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AFRPL-TR-72-60

# AMBIENT TEMPERATURE CURE CATALYSTS FOR CARBOXY BINDER SYSTEMS

R. E. Foscante C. Weis W. Wiegand Midwest Research Institute

H. Gilbert T. Comfort Allegany Ballistics Laboratory Hercules, Inc.

#### **TECHNICAL REPORT AFRPL-TR-72-60**

JUL 13 1972

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#### FOREWORD

This technical report relates work performed under Contract F04611-70-C-0018, Air Force Rocket Propulsion Laboratory, Edwards, California 93523, for the development of ambient cure catalyst systems for solid propellants based on carboxyl containing binders. The catalyst screening and chemistry studies were performed in the Physical Sciences Division of Midwest Research Institute, Kansas City, Missouri, and the propellant processing and evaluation studies were conducted by Hercules, Inc., Allegany Ballistics Laboratory, Cumberland, Maryland. The Air Force Project Engineer was Captain D. F. Clark, USAF, AFRPL.

Personnel at MRI contributing to the program were: Dr. A. D. McElroy, Program Supervisor from January 1970 to November 1972; Dr. R. E. Foscante, Project Leader and Program Supervisor for the final period of the program; Mrs. C. Weis, Assistant Chemist; and Mr. C. J. Wiegand, Senior Chemist. Those contributing at ABL were: Mr. H. Gilbert, Program Supervisor; Dr. T. Comiort, Project Leader; and Mr. C. Dickey.

The inclusive dates of the program technical effort were January 1970 to March 1972. No classified material is contained in the report nor was any classified information abstracted from classified sources.

Publication of this report does not constitute Air Force approval of the findings and/or conclusions stated herein. It is published only for the exchange and stimulation of ideas.

This technical report has been reviewed and is approved for distribution.

CHARLES R. COOKE CHIEF, SOLID ROCKET DIVISION

#### ABSTRACT

An extensive series of organometallic compounds were screened to determine their utility as catalysts for the ambient temperature cure of carboxylic acid terminated polymers with epoxide and aziridine curatives. Chromium salts, specifically chromium 2-ethyl hexanoate, were shown to promote the desired cure reaction efficiently and stoichiometrically at 1% and less levels at 25°C in less than 7 days in both model and binder cure studies. The presence of other propeilant components (e.g., ammonium perchlorate, aluminum, iron oxide, antioxidants, Surfactants, plasticizers, etc.) and certain contaminants (water, perchloric acid) showed no significant or uncontrollable effects on the course of the Cr catalyzed reaction of both model and binder systems. Catalytic trends appear similar in both aziridine and epoxide systems. Reaction mechanisms and catalyst design parameters were developed and are discussed.

Using the catalyst system selected by Midwest Research Institute, composite propellants based on several different carboxy binder systems were successfully manufactured and cured at ambient temperature (75°F) by ABL. A 6-month surveillance program proved that chromium-octoate-catalyzed propellants had completely cured at ambient and did age as well as 140°Fcured control propellants. The Telagen-epoxide ambient-cured binder system, which was the prime candidate in this program, is deemed ready for scale-up beyond the 40-lb batch size, 85.5% total solids propellant demonstrated in this program. GEDING PAGE BLANK-NOT FILMED."

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#### INTRODUCTION

#### A. Background

Some of the well known advantages of the solid propellant are: simplicity of design, instant availability (compared to cryogenics), and high density. In simplest terms, the composite solid propellant consists of a polymeric matrix containing the other ingredients (oxidizer, fuel) which participate in the combustion process. The curing of the propellant is actually an <u>in situ</u> polymerization process, the results of which control the mechanical, physical, and often combustion properties of the final propellant.

Stresses and strains which may lead to the formation of cracks in the propellant must be avoided. High strains in the port area may result from shrinkage due to a decrease in polymer volume on cure and temperature drop upon cool-down of the motor from the 130° to 160°F cure temperature. The shrinkage due to cool-down may be avoided by using a catalyst to drive the cure reaction at a reasonable rate at ambient (70°F) temperature.

Ambient cure also gives the propellant a comparatively low stressfree temperature (80°F vs. the usual 160°F). The advantage of an 80°F stress-free temperature is lower thermal strains upon exposure of the motor to cold temperatures and only mild compressive forces on the grain at high temperatures.

The general types of binder systems of interest on this program are the carboxylic acid terminated prepolymers cured with either epoxides or aziridines. Currently, elevated temperatures are required to effect the cure of propellants based on these binders.

The interactions which can affect the mechanical and physical properties and aging characteristics of a propellant can be summarized as follows:

1. Homopolymerization or rearrangement of curatives (especially significant for the aziridines).

2. Side reaction of the curative during cure with other ingredients, notably the oxidizer.

3. Postcure reactions due either to side products generated in 2, or to slow reaction of functionalities of the cured propellant with other components (leading to hardening of the propellant).

4. Chemical or thermal instability of the cure linkage leading to cure reversal (softening of the propellant).

5. Cracking of the propellant during cool-down from cure due to the large difference in the coefficient of thermal expansion between the case and the propellant.

Many of these problems can be either eliminated or mitigated by catalyzing the cure reaction. Reaction rates can be accelerated at lower (or ambient) temperatures, thereby eliminating the cool-down and its attendant differential volume change. The main cure reaction can be promoted in the correct direction at the expense of side reactions and homopolymerization. Those postcure reactions proceeding from side products would then be controlled. Of course, another benefit would arise from the cost savings involved in curing propellants at ambient temperature.

Some problems may arise, however, in utilizing an ambient cure process. The lower the temperature, the greater is the slurry viscosity of the formulation. In certain systems ambient mixing and casting may be difficult (especially those of high solids loading). If the temperature is raised to increase fluidity, the presence of a rate accelerator in the mix will cause pot life problems.

Cure reversal is another potential problem in a catalyzed system. There are instances in which a catalyst will assist in cleavage of the bond it helped form (e.g., as may occur in the urethane system). In addition, the cure catalyst may promote side reactions between other functionalities in the propellant during storage (e.g., oxidative cross-linking of double bonds).

To recapitulate, the probable benefits of an ambient temperature propellant cure catalyst system are several, including the following: reduction in the liklihood of undesirable physical and chemical phenomena during cure; increased control of the specificity of cure chemistry, and therefore, of the properties of the cured propellant; a lower stress-free temperature, increase in the variety of binder systems amenable to cure; reduction in the complexity and cost of formulation and cure; and a better opportunity to successfully handle advanced propellant ingredients.

#### B. Objectives

The overall objective of this program is the development and demonstration of ambient temperature cure catalyst systems for propellants containing the carboxylic acid-epoxide and aziridine based binder. The specific approach to the problem involves the following steps. 1. A survey of the literature, to define the state of the art in this area and identify useful information from related fields.

2. Select a series of candidate eatalysts giving desirable reaction rate and stoichiometry.

3. Evaluate those candidates in the presence of propellant components and potential contaminants.

4. Develop and demonstrate processing procedures to utilize the ambient temperature cure catalyst.

5. Evaluate the aging characteristics of solid propellants formulated with these cure catalyst systems.

#### C. Approach

The literature survey indicated the types of materials most likely to be effective as catalysts for the epoxide-carboxylic acid condensation reaction. The selection procedure involves screening the potential catalysts in a model reaction system to obtain relative rate and reaction direction data. The model system approach was taken because it provides a rapid and simplified method for evaluating large numbers of materials. In addition, these reaction studies can be designed so that the effects of selected reaction parameters on rate and stoichiometry can be evaluated. 2-Ethyl hexanoic acid (EHA) and allyl glycidyl ether (AGE) were selected as model reactants; gas-liquid chromatography was employed to monitor the rate and stoichiometry of consumption of reactants and the formation of products. In addition, selected propellant components and contaminants were introduced into the model system and their effects on reaction rate and stoichiometry were determined.

Those materials Surviving the screening studies were then evaluated as cure catalysts in the octual binder system in two types of studies: (1) reaction chemistry through the use of an infrared method to follow the course of the cure reaction; and (2) gumstock and low solids-loaded binder studies as a prelude to processing evaluation using conventional physical property measurements. The effects of propellant ingredients and contaminants on the rate and direction of the catalyzed cure reaction are also under study in these binder studies.

Ambient temperature processing procedures are evaluated and developed in terms of processability and cure characteristics in a series of propellant mix studies ranging from 30 g. to 30 lb. Finally, the aging characteristics of propellants cured with the candidate catalysts (and in some cases with contaminants present) are evaluated in both model and propellant surveillance studies.

#### CATALYST SCREENING (EPOXIDE SYSTEM)

#### A. Selection of Catalysts for Initial Screening

The characteristics of a good cure catalyst are apparent from previous discussion. Notably, such a material must: (1) catalyze only the main reaction; (2) be stable in the propellant environment; (3) be effective in low concentration; (4) control the reaction at a rate conducive to good pot life; (5) be highly soluble in the binder; (6) be intrinsically stable (it must not itself decompose); (7) not contribute to reversal of the cure reaction; (8) be nonreactive with components of the formulation; and (9) require no special handling.

The literature indicates that a wide number of materials (of the mineral and Lewis acid type) have been used to accelerate the opening of the epoxide ring and facilitate condensation of the epoxide with nucleophilic species. The major similarity of these catalytic materials is the fact that they are electrophilic in nature, that is, they are capable of attracting electron rich atoms or groups to themselves. Many of these materials are not suitable for use in propellants because they may catalyze the wrong type of reaction (e.g., epoxide homopolymerization instead of carboxylic acid-epoxide addition esterification) or they may be ineffective or incompatible in the propellant environment. Catalysis by organometallics takes precedence in propellant chemistry as a means of catalyzing the cure reaction because of the apparent efficiency, and ease of handling, storage, and use of these catalysts.

#### B. Reaction Paths and Experimental Design

Possible reaction paths for the carboxylic acid-epoxide reaction include:

II.



Long time or elevated temperatures required.

When R' = R", C-1 = C-2
Where R' ≠ R" substitution occurs preferentially on the least
substituted carbon.

Ф - С-ОН / М 0 + О-С ~ -С-О-С-С-R'' R' Н

Reaction Path A is a possibility only in the catalyzed systems under the low temperature conditions of this study. (The epoxides used are stable at ambient and test temperatures.) Reaction B is an extremely slow reaction, and is mentioned here for the sake of completeness (for the screening studies, stock solutions of acid and epoxide are prepared in advance of most determinations; hence, their reactivity, if any, during storage must be known). Path C is the main and desired reaction route, that is, to compound C (where R' - R", C-l and C-2 are identical). For unsymmetrical epoxides (as are found in the curatives) addition of the carboxyl group will occur on the least substituted carbon (Ref. 1). When R' = H and R" # alkyl, C-1 will be the favored product; that is, the main reaction product will be a 3-hydroxy ester with the hydroxyl group in a secondary configuration on the main carbon chain (in the binder the hydroxyl group will be in a secondary position along the polymer backbone). The unfavored product would contain the alcohol group as a primary substituent on a side chain (C-2 where  $R^* = H$ ,  $R^{**} = alkyl group$ ). Path D represents a route which could conceivably be a competitive reaction to Route C during the cure reaction (depending on the catalyst present) and also be a postcure reaction situation through condensation of unreacted carboxylic acid starting material with the hydroxyl group of the product ester.

Hence, the ideal eatalyst will be either specific to the acid-epoxide condensation or will direct the reaction through Route C to product C-1 (where  $R' \neq R''$  and R' = H) sufficiently fast so that Path D is eliminated because the least reactive product (secondary alcohol along the polymer backbone should be more sterieally hindered than the primary hydroxyl of the unfavored ester and hence, less reactive, if, indeed, such a condensation were to occur) is formed and acid is consumed preferentially in Path C.

In light of this discussion, the data required from the catalyst screening experiments are as follows: (1) relative rates of disappearance of aeid and epoxide and the stoiehiometry of the consumption; (2) extent to which a given catalyst will eause homopolymerization of the epoxide; (3) the number, relative abundance, and identity of products formed; and (4) the extent to which reaction of starting acid and product hydroxy ester is possible. The above data items can be derived <u>directly</u> from the gas chromatographie screening runs (except for the identity of the products); Items 2 and 4 are the subject of additional correborative studies.

#### C. Catalyst Screening Studies

The effects of metal type, ligand nature, and temperature on the rate and stoichiometry of the acid-epoxide reaction were studied in an experimental series utilizing 2-ethyl hexanoic acid (EHA) and allyl giycidyl ether (AGE) as models. Table I lists the materials evaluated in this series. The results of these screening experiments have suggested three categories of catalytic effectiveness for the acceleration of the epoxide-acid reaction. The chromium salts give complete reaction in less than 2 days at 50°C and less than 7 days at 25°C. Iron salts, specifically, ferric octoate, were shown to accelerate the AGE-EHA reaction at a rate less than the analogous chromium salts, but still at a rate which is faster than most other materials and within the desired performance range. The value of the intermediate performing catalysts was assessed as follows: (1) should the chromium salts have proven to be too fast for good pot life behavior, a slower catalyst would be required; (2) should cure reversal during aging be indicated, the intermediate group would prove a mild compromise rate aecelerator.

#### TABLE I

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# SUMMARY OF CATALYST SCREENING STUDIES

	Catalyst					
	Level	Temperature	% Reaction <sup>a</sup> /			
<u>Catalyst</u>	(wt. %)	(°C)	7 Days	<u>14 Days</u>	30 Days	
CrOct	1	25	99	100	100	
	1	50	100	100	100	
CrOleate	1	25	05	100	100	
	1	50	100	100	100	
CrNapthenate	1	25	05	100	100	
	1	50	100	100	100	
Cr-neo-decanoate	1	25	97	ίuô	100	
	1	50	100	100	100	
Cr(AA) <sub>3</sub>	1	25	< 1.0	_	*	
-	1	50	5.5	10.7	72.2	
MnOct	2	25	-	2 5	۰ <b>.</b> ۲	
	2	50	8.3	22.2	23.2 58.3	
CoOct	1	25	11 3	_	35.0	
	1	50	23.6	-	48.7	
NiOct	1	25	7 8	-	0 0	
	1	50	19.8	-	20.1	
ZrOct	1	25	< 1.0	_	< 1.0	
	1	50	6.8	-	10.0	
Zr(AA) <sub>3</sub>	1	25	< 1.0		3 5	
	1	50	7.2	-	15.2	
CuOct	1	25	< 1.0		5 9	
	1	50	< 1.0	-	38.2	
Co(AA) <sub>2</sub>	1	25	12.0	-	48.0	
	1	50	27.7	52.7	83.8	
Co (AA) 3	1	25	< 1.0	-	-	
	1	50	11.0	-	23.5	

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# TABLE I (Continued)

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	Catalyst				,
Cateline /	Level	Temperature		% Reaction	<u>a/</u>
catalyst	<u>(wt. %)</u>	(°C)	<u>7 Days</u>	<u>14 Days</u>	<u>30 Days</u>
Fe(AA) <sub>2</sub>	1	25	< 1.0	-	44.4
	1	50	25.0	55.5	77.8
Fe (AA) 3	1	25	< 1.0	13.8	33.3
	1	50	15.7	-	61.7
Ce (AA) 3	1	25	0	0	0
	1	50	0	< 1.0	29.4
Ní (AA) 2	1	25	3.7	_	22.5
	1	50	9.8	-	62.7
Mn (AA) 3	1	25	5.0	-	41.5
-	1	50	23.6	38.8	> 99.0
SnOct	1	25	0	-	< 1.0
	1	50	< 1.0	5.7	13.8
V (AA) 3	1	25	0	-	< 1.0
	1	50	< 1.0	-	36.1
	2	25	0	-	2.5
	2	50	4.0	-	30.6
V0 (AA) 2	1	25	0	-	< 1.0
	1	50	< 1.0	-	< 1.0
	2	25	0	-	16.7
	2	50	< 1.0	-	58.7
VONapthenate	1	25	0	-	< 1.0
	1	50	< 1.0	-	2.0
	2	25	0	_	< 1.0
	2	50	< 1.0	-	2.3
FeOct	1	25	35.7	-	773
	1	50	75.0	-	84.0
	2	25	51.3	-	83 3
	2	50	81.3	-	86,1
In(AA)4	1	25	< 1.0	< 1.0	30.0
	1	50	-	36 1	1.1. 0

#### TABLE I (Concluded)

	Catalyst	Temporaturo	% Reaction		
Catalyst	<u>(wt. %)</u>	(°C)	7 Days	14 Days	30 Days
V-neo-decanoate	2	25	< 1.0	) en	8.9
	1	25	< 1.0	-	0
	1	25	< 1.0	-	5.3
Dibutyltin diacetatc	1	25	0	-	< 1.0
	1	50	0	82.	< 1.0
Tetra-n-butyltin	1	25	< 1.0	-	5.7
	1	50	10.5	-	20.3
Triphenyllead chloride	1	25	0	0	-
	1	50	0	t <u>c</u> /	-
Diphenyllead dichloride	1	25	0	0	-
	1	50	0	t	-
Tetraphenyllead	1	25	0	0	-
	1	50	0	t	-
Lead octoate	1	25	0	0	-
	1	50	0	t	-
Tetraphenylgermanium	1	25	0	0	-
	1	50	0	t	-
Cobaltocene	1	50	55.5/47.2	2 <u>b</u> / 66.6/-	÷

<u>a</u>/ Percent reaction is given as the average of the percent composition of the reactants within error limits.

b/ Deviation in stoichiometry noted; epoxide/acid.

