal and Primer-Metal (CM) Failure	. 12
Cement-Primer (CP) Failure	. 12
Combination Failure	. 12
Remedies to Combination Failures	19
	. 10
Types of Failures	. 14

INTRODUCTION

This brochure is a reference guide to elastomer vulcanization bonding. It provides designers and manufacturers with recommended operations for producing high-quality, elastomer-metal assemblies.

Prior to the 1940s, the primary methods for attaching rubber to metal were based either on brass plating of the metal or on "ebonite," as an adhesive material.

Brass plating enabled sulfur-vulcanizing natural rubber (NR) to adhere spontaneously during the vulcanization cycle. Brass plating was the most widely used technique in the years between World War I and World War II. However, it had several shortcomings: the elastomer had to be NR, the brass plate had to be fresh, the adhesion was sensitive to compound variations and processing conditions, and the finished bonds were often undermined by aggressive service environments. Despite these shortcomings, NR-brass plate assemblies were superior to those bonded with ebonite. The latter offered virtually no service heat resistance and also suffered disbonding under mildly corrosive conditions.

In the late 1930s and the early 1940s, proprietary cements became available which were based on resins other than ebonite. These organic solvent formulations continued to show many deficiencies: processing restrictions, poor layover characteristics and inadequate environmental resistance in service. These proprietary cements did broaden the scope of synthetic elastomers and compound variations that were viable options for production of assemblies. The inadequacies of the proprietary cements and the brass plating technique prompted Lord Corporation to initiate broad investigations of the science and technology of elastomer adhesion. This research lead to the development of specially synthesized resins and cross-bridging agents. The resulting technology dramatically raised the performance level of vulcanization bonding.

In 1956, Lord commercialized an adhesive system consisting of a primer and covercoat that produced elastomer-tearing bonds with virtually all commercially available elastomers. This "quantum jump" in technology provided a broad processing latitude for bonding and drastically reduced scrap levels. The primer and covercoat system enabled bonded assemblies to have better environmental service performance.

Continuing research at Lord has produced a growing line of elastomer-bonding formulations that can accommodate all commercially available elastomers, diverse metals and rigid plastic substrates. Lord adhesives have earned worldwide preeminence for their superior consistency, reliability, versatility and ease of use.

New developments have focused on improved bonding performance and environmental acceptability of the formulations due to recent air quality (emissions) restrictions, OSHA regulations and other governmental mandates. These new adhesive systems use water as their medium rather than volatile organic solvents.

THE ELASTOMER AND THE ADHESIVE – CHOOSING THE RIGHT COMBINATION

Production of high-quality bonded parts begins with two choices: the elastomer and the adhesive system.

The first choice is the elastomer. The gumstock type and the details of its formulation will be predicated by the intended function of the bonded assembly. Thus, the rubber for a highly engineered automobile engine mount will be selected for its dynamic performance in controlling vibrations and for its ability to endure under-the-hood operating conditions. Conversely, the elastomer for an engine seal must provide superb resistance to attack by engine fluids.

The second choice is the adhesive system. The adhesive system must provide an excellent bond under the specified vulcanization conditions, as well as maintain its bond under service conditions.

Elastomers

Natural rubber and many synthetic elastomers make up the range of rubber polymers that are available for fabrication. Factors to consider when selecting the elastomer are performance requirements of the part, ease of mixing, processing and molding.

The largest percentage of vulcanizationbonded assemblies makes use of:

- natural rubber (NR)
- styrene-butadiene copolymers (SBR)
- polychloroprene (CR)

• acrylonitrile-butadiene copolymers (NBR) Other commonly used synthetic elastomers include:

- ethylene-propylene-diene monomer (EPDM)
- butyl rubber (IIR)
- isoprene, synthetic (IR)
- polybutadiene (BR)
- chlorosulfonated polyethylene (CSM)
- polyacrylate (ACM)
- ethylene-acrylate ester types (AEM)
- various curable polyurethanes (AU or EU)

High and ultra high-performance elastomers are specified where durability and extreme service conditions are mandated. These include various fluoroelastomer (FKM) and silicone (MQ) types, and hydrogenated NBR (HNBR). For their special dynamic capabilities, elastomeric copolymers from propylene oxide and epichlorohydrin are sometimes selected.