 $\underline{c}$  /  $t \approx Trace$ .

The bulk of the other materials examined performed poorly as rate accelerators for the epoxide-carboxylic acid reaction. Elevated temperature is required to bring these reactions close to completion in 30 days. The ambient temperature runs had reacted only to the extent of 10 to 15% in 10 days. In general, the chelates (AA) are better catalysts than the salts for members of this group. There appears to be an induction period for some of these materials, especially chromium acetylacetonate. Mechanistically, this induction period may be the time required for the chelate to "open" and allow coordination of the reactants. Several of the materials falling into the "poor" category were reexamined with more attention being paid to examining the induction period. The results of the duplicate studies were in agreement with the previous results for the materials examined, i.e., they are poor catalysts, especially at ambient temperature. Doubling the amount of the materials used, e.g., manganese octoate, made no difference in the reaction rate at ambient and increased it only slightly at 50°C.

A series of vanadium and vanadyl ( $VO^{+2}$ ) derivatives were also shown to be poor rate enhancers for the AGE-EHA reaction.

Stannous octoate is reported to be a good catalyst for the anhydrideepoxide copolymerization reaction at low and ambient temperatures. In the AGE-EHA system (as well as Telagen CI and ERLA-0510) it is a poor reaction promoter. Several other tin compounds were investigated to determine if this class of materials (nontransition) can be as effective in the carboxylic ester system as they are in the urethane. The results, once again have been disappointing. Those materials examined displayed poor rate enhancement properties.

To recapitulate, chromium salts were shown to be the best rate enhancers for the epoxide-acid reaction, with iron salts providing an intermediate category of catalysts worthy of continued evaluation. All other materials tested, including chelates and salts, are poor rate promoters at ambient temperature compared to the first two categories. As will be discussed later, these results were also verified in the binder studies. The chromium and iron salts were then subjected to more detailed kinetic analysis as described below.

#### D. Baseline Interactions in the Model Epoxide Acid Systems

The intent of the model systems studies was to provide a rapid and simplified means for screening catalysts in a relative sense insofar as they affect the rate and stoichiometry of the reaction. In other words, the desired result of these screening studies was to examine only the reaction of the cure functionalities and any alteration in reaction rate

and/or direction should be due to the catalysts and other known variables selected for evaluation. To determine those parameters within the model reaction system which have significant effects on the rate and direction of the main reaction, a single factorial experiment was designed to establish the existence of any parameter interplay. Model compounds of several different structures were employed: 2-ethyl hexanoic acid, octanoic acid (OA), allyl glycidyl ether (AGE) and epoxy butane (EB). The results of this study are shown in Table II. The responses used for the factorial experiments were percent reaction occurring in a 24-hr. period and the stoichiometric factor as defined in Footnote B, Table II. It was anticipated that only temperature and the presence of catalyst would affect the rate and direction of the reaction. Indeed, this was shown to be the case. Using the Yates method for analyzing the results of a factorial experiment (Ref. 2), the one most significant parameter (within the context of the matrix design used) for determining reaction rate is the presence or absence of catalysis. Temperature is the most important secondary parameter; however, compared to catalysis it is not statistically significant (once again, this conclusion is based on the statistical evaluation of the total matrix shown in Table II). When the experiment is reconsidered in light of the factors of major importance in the catalyzed system, examination of data in Table 11 shows that temperature is the most significant parameter. Acid structure appears to be affecting reaction rate also, though based on the data available this cannot be stated unequivocally. When octanoic acid is the substrate rather than 2-ethyl hexanoic acid, reaction goes approximately 20% faster at room temperature. At 50°C the rates follow the same trend but are too close in value to be used as the basis for any firm conclusion. Uncatalyzed, AGE-EHA react at 100°C; in 48 hr., 13.8% reaction has occurred with a stoichiometry of 1.

Reaction stoichiometry appears dependent on the type of acid used, the alpha-substituted EHA giving only one product while <u>n</u>-octanoic acid consistently gave a minor product amounting to nearly 3% of the product in some cases.

Since the relationship of acid and epoxide structures to reaction rate and direction is one of obvious importance, in both a mechanistic and a practical sense, further study was given to the rate and stoichiometric parameters in the solution reaction system. The results of the initial factorial experiment suggest that reaction stoichiometry is a function of acid type rather than a catalyst parameter at least in the chromium salt system). Table III and Figure 1 show the reaction of a series of experiments designed to change the reaction rate in the octanoic acid-AGE system, by either thermal or diffusional means. Only one side product was observed and its amount was always less than 2% of the total yield. These values are within the range of experimental error and no significance can be attributed to variations among them. TABLE II

# BASELINE INTERACTIONS FOR THE EPOXIDE-CARBOXYLIC ACID REACTION

ane OA		EB/OA/RT/Cat 0.975 50.4 (0.025)	EB/OA/RT/No 0	EB/OA/50/Cat 0.964	98.5 (0.016) EB/0A/50/No -
Epoxy Buta	EB/EHA/RT/Cat 43.2 1	EB/EHA/RT/No 0 -	EB/EHA/50/Cat	92 I EB/EHA/50/No 0 -	
	idyl Ether <u>OA</u>	AGE/OA/RT/Cat 0.973 48.0 (0.027)	AGE/OA/RT/No O -	AGE/OA/50/Cat 0.990	99.3 (U.UIU) AGE/OA/50/No 0 -
Allyl Glyci EHA	AGE/ENA/RT/Cat 7. RXN <u>a</u> / Stoich <u>b</u> / 40.2 1	AGE/EHA/RT/No 0 -	AGE/ENA/50/Cat	95.9 L AGE/EHA/50/No C -	
Curing Agent Acid Structure Temperature (°C) Catalysis	Catalysis	CrOct Present	Absent	CrOct	Present Absent
	Temperature (°C)	ě	0		50

Fercent RXN in 24-hr. period as measured by the average consumption of starting material within experimental error limits.

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Stoichiometry is determined by the number of products noted in GC; product composition is determined value of 1, a mixture of 98% major product, and 1.5 and 0.5% minor products, respectively, would be reported as an overall stoichiometry of 0.98 with the amounts of secondary products noted in relative to total area for products only. Hence, only one product would give a stoichiometric purentheses, e.g., 0.980 (0.015, 0.005). <u>آم</u>

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TABLE :	II	I
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	RATE-SI	RATE-STOICHIOMETRY STUDY FOR OCTANOIC			
Catalyst Level (wt. %)	Agitation <sup>a</sup> /	Temperature (°C)	Reaction <u>Time (%)</u>	<u>Stoichiometry</u> <sup>b/</sup>	
1	None None None None None None	4	4 hr. 5.5 5 hr. 8.3 23 hr. 16.6 48 hr. 38.8 125 hr. 66.6 197 hr. 79.1 12 days 91.6	0.984 (0.016)	
2	Shaker Shaker Shaker	25	5 hr. 40.3 25 hr. 99+ 52 hr. 100	0.980 (0.020)	
1	Shaker Shaker	25	4.5 hr. 19.4 22.0 hr. 77.7 48.0 hr. 94.4	0.986 (0.014)	

 $\underline{a}$  / In the standard experimental procedures the reaction systems are neither stirred nor agitated.

b/ With recorder sensitivity at maximum only one side product was noted; values noted in table were derived from data taken under normal operating conditions.



The results of this experiment do indicate that product distribution under low temperature conditions is not a function of reaction rate.

The OA-AGE condensation reaction catalyzed by CrOct was then run under toluene reflux conditions, i.e., 110°C. In this situation only one side product was noted; however, it comprised 22% of the total yield. These results indicate that the reaction sequence leading to the formation of the side product is a higher energy process than that leading to the main product.

Acid molecular weight may also be a key factor controlling both reaction rate and stoichiometry in solution. The smaller the acid, the greater would be its diffusional rate and hence reaction rate. In addition a smaller acid (than OA or EHA) would have a greater tendency to overcome steric hindrance at various reaction sites. Hence, for an acid of molecular weight lower than AGE (or OA), the rate of conversion should be faster and a greater number of products should form. This was found to be the case for propionic acid (PA) Table IV). Figure 2 shows a comparison of the reaction rate profiles for PA-AGE under various conditions and that for EHA-AGE. Under comparable conditions, the propionic system undergoes complete reaction significantly faster than the ethyl hexanoic mixture. Reaction stoichiometry is good with the major component forming 99% of the product. A widely varying number of side products is formed under the different conditions described with the greatest multiplicity being produced at elevated temperature (see Table IV). It should be noted that these products are present as trace quantities and their total in any given PA system was always less than 1%. This trend was also verified with valeric acid (C5). In this case a total of two side products in less than 1% total quantity was noted.

Figure 3 is a schematic representation of the direction which can be taken by the catalyzed reaction of a monofunctional epoxide with monofunctional carboxylic acid. Compounds I and III are the expected major products of the reaction, that is, epoxides should be attacked at the least hindered carbon. Side products formed through attack on the other epoxide carbon are II and IV. Continued reaction of the hydroxyl group of the product ester with acid would lead to Compounds Ia, IIa, IIIa, and IVa. The results reported in Table II show that EHA gives only the major product while OA gives a mixture of products containing approximately 3% minor ingredient, which, for purposes of this discussion, will be assumed to be IV. The simplest explanation for this behavior is in terms of steric effects. EHA with its ethyl group adjacent to the reacting functionality will be directed toward the least crowded reaction path, i.e., the unsubstituted carbon of the epoxide. These steric effects may take on added importance since the reaction occurs through a catalytic intermediate. Octanoic acid does not possess this steric limitation and hence, a minor amount can "fit into" the reaction site around the substituted carbon of the epoxide.

TΑ	BLE	1 I	V

	RATE DATA F	OR PROPIONIC	ACID - AGE	
	REACTION	a/ CATALYZED	BY CrOct	
Catalyst				
Level	Temperature	Time	Reaction	. /
(wt. %)	(°C)	<u>(hr.)</u>		<u>Stoichiometry</u> <sup>b</sup> /
1	4	3.8	22.4	-
	4	22.5	36.7	*
	4	52.0	40.8	-
	4	144.0	75.5	0.99 + 2 minor
1	25	4.25	32.6	-
	25	28.0	88.7	0.99 + 3 minor
1	50	2.5	59.1	-
	50	5.5	86.7	-
	50	30.0	100.0	0.99 + 9 minor
2	25	4.5	54.0	-
	25	28.5	109.0	0.99 + 2 minor
2	50	2.5	91.8	-
	50	5.5	100.0	0.99 + 4 miror

 <u>a</u>/ Reaction system consisted of equimolar amounts of acid and epoxide and an amount of toluene equal to the weight of both reactants.

b/ Reaction stoichiometry - only one side product is observed with recorder operating under normal conditions. At maximum sensitivity, a multiplicity of side products can be found; these, however, account for less than 1% of the total product.

1 - 1 A 4 - 1 - 2







Should reaction continue through esterification of the product hydroxyl group, these steric factors would continue to govern product distribution. Ethyl hexanoic acid would be less inclined to react with its primary product because the hydroxyl group in question is in a secondary position along the carbon backbone, i.e., somewhat hindered. The primary, side chain hydroxyl group (as in II and IV) would seem to be a better target for continued reaction. Compound IV has not been observed; hence this continued reaction may not occur in the EHA system. For octanoic acid, both Ia and IIa are possible. Compound IIa may be sterically favored, but on a statistical basis more of Ia may form (since there is an abundance of I compared to II). The data reported in Table II show that only two products have thus far formed. The multiplicity of products formed when propionic acid or valeric acid is substituted for the Cg acids used in the factorial study tends to support this mechanistic hypothesis. Both propionic and valeric acids are unhindered and smaller than either OA or EHA, and would be even less subject to storic control. Hence, a greater number of side products can result.

Homopolymerization of the epoxide is another factor which was evaluated to insure accurate interpretation of the results from the model systems studies. A series of potential catalysts (CrOct, CrOleate, Crneo-decanoate, MnOct, Fe(AA)<sub>3</sub>, Cr(AA)<sub>3</sub>, NiOct) were placed in the reaction system less acid, i.e., allyl glycidyl ether or epoxy butanc in toluene, and subjected to 50°C. Compositions were monitored periodically by GC. No evidence was found for homopolymerization, i.e., no new products formed nor was there a decrease in the amount of epoxide content during a 14-day period. It was concluded that homopolymerization of the <u>model</u> epoxides in the presence of the catalysts does not occur under screening test conditions, i.e., 25° and 50°C in toluene solution.

Several conclusions about the nature and value of the solution condensation reaction can be made: (1) the reaction of AGE and EHA is stoichiometric and this system is suitable for the comparative screening of potential catalytic materials; (2) side reactions in the simplified model system are relatively unimportant; (3) there is a delicate balance among such parameters as acid molecular weight, carboxyl group environment, and product stoichiometry; and (4) the nature of the epoxide group, at least in the model system studies, appears to have little or no effect on the outcome of the reaction.

#### E. Summary of Catalyst Screening and Selection Studies

A large number of organometallic materials were screened in model system studies to assess their utility as potential cure catalysts for the carboxylic acid-epoxide binder. The materials fall into three categories with catalytic activity. Chromium salts (e.g., octoate, neodecanoate, oleate) were found to be excellent catalysts for converting carboxylic acid and epoxide to ester. They gave complete stoichiometric reactions at room temperature in less than 7 days (in model systems). An intermediate group, composed of iron salts, gave approximately 70% reaction completion in 14 days at room temperature with 99% stoichiometry at this level of reaction. There is some question about the overall utility of the iron salts as cure catalysts because of the rate plateau observed and the deviation from stoichiometry observed during the later stages of reaction. All of the other materials evaluated as potential catalysts for the epoxidecarboxylic acid reaction were found to be poor rate accelerators at room temperature. In general, to reach 50% conversion a time period of 30 days at 50°C was required.

Hence, the chromium salts were selected for further development with CrOct as the major candidate. This material shows a good balance between solubility and chromium content, and it is commercially available at a reasonable cost. The chromium salts in general have thus far met the criteria for a cure catalyst in the epoxide-carboxylic acid binder system. They are catalytically active (once again especially CrOct) in low concentrations and give the desired reaction stoichiometry. As the result of this evaluation, CrOct was selected for evaluation in binder gumstocks and processing studies at ABL.
#### CATALYST DESIGN AND REACTION MECHANISMS STUDIES

#### A. <u>Objective</u>

The objectives of these studies are to understand the factors controlling the behavior of the cure reaction (i.e., rate and direction) and the utilization of this information to optimize cure catalyst design. The mechanism and kinetics of the cure reaction is affected by such parameters as the nature and structure of the catalyst, the structures and steric environment of the reacting functionalities, temperature, contaminants, and type and level of solids loading.