Part designers are beginning to use meltprocessable or thermoplastic elastomers for assemblies whose main function is cushioning or shock control. These elastomers include various polyolefinics (TPO), styrene-butadiene block copolymers, and thermoplastic polyurethanes. These materials are atypical for bonded assemblies as they do not require vulcanization, but they are easy to process, and waste can be recycled. End-uses generally require service at ambient temperatures.

Many of the above mentioned elastomers have features which satisfy specific end-use requirements: oil and organic fluid resistance, heat resistance, resistance to chemical attack, high strength, superior dynamic properties, and/or ease of processing. These elastomer properties are summarized in the accompanying chart (Chart 1).

Compounding Effects

Data generated by Lord technical service laboratories, combined with customer input, provides the information needed for understanding compounding variables and bonding. These formulation guidelines pertain mainly to the non-polar diene elastomers: EPDM, IIR, and NR, and to a lesser extent, the easier to bond and more polar types, such as CR and NBR.

The following compounding ingredients, cure system, fillers, extender oils/plasticizers, and antidegradants all affect "bondability" to varying degrees. The effects of these ingredients are listed below:

Sulfur Levels — The amount of sulfur in the compound has a significant role: sulfur levels of one p.h.r. or higher have a favorable effect on bondability, whereas little or no sulfur results in a compound that is more difficult to bond.

Accelerators — Of the more commonly used accelerators, MBT generally allows good bondability. ZDMC and the ultra-accelerators such as TMTD detract from bondability, particularly in "EV" or "semi-EV" cure systems. The prevulcanization inhibitor (PVI) is often added to fast-vulcanizing stocks to increase processing safety. However, when using ultra-accelerators, high levels of PVI in NR formulations are detrimental to bonding. Amounts of PVI below 0.15 p.p.h. usually allow satisfactory bonding.

Fillers — The type and amount of filler is critical. Compounds with 40 to 80 p.h.r. of carbon black are easier to bond than those with lower black levels. Non-black fillers, such as clays and silicas, also facilitate bonding.

Waxes and Oils — Waxy or oily compounding ingredients that migrate to the vulcanizing elastomer surface cause bonding difficulties. These include low molecular-weight polyolefin auxiliaries, (i.e. low-melting polyethylene and polypropylene processing aids/lubricants), aromatic oils and fatty acid esters (i.e., ricinoleates). Naphthenic or paraffinic oils are less problematic.

Phthalate Ester Plasticizers — Although phthalate esters such as dioctyl phthalate are often recommended for maintaining the mechanical properties of polyolefinic elastomers (EPDM and IIR) in low-temperature, end-use applications, they are detrimental for bonding. Using phthalate esters can compromise the bondability of NBR stocks. However, incorporating high surface area, inorganic fillers, such as silicas, can sometimes neutralize the negative effects of phthalate ester plasticizers. **Anti-ozonants** — High levels of antiozonants and certain antioxidants, particularly the p-phenylene diamine type, may detract from bondability.

Non-diene Elastomers — Elastomers not cured with sulfur and accelerators are easier to bond through inclusion of high surface area fillers. They become more difficult to bond when compounded with certain oils, plasticizers and waxes.

Elastomer Blends

Blends of two or more gumstocks (e.g., NR-SBR mixtures, NBR mill-mixed with IR) are chosen so the most desirable features or properties of each component are available. Blends are also selected in an effort to improve raw material economics, without compromising finished part quality.

Elastomer blends are almost always heterophase systems, i.e. dispersions of one type of elastomer in a continuum or matrix of the other. This heterogeneity is because most elastomer pairs are not mutually soluble. Blending results in less-than-uniform distribution of the compounding ingredients, which often causes one of the elastomers to be preferentially vulcanized by the sulfur and accelerators.

The overall effects of elastomer blending can impact bondability and adhesive selection. For example, blends of NBR and NR will be more difficult to bond than compounds comprised entirely of nitrile elastomer.

CHART 1: ELASTOMER PROPERTY EVALUATION

	ELAST	OMER	S												
LEGEND 1 Excellent 2 Good 3 Fair	Natural Rubber	Isoprene, Synthetic	Styrene-Butadiene Copolymer	Polybutadiene	Polychloroprene	Ethylene-Propylene-Diene Monomer	Butyl Rubber	Chlorosulfonated Polyethylene	Polyurethane	Acrylonitrile-Butadiene Copolymer	Hydrogenated Nitrile	Polyacrylate	Ethylene-Acrylate Ester	Silicone	Fluoroelastomer
4 Poor	NR	R	SBR	BR	CR	EPDM	IIR	CSM	AU/EU	NBR	HNBR	ACM	AEM	MQ	FKM
PHYSICAL PROPERTIES		•		•	•	•						•	•		
Tensile Strength	1	1	2	2	1	2	2	2	1	2	2	3	2	4	3
Elongation	1	1	2	2	1	2	1	2	1	2	2	3	3	1	3
Compression Set	3	3	3	3	2	2	3	2	2	2	2	2	2	1	1
Resilience	1	1	2	1	1	2	4	2	1	2	2	3	3	2	4
Electrical Resistivity	1	1	1	1	2	1	1	1	2	3	3	2	2	1	2
MECHANICAL RESISTANCE															
Tear	1	1	2	2	1	3	2	1	1	2	2	3	3	3	3
Abrasion	1	1	1	1	1	2	2	2	1	1	1	3	3	4	3
Cut Growth	1	1	2	3	2	2	1	2	2	2	2	3	3	4	4
TEMPERATURE															
Heat Resistance	4	4	3	4	3	2	2	3	3	3	2	2	2	1	1
Low-Temperature Resilience	2	2	3	1	3	3	3	3	3	3	3	4	3	1	4
SERVICE PERFORMANCE															
Water	1	1	2	1	2	1	1	2	2	2	2	3	2	1	2
Acid	2	2	2	2	2	1	1	2	3	2	2	3	3	4	2
Alkali	2	2	2	2	2	1	1	1	3	2	2	3	3	4	2
Aliphatic Hydrocarbons	4	4	4	4	2	4	4	2	2	1	1	1	2	2	1
Aromatic Hydrocarbons	4	4	4	4	4	4	4	1	3	2	2	3	3	3	1
Chlorinated Solvents	4	4	4	4	4	4	4	1	2	4	4	4	4	3	3
Ketones	4	4	4	4	4	2	2	4	4	4	4	4	3	3	4
Alcohols	2	2	2	2	2	2	1	2	2	1	1	4	4	2	2
Lubricating Oils	4	4	4	4	2	4	4	4	2	1	1	1	2	2	1
Synthetic Oils	3	3	4	3	4	3	3	4	4	2	2	2	2	3	2
Hydraulic Fluids	4	4	4	4	3	2	3	3	4	3	2	2	2	4	2
Fuels	4	4	4	4	3	4	4	3	2	1	1	1	2	3	1
Weather	3	3	3	3	2	1	1	1	1	2	1	1	1	1	1
Oxidation	4	4	3	4	3	2	2	3	3	3	2	2	2	1	1
Ozone	4	4	4	4	2	1	1	1	1	4	1	1	1	1	1

Why Choose Chemlok[®] Adhesives?

Easy to Use — Lord adhesives wet and spread evenly on prepared metal substrates. They are tolerant of minor amounts of surface contaminants on both metals and elastomer compounds. They accommodate variations in compound formulations and in vulcanization processes.

Elastomer Compatibility — The adhesive must be compatible with the gumstock in providing bond qualities of strength, elongation, dynamic fatigue resistance and durability under expected service conditions. Depending on the elastomer compound selected and the specified vulcanization process, there will be one or more Chemlok adhesive offering this compatibility. This is especially true of the versatile adhesives that bond NR and most of the sulfur-vulcanizable synthetic diene elastomers.

Specialty Elastomers — Many of the highperformance elastomers (i.e., the non-diene types that are not vulcanized by sulfur) require adhesive bonding chemistries that are different from the broad-purpose Chemlok formulations. These formulations include Chemlok adhesives for peroxide-curing silicone elastomers, ionically-curing fluoroelastomers and castable polyurethane types. These Chemlok adhesives for specialty elastomers can often be applied in one coat.

Additional Considerations

The finer points of adhesive selection include considerations regarding the design of the part, the molding method and the compound formulation.