#### B. Chromium Salts Studies

Based on the results of the screening studies, the chromium salts and soaps were selected for a more detailed evaluation of their chemistries as catalysts, their behavior in the presence of other propellant components and contaminants, and their behavior in binder and propellant systems. Of the commercially available chromium salts listed in Table V, chromium 2-ethyl hexanoate gave the greatest degree of rate enhancement. Examination of Table V gives an insight into the reasons for the differences in rates among the various chromium salts. It can be seen that the octoates as compared to the other commercially available chromium salts shown in Table V have higher percentages of chromium metal present per mole of material. Rate profiles for this and other chromium salts as functions of temperature and catalyst loading are shown in Figures 4, 5, and 6. The rate behaviors of the various chromium salts follow the same general pattern. As would be expected, the slowest segment of the reaction occurs during the last 10% conversion. The difficulty of this final conversion increases with decreasing amounts of catalyst. The reaction rate during the latter stages of the conversion is diffusion controlled. Hence, rates become increasingly slower as more of the reactants are consumed. Diffusion of curative and catalyst is an even more important rate parameter in the binder system than in the solution reaction. Continued reaction creates a aolid matrix and the hindrance to reactant diffusion is drastically increased compared to the situation in the test solution. The catalysis of the final segment of the reaction should be enhanced by a catalyst of low molecular weight, which would be better able to diffuse throughout the matrix.

#### TABLE V

#### CHARACTERISTICS OF CHROMIUM SALTS

	Structure	<u>Wt. % Cr</u> ª/	Solubility <u>b</u> ' <u>Characteristics</u>
Oleate <sup>c/</sup>	C <sub>8</sub> H <sub>17</sub> CH=CH(CH <sub>2</sub> )7CO <sub>2</sub> 3Cr	5.7	Excellent
ne Decanoate <u>c</u> /	(CH <sub>3</sub> (CH <sub>2</sub> ) <sub>5</sub> C(CH <sub>3</sub> ) <sub>2</sub> CO <sub>2</sub> ) <sub>3</sub> Cr	9.2	Excellent
Decanoate	(CH <sub>3</sub> (CH <sub>2</sub> ) <sub>8</sub> CO <sub>2</sub> ) <sub>3</sub> Cr	9.2	Acceptable
Octanoate	(CH <sub>3</sub> (CH <sub>2</sub> ) <sub>6</sub> CO <sub>2</sub> ) <sub>3</sub> Cr	10.7	Acceptable
2-Ethyl Hexanoate <u>C</u> /	$(CH_3(CH_2)_3CHCO_2)_3Cr$ $c_2H_5$	10.7	Goud
Heptanoate	(CH <sub>3</sub> (CH <sub>2</sub> )5CO <sub>2</sub> )3Cr	11.9	Good-Acceptable
Hexanoate	(CH <sub>3</sub> (CH <sub>2</sub> ) <sub>4</sub> CO <sub>2</sub> ) <sub>3</sub> Cr	13.2	Poor
Acetate <mark>c</mark> /	(CH <sub>3</sub> CO <sub>2</sub> ) <sub>3</sub> Cr	22.7	Insoluble
Formate <sup>c</sup> /	(HCO <sub>2</sub> ) <sub>3</sub> Cr	27.8	Insoluble

- <u>a</u>/ Stoichiometric amount; weight percent Cr in compound. Soaps are also available containing diluent or residual acid; in that case Cr content is given as the percent of the total sample (including diluent).
- $\underline{b}$ / The evaluation of salt solubility is qualitative, based on ease of handling and solution of the catalyst in polymers as well as common solvents such as toluene, ethanol, acetone, methylene chloride, and tetrahydrofuran.
- c/ Commercially available.



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Figure 6 - The Effect of Temperature and Catalyst Quantity on Cr Oleate Catalyzed Reaction of AGE and EHA

These results suggest at least three key rate determining parameters for the chromium soap catalysts: (1) chromium content of a given compound (the higher the metal content the faster the conversion, e.g., CrOct vs. CrOleate); (2) solubility of the Cr material in the reaction medium (binder mix or solution); and (3) as will be discussed below, steric environment around the carboxyl group of the organic portion of the metal salt. Table V shows some of the structural characteristics of a number of Cr salts.

Theory would suggest that the steric environment within the catalyst itself may have an effect on the conversion rate of the acid-epoxide reaction. Table VI lists rate data for the reaction of AGE and EHA catalyzed by chromium heptanoate and hexanoate. The chromium heptanoate analyzed at 8.45% chromium (11.80% is theoretical), indicating the sample to contain 28.3% free heptanoic acid.

#### TABLE VI

	<u>Catalyst</u>	Catalyst Level (wc	Temp. (°C)	Time (hr.)	<u>% Rxn</u>
Cr	Heptanoate <u>a</u> /	1	25	19	42.6
				22	50.0
				43	83.8
				48	97.1
		1	5Û	1	13.2
	•			3.8	70.5
				5.8	82.4
				23	··}.0+
$\mathbf{Cr}$	Hexanoateb/	1	25	21	13.2
				45	27.9
				69	45.5
				30 days	100.0
		1	50	3	33.8
				4	38.2
				6	47.1
				23	99.0+

#### CATALYSIS BY Cr HEPTANOATE AND HEXANOATE

<u>b</u>/ Sample contained 43% titratable acid; hence, Cr content is 5.67%.

<sup>&</sup>lt;u>a</u>/ Sample contained 71.7% salt and 28.3% free acid; hence, catalyst sample contained 8.46% Cr.

Table VII shows a comparison of rate data for identical AGE/EHA reaction systems catalyzed by chromium octoate and chromium heptanoate. Consider the following relationships:

Catalyst ratio:  $\frac{\text{CrHep}}{\text{CrOct}} \equiv \frac{7.1}{9.2} = 0.7/1$ 

Time to 50% conversion:  $\frac{\text{CrHep}}{\text{CrOct}} = \frac{22}{20} = 1.1$ ; inverse 0.91

Comparing the ratios for a situation of equal 50% conversion times, 0.845 mole of chromium heptanoate would equal 1 mole of chromium octoate in catalyzing ability. This exercise may oversimplify the kinetic situation; however, suffice it to say that chromium heptanoate gives better rate acceleration than chromium octoate on a mole-for-mole basis in a completely miscible system.

#### TABLE VII

#### RELATIONSHIP OF CATALYST QUANTITY AND RATE FOR Cr OCTOATE AND HEPTANOATE

<u>Catalyst</u>	Time to 50% Conversion (hr.)	Amount of Catalyst Present (moles)	Grams <u>Cr Present</u>
CrOct	20	9.2 x 10 <sup>-5</sup>	0.0047
Cr Heptanoate	22	$7.1 \times 10^{-5}$	0.0037

The difference in activity displayed by the two catalysts may be due to the presence of the ethyl group at the two-position in the CrOct organic ligand which would hinder (relative to the unsubstituted situation) the approach of the reactant carboxylic acid group to the chromium nucleus for activation purposes. Hence, a given acid can reach the chromium easier in chromium heptanoate than it can in chromium octoate (the octoate group being 2-ethyl hexanoate).

The low purity of the bexanoate sample precluded a similar comparison to other chromium catalysts.

#### C. <u>Substrate Steric Parameters</u>

Early work on this program suggested that reaction rate and direction were functions of acid type rather than of catalyst parameters.\* Further studies have supported the acid structure rate contention; however, the nature of the organic group in the metal catalyst has also been shown to alter conversion rate.

Concerning acid steric control, two mixed-acid systems catalyzed by a 1% loading of CrOct were examined: (1) equimolar amounts of octanoic acid and 2-ethyl hexanoic acid were reacted with an equivalent amount of allyl glycidyl ether (equivalent amounts of total acid and epoxide); (2) equimolar amounts of 2-ethyl hexanoic and propionic acids with an equivalent amount of allyl glycidyl ether. The rate data for these reactions are given in Table VIII, and Figures 7 and 8 show rate profile plots for these results.

The least sterically hindered acids (OA, PA) and the lower molecular weight acid (PA) always reacted faster than the  $\alpha$ -substituted EHA. The most revealing comparisons result from an examination of the conversion data for the OA and EHA experiment where molecular weight is not a competitive parameter. Table IX shows the data for time to 50% conversion of the two acids with AGE in the presence of CrOct.

Analysis of these data shows correlation of reaction rate with steric effects. In the mixed acid system OA reaches the 50% consumption point in 5 hr. When OA is the only acid, 2% catalyst loading is required for the same equivalence level ( $CO_2H$ /epoxide) to give 50% conversion of OA in the same time period. Hence, the steric factors are compelling enough to virtually exclude EHA from activation by the catalyst until OA is consumed (hence, OA "sees" the catalyst level as 2% instead of 1).

<sup>\*</sup> Q-Substituted acids give slower reaction rates than unsubstituted ones but give only one product while the unsubstituted acid gives a side product amounting to 2-5% depending on conditions.

TABLE	VI	II
-------	----	----

Temperature (°C)	Catalyst Level	Time (hr.)	<u>%</u>	Reaction	<u>•</u> /
PA/EHA Series			AGE	PA	EHA
25	1	5	36.5	50.0	30.9
		24	85.4	95.2	76.7
		31	86.6	97.6	76.2
		49	95.1	100	90.5
50	1	1.5	46.3	62.0	35.7
		3	76.8	90.5	64.3
		6	87.8	95.2	80.9
		24	99 <del>1</del>	00ء	99+
25	0.4	4.5	9.8	28.5	0
		26.5	45.1	61.9	35.7
		50.0	69.5	80.9	61.9
		76.0	81.7	100	69.0
		9 days	99+	100	99+
50	0.4	1	10.9	23.8	0
		2.5	14.9	28.5	0
		29	95.1	100	90.5
		53	98.7	100	97.6
		72	99+	100	99+
			AGE	<u>OA</u>	EHA
OA/EHA Series					
25	1	5	30.5	44.4	11.1
		25	73.6	91.6	55.5
		31	80.5	94.4	66.6
		52	92.7	100	85.5
		74	97.2	100	ે⊸ 4
50	1	2	47.2	66.0	22.2
		4	70.8	91.5	50.0
		25	99+	100	99+
25	0.4	26	41.6	55.5	16.6
		48	59.7	77.7	44.4
		51	62.5	83.3	50.0
		76	75.0	91,0	63.5
		9 days	94.4	100	88.8
50	0.4	2	22.2	27.7	16.6
		4	41.6	/2.2	10.0
		24	91.8	100	83.3
		29	A7.0	100	00.1 ga c
		UC ( 0	94.4 65.3	100	00,0
		+0 73	7/.4	100	100

REACTION OF AGE WITH MIXED ACIDA/ CATALYZED BY CrOct

a/ Reaction mixtures consisted of equivalent amounts of epoxide and acid functionalities in the following molar ratios: hACE to 500 to 500

1 AGE/0.5PA/0.5EHA and 1 AGE/0.50A/0.5EHA

<u>b</u>/ Percent reaction is given as the percent conversion of the given reactant; % AGE = % CONVERSION

$$7 \text{ AGE} = \frac{7 \text{ Acid}_{A} + 7 \text{ Acid}_{B}}{2}$$



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Rea <u>Sys</u>	ction tem <u>a</u> /	Temp. <u>(°C)</u>	Catalyst Level <u>b</u> / Wt. %	ד -	ime to 50% Conversion (hr.)
AGE +	OA	25	1		11.0
		25	2		5.5
AGE +	EHA	25	1	OA	20.0
AGE +	EHA/OAC/	25	1	EHA	22.5

#### RATE DATA FOR CONVERSION OF VARIOUS ACIDS WITH CrOct

a/ Equivalent amounts of acid and epoxide in toluene solution.

b/ Catalyst level in wt. % of catalyst based on total reaction solution (including tolugne).

<u>c</u>/ Reaction mixture consisted of equivalent amounts of epoxide and acid functionalities in the following molar ratios: 1 AGE/0.5 EHA/0.5 OA.

The following conclusions may be drawn concerning substrate acid structure rate effects: (1) where acid molecular weight is not a competitive parameter, as in the OA-EHA situation, steric crowding about the carboxylic acid group is a rate as well as stoichiometry controlling factor; (2) this conclusion can be extrapolated to the polymer situation, that is, where different types of acid groups are present in the prepolymer these will be consumed at different rates, and since binder reactions do not go to completion, the identity of the unreacted group can be predicted; (3) the nature of the catalytic intermediate (in at least the Cr system) is such that  $\alpha$ -carbon substituents in the reactant acid are in proximity to the activation site (further supporting the contention that it is the acid which is activated during the catalysis step); (4) in terms of catalyst design, less bulky substituents in the organic portion of the catalyst may facilitate the reaction rate (a point discussed below); and (5) metal catalysis does not level the reaction rates of different carboxylic acid groups. Hence, in addition to acting as a carrier for the metal, the ligand plays a role in controlling reaction rate.

#### D. Variable Substrate Concentration Studies

To gain further insight into the mechanism of the catalyzed conversion and to determine the nature of those reactions which may be important in a post-cure sense, studies were undertaken in which the initial amounts of reactants were varied. 1. Excess epoxide: Three separate systems were examined: (1) EHA and twice the number of equivalents of AGE, (2) OA and twice the equivalents of AGE, (3) separately prepared ester from AGE-EHA with an equivalent amount of AGE. The results of the reactions are shown in Table X. In the presence of CrOct, the carboxylic acid-epoxide addition reaction is exclusive at 25, 50 and most probably 100°C. This can be seen from the GC analysis in which acid and epoxide are consumed at equivalent rates until all acid is consumed and no additional products are noted up to this point relative to behavior of these two materials in the standard reaction system (equivalent initial amounts of reactants). The rates of reaction are those expected for the catalyst/reactant (acid) molar ratio used. These results tend to support the notion that the key catalytic interaction involves the carboxylic acid and the chromium salt.

Beyond the total acid consumption point, postesterification epoxide consumption (with catalyst present) is very temperature dependent. Epoxide loss does occur to a limited extent at 50°C (~ 15-20%) in 40 days. The fact that only trace products were noted in the GC is most likely due to the fact that they are of high molecular weight and not sensitive to the method of analysis used. At 100°C nearly complete epoxide loss is noted, accompanied by polymerization of the reaction system. Postesterification reaction at 25°C (in presence of Cr) is minor (~ 13% in 40 days). The control experiment (ester/AGE/no catalyst/100°C) showed only minor epoxide consumption (11% in 35 days) while similar reactions with Cr catalyst present resulted in polymerization and high (85%) epoxide loss in the same time interval.

Several conclusions may be drawn: (1) in the presence of diffusible (excess or residual) epoxide, further reaction between epoxide and ester product will occur (unreacted epoxide will be present in a propellant matrix, however, it will be attached to a polyfunctional curative incorporated into the binder network and will be nondiffusable or only slightly so in a reactivity sense); (2) the results (exclusive acid-epoxide reaction until total consumption of acid) support the suggestion that the key catalytic interaction involves carboxylic acid and the chromium species; (4) chromium catalyst promotes the postesterification reaction of epoxide and hydroxyl group of the product, and (5) this postesterification reaction is important only at elevated temperatures (100°C) and does not appear significant under ambient conditions.

2. Excess acid studies: The effect of excess acid on the rate and direction of the CrOct catalyzed cure reaction was examined in model systems employing EHA-ACE mixtures 1.5:1 and 1.2:1 equivalent proportions at 25° and 50°C. As in the excess epoxide case, reaction is stoichiometric, i.e., acid and epoxide disappear at equal rates. Unlike the excess epoxide situation, however, no post-esterification reaction occurs in 35 days after completion of the main reaction at either 25° or 50°C. Rate patterns (see Figure 9) are within experimental error of the equivalent acid-epoxide reaction.