Part Design — The design or geometry of the assembly will influence bonding and how well that part will stand up in service. Fluid engine mounts (i.e., those with contained liquids for back-and-forth movement through orifices or baffles) may place atypical demands on the environmental resistance of the cured adhesive. If the elastomer-metal interface is exposed to a confined fluid such as hot glycolwater mixture, the adhesive system will need to withstand this particular service exposure (i.e., Chemlok 259 adhesive in conjunction with Chemlok 207 primer).

Molding Method — The molding method will affect the tendency for undesirable wiping or sweeping of the adhesive. This phenomenon

sometimes results when a molten elastomer compound moves across the adhesive-coated metal part surfaces prior to vulcanization. Under these conditions, some adhesives can be swept away from the interfaces where they are needed. Chemlok 220 adhesive has good resistance to sweeping or wiping. Chemlok 234B adhesive, by contrast, should not be selected if sweeping is apt to be problematic.

Prebaking — Prebake tolerance is the adhesive's ability to withstand high-temperature exposure before it contacts the vulcanizing elastomer. Bonding is dependent on chemical reactions that occur at elevated temperatures between the adhesive and the vulcanizing elastomer compound (i.e., across the adhesive-elastomer interface). If chemical reactions begin in the adhesive before elastomer contact, a significant amount of the adhesive's bonding capability can be lost. Sublimation of key adhesive ingredients from the adhesive-coated metal surface can cause the adhesive to lose some of its bonding activity. Sublimation may result in migration of highly reactive species to the compound surface. Pre-reaction of key ingredients can then occur at the outer layer of the elastomer, before adhesive-elastomer contact.

Specific Formulation for Specific Needs — The newer aqueous adhesives and the fastbonding adhesives (i.e., Chemlok 252X adhesive) are well suited for brief, hightemperature vulcanizations. The more reactive, faster-bonding Chemlok adhesives (i.e., Chemlok 8560S or D adhesives) are best for difficult-to-bond stocks.

Further Reading

For a comprehensive listing of available Chemlok adhesives, refer to "Chemlok Adhesives Selector Guide," (PB2015). This easy-to-follow guide recommends adhesives for elastomer types, metallic, ceramic, plastic and textile fiber substrates. More detailed information on the aqueous bonding products is presented in the brochure "Chemlok Adhesives — A Guide to Handling and Application," (SI 3018).

OVERVIEW OF THE BONDING PROCESS

CHART 2: ELASTOMER BONDING FLOWCHART

There are four stages to the bonding process:

- 1) Substrate Preparation
- 2) Primer/Adhesive Application
- 3) Elastomer Preparation
- 4) Molding/Curing/Finishing

The flowchart (Chart 2) at right depicts the steps of the bonding process and lists factors to control in each step.

SUBSTRATE PREPARATION

CHEMICAL TREATMENT

- 1. Chemical Cleaning Removal of inorganic surface contaminants (Caustic or Acid)
 - A. Concentration
- B. Temperature
- C. Cleanliness

2. Rinse

- A. Cleanliness
- B. Thorough Agitation
- 3. Conversion Bath
 - (Zn of Fe Phosphate)
 - A. Concentration
 - B. Temperature
 - C. Crystal Size
 - D. Film Weight
 - E. Dispersal
 - F. Residence Time
- 4. Rinse
 - A. pH
 - B. Temperature
 - C. Concentration
- 5. Air Dry

MECHANICAL TREATMENT

- Vapor Degreasing *Removal of oil/grease surface contaminants* A. pH Stabilization
- B. Cleanliness
- C. Volume
- D. Temperature
- E. Treatment
- 2. Mechanical Treatment Removal of inorganic surface contaminants (Grit or Sand)
 - A. Cleanliness of media B. Avoid Fe With Non-
 - Fe Alloys C. Thorough Particle
 - C. Thorough Particle Removal
- 3. Vapor Degreasing Removal of oil/grease surface contaminants
 - A. pH Stabilization
 - B. Cleanliness C. Volume
 - C. volume
 - D. TemperatureE. Treatment
 - E. Heatmen