#### TABLE X

- Participation of the Partici

## DATA SUMMARY

## Excess Epoxide Experiment

A. Early React	ion Data				bservation
			Temperature	Product Formation	Epoxide Consumption Beyond Complete
System <sup>a</sup> /	<u>Catalyst</u>	Time	<u>(°C)</u>	(%)	Esterification
EHA (1)/ACE (2)	1% CrOct	6.8 hr.	25	32	-
		25.5 hr.	25	100	0
		7.0 days	25	100	13.0
		2.5 hr.	50	100	-
		24.5 hr.	50	100	0
		7.0 days	50	100	8.6
		1.0 hr.	100	100	8.6
		2.0 hr.	100	100	8.6
		3.0 hr.	100	100	8.6
		7.0 days	100	100	21.7
		13.0 days	100	100	43.4
OA(1)/ACE(1)	1% CrOct	6.5 hr.	25	58	-
		26.0 hr.	25	100	6.7
		9.0 days	25	100	8.5
		2.8 hr.	50	100	1.3
		25.0 hr.	50	100	6.2
		7.0 <b>day</b> s	50	100	13.0
		1.0 hr.	100	100	2.2
		2.0 hr.	100	100	6.7
		3.0 hr.	100	100	11.6
		20.0 hr.	100	100	13.0
		o.0 davs	100	100	24.6
		13.0 days	100	100	29.6
EHA Prod/ACE	1% CrOct	3.0 hr.	100	-	5.8
		20.0 hr.	100	-	44.7
		6.0 days	100	-	11.7
		13.0 days	100	-	14.7
		4.3 hr.	50	-	0
		25.4 hr.	50	-	0.8
		7.0 days	50	-	2.0
		6.0 hr.	25	-	0
		27.0 hr.	25	-	0
		9.0 days	25	-	0

 $\underline{a}$  / Numbers in parentheses refer to relative molar ratio of the two components.

## TABLE X (Concluded)

# B. Surveillance of Above Experiments

				Epo	xide Consump	tion Beyond
	Catalyst	Temperature	Time		Ester Form	ation
System	<u>(Wt. %)</u>	<u>(°C)</u>	Days	<u>G.C.</u>	<u>Titration</u>	Comment
EHA(1)/	1% CrOct	25	7	13.0	-	
ACE(2)		25	22	14.0	-	No side prod.
		25	41	14.0	13.8	No side prod.
		50	7	8.6	-	
		50	22	15.3	-	
		50	40	20.5	22.3	Trace side prod.
		100	13	43.4	-	
		100	16	56.5	-	
		100	40	-	90.0	Polymerized
QA(1)/	1% CrOct	25	9	8.5		
ACE(2)		25	22	13.0	-	
		25	41	18.5	13.0	No side prod.
		50	7	13.0	-	
		50	22	15.0	-	
		50	40	-	17.7	
		100	13	29.0	-	
		100	16	80.0	-	2 major side prod.
		100	20	85.0	-	Polymerized
		100	40	-	92.6	Polymer
EHA Prod/	1% CrOct	28	9.0	0	-	
ACE		<b>2</b> 5	20.0	2.0	-	
		25	41.0	4.8	6.4	
		50	7.0	2.0	-	
		50	22.0	2.8	-	
		50	40.0	13.0	14.7	
		100	13.0	14.7	-	
		100	16.0	27.7	-	
		100	20.0	<b>3</b> 5.0	-	Side prod.,
		100	40.0	_	100 0	brown, viscous Bolumerized
		100	40.0		100.0	rorymerraed
Control:	No	100	1.0	0	-	
EHA Prod/	catalyst	100	5.0	0	-	
AGE		100	8.0	2.8	-	
		100	9.0	3.5	-	
		100	30.0	-	11.1	Slight yellow- ing, no polymer- ization

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3. <u>Rate dependence on catalyst concentration</u>: The dependence of reaction rate on catalyst quantity is shown in Figure 10. The data from an experimental series in which catalyst concentration was the only reaction variable are shown in Table XI. Initial concentrations of reactants (epoxide and acid) are the same; the reaction is stoichiometric, i.e.,  $-d[CO_2H]/dt = -d[epoxide]/dt = +d[product]/dt$ . The rate is expressed as the number of moles of product formed per hour at 50% conversion. Figure 3 is a log-log plot\* of the reaction rate versus the ratio of molar concentrations of catalyst to reactant at fixed initial reactant concentrations. The plot is linear with a slope of one indicating first order dependence of reaction rate on catalyst concentration. Mechanistically, these results suggest the existence of a reactant-organometallic intermediate in the rate-controlling step for the conversion and are in agreement with the spectroscopic evidence indicating coordination of the chromium salt with carboxylic acid.

\* The log-log plot is based on the following derivation:

 $A + B + Cat \longrightarrow Products$   $\frac{d[c]}{dt} = k [acid] [epoxide] [cat]$  k, [acid] and [epoxide] are constant  $\vdots k [acid] [epoxide] = K$   $\frac{d[c]}{dt} = K [cat]$   $let \frac{d[c]}{dt} = R$   $R = K [cat] \times$  log R = x log [cat] + log K

The relationship is now expressed in straight line form, and x is the slope.



TA BLE XI

March 1. States

# KINETIC ANALYSIS RATE DEPENDENCE ON CATALYST CONCENTRATION

	Moles of	Components	Molar Ratio Catalyst	Log Molar	Rate <u>a</u> / d[c]/dt	I.oe
NIE	AGE	CrOct	Reactant	Ratio	Moles/hr x 10 <sup>4</sup>	Rate
0.0086	0.0086	4.60 x 10 <sup>-5</sup> (0.5) $\frac{b}{b}$	0.0053	$\frac{-}{2.2758}$	1.43	$\frac{-}{3.38448}$
0.0086	0.0086	$9.24 \times 10^{-5}$ (1)	0.0107	$\overline{1.9706}$	2,15	3.6676
0.0086	0.0086	$1.85 \times 10^{-4}$ (2)	0.0215	T.6676	5.73	<u>3</u> .2419
0.0086	0.0086	$3.69 \times 10^{-4}$ (4)	0.0429	$\overline{1.3675}$	15.10	$\overline{2}.8211$
0.0086	0.0086	$5.54 \times 10^{-4}$ (6)	0.0644	T. 1911	24.00	<u>7</u> .6198
0.0086	0.0086	$9.24 \times 10^{-4}$ (10)	0.1074	0.9690	43.00	$\overline{2.3666}$

Rate = moles of product formed per hour in 0-50% reaction range.

Number in parentheses refers to catalyst levels in reaction system expressed as weight percent. <u>e la</u>

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#### E. Ferric Octoate Studies

From the results of the screening studies, ferric octoate appeared to be a good rate accelerator for the AGE-EHA reaction. However, a more detailed examination of its rate acceleration behavior shows that after approximately 60 to 70% conversion, a rate plateau is reached. This effect is noticeable at both 1 and 2% by weight catalyst levels, and elevated as well as ambient temperature. This thermal effect is strongly suggestive of a loss of catalytic activity by the iron compound. Indeed, examination of the sample vials shows a gelatinous precipitate present in the 2% sample and a color change in the 50°C runs. Of the msterial in solution, six minor products (each in trace quantities) 'amounting to approximately 2% of the totsl product were noted. The rate data for FeOct as a catalyst are shown in Figure 11; Table XII displays the results of the various reaction series run to examine the FeOct rate phenomena and to check its reproducibility. Analysis of the initial series after 60 and 88 days shows nearly total consumption of the epoxide while 20 to 30% of the acid remains unreacted. The side product formed by the extra consumption of epoxide is the precipitate observed previously, since GC analysis of the dissolved material shows no eccraordinary deviations in reaction direction.

The side reaction behavior was noted also in solutions contaminated with water. Again, epoxide was consumed faster than acid and the threshold for this deviation in stoichiometry appears at 60 to 65% conversion.

There are at least two possibilities which would explain the side reaction: (1) an epoxide homopolymerization reaction which becomes important during the later stages of cure where the rate of ester formation slows down due to the decrease in reactant concentration and ensuing importance of diffusion; (2) reaction of epoxide with the catalyst itself due to the decrease in the amount of activated reactant acid.

An exploratory study was made using a mixed FeOct/CrOct catalyst (0.5/ 0.5% weight). The reaction behavior (that is, rate and stoichiometry) observed was that expected of the CrOct catalyst at the 0.5% level with a slight acceleratory assist from the FeOct (see Table XIII), thereby confirming the dominance of the catalytic effect of Cr even in the presence of potential reaction-promotion competitors.



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## TABLE XII

## FeOct STUDIES

		Leve 1	Temperature	Time	Reaction	
Å.	Rerun Series	2%	RT	2 hr. 26 hr. 3 days 6 days	> 1 4.8 32.9 52.9	
				30 days	Acid 65.2 Epoxide 77.7	
		2%	50°C	2.5 hr.	> 1	
				1 days	60.0	
				6 days	70.0	(epoxide 74.2)
				30 days	[ Acid 72.2	(0)000000000000000000000000000000000000
				50 0095	Epoxide 91.6	
в.	Initial Series	2%	50°C	60 days	Acid 69.4	
					Epoxide 97.2	
		2%	RT	88 days	Acid 75.0	
					Epoxide 97.2	
		1%	50°C	60 days	Acid 69.4	
					Epoxide 91.6	
		1%	RT	88 days	Acid 66.6	
					Epoxide 79.1	
С.	H <sub>2</sub> O Effect Studies <u>Catalyst Level</u>					
	2%	18 µ1.	50 <b>°C</b>	2.5 hr.	> 1	
				24 hi.	28.5	
				3 days	55.5	
				6 days	61.1	
				33 days	Epoxide 95.8	
	27.	18 µl.	RT	2 h <b>r</b> .	>1	
				26 hr.	19.4	
				3 days	34.7	
				6 days	44.4	
				33 days	∫ Acid 72.2	
					Epoxide 77.7	

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## TABLE XIII

## MIXED METAL OCTOATE CATALYST STUDIESa/

<u>Cemperature</u>	Time	Reaction (%)	Analogous 0.5% CrOct Reaction
RT	l hr.	t	
	1 day	47.2	(43.9)
	3 days	86.1	(80.2)
	6 days	91.6	(94.3)
50 °C	25 hr.	30.6	
	1 day	77.7	
	3 days	99+	
	6 days	100	

a/ Catalyst in this case was an equal weight mixture of CrOct and FeOct, that is, 0.5% CrOct and 0.5% FeOct by weight of the total reaction solution.

#### EXTRA INGREDIENT EFFECTS ON THE CHROMIUM SALT SYSTEMS

#### A. Propellant Components

The chromium salts are excellent catalysts for the carboxylic acidepoxide reaction. To be successful as cure catalysts, they must demonstrate the same type of catalytic behavior in the propellant environment; i.e., they must promote the esterification reaction in the presence of other potentially reactive ingredients. CrOct, CrOleate, and Cr-neo-decanoate were selected for this evaluation, the octoate being the prime candidate while the other two are representative of the other types of chromium salts available.

1. <u>Ammonium perchlorate</u>: The introduction of 5% by weight of ammonium perchlorate into the standard test system (ACE-EHA-toluene-CrOct) does not alter the rate or stoichiometric outcome of the condensation reaction at either ambient temperature or 50°C. These data are presented in Table XIIIa and plotted relative to the behavior of the reaction system without AP in Figure 12. All points fall on their respective temperature plots. It is concluded that AP, in this weight percent range at least, does not alter the outcome of the solution reaction. The completed reactions containing ammonium perchlorate, catalyst, and product ester were set aside and monitored throughout the course of the program to determine the long-range effects of AP in the presence of CrOct on the stability of the ester linkage.

2. Additional propellant solids: Table XIV lists several other propellant ingredients whose effect on the catalyzed AGE-EHA reaction was examined. One surfactant and two antioxidants were included: Roccal (a quaternary ammonium salt), AO-2246, and phenyl-B-naphthylamine, respectively. The results of these experiments are displayed in Figures 13 and 14. No alteration in reaction direction was noted for the reactions catalyzed by CrOct and Cr-neo-decanoate. Considering the CrOct case first, the rate of ester formation was only slightly accelerated by the presence of the extra ingredient, and in many cases, it is doubtful whether the differences relative to the baseline cure (CrOct alone) are real. Only phenyl-B-naphthylamine appears to give a definite reaction rate increase (12% more conversion after 24 hr.).

An acceleratory effect by the antioxidants would not be surprising, especially at the concentration levels being used (0.5% by weight, or half as much by weight as the cure catalyst). The antioxidants are amines, and hence have catalytic properties toward the acid-epoxide reaction.

## TABLE XIIIa

	Temperature	Time	Reaction	
		lime	(/_)	Stolchlometry
A. 0.1% catalyst	25	1 day	13.8	1
level; <u>b</u> /	25	2 days	16.6	1
5% AP <u>b</u> /	25	3 days	26.3	1
	25	7 days	41.6	1
	25	10 days	48.6	1
	25	17 days	70.8	1
	50	6.5 hr.	13.8	1
	50	24.0 hr.	38.8	1
	50	5 days	77.7	1
	50	9 days	83.3	1
B. 1.0% catalyst	25	5.5 hr.	25.0	1
level;	25	24.0 hr.	62.5	1
5% AP	25	48.0 hr.	80.5	1
	25	4 days	94.4	1
	25	7 days	100.0	1
	50	2.5 hr.	41.6	1
	50	5.0 hr.	62.0	1
	50	6.5 hr.	69.4	1
	50	24.0 hr.	97.2	1
	50	48.0 hr.	100.0	1

## THE EFFECT OF AMMONIUM PERCHLORATE ON THE CrOct CATALYZED REACTION OF AGE AND EHA<sup>a/</sup>

a/ Standard toluene stock solution.

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b/ Weight percent based on weight of stock solution used.



#### TABLE XIV

#### EXTRA-INGREDIENT COMPATIBILITY

Ingredient and	Catelyst and				
Concentration Level (wt. % of total <u>mixture</u> )	End Use of Material	Concentration Level (wt. % of total mixture)	Tims	Temparature (°C)	Percent Reaction
0.51.2000	aurfectant	17 CrOct	15 min	25	> 1
0.3% Roccar		14 0000	24 hr.		65 7
			6 days		99 +
0 FN B 1		17 Chanadada	5 5 hr	25	> 1
0.5% Roccal		1% CE-NBO-GEC	5	• ·	38.5
			S P P P		77.1
			: leve		84.2
			5 T <b>Y</b> B		85.7
		18 0-0.0	7 5 6 .		61.6
0.5% phen/1-8-N	entioxident	1% Groet	1.5 nr.	15	36.3
			6 Java	6.3	99.4
			7 days		100
				16	11.4
0.5% phenyl-g-N		1% Cr-neo-dec	) hr.	43	11,4
			24 hr.		40,0
			4 days 5 days		97.1
			6 deve		90.0
			7 deys		91,4
18 43		18.0-0-1	1 h-	75	21
17, 81	tuel	1% Croci	1 nr. 25 hr	23	68.5
			6 dave		99 +
			7 days		100
1* +1		17 0	6 hr	15	10.0
14 AI		14 Graneoadec	4 nr. 76 hr	4.2	20,0
			20 HEL 6 dave		77.1
			5 devs		84.3
			6 deye		88,6
17 oxemid	ballistic	17. CrOct	l.hr.	25	11.4
	modifler		24 hr.		65.1
			6 days		99.+
			7 dava		99 +
17. oxemid		1% Cr-neo-dec	7 hr.	25	12.0
			24 hr.		.2.9
			4 dæyø		82.6
			5 deys		83
			6 deys		87.1
0.5% A0-2246	antioxidant	1% Cr-neo-dac	6.5 hr.	25	17.1
			24 hr.		5-1.2
			4 deys		82.9
			5 days		40.0
			7 джун		۹1 ~
0,5% Fe <sub>3</sub> 0 <sub>3</sub>	ballistic	No cetalyst	33 dave	25	> :
<b>A</b> 3	modifier	No catalyst	t4 deye	50	15.2
		No cetalyst	33 dave	50	19.~
1.02 Fr301		No cetelyst	33 days	25	• 1
÷ 2		No cetalyst	14 deve	50	9,7
		No catalyst	33 dave	50	19.8
1.02 80.01		17, CrOct	1 hr.	25	ø
< )		-	1 h <b>r</b> .	25	12.2
			4 hr.	25	27.*
			5 hr.	25	41 в
			26 hr.	25	66.2
			4 de ys		99 <del>v</del>
			6 deye	2.5	44 •
			11 dava	25	10012 mt

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However, even at these high concentration levels compared to the actuel propellant situation, the antioxidants have no significant effect on the rate and direction of the cure reaction as compared to the effect of the catelyst itself.

The same can be said for the presence of eluminum, iron oxide, and oxamid; as expected, they have no intrinsic effect on the reaction (though iron oxide by itself at elevated temperatures and after extended time periods displays some slight catalysis for the reaction).

In Figure 14, the rate data are plotted for the Cr-neo-decanoate system. The acceleratory effect of the extra ingredient, once again, is minor but more pronounced than in the CrOct catalyzed reaction. This is not unexpected since Cr-neo-decanoate is a slower catalyst than the octoate (the neo-decanoate sample used here has roughly half the amount of chromium as the octoate) and, the extra contribution to the conversion rate by the other components is more noticeable.

#### B. <u>Contaminants</u>

The extra-ingredient compatability survey was extended to include the following potential contaminants: water, perchloric acid (which may be introduced by the oxidizer), nitric acid, ammonia, and <u>n</u>-propyl nitrate (as a model for nitrate ester).

1. Water: The effect of water on the model system is revealing in a mechanistic sense. Standard CrOct reaction systems (containing 0.01 mole of acid and epoxide) were contaminated with 9, 18, and 90  $\mu$ l. of water. These water concentrations correspond to 0.0005, 0.001, and 0.005 mole of water, respectively. The rate data are listed in Table XIVa and plotted in Figure 15, relative to the rate profile for the reaction without added moisture. It can be seen that there is an initial rate enhancement corresponding to the amount of water present. The concentration effect diminishes in importance as the reaction proceeds, and water contamination does not significantly change the total time required for a complete reaction, nor does it alter the stoichiometry.

To provide a baseline for isolating the specific effect of water on the reaction, the behavior of the epoxide alone, and epoxide and catalyst with water, were examined. No reaction was detected between AGE and water after 10 days at 50°C. Very slight reaction was detected (less than 1%) under similar conditions when 1% CrOct was present in the system. Hence, water appears to act as a co-catalyst with CrOct in accelerating the acid epoxide condensation reaction.

#### TABLE XIVa

Mater Content	Temperature (°C)	Time <u>(hr.)</u>	Reaction <u>c</u> / (%)
9	25	3.5	30.5
		24.0	66,6
		50.0	87.5
9	50	2.0	41.6
	2.2	5.0	75.3
		24.0	94.4
		48.0	99.0 <del>+</del>
18	25	4.0	47.2
		26.0	62.5
		52.0	87.5
18	50	3.0	41.6
		5.0	61.1
		24.0	93.0
		48.0	99.0+
90	25	4.0	55.5
		26.0	66.0
		53.0	87.5
90	50	6.0	72.2
		25.0	93.0
		48.0	99.0+

## EFFECT OF WATER ON THE CrOct CATALYZED REACTION OF AGE AND EHAa/

 $\underline{a}$ / Standard reaction system, 1% catalyst level.

<u>b</u>/ The test solutions contained 0.01 mole of reactants; 9  $\mu$ l. of H<sub>2</sub>0 = 0.0005 mole, 18  $\mu$ l. = 0.001 mole, 90  $\mu$ l. = 0.005 mole.

<u>c</u>/ One trace product noted in GC with recorder at maximum sensitivity; major component in excess of 99% of product mixture.



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Summary	÷
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	System*					
AGE Mole	H20 Mole	Catalyst (wt. %)	Temperature (°C)	Time <u>(days)</u>	Resulc	
0.01	0.005	-	50	10	No reaction	
0.01	0.005	1	50	10	1% loss of AGE; product detected	

A trace product is formed at  $50^{\circ}$ C when water is added to a mixture of AGE and EHA without catalyst. The carboxylic acid is most likely acting as a protonic catalyst for glycol formation. The trace product has the same GC retention properties as that observed in the AGE/H<sub>2</sub>O/CrOct system. Hence, water does aet as a eo-eatalyst in the system for conversion of epoxide and carboxylic ester, and does not lead to significant side product formation. This effect is most pronounced during early stages of reaction when the ratio of catalyst to reactant is high.

More significant, however, is the impact these observations have on the study of the mechanism of the reaction. The rate controlling step in the conversion sequence is obviously affected by the presence of water. Information from the literature and evidence from this study (discussed later) indicate the catalytic activation step involves interaction between catalyst and acid to give the carboxylate anion which, in turn, attacks the epoxide ring (Ref. 3). The presence of water would be expected to facilitate this ionization step. The chromium catalyst could be depicted as consisting of an inner sphere containing the primary metal-oxygen bond, and an outer sphere in which bonding forces would be weaker but still sufficient to hold solvent molecules or nucleophilic materials such as a carboxylic acid. This outer sphere may be the activation site; water

In toluene solvent.

## TABLE XV

## THE EFFECT OF H<sub>2</sub>O ON THE COMPONENTS OF THE EHA/AGE/CrOct SYSTEM

Water				
	Content	Temperature	Time	Reaction
Reactants	<u>(µl)</u>	(°C)	(days)	(%)
AGE - no catalyst	9	50	6	
·	9	50	12	ND
	9	50	26	NR
ACF + 1% C=0.5	0			
AGE 4 1% CLOCE	9	50	1	Т
	9	50	6	т
	9	50	12	> 1
	9	50	16	> 1
	9	50	26	> 2
AGE/EHA - no catalyst	9	50	16	> 1
AGE/EHA - no catalyst	0	50	6	NR
Completed Reactions				
AGE/EHA/CrOcta/	9	50	20	100 NG
	18	50	20	100 NC
	90	50	28	100 NC 100 NC
	0			
	9	25	28	100 NC
	18	25	28	100 NC
	30	25	28	100 NC
Product - no catalyst	9	50	6	NC
	9	50	16	NC
	9	50	26	NC
Product + 1% CrOct	9	50	1	NC
	9	50	6	NO
	9	50	16	NG
	9	50	26	NC
	-	2.4	2.0	193.

NC = No Change.

NR = No Reaction.

<u>a</u>/ Surveillance.

present in the system would be bonded here (akin to solvation). The activation of the acid probably involves ionization to carboxylate anion by release of the proton. Water would facilitate this step. Mechanistically, the following steps are postulated.

$$-\operatorname{CO}_{2}H + M \xleftarrow{k_{1}} \left[-\operatorname{CO}_{2}-MH^{+}\right]$$

$$-\operatorname{CO}_{2}-MH^{+} + A \xrightarrow{k_{3}} -\operatorname{CO}_{2}-CH_{2}-C^{+} + M$$

The effect of water on the rate of reaction in the model system was further explored using CrOleate and neo-decanoate as catalysts. Effects similar to those noted above for the octoate system were noted for the oleate and neo-decanoate catalyzed reactions as shown in Figures 16 and 17 respectively. Data tabulation is given in Tables XVI and XVII. Reaction is enhanced during initial stages, though not as sharply as in the CrOct case. The extent of initial rate enhancement appears to be independent of the actual amount of water present; this is unlike the CrOct situation. Since both Cr-neo-decanoate and CrOleate at the weight percent used have approximately half the amount of metal as compared to CrOct, the lowest water level (9  $\mu$ L or 0.0005 mole) may be beyond the critical interactantratio-point for the effect; that is, the CrOleate and neo-decanoate systems are "swamped" even at 9  $\mu$ L of water.

2. <u>Ammonia</u>: This material was introduced to the model cure system as ammonium hydroxide. The control experiment indicates reaction promotion by the base, especially at 50°C. The extent of catalysis at 25°C is insignificant relative to the normal rate promotion by CrOct. When both ammonium hydroxide and the Cr catalyst are present in the same system, the extent of reaction is significantly greater in a 24-br. period than for the organometallic alone, i.e., 77.7% vs. 56%. The acceleration is due only partly to the amine, however, since the amount of water present in the system will increase the conversion in 24 hr. by ~ 10% over the Cr control rate. Side products were insignificant in quantity (two traces) indicating the ammonia to be catalyzing the desired esterification. (See Table XVIII.)

3. HNO3: Nitric acid has no significant effect on the rate or stoichiometry of the catalyzed reaction and does not catalyze the main reaction itself. A slight yellowing of the control solutions was noted, especially at 50°C, but no product formation was noted at 25°C and ~ 5.0% at the elevated temperature. (See Table XVIII.)

4. <u>RCNO2</u>: The nitrate ester functionality has no effect whatever on the rate or stoichiometry of the acid-epoxide reaction.




# TABLE XVI

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Mater Content (µl)	Temperature (°C)	<u>Time</u>	Reaction(%)
9	50	2 hr.	28.4
9	50	25 hr.	83.8
9	50	3 days	99+
9	50	18 days	100
18	50	3 hr.	22.9
18	50	25 hr.	83.8
18	50	3 days	99+
18	50	18 days	100
90	50	2 hr.	22.9
90	50	26 hr.	78.4
90	50	3 days	99 <del>+</del>
90	50	18 days	100
9	25	22.5 hr.	35.2
9	25	90 hr.	83,8
9	25	13 days	99+
18	25	23 hr.	37.8
18	25	90 hr.	75.7
18	25	120 hr.	81.0
18	25	14 days	<del>99+</del>
90	25	24.5 hr.	37.8
90	25	91 hr.	70.3
90	25	120 hr.	78.4
90	25	13 days	91.8

# EFFECT OF WATER ON THE REACTION OF AGE/EHA CATALYZED BY CR-NEO-DECANOATE

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# TABLE XVII

Water Content	Temperature			Reaction
<u>(µl)</u>	<u>(°C)</u>	<u>T</u>	ime	(%)
9	50	2	h	27 5
9	50	2		27.5
9	50	20	nr.	84.5
2	50	5	days	99+
9	50	18	days	100
18	50	3	hr.	24.2
18	50	25	hr.	85.0
18	50	3	davs	99+
18	50	18	days	100
90	50	2	hr.	23.4
90	50	20	hr.	79.5
90	50	3	days	99+
90	50	18	days	100
9	25	7	hr.	21.5
9	25	3	davs	77.2
9	25	4	days	83.5
9	25	. 6	days	89.9
9	25	18	davs	100
	•••	10	days	100
18	25	7	hr.	21.4
18	25	3	days	71.2
18	25	4	days	81.5
18	2.5	6	davs	89.7
18	25	18	days	100
90	213	7	hr.	19.1
90	25	3	days	73.4
90	25	4	days	80.6
90	25	6	days	86.5
90	25	10	days	97.2
90	25	18	days	100

# THE EFFECT OF WATER ON THE REACTION OF AGE/EHA CATALYZED BY CROLEATE

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EFF	SCT OF SELECTED CONTAMI	NANTS ON THE CATALYZE	D REACTION OF AU	E AND ERR
<u>Contaminant</u>	Catalyst Level (%)	Temperature (°C)	Time <u>(days)</u>	Percent Reaction
0	1	R.T.	24 hr.	56.0
H20 <u>#</u> /	1	R.T.	24 hr.	65.0
HNO 3	1	R.T.	1	58.3
			4	96.1
			7	98.3
	0	R.T.	11	< 1
	1	50	1	97.2
			4	97.2
			7	99.1
	0	50	11	5.0
NH40Ha/	1	R.T.	1	77.7
			4	91.6
			7	94.4
			11	97.5
	0	R.T.	1	5.5
			7	8.5
			11	16.6
	1	50	1	94.4
			4	99+
			7	160.0
	0	50	1	38.8
			4	52.7
			7	52.7
			11	63.8
rono2ª/	1	R.T.	1	58.7
			4	94.4
			/	188.0
	0	<b>R</b> . <b>Τ</b> .	11	S.0
	1	50	1	97.2
			4	99+
			7	100.0
	¢	50	11	0.0
HC104		R.T.	1	60.0
(9 µl. of 70%			4	. 94.0
HC104)			7	100.0
	0	R.T.	11	0.0 (trace product)
	1	50	11	< 2.0 (4 trace products)
	0	50	11	< 1.0 (4 trace products)

Contaminant level was 0.1 to 1 molar ratio of contaminant to reactant (i.e., contaminant level  $\pm /$ actually exceeds catalyst amount).

5. Perchloric acid: As in the nitric acid case (in the concentration range studied) perchloric acid displayed no significant cure alteration effect in Cr system. A blank series was run first, i.e., reactants less catalyst employing three different concentration levels of the acid (9, 18, 90  $\mu$ l. in the standard 5 cc. reaction sample). These amounts of acid would be representative of a catastrophic decomposition situation in the propellant. At both 25° and 50°C, four distinct products were observed in the gas chromatogram for the 18-µ1. and 90-µ1. samples, all in trace quantities. Only at 50°C and 90 µ1. HClO4 did the total amount of these products exceed 2% of the reactants. The nature of these reactions are speculative: oxidation, acid catalyzed condensation, and homopolymerization are possibilities. With 1% CrOct present in addition to the contaminants, reaction catalysis appeared normal at both 25° and 50°C at the two lower  $HC10_{1/2}$  levels, with side reactions unchanged from the uncatalyzed systems. At the highest contaminant level, cure reactions occur, but numerous side reactions are noted by the gas chromatograph.

Conclusions possible at this point concerning the presence of HClO4 in a formulation include: (1) trace levels of HClO4 (that is, realistic levels) are not likely to affect catalytic effectiveness of the Cr catalyst (specifically CrOct); and (2) as expected, HClO4 undergoes a number of reactions with organic material regardless of the presence of the catalysts, and in this sense will be detrimental to cure.

6. <u>HF</u>: HF (introduced as the aqueous solution) in an amount equivalent to a 1% loading of CrOct showed very definite epexide consuming properties. In an uncatalyzed control, all epoxide was consumed at ambient temperature in 7 hr. EHA remained unaffected. The HF promoted reaction competes successfully with the CrOct catalyzed esterification reaction in terms of epoxide demand; only 44% of the total theoretical amount of product ester has formed when all epoxide has been consumed. The gas chromatograph indicates four side products due to HF promotion. A different effect, discussed below, was observed by ABL in propellant systems. (See Table XIX.)

7. <u>N20</u>: N20 showed no catalytic properties in the uncatalyzed control and its presence did not alter the rate and stoichiometry of the CrOct promoted reaction.

#### C. <u>Amine Catalysis Studies</u>

Since many propellant additives, e.g., processing acids and antioxidants, are amines or amine salts, an examination of their potential catalytic properties in the acid-epoxide system was undertaken. The objective of this study was to identify amine types which would interact with the cure functionalities to the detriment of the main reaction; in a quantitative sense, reaction rates (e.g., epoxide consumption) relative to the main reaction were determined.

#### TABLE XIX

	http://www.		
<u>Catalyst</u>	Time (hr.)	% Epoxide Consumed	% Acid Consumed
0	5	97.0	0.0
	25	100.0	0.0
1% CrOct	5	27.9	5.0
	25	30.8	7.0
	8 days	100.0	44.1

### EFFECT OF HF ON EPOXIDE-ACID REACTIONA/

a/ HF amount equivalent to 1% CrOct loading.