4. Air Dry

ELASTOMER PREPARATION Elastomer

- A. Properties
- B. Bondability
- C. Mixing
- D. Dispersal
- E. Cure Rate

Bond Design

- A. Radii
- B. Interface Cut-off
- C. Stresses

PRIMER/ADHESIVE APPLICATION

- 1. Primer Application
- A. Primer Choice
- B. Agitation
- C. Evenness of Application
- D. Thin Coats
- 2. Hot Air Dry
- 3. Covercoat
 - A. Adhesive Choice
 - B. Temperature
 - C. Viscosity
 - D. Agitation
 - E. Evenness of Application
 - F. Avoidance of Precure
- 4. Warm Air Dry
- 5. Protection from Contamination

MOLDING/CURING/FINISHING

- Finished Part:
 - A. Mold Design
- B. Loading Time
- C. Temperature
- D. Pressure
- E. Mold Integrity
- Avoid the following:Bond-Destroying Solvents
- Excessive Heat
- Excessive Stress

SUBSTRATE PREPARATION

Proper surface preparation is essential to achieving maximum bond strength. Use Chart 3 to determine the appropriate surface-cleaning procedure. This chart lists guidelines and recommendations for metallic and nonmetallic surfaces.

During the surface preparation, there are certain control parameters which need to be taken into account. These process control checkpoints are listed in Chart 4.

Removal of Oily Contaminants — remove cutting-oils, die lubricants, and other oily contaminants by solvent degreasing. Solvent degreasing consists of three steps:

- 1) Place metals in boiling solvent.
- 2) Rinse in cool solvent.
- 3) Place substrates in solvent vapor until condensation of vapor on metal has stopped.

Degreaser solvents normally used are trichloroethane and perchloroethylene. It is important to maintain clean solvent baths and control pH and temperature. Alkaline detergent scouring can be used in lieu of solvent degreasing. The alkaline bath must have temperature and concentration controls and have an overflow system. Cold and hot water rinse tanks are required to ensure removal of alkali and detergent traces.

Removal of Insoluble Materials — remove scale, rust or other oxide coatings by mechanical or chemical treatments.

Mechanical Treatments include blasting, abrading, machining or grinding. Clean grit or abrasives must be used. Vapor degreasing before and after blasting is preferred. These methods remove dry soil and corrosion, increase the surface area and provide an active surface for bonding.

Chemical Treatments include phosphatizing, acid-pickling, conversion coating or electroplating to provide a clean surface. Treatment solutions must be controlled. Rinse water and drying air must remain pure. Further details are provided in Lord Bulletin 7101, "Preparation of Substrates for Bonding."

Maintaining Prepared Substrates —

prevent exposure to dust, moisture, chemical fumes, mold sprays and other contaminants. Apply the primer as soon as possible after surface preparation.

CHART 3: SURFACE PREPARATION SUBSTRATE | MECHANICAL TREATMENT **CHEMICAL TREATMENT** Metals Steel, Low Carbon Phosphate Blast with clean, sharp 40-Steel, Stainless mesh steel grit, sand or Acid etch aluminum oxide grit. Aluminum and Magnesium Blast with clean, sharp 40-Chromate conversion Brass and Copper mesh aluminum oxide grit Ammonium persulphate etch Zinc and Cadmium Phosphate or chromic acid or sand. Titanium HF acid pickle Plastics Nylon Lightly blast, using sand Clean with alkaline cleaner Cured Phenolic or aluminum oxide grit. or solvent Cured Epoxies Polycarbonate NA Teflon[®] Sodium-napthanate process Delrin® NA DuPont[®] satinizing process

PROCESSING	ELEMENT FOR CONTROL	CONTROL PARAMETERS ¹		
Chemical Cleaning				
Alkaline (may replace	Alkaline bath	Provide overflow system or		
degreasing)		vigorous clean water flush		
Conversion Coating				
Zinc or Iron phosphate	Chemical bath	Maintain concentration,		
		temperature, cleanliness ²		
	Metal parts	Provide adequate residence		
		time, thorough rinsing		
		between baths		
Mechanical Cleaning				
Vapor Degreasing	Chemical vapor cleaning	Maintain pH, volume,		
Grit Blasting or Sand	bath	cleanliness, temperature ³		
Blasting				
¹ See text for specific recommen ² To ensure proper crystal struct ³ To ensure proper deposited film	dations. ure and deposited film weight. 1 weight.			