Two Levis bases, pyridine and triethylamine, were selected to provide points of comparison between amine catalysis and organometallic catalysis of the acid-epoxide reaction. The results are shown in Table XX; even at 50°C the amines are significantly less effective than Cr salts as catalysts for the solution epoxide-acid reaction. The results reported above for ammonia also agree with this conclusion.

#### TABLE XX

#### AMINE CATALYSIS STUDIES OF EHA/ACE MODEL SYSTEM AT 50°Ca/

	% Reaction				
Time	Pyridineb/	<u>Triethylamineb</u> /	<u>Cr0ct</u>		
2 hr.	< 1	< 1	50		
22 hr.	2.5	< 1	<b>9</b> 8		
4 days	30.7	7.7	100		
7 days	34.6	23.1			
19 days	66.6	76.9			
27 days	65.6 <u>c</u> /	76.9 <u>°</u> ∕			

a/ Standard model reaction system used.

b/ Amounts of amine catalyst used were equal on a molar basis to a 1% loading of CrOct, i.e., 9 x 10<sup>-5</sup> moles, or 0.0073 g. pyridine and 0.0093 g. triethylamine.

c/ Stoichiometry of 0.99 noted; one trace side product detected.

An interesting problem arose during propellant formulation studies at ABL: a number of mixes containing the antioxidant TDPA (thiodiphenylamine) did not cure. Mixviscosity increased significantly and no cure was achieved-an unusual occurrence since TDPA has been used successfully in numerous similar ABL propellants cured at 140°F. To determine whether the effect is due to a chemical interaction, the series of experiments was undertaken; those in model systems are listed in Table XXI.

#### TABLE XXI

#### MODEL SYSTEM STUDIES

	TDPA	Cr0ct		
Reactant Combination	<u>% by Wt.</u>	<u>% by Wt.</u>	Temperature	<u>Observation</u>
EHA/AGE	0.5	0.0	R.T.	No reaction or
	0.5	0.0	50	change in 7
				days
EHA/AGE	1.0	0.0	R.T.	
	1.0	0.0	50	
EHA/AGE	2.0	0.0	R.T.	
	2.0	0.0	50	$\checkmark$
EHA/AGE	1.0	1.0	R.T.	Normal rate and
	1.0	1.0	50	stoichiometry
				behavior

No adverse or unexpected effect due to TDPA was noted in the model studies. The uncatalyzed solution studies at both R.T. and 50°C showed the typical amine effect--only a slight product formation (trace amount, less than 1%) in the presence of the antioxidant after a duration of at least 7 days. With CrOct present along with TDPA, the normal catalyzed reaction behavior is observed for EHA/ACE at both R.T. and 50°C.

Reaction hehavior was examined in the polymer system and the results from those experiments are given in pages 77-79.

#### D. Summary of Contaminant and Extra-Ingredient Effects

No evidence has been found of catalyst poisoning in the Cr system by common propellant ingredients and contaminants. Oxidizing acids such as  $HNO_3$  and  $HClO_4$  (at realistic contaminant levels) do introduce trace amounts of side products; however, these side reactions are insignificant in a rate and stoichiometric sense, when compared to the course of the main esterification reaction. Amines are co-catalysts for the esterification reaction but, again, they are poor at ambient temperature compared to CrOct. The presence of an amount of amine in a system equimolar to the Cr catalyst will result in an acceleration of the conversion rate during earlier reaction stages equal to no more than 10% of the Cr rate. The presence of water in the system causes a similar rate effect in the model systems, though in the binder situation this rate acceleration effect is not as noticeable.

#### CATALYST EVALUATION IN BINDER SYSTEM

Following the selection of candidate catalysts from the model studies, the evaluation was expanded to actual polymer systems. The purpose of this work was: (1) to correlate the results of the model system studies with the actual binder situation; (2) to relate catalytically controlled chemical conversion behavior to physical effects, such as gelation, pot life. catalyst levels, total cure time, etc.; (3) to study effects that could not be determined by GC, for example, to examine the effects of high molecular weight materials such as plasticizers and surfactants on the outcome of the catalyzed cure reaction. Infrared spectroscopy was applied to the problem of following the conversion of Telagen CT to ester by observing the decrease in absorption at 5.9  $\mu$  (carbonyl stretching frequency for acid) and the increase in the peaks at 5.8  $\mu$  (carbonyl stretching frequency for ester) and 2.9  $\mu$  (OH stretching vibration) (Refs. 3 to 5).

A description of the experimental method is given in the experimental section along with a discussion of the advantages and disadvantages of the procedure.

Quantitative determinations are made by reference to an empirical absorbance ratio-concentration curve. In some case, where only relative comparisons were desired of general reaction trends, the data are presented as plots of absorbancy versus time.

#### A. Binder Cure Behavior in Cr Salt Catalyzed Systems

It was desired to determine the effect of free (residual) acid in the catalyst on the overall trend of the reaction. Figure 18 shows a comparison of the rates of reaction for Telagen CT and AGE catalyzed by GrOct and CrOleate. The quantities of active ingredient, Cr, are the same for both reactions; 2-ethyl hexanoic acid was added to the 10.5% GrOct to compensate in kind for the additional oleic acid present in the 4.2% GrOleate. The patterns for both reactions are very similar, though the octoate catalyzed reaction appears to proceed faster initially to a rate plateau, while the oleate goes farther toward completion in the same time period. This difference in rates is most likely due to differences in solubility between the GrOct and CrOleate; the oleate is more soluble in organic materials than the octoate. The addition of the compensatory acid to the GrOct does not appear to change the reaction pattern or rate of reaction relative to the same catalyst without the free acid (10.5% CrOct) as long as the same amount of active metal is used.

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The conversion profile for the reaction of Telagen CT and ERLA 0510 catalyzed by CrOct at the 1% by weight level is given in Figure 19. Reaction is about 75% complete in 5 hr. The rates of disappearance of acid and appearance of alcohol and ester are equal during this stage. Beyond this point acid is consumed at a faster rate (i.e., loss of abosrbance at 5.9 u) than either ester or alcohol is formed. At 100 hr., the amount of acid consumed equals the amount of ester formed. This pattern is experimentally reproducible, hence it is a real effect and not due to error in the method. The alcohol present after 100 hr. is about 5% less than theoretical, the difference in this case is the method. The rapid decrease in acid can be explained in terms of a shift in kinetics and the resulting survival of metal-coordinated carboxylate. After approximately 75% reaction, diffusion of reactants is extremely slow. Diffusion of the catalyst will be faster than epoxide since, at this stage, the latter being polyfunctional will have its mobility significantly reduced through incorporation of one or more of its epoxides into the fixed binder matrix. Therefore, catalyst diffusion and coordination with carboxyl will be rapid compared to the condensation, and this catalytic intermediate will exist as such for a longer time period. Hence, there will be a difference in the rates of disappearance of carboxylic acid and appearance of ester. This point will be dealt with in more detail below.

Of significance in a processing sense, 1.0% loading of CrOct leads to 75% conversion in 5 hr. This may be too fast for good pot life characteristics. Studies were therefore undertaken using 0.2, 0.4, and 0.8% loadings of CrOct. Figure 20 shows the results of this experimental series in terms of the rate profile (conversion of carboxylic acid) for the cure of Telagen CT with ERLA 0510. At the 0.8% catalyst level the rate transition point (discussed below in terms of gelation) appears to begin at around 6 hr., while with 0.4% catalyst inflection point on the curve occurred at 20 hr. The rate of disappearance of COOH for the reaction catalyzed by 0.2 and 0.1% CrOct, respectively, do not display any sharp shift in the time range of the experiment, a point in agreement with 30 g. propellant mix studies (discussed later). The minimum catalyst level for cure in a reasonable time period appears in the 0.4% range; results from ABL propellant processing studies indicate that this range is indeed 0.37 to 0.4% catalyst level.

Figure 21 displays the decrease in acid carbonyl content with time for an equivalent mixture of Telagen CT and CrOct. This plot is a demonstration of the "precoordination" effect; as the amount of carboxylic acid decreases relative to the amount of catalyst present, the more pronounced is the effect. Hence, it is reasonable to propose that the deviation in the rate of disappearance of acid and appearance of ester noted in the above studies is due to coordination of unreacted acid with catalyst.



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At this point some comment should be made concerning the reasons for the character of the reaction conversion curves. In general, the chemical reaction behavior of polymers is essentially the same as that for low molecular species with the exception of systems in which diffusion rates are low (Refs. 5 and 6). Where gelation occurs, reaction continues only slightly beyond the gel point and a complete reaction does not occur. French and co-workers (Ref. 5) have demonstrated that the maximum extent of polymer reactions is dependent on functionality and suggest that the limiting factor in determining the final extent of reaction depends on the geometry of the solid matrix formed by the cure reaction (functionality dependence) and not on thermodynamics or kinetics.

This effect becomes apparent when results from the model system studies are compared with those from the binder reactions. The solution reactions go to completion, while the polymer binder cure reaction consumes only 75 to 90% of the starting materials. Based on the French hypothesis, then, the extent of reaction (but not rate) is not controlled by the catalyst and is a function only of prepolymer and curative structure. Time to gelation and rate of increase in binder viscosity during cure are catalyst dependent. Since these parameters are of critical importance to processing, catalyst selection is crucial to the success of that procedure. In simplest terms, gelation occurs when cure linkage (especially cross-linking) formation has reached a point where a sufficient number of cross-links exist to give a three-dimensional matrix with mechanical integrity (i.e., solidification occurs). Reaction of the remaining cure functionalities becomes increasingly dependent on diffusion and the rate of ester formation will be curtailed sharply. Referring to Figure 19, Point A is most likely the beginning of the gelation process and Point B may be the gel point (this cannot be established without direct correlation with actual gel point determination studies). A-C represents the "precoordination" range for unreacted carboxylic acid groups with the catalyst -- the only truly diffusable species present in the system at this point.

The effect of other binder parameters (e.g., mixed curative, plasticizer) were investigated in the Telagen CT system at both 1.0 and 0.4% CrOct levels as shown in Table XXII and Figure 22. At the 1% CrOct level, the reaction rate profiles are identical regardless of the nature of the curative (all tri-epoxide vs. an equivalent mixture of di- and tri-epoxide) or the presence of plasticizer. When 0.4% CrOct is used to catalyze the reaction of the mixed epoxide and Telagen CT, the presence of plasticizer does not alter the conversion pattern. There appears to be a slight increase in the rate of conversion for the mixed curative system compared to the all tri-epoxide (though the differences in values are precariously close to the experimental error of the method).

#### TABLE XXII

# PARAMETER STUDY FOR AMBIENT CURE OF TELAGEN CT

Curative (meg. ratio)	Catalyst Level <u>(wt. + % CrOct)</u>	Plasticizer Level (wt. + % polybutene)	Time (hr.)	% Reaction
DEP 332 (0 5) + EPIA 0510 (0 5)	1.0	5.0	1	3.0
$0.00 \pm 0.00 \pm 0.00 \pm 0.000 = 0.000$	1.0	5.0	2	57 5
			4	74.2
			10	85.0
			24	85.0
			49	87.8
			14 days	89.0
			I4 Gays	05.0
DER 332 (0.5) + ERLA 0510 (0.5)	0.4	5.0	2	30.3
			4	39.3
			10	66.6
			24	82.0
			48	82.0
			4 days	89.0
			14 days	89.0
DEP 332 (0 5) + EPIA 0510 (0 5)	1.0	0.0	1	16.6
DER 332 (0.3) + ERLA 0310 (0.3)	1.0	0.0	2	54.5
			4	72.7
			10	83.3
			24	85.0
			48	85.0
			14 days	90.1
	<u> </u>		,	• •
DER 332 $(0.5)$ + ERLA 0510 $(0.5)$	0.4	0.0	1	9.0
			2	24.7
			4	56 1
			26	70.1
			24	72+7 8/ 8
			9 dawe	95.0
			14 davs	90.0
			14 days	,0.0
ERLA 0510 (1.0)	1.0	5.0	1	5.2.7
			2	17.6
				76.0
			10	81.8
			24	85.0
			48	85.C
			8 days	85.0
			14 days	88.0
FRIA 0510 (1.0)	0.4	5.0	1	13.6
0/00 \/	- • •		2	39.4
			4	41.0
			10	63.6
			24	82.0
			48	83.2
			8 davs	87.0
			14 days	89.0

10.00

Å



An interesting observation was made when the effect of plasticization on the rate of cure of Telagen CT with all tri-epoxide curative was studied. Figure 22 shows a comparison of rate patterns for plasticized reactions at both 1.0% and 0.4% catalyst levels. No effect is noted at the 1% level; while the 0.4% CrOct reaction is accelerated by the presence of plasticizer during early stages of cure (up to approximately 70% conversion) compared to the unplasticized case.

Several propellant components, listed in Table XXIII, were added to Telagen-ERLA mixtures to determine whether their presence would alter the CrOct catalyzed cure reaction. The data are plotted in Figure 23 relative to the rate data for the binder mixture without the ingredients. At the 1% catalyst level, AP and water had no effect on the rate of the cure reaction. The presence of lecithin or  $Fe_2O_3$  tends to slow the reaction during early stages of cure, though the total time to 90% conversion is unaffected. At this writing, the 0.4% catalyst level studies of the lecithin and Fe<sub>2</sub>O<sub>3</sub> effects are under way. At the 0.4% catalyst level, again, AP and water do not alter the rate behavior of the cure reaction.

The binder cure rate studies reported above employ an equivalent mixture of carboxylic acid and epcxide. Since E/C\* ratios other than one are often employed in formulation, the effect of this parameter was studied in the Telagen CT/ERLA 0510 system. Table XXIV summarizes this experiment.

Excess epoxide gives a higher cure rate than in the analogous excess Telagen system. Cure is complete in a shorter time period (2 days vs. 7 days). The final reaction segment is much shorter than in the excess Telagen situation. Ultimate reaction is slightly higher, approximately 4%, in the excess epoxide system.

In summary, the ambient temperature cure behavior of carboxylic acid prepolymer with epoxide remains essentially unaffected by the presence of binder and propellant components, i.e., the total cure times appear unchanged. Diffusion of reactants and catalyst is a rate controlling parameter as evidenced by the differences in rate observed between di- and triepoxide curatives, and in the presence of plasticizer. These observations are in agreement with the hypothesis made above concerning chemical conversion behavior and its relation to physical effects.

<sup>\*</sup> E/C - epoxide/carboxyl equivalence ratio.

# TABLE XXIII

	Contraction of the second second second	Reaction	
Ingredient (% by wt.)	<u>Catalyst Level (% by wt.)</u>	Time (hr.)	(%)
AP (5.0)	1.0	1	28.8
		2	60,6
		4	72.7
		11	81.8
		24	83.3
		48	84.0
		8 days	90.0
AP (5.0)	0.4	2	18.1
		4	31.8
		11	56.0
		24	78./
		48 5 deue	81.8
		Jaays	85.0
		8 days	86.0
H <sub>2</sub> O (0.04 meq.)	1.0	1	27.2
		2	50.0
		4	68.1
		11	81.8
		24	82.0
		5 d <b>a</b> ys	86.3
H <sub>2</sub> O (0.04 meq.)	0.4	1	13.6
		2	21.2
		4	33.3
		11	66.6
		24	78.7
		48	82.0
		5 days	89.0
Lecithin (0.5)	1.0	2	31.8
		4	51.5
		11	75.7
		24	78.7
		48	81.8
		5 days	83.0
Fe <sub>2</sub> 0 <sub>3</sub> (1.0)	1.0	2	27.3
		4	46.9
		11	75.8
		24	78.8
		48	81.0
		5 days	84.0

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# EXTRA-INGREDIENT EFFECTS ON AMBIENT CURE OF TELAGEN CT WITH ERLA 0510 CATALYZED BY CrOct



#### TABLE XXIV

Telagen/ERLA 0510	Temperature (°C)	Time <u>(hr.)</u>	<u>% Rxn</u>
1/0.75	25	2	26.4
		10	51.9
		23	59.0
		48	61.2
		3 days	62.7
		7 days	64.6 - cured
		10 days	64.6
0.75/1	25	2	22.0
		10	59.3
		23	68.3
		48	68.6 - cured
		3 days	68,6

#### EFFECT OF VARYING E/C RATIO ON CURE REACTION

#### B. Cure Processing Problem - TDPA

In the propellant formulation phase of the program, ABL encountered processing difficulty when the antioxidant TDPA was incorporated in certain mixes. The studies described below were an attempt to isolate and identify the components which were interacting to produce the effect.