ADHESIVE APPLICATION

Thoroughly mix pigmented adhesives prior to and during application. Evenly apply the primer and allow to completely dry before top-coating. A thin coat of primer is preferred, as heavy coats can lead to solvent entrapment and subsequent bond failure during molding. Maintain film uniformity by controlling the temperature and viscosity of the wet adhesive or primer. When applying more than one coat of adhesive, allow adequate time and temperature between coats to ensure complete solvent evaporation.

Apply the primer or adhesive by dipping, spraying, brushing, roll coating or tumbling. The choice of application method depends on size, shape of parts and the number of pieces being coated. Listed to the right are the features of the five application methods: **Dipping** — used for solvent and aqueous adhesives. Dipping accommodates both large and small production runs, depending on the level of automation.

Spraying — provides the highest level of bond performance and the most rapid evaporation rate of carrier solvent.

Brushing — recommended only for solvent adhesives. Useful for small runs or production which is not continuous.

Roll Coating — provides an excellent method of coating large flat areas as well as cylindrical objects.

Tumbling — economically coats the parts in a revolving barrel. The adhesive can be dried by either discharging the parts into drying trays or by circulating warm air through the tumbling drum.

Precise guidelines for control of the application processes can be found in the brochure "Chemlok Adhesives — A Guide to Handling and Application," (SI 3018).



MOLDING AND FINISHING OPERATIONS

Molding is the most important step in the bonding procedure; any variation in the individual molding parameters can result in bond failures or a high scrap rate. When designing the mold, make provisions for easy loading of the adhesive-coated metals as well as for easy removal of the vulcanized part.

Place the adhesive-coated metal and rubber compound in the mold cavity. Use the correct time, temperature and pressure to form a quality bonded assembly. Periodically check the mold cavity temperatures by using thermocouples, pyrometers, Tempilsticks[®] or selective melting point wax pencils. Leaky molds, temperature variations, lack of curing or overcuring will adversely affect the bond integrity.

The ideal bonding environment occurs when the elastomer is under maximum pressure and at a minimum viscosity during vulcanization and curing. To obtain these conditions, follow the specified time and temperature requirements of the elastomer being cured. Chart 5 lists the process control checkpoints for molding and finishing operations.

Molding Methods

There are three techniques of molding: transfer molding, injection molding and compression molding. Transfer and injection molding comprise the majority of all manufactured rubber-to-metal parts. Listed in Chart 6 are typical conditions imposed for satisfactory vulcanization bonding.

CHART 5: PROCESS CONTROL CHECKPOINTS FOR MOLDING AND FINISHING OPERATIONS

PROCESSING STAGE	ELEMENT FOR CONTROL	CONTROL PARAMETERS
Assembly Design	Metal-Elastomer Configuration	Avoid sharp radii, abrupt elastomer-metal cutoff and stresses at elastomer-metal interface.
Molding	Mold	Strive for a relatively low cure rate. Control temperature and pressure. Minimize loading time. Maintain mold integrity.
Finishing Operations	Deflashing Plating Coating	Avoid high-solvent paints, high- solvent rust preventatives, acetone exposure and excessive heat and stress.

CHART 6: TYPICAL MOLDING PARAMETERS								
	MOLD	TRANSFER OR	CURE	COMPOUND (RUBBER)				
	TEMPERATURE	Injection time	Time	TEMPERATURE				
Transfer	149°C (300°F) to	20 seconds to	10 to 30	82°C (180°F) to				
	171°C (340°F)	2 minutes	minutes	93°C (201°F)				
Injection	165°C (329°F) to	10 to 20	3 to 10	99°C (211°F) to				
	188°C (370°F)	seconds	minutes	121°C (249°F)				
Compression	149°C (300°F) to 171°C (340°F)	Not Applicable	10 to 30 minutes	82°C (180°F) to 93°C (201°F)				

Finishing Operations

It is often necessary to perform additional treatments to bonded parts. Common bond failures associated with these additional treatments:

- **Deflashing with dry ice or nitrogen** failures between the metal and rubber materials when large loads are in the tumbler at too low of a temperature for an extended period of time.
- Wire brushing, grinding or machining failure of the bonded part due to heat build-up.
- **Electroplating** failures when current densities are too high.
- **Painting** failure when the adhesive does not resist the solvents in the paint.

TROUBLESHOOTING

The American Society for Testing and Materials (ASTM) provides a set of detailed symptom descriptions for bond failures. These descriptions can be used to assess the problem and effect swift, corrective action. (The terms, "elastomer" and "adhesive," in this brochure can be interpreted as "rubber" and "cement," respectively.)

Covering approximately 80% of all bond failures, the four basic ASTM designations are:

- R failure in the rubber. This classification can be further broken down into additional sub-classifications as described below.
- ${\bf RC}$ failure at the rubber-cement interface.
- **CM**—failure at the cover cement and metal; or primer and metal interface.
- **CP** failure at the cover cement-primer interface.

Rubber (R) Failures

Commonly used industry designations for types of rubber failure include:

SR - Spotty Rubber: Appears on the metal surface looking like splattered rubber. Often caused by a metal surface contaminated with dust or other foreign deposits prior to bonding. An SR break can also be caused by ultrafast drying of adhesive as it leaves the spray nozzle (cobwebbing).

TR — *Thin Rubber*: An even, but very thin rubber residue on the metal surface. Usually occurs with butyl or rubber stocks that are very highly oil extended. The oils migrate to the RC interface and provide a bond layer that is part adhesive, part oil and part rubber. This weak layer easily fails when the part is stressed.

HR — *Heavy Rubber*: A thick or heavy layer of rubber remaining on the metal surface indicates an excellent bond. The stock fails because it is stressed beyond its cohesive strength. *SB* — *Stock Break*: A failure of the rubber that makes the elastomer appear to have been folded back on itself and then broken off. The break is jagged and at a sharp angle to the metal surface.

Rubber-Cement (RC) Failures

Separation between rubber and cement or adhesive is usually characterized by a relatively glossy, hard surface on the metal with little or no rubber visible.

Common causes of RC failure are precuring of the adhesive or rubber before the rubber comes in contact with the adhesive, inadequate film thickness, low-molding pressure or temperature, under-cure and migration of plasticizers, oils and other incompatible compounding ingredients.

Cement-Metal and Primer-Metal (CM) Failures

A clean separation between metal and adhesive indicates that no adhesion has occurred. This may be due to several factors. Oil, dirt, dust or other foreign matter on the metal surface may have prevented adhesion from taking place. Environmental factors affecting the metal surface may have caused under-bond separation.

Too-rapid evaporation of solvents in the adhesive may result in ultrafast drying of the adhesive as it leaves the spray nozzle (cobwebbing). Flow of the elastomer stock during bonding may cause displacement of the adhesive from the metal.

Cement-Primer (CP) Failure

Separation at the cover cement-primer interface is easily detected if primer cement and cover cement are of different colors. Such a failure is invariably due to contamination of the primer or plasticizer migration from the elastomer.

Combination Failure

Combination failures can occur when cement-metal, rubber-cement and rubber failures are found on the same part. Consult the charts (Chart 7 and Chart 8) for remedies to combination failures.

REMEDIES TO COMBINATION FAILURES

CHART 7: CEMENT-METAL FAILURE				
POSSIBLE CAUSES Poor metal surface, oil or powdery residue evident	REMEDY Better chemical or mechanical cleaning. Check degreasing operations.			
Adhesive dried before reaching metal (cobwebbing)	Adjust adhesive to proper viscosity. Add higher boiling diluent. Reduce atomization pressure. Reposition gun to ensure adhesive spray coats the targeted area(s).			
Solvent or water entrapped in adhesive film	Increase time between adhesive application and bonding.			
Poor metal preparation — leading to CM failure in service	Check surface preparation. Use a primer on clean metal and follow with covercoat. Prebake primer 10-15 minutes at 138°C (280°F) to 149°C (300°F).			
Galvanic decay or sacrificial metal acitivity	 Avoid dissimilar metal contact upon installation of finished bonded parts. Avoid dissimilar metals in the design of bonded parts. Avoid dissimilar metals in abrasive cleaning (i.e., steel grit for cleaning brass, copper and aluminum. 			
Contamination of treated metal part before adhesive application	Cover parts in storage. Eliminate possible contamination from nearby equipment and/or operations.			