Results for gumstock and filled binder cure studies are shown in Table XXV.

The Telagen CT-ERLA 0510 gumstocks with 0.4 and 0.5% CrOct and 1% TDPA showed normal reaction behavior relative to the reaction profile at these catalysts levels without the antioxidant. Hence, TDPA does not appear to interfere directly with either of the reactants or the action of the catalyst on the cure functionalities.

The addition of AP to the gumstock mixture does not change the cure behavior. The samples loaded with 50% AP and 20% Al also appeared to cure completely in approximately 10 days (the observations in this case are only qualitative since no measurement of cure was made other than visual and handling observations).

A closer examination was made of the possibility of interaction of TDPA antioxidant with the specific curative ERLA-0510 (previous quantitative work had been done in the model test solution). Epoxide consumption in TDPA/ERLA-0510 mixtures was measured by titration of the epoxide.

## TABLE XXV

#### TDPA BEHAVIOR IN BINDER

Binder Composition	Solids Loading		
<u>% by Wt. of Binder</u>	% by Wt. of Total	Temperature	<u>Observation</u>
Telagen CT/ERLA 0510/0.5% CrOct/1% TDPA	0	R.T.	Normal rate and cure behavior
Telagen CT/ERLA 0510/0.4%	0	R.T.	ļ
CrOct/1% TDPA Telagen CT/ERLA 0510/0.5% CrOct/1% TDPA	63	R.T.	
Telagen CT/ERLA 0510/0.4% CrOct/1% TDPA	63	R.T.	00
Telagen CT/ERLA 0510/0.5%	50-AP	R.T.	
CrOct/1% TDPA	20-A1		
Telagen CT/ERLA 0510/0.4%	50-AP	R.T.	$\vee$
CrOct/1% TDPA	20-A1		

Table XXVI shows a summary of the results of the experimental runs.

#### TABLE XXVI

#### EPOXIDE DEMAND-ERLA-0510 (LOT 11)

	System	TDPA Level Wt. % Based on Curative	Temp. (°C)	Time (hr.)	Epoxide Analysis (meq/sample)	Comment
1.	ERLA-0510	0	25	0	11.33 ± 0.20	Based on 8 de-
2	paseline	10	25	27	11 00	Na shuras
Ζ.	EKLA-0510/	10	25	24	11.29	No change
	TD <b>PA</b> in	1	25	24	11.19	No change
	Chloroform Soln.	0.5	25	24	11.56	No change
		1	50	24	11.19	No change
3.	ERIA-0510/	10	25	5 days	10.11	10.7% consumption
	TDPA Neat	1	25	5 days	11.09	~ 1% Rxn
	Mixture	0.5	25	5 days	11.01	∼ 1% Rxn

As the weight ratio of TDPA to curative increases, there is an increased epoxide demand in the system. This conclusion appears to hold true only in the bulk (neat) mixture situation and not in solution. In addition, these observations agree with the propellant cure data from ABL. The nature of the epoxide consumption is most likely a homopolymerization reaction; such an occurrence would explain the increase in viscosity and concomitant decrease in cure rate (homopolymerization of a polyfunctional material such as ERLA-0510 would undoubtedly lead to significant viscosity increases). To complicate matters further, samples from a new lot of ERLA 0510 do not exhibit the consumption behavior in the presence of TDPA as noted of the previous lot. These data showing no epoxide loss are summarized in Table XXVII.

#### TABLE XXVII

#### EPOXIDE DEMAND BY TDPA AT 25°C (ERLA 0510 Lot 9)

	Milliedulvalents Epoxide Kemaining					
	(ERLA-0510) Sample					
<u>25% TDPA</u> a/	12.5% TDPA	10%_TDPA	5% TDPA			
$11.50 \pm 4$	12.50	12.60	13 <b>.2</b> 0			
11.50	12.30	12.42	13.14			
11.91	12.30	12.53	13.00			
11.17	12.51	12.34	13.04			
11.63	12.04	12.36	13.48			
	<u>25% TDPA</u> <sup>2</sup> / 11.50 ± 4 11.50 11.91 11.17 11.63	$\begin{array}{r c c c c c c c c c c c c c c c c c c c$	MITTEQUIVATENT'S EPONIAE Remaining (ERLA-0510) Sample25% TDPA <sup>a</sup> 12.5% TDPA10% TDPA11.50 $\pm$ 412.5012.6011.5012.3012.4211.9112.3012.5311.1712.5112.3411.6312.0412.36			

a/ Wt. % TDPA of total sample (TDPA + ERLA 0510).

Conclusions are as follows: (1) TDPA does not appear to interact with the carboxylic acid or epoxide groups to any appreciable extent for a length of time equal to the duration of the processing period; (2) TDPA does not interfere with the activity or CrOct as a catalyst for the epoxide-acid condensation reaction in either the model or binder situation; (3) the presence of AP at a moderately high loading (63%) does not alter the above conclusions; (4) aluminum in the formulation containing TDPA does not alter the cure behavior of the system. The problem may be unique to the lot of ERLA used in the propellant formulation study.\*

<sup>\*</sup> A sample from this lot homopolymerized during storage for no apparent reason.

#### C. Evaluation of CrDIPS System

CrLIPS\* is a catalyst system recommended for the all-epoxide cure of CT binders by the Explosives Research and Development Establishment, Waltham Abbey, England. The sample used in this study was supplied by AFRPL. Compositional data and a recommended formulation follow:

Catalyst: CrDIPS in IDP (isodecylpelargonate)-1.6% wt. Cr content.

Formulation:	AP 300 µ	39.5% wt.
	AP 30 μ	39.5
	Α1 15 μ_	5
	ERLA 0510	
	DER 332 >	14
	HC 434	
	CrDIPS in IDP	2

The catalyst (including IDP plasticizer) level in the binder is, therefore, 12.5%. In our evaluation, we used a 10% level of CrDIPS/IDP in Telagen CT-ERLA 0510 binder mix. The active metal content is 0.16%; our 1% CrOct catalyzed systems contain 0.108% Cr.

Table XXVIII shows the results for a binder cure study using CrDIPS. It is effective at both ambient and 50°C, at least as effective as a similar loading of CrOct.

Figure 24 is a comparison of the effectiveness of a 5% loading of CrDIPS in the epoxide and aziridine cure of Telagen CT. In general, this catalyst appears more effective for the epoxide system than for the imine in that a higher extent of conversion is achieved. The chart below summarizes the catalytic behavior of CrDIPS in the Telagen system and compares it with CrOct.

Catalyst Loading (Wt. % in binder)	Wt. % Cr Metal	Time to 50% Conversion	
CrDIPS (10) <u>a</u> /	0.16	20 hr.	
CrDIPS (5)	0.08	2.5 days	
CrOct (1)	0.11	1.5 days	

a/ CrDIPS in IDP - 1.6% wt. Cr content.

\* Chromium 2,6-diicopropyl salicylate.

# TABLE XXVIII

# CATALYSIS BY CrDIPS IN IDPa/

Catalyst Level	Temperature	Time	
<u>(Wt. %)</u>	(°C)	<u>(hr.)</u>	<u>% Rxn</u>
10	Ř.T.	2.5	13.8
		4.5	24.6
		22.0	58.0
		27.0	64.0
		2 days	Cured (67)
10	50	2.5	60.5
		4.5	Cured (66)
5	R.T.	3.5	16.7
		27.0	55.1
		48.0	63.0
		7 days	Cured (67.6)
5	50	3.5	59.2
		6.5	66.5
		22.0	Cured (67)

<u>a</u>/ Cr content of catalyst mixture is 1.6%. Binder system: Telagen CT/ERLA 0510.



Figure 25 shows the effect of AP and Ammonyx T on CrDIPS catalyzed cure to Telagen CT and ERLA C510. The cure reaction pattern is unaffected by the presence of these ingredients.

As a result of these cure reaction rate observations, CrDIPS was selected for further screening at ABL of its performance in gumstocks and propellants.

#### D. PBAN Cure Studies and the Ammouyx T Question

ABL reported cure difficulties with PBAN polymer. The latter contains 4% by weight Ammonyx emulsifier. Two types of experiments were run to determine the nature of the interaction of the additive with the cure process: (1) GC model studies, and (2) binder cure studies with and without solids. Table XXIX summarizes the results of the model studies.

1. <u>Ammonyx effect in model systems</u>: The presence of Ammonyx at 1 and 4% leels in the model system has a slight acceleratory effect on the CrOct catalyzed reaction of AGE and EHA. The uncatalyzed controls (no CrOct) show a low level of reaction catalysis, especially at 50°C (93.6% conversion in 28 days at 4% Ammonyx T level). Hence, Ammonyx T displays the usual amine co-catalysis effect.

2. <u>Ammonyx effect in Telagen CT epoxide system</u>: Polymer cure studies were run in the Telagen CT/ERLA 0510 system in the presence of Ammonyx T and AP. These data are shown in Table XXX. The results are somewhat surprising; in the presence of 4% Ammonyx T, cure occurs at the same rate for both 1 and 0.5% catalyst loading. The presence of 5% AP in the sample leads to a 10% suppression of the rate of ester formation. These reactions were rerun to determine reproducibility.

Figure 26 shows the effect of AP and emulsifier level on cure behavior at 1 and 0.5% CrOct loadings. In this series, the presence of the Ammonyx T accelerated cure at both catalyst levels. For example, the time to 50% conversion for 1 and 0.5% CrOct loadings are 1.5 and 6.5 days, respectively. With the emulsifier these values become 0.5 day at 1% CrOct and 0.75 day at 0.5% catalyst level. One reaction containing 0.5% CrOct, 1% AP and 4% Ammonyx T behaved in an unexplainable fashion, that is, cure followed the pattern displayed by the other reactions until 40% conversion and then slowed considerably. Ultimate extent of reaction achieved was only 58% as compared to 70-73% for the others.

The presence of the emulsifier does accelerate and in some cases, alter, catalyzed reaction possibly by increasing the dispersion of the CrOct throughout the system and by solubilizing other potential trace reactants which may be present as contaminants in the other propellant components.



ALL STREET

Weight % <u>Ammonyx T</u>	Catalyst Level <u>a</u> /	Temperature (°C)	Time (hr.)	% Rxn	Stoichiometry
1	1	R.T.	5	23.5	1
			23	64.7	1
			48	32.4	1
			5 days	100.0	1
1	1	50	3	55.9	1
			23	97.9	1
4	1	R.T.	5	26.5	1
			23	67.6	1
			49	99.1	1
4	1	50	3.5	55.8	l
			24	95.0	1
1	0	R.T.	28 days	8.8	1
1	υ	50	51	26.5	1
			13 days	35.4	1
			28 days	55.9	1
4	0	R.T.	5 days	11.6	1
			13 days	17.6	1
			28 days	23.5	1
4	0	50	l day	8.8	1
			2 days	20.6	1
			5 days	44.1	1
			13 days	76.8	1
			28 dave	93.6	

# TABLE XXIX EFFECT OF AMMONYX TA ON REACTION OF EHA AND AGE

a/ Cetyl-dimethylbenzylammonium chloride.

# TABLE XXX

Statistics of the

# THE EFFECT OF AMMONYX T ON CURE OF TELAGEN CT WITH ERLA 0510

Catalyst Level	AP Level	Ammonyx Level	Temperature (°C)	Time (hr.)	<u>% Rxn</u>
1	0	4	R.T.	17.5	59.8
				23.0	61.4
				42.0	61.8
				6 days	Cured (65)
0.5	0	4	R.T.	17.5	58.5
				23.0	63.0
				42.0	64.6
0.5	5	4	R.T.	4.0	22.7
				17.0	43.0
				23.0	51.1
				42.0	51.5
				6 days	52.5

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3. <u>PBAN cure studies</u>: An infrared calibration curve was prepared for monitoring the change in ester and acid functionalities with time for the CrOct catalyzed cure of PBAN and epoxide. These data are summarized in Table XXXI. In general, the acid-ester conversion reaction proceeds as expected as a function of catalyst quantity and temperature. A 3% loading of AP tends to slow the 1% CrOct catalyzed reaction, as does Fe<sub>2</sub>O<sub>3</sub>.

The IR spectra for the PBAN cure systems show evidence of a side reaction which occurs during later stages of cure. An absorbance at 6.3 -6.4  $\mu$  appears after approximately 60% acid conversion and has continued to grow. Functional groups of interest absorbing in this region include:  $CO_2^2$ , NH<sub>3</sub><sup>+</sup> (deformation), NH<sub>2</sub><sup>+</sup> (deformation). This phenomena did not occur in the Telagen-epoxide-Ammonyx T\* mixes; hence, we attribute it to a PBAN interaction. The interaction could involve the nitrile of PBAN, the quarternary ammonium salt (Ammonyx T), and the cure catalyst. The decreased availability of carboxylic acid (with which CrOct will preferably coordinate) at this stage of cure (> 60%) would then allow the side reaction to become increasingly important. The net effect would be to prolong the time for cure to a state giving desired mechanical properties.

<sup>\*</sup> Cetyl-dimethylbenzylammonium chloride.

# TABLE XXXI

# PBAN/ERLA 0510 CURE STUDY

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Catalyst Level (% CrOct)	Temp. (°C)	Time <u>(Days)</u>	Extra Ingredient (%)	% Reaction Acid Consumed	Comment
1	25	(5 hr.)		27.3	No side rxn
		1		78.4	Side rxn beginning
		2		79.6	Side rxn continuing
		5		79.6	Side rxn continuing
1	50	(5 hr.)		76.2	Side rxn
		1		76.2	Side rxn
		2		76.2	Side rxn
		5		77.0	Side rxn
0.5	25	(6 hr.)		14.6	
		1		49.1	Side rxn beginning
		2	<b>*</b> =	62.3	Side rxn continuing
		5		78.9	Side rxn continuing
0.5	50	(6 hr.)		49.6	Side rxn
		1		60.8	Side rxn continuing
		2		70.5	Side rxn continuing
		5		76.8	Side rxn continuing
1	25	(5 hr.)	3% AP	26,2	
		1		63.1	Side rxn apparent
		2		78.6	Side rxn continuing
		5		79.0	Side rxn continuing
1	50	(5 hr.)	3% A.P	51,2	
		1		78.2	Side rxn apparent
		2		78.2	Side rxn continuing
		5		78.2	Side rxn continuing
1	25	(6 hr.)	1% Fe <sub>2</sub> 0 <sub>3</sub>	21.7	
		1		78.8	Side rxn apparent
		2		81.4	Side rxn continues
		5		83.7	Side rxn continues
1	50	(6 hr.)	1% Fe <sub>2</sub> 0 <sub>3</sub>	78.7	Side rxn apparent
		1	-	79.8	Side rxn continues
		2		79.8	Side rxn continues
		5		79.8	Side rxn continues

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#### AZIRIDINE - ACID CATALYSIS STUDIES

VI.

#### A. Introduction

Like the epoxide, the aziridine group is also a strained, polar threemembered ring which undergoes reactions of the type similar to the epoxides, i.e., nucleophilic addition. Unlike the epoxide, however, aziridine reactivity is much more affected by substituents on and adjacent to the ring. Aziridines show a greater tendency than the epoxide to undergo rearrangement and homopolymerization reactions. Aziridines containing the P-N bond, while displaying acceptable cure reaction chemistry, tend to undergo bond cleavage

at elevated temperatures. Acyl aziridines  $(-C-N_1)$  are prone to rearrange to oxazolines which, in turn, will also react with carboxylic acid but at a slower rate than the parent aziridine. The development and evaluation of aziridine curatives have been the subjects of an Air Force sponsored program (Ref. 8); based on results of that program and recommendation from the Air Force, two curatives were selected for primary use in our catalyst demonstration studies: HX-868 (1,3,5-Benzene-tris-carbonyl-2-ethylaziridine), and a new curative supplied by ESSO, 1,3,5-Benzene-tris-carbonyl-2,3-dimethylaziridine (AcAz).