CHART 8: RUBBER-CEMENT FAILURE					
POSSIBLE CAUSES	REMEDY				
Precure of rubber	Use fresh stock. Reduce cure acceleration. Check rubber mixing and compound age before bonding.				
Precure of adhesive (cement)	Accelerate loading cycle. Reduce molding temperature.				
Low mold pressure	Check mold pressure. Check mold for leaks. Increase amount of rubber in mold.				
Incorrect molding cycle	Check mold and platen temperature. Recheck cure rate of stock.				
Low adhesive film thickness	Use heavier or multiple coats of adhesive.				
Bloom or migration from rubber	Substitute more compatible ingredients. Reprocess or freshen stock.				
Contamination of adhesive coated parts	Check unbonded specimens for oil or dust. Avoid over-spray or excessive use of mold release agents. Check for possible contamination from nearby equipment and/or operations.				
Adhesive-rubber incompatibility	Select proper adhesive for elastomer and application.				
Wiping or sweeping of adhesive	Slow transfer rate. Reduce molding temperature. Reduce topcoat dry film thickness.				

TYPES OF FAILURES

FIGURE 1: RUBBER FAILURE



Photos at right illustrate types of rubber-bond failures: thin rubber (TR), heavy rubber (HR) and stock break (SB).



THIN RUBBER (TR) HEAVY RUBBER (HR)

STOCK BREAK (SB)

FIGURE 2: RUBBER-CEMENT FAILURE



Separation at rubber-cement interface. Photo at right shows appearance of the part after such separation — relatively glossy, hard surface with little or no rubber visible.



RUBBER-CEMENT (RC)

FIGURE 3: CEMENT-METAL OR PRIMER-METAL FAILURE



Separation at the primer-metal interface, or cement-metal interface for a one-coat adhesive, results in a part with a clean metal surface as depicted at right. Both are referred to as CM failure.



CEMENT-METAL (CM) One-coat system

FIGURE 4: CEMENT-PRIMER FAILURE



Separation at cement-primer interface. Photo at right shows appearance of the part after cement-primer failure caused by lack of time for solvents in primer to evaporate before cover cement is applied.



CEMENT-PRIMER (CP)



Cross-sectional view shows typical rubber-tometal failure in which 60% of bond area shows failure in rubber (60R), and 40% failure between cement or primer and metal (40CM). Appearance of failure is seen in photo at right.



COMBINATION

Values stated in this bulletin represent typical values as not all tests are run on each lot of material produced. For formalized product specifications of specific product end uses, contact our Customer Service Department. Information provided herein is based upon tests believed to be reliable. Inasmuch as Lord Corporation has no control over the manner in which others may use this information, it does not guarantee the results to be obtained. In addition, Lord Corporation does not guarantee the performance of the product, the results obtained from the use of the product or this information where the product has been repackaged by any third party including, but not limited to, any product end-user. Nor does the

company make any express or implied warranty of merchantability, or fitness, for a particular purpose concerning the effects or results of such use.

Cautionary Information

Before using this or any other Lord product, refer to the product Material Safety Data Sheet (MSDS) and label for safe use and handling instructions.

For additional information, contact: Lord Corporation Chemical Products Division 2000 West Grandview Blvd. P.O. Box 10038 Erie, PA 16514-0038 1-800-CHEMLOK, ext. 3211 FAX: 814/864-3452 www.chemlok.com

Chemical Products Division Worldwide



Lord Corporation P.O. Box 10038 Erie, PA, USA 16514-0038 1-800-CHEMLOK

Lord Corporation (Europe) Ltd. Stretford Motorway Estate Barton Dock Road Stretford, Manchester England M32 0ZH +44 (161) 865-8048

Lord Industrial Ltda. Via Anhanguera, KM. 63,5 Distrito Industrial 13.200 Jundiai, Sao Paulo Brazil +55 (11) 7392-7755

Lord Far East, Inc. 2F, Yoyogi Center Building 57-1, Yoyogi 1-Chome Shibuya-ku, Tokyo Japan 151 +81 (3) 3378-9011

Cautionary Information: Before using this or any other Lord product, refer to the product Material Safety Data Sheet (MSDS) and label for safe use and handling instructions.



Lord and Chemlok are registered trademarks of Lord Techmark, Inc., a subsidiary of Lord Corporation. Teflon is a registered trademark of E.I. Dupont De Nemours, Inc., and Delrin is a registered trademark of E.I. Dupont De Nemours, Inc. Tempilstick is a registered trademark of Tempil Co.