In addition, MAPO and MAPS were used in certain experiments to provide a baseline for rate behavior since they are well-known and well characterized curatives.

#### B. Catalyst Selection Studies

A major contention of the mechanistic scheme proposed above for the catalysis of the acid-epoxide cure involves activation of the acid so as to make it more nucleophilic. Such a step facilitates attack on the ring of the curative (epoxide or aziridine) by the carboxylic acid and gives rise to the resulting ester. It is reasonable to assume that the mechanisms for epoxide and aziridine cures are similar in this respect. Thus, the information gathered from the epoxide studies was used as a starting point for catalyst selection for the aziridine system.

With this possible similarity of curing mechanisms in mind, several binder mixes were made using CrOct as catalyst and one of several aziridines as curatives (MAPO, MAPS, AcAz). In all cases, at a 1% CrOct loading, cure to a tough, nontacky binder was achieved at 25°C in less than 9 days. Figure 27 displays these results comparatively. The reaction behavior resembles that of the epoxide system, with several marked differences. First,



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thermal catalysis is an important parameter; at 50°C, there are no significant differences in cure rate of the AcAz sample attributable to the different catalyst concentrations (0.5 and 1% CrOct). In fact, an uncatalyzed control (AcAz) at 50°C had reached 80% completion in 6 days while the analogous catalyzed reactions were only at the 90% level. At 25°C, the difference in conversion rates between 0.5 and 1.0% catalyst level is not as dramatic as in a similar epoxide Telagen CT system. IR spectra show evidence of minor side reactions during cure, probably oxazoline formation; this question is discussed in detail below. Another unusual result is the high extent of reaction noted for AcAz/Telagen CT, i.e., 90%. As previously noted in both our studies and in the literature, 65 to 80% (depending on the individual ingredients) is usually the conversion achieved in three-dimensional matrix forming reactions.

Several organometallics of different structural characteristics were evaluated for catalytic activity in the Telagen CT-acylaziridine (ESSO) system. Only those mixtures containing CrOct or FeOct cured at ambient temperature; as in the epoxide system all other additives failed to catalyze the reaction in the desired time length. These results are summarized in Table XXXII.

#### TABLE XXXII

#### THE EFFECT OF SELECTED ORGANOMETALLICS ON THE AMBIENT CURE OF TELAGEN CT WITH ACYL AZIRIDINE<sup>b</sup>/

#### Catalysta/

#### **Observation**

Ferric Octoate

Ferric Acetylacetonate Ferrous Acetylacetonate Chromium Acetylacetonate Stannous Octoate Nickel Octoate 50% reaction in less than 7 days; cure in 10 days No cure No cure Insoluble, no reaction No cure No cure

<u>a</u>/All catalysts at the 1.0% by weight of binder level. b/ ESSO product.

In summary, chromium octoate and to a lesser extent ferric octoate do accelerate the reaction of aziridine curatives and Telagen CT at ambient temperature. Visual cure is noted (i.e., sample is elastic and nontacky) at 7 and 12 days (~ 50% conversion) at 1.0 and 0.5% CrOct levels, respectively.

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ABL reports CrOct to be less effective as a catalyst in the HX-868/ Telagen CT system than in the analogous epoxide system. We agree with ABL's evaluation based on the results of our studies with AcAz, which is reported to be a better curative in terms of stability, reactivity and stoichiometry than HX-868. The problem is rooted in the differences between the two classes of curatives. Epoxides are more stable instrinsically than aziridines and give more controllable and stoichiometric reactions with carboxylic acids.

Two conclusions are possible: (1) the catalytic trends are similar in both epoxide and aziridine systems. Aspects of the mechanism postulated for the cure of acid-epoxide systems are apparently applicable to the aziridine-acid reaction also (i.e., the same materials are effective as catalysts suggesting activation of the acid through coordination with the catalyst and attack by this intermediate on the curative); (2) differences in reactivity due to aziridine structure still exist in the catalyzed reaction, supporting further the acid activation mechanism in the organometallic reaction (i.e., if the aziridine were involved in the catalytic activation step a leveling effect on the rate <u>might</u> be expected throughout a series of aziridines of different structures since catalysis of the aziridine ring opening involves interaction at the nitrogen and not at the substituted carbons).

At this point chromium and iron salts (especially the octoates) were recommended for further evaluation in Phases III and IV (ABL) as ambient cure promoters in the aziridine cure system.

### C. Side Reaction Problem

Figure 28 shows a comparison of the initial infrared spectrum for a mixture of Telagen CT, AcAz and chromium octoate with that for the same mixture after 13 days of reaction at room temperature. As expected, the carbonyl absorption due to carboxylic acid at 5.9  $\mu$  (2) decreases as ester carbonyl at 5.8 µ grows (1). Based on the disappearance of acid, reaction is approximately 70% complete after 13 days. The carbonyl absorption due to the acyl aziridine curative (3) also decreases with time as the 6.4  $\mu$  amide peak grows (4). The peak at 6.1  $\mu$  (5) due to unsaturation in the polymer (which should be an invariant in the spectrum) also shows an increase as reaction proceeds. In addition, several other absorbances grow, especially at 8.5 (7) and 8.9  $\mu$ (8). Considering the 6.1  $\mu$  peak growth first, two possibilities come to mind: (1) olefin formation through opening of the aziridine ring during cure or through elimination of acid from the product amide; (2) -C=N- absorption in exazeline which could be formed under reaction conditions. Figure 29 displays the various reaction paths which can occur in the aziridine-carboxylic acid system.



Workers at ESSO report (def. 8) acid elimination in the sulfonyl aziridine system to the extent of 40% of the product. However, they also claim excellent stoichiometry for the 2,3-dimethylaziridine-acid reaction. There are three major differences between these studies: temperature (ESSO-150°F, this study--25°C), the presence of the chromium catalyst, and the nature of the reaction system (ESSO--model, this study--gumstock).

Returning to the IR study, the increase in intensity at 6.1  $\mu$  can be due to either olefin formation or the -C=N- linkage in oxazoline. The absorptiens at 8.5 and 8.9 are more clearly assignable to the -C-O- stretching vibration as found in the oxazoline. Hence the latter structure tends to explain major changes in the spectrum, which the olefin alone does not. One further point--if olefin formation is indeed occurring, it is unlikely that it would be formed through cure reversal (Figure 29, rxn 1) since acid is consistently consumed; no evidence has been found (in the IR) for carboxylic acid reformation. Rearrangement of the aziridine under the influence of the cure environment (rxn 2) would be a more logical explanation for olefin formation. The fact that these peaks do not appear in the IR spectrum of the Telagen CT-MAPO or MAPS reactions also suggests that the presence of these absorptions in acyl aziridine reactions is due to oxazoline formation, since neither the phosphine oxide nor the sulfide aziridine can rearrange to an oxazoline structure.

The control (uncatalyzed) AcAz-Telagen CT reaction run at 50°C also shows the same extra peak growth as catalyzed runs, indicating that the transformation in question is characteristic of acyl aziridine-carboxylic acid reaction. (The 8.8 to 8.9  $\mu$  peak is conspicuously absent from the spectra of the MAPO and MAPS cured polymers indicating the absorption in the AcAz system to be due to some group other than -C-O- in the ester formed from carboxylic acid attack on the imige ring.)

In summary, oxazoline formation may be occurring during the CrOct catalyzed cure of AcAz and Telagen CT, though olefin formation is also remotely possible.

£ (5)etc. 0 H R' CH2 0 → R-C-N-CH-CH + H0-C-R''' -0-C-R " elimination, R" = CH3 2 പ  $R-\ddot{C}-NH-CH+CH = CH_2$ R-C-N-C-C-2 0= R-C-OH (desired product) 0 H R' R'' 0 R-C-N-CH-CH-O-C-R''' R-C-K Γ<sup>H</sup> R-C-K ¢-R" - H ч Ж R-C-N 0 R-Q homopolymerization olefin formation via ring opening о к...- сон oxazoline formation [cat]? - 8 R-0=0 R-0=0

Figure 29 - Possible Reaction Paths in the Telagen CT-AcAz Cure System

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### D. Aziridine Catalysis Studies

The effectiveness of CrOct and FeOct as catalysts for the aziridine cure system was explored further using Telagen CT and the curatives AcAz\* and BITA.\*\* In addition, the effect of AP,  $Fe_2O_3$ , and water on imine cure was examined. Table XXXIIa lists the reaction rate data from these experiments.

In general, both organometallics are effective as cure catalysts, in that they promote the conversion of acid and imine to ester at ambient temperature in acceptable time periods. The reaction appears stoichiometric in a practical sense, that is, acid consumption proceeds to a level expected in a cure reaction (60-80%) and absorbances noted in the IR spectra do not indicate significant or appreciable side reaction of curative (oxazoline formation).

Figure 30 displays the cure conversion behavior of the Telagen CT-BITA system. As previously reported, the reaction rate is quite temperaturedependent. In fact, rates are identical for the catalyzed 25°C mix and the uncatalyzed 50°C run. A 1.0% loading of CrOct will promote the reaction to the 60% level in approximately 30 hr. while 8 days are required for the uncatalyzed gumstock cure. Hence, CrOct is an effective catalyst for the BITA curative system.

Figure 31 is a similar display for the results from the study of Telegen CT-BITA cure promoted by 1% FeOct. Conversion rates and thermal effects are identical to the CrOct system. A plausible interpretation follows. In the epoxide system, the reaction may be truly nucleophilic. That is, the ring opens only (or predominantly) through attack by the activated acid. Activation of the carboxylate (concentrating electronic charge by proton removal and coordination) is of critical importance in epoxide catalysis. In contrast, the aziridine ring opens readily and the carboxyl group is acidic chough by itself to undergo condensation with the imine. The organometallic thus merely facilitates a reaction which does not require the extensive activation needed in epoxide-carboxyl cure.



### TABLE XXXIIa

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## <u>CURE OF TELAGEN CT</u> WITH SELECTED AZIRIDINES

	Catalyst	Extra Component			Meq. CO <sub>2</sub> H	
<u>Curative</u>	(wt. %)	(wt. %)	Temperature	Time	Present	<u>% Rxn</u>
BITA	1% CrOct	-	R.T.	Initial	0.306	0.0
				24 h <b>r</b> .	0.128	58.2
				3	0.112	63.4
				4	0.113	63.1
				7	0.101	66.9
				10	0.10	67.3
BITA	1% CrOct	-	50	Initial	0.306	0.0
				2	0.190	37.9
				6	0.130	57.5
				19	0.110	64.1
				26	0.100	67.3
				43	0.099	67.6
				3 d <b>a</b> ys	0.090	70.5
				4	0.100	67.3
				7	0.089	70.9
				10	0,86	71.8
BITA	0	-	R.T.	Initial	0.308	0.0
				1 day	0.206	33.1
				3	0.168	45.4
				4	0.143	53.6
				7	0.128	58.4
				9	0.109	64.6
				10	0.108	64.9
BITA	0	-	50	Initial	0.308	0.0
				3 hr.	0.264	14.2
				7	0,205	33.4
				20	0,150	51.3
				27	0,128	58.4
				43	0,124	59.7
				3 d <b>a</b> ys	0.113	63.3
				4	0.109	64.6
				7	0.108	64.9
				8	0.106	65.5
				10	0.105	65.9

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	Catalyst	Extra Component			Meq. CO <sub>2</sub> H	
<u>Curative</u>	<u>(wt. %)</u>	(wt. %)	Temperature	Time	Present	<u>% Rxn</u>
AcAz	1% CrOct	-	R.T.	Initial	0.314	0.0
				5.5 h <b>r.</b>	0.308	1.9
				2.5	0.250	20.4
				3 days	0.234	25.4
				6	0.204	35.0
				8	0.184	41.4
				9	0.180	42.6
AcAz	1% CrOct	-	50	Initial	0.314	0.0
				2 hr.	0.282	10.2
				7	0.249	20.7
				l day	0.172	45.2
				2.5	0.111	64.6
				3	0.084	73.2
				6	0.056	82.2
				9	0.047	85.0
AcAz	I% FeOct	-	<u>R.T.</u>	Initial	0.330	0.0
				4 hr.	0.316	4.2
				2.5 days	0,178	46.1
				3	0.188	43.0
				6	0.177	46.4
				8	0,172	47.9
			_	9	0,173	47.6
ACAZ	1% FeOct	-	50	Initial	0,330	0.0
				4 hr.	0.285	13.6
				7	0.257	22.1
				l day	0,253	23.3
				2.5	0.210	36.4
				3	0.167	49.4
				6	0.135	59.1
				9	0,118	64,2
BITA	1% FeOct	-	R.T.	Initial	0,334	0.0
				l day	0.132	60,4
				3	0.116	65.3
				4	0,120	64.1
				7	0.113	66.2
				9	0.125	62.5
				10	0,116	65.3

# TABLE XXXII (Continued)

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	1% FeOct			lime	Present	<u>% Rxn</u>
BITA		-	50	Initial	0.334	00
				2 hr.	0.174	47 9
				4	0,120	64 1
				7	0.132	60 /
				24	0.114	65.8
				28	0.110	67 1
				2 days	0.110	67.1
				3	0.104	68.8
				4	0.100	70.1
				7	0.102	69.5
				8	0.074	77.8
				10	0.098	70.7
AcAz	0	5% AP	R.T.	Initial	0 382	0.0
				l dav	0.309	10.0
				3	0.328	14 1
				4	0.300	21 4
				7	0.300	21.4
				9	0.280	26 7
				10	0.282	26.2
ACAZ	0	1% Fe <sub>2</sub> O <sub>3</sub>	R.T.	Initial	0.340	0.0
				l day	0.311	8.5
				3	0.312	8.2
				4	0.300	11 7
				7	0.296	12 9
				9	0.289	15.0
				10	0.286	15.8
AcAz	1% CrOct	2 µl H <sub>2</sub> O	R.T.	Initial	0.327	0.0
		2		4 hr.	0.298	8.8
				2 day	0.268	18.0
				3	0.236	27.8
				6	0,210	35.7
				8	Ů.197	39.7
				9	0.197	39.7

# TABLE XXXII (Continued)

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	Catalyst	Extra Component			Meq. CO <sub>2</sub> H	
<u>Curative</u>	<u>(wt. %)</u>	<u>(wt. %)</u>	Temperature	Time	Present	<u>% Rxn</u>
AcAz	1% CrOct	5% AP	R.T.	Initial	0.319	0.0
				5 hr.	0.298	6.5
				2 days	0.262	17.8
				3	0.231	27.6
				6	0.210	34.2
				8	0.198	37.9
				9	0.166	47.9
AcAz	1% CrOct	5% AP	50	Initial	0.319	0.0
				5 hr.	0.256	19.7
				l day	0.179	43.9
				2	0.115	63.9
				3	0.090	73.7
				5	0.060	81.2
				7	0.053	83.4
				9	0.044	86.2
AcAz	1% CrOct	1% Fe <sub>2</sub> 0 <sub>3</sub>	R.T.	Initial	0.300	0.0
				2 days	0.270	10.0
				3	0.259	13.6
				6	0.231	23.0
				8	0.238	20.7
				9	0.210	30.0

# TABLE XXXII (Concluded)

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Figure 32 displays the rate patterns for the AcAz cure system catalyzed by the iron and chromium organometallics. At ambient temperatures, reaction is slower for both catalyst systems. Two factors may explain the difference between the two aziridine curatives: (1) AcAz is intrinsically more stable than BITA and should be less disposed to reaction; (2) possibly the most important difference is solubility--BITA is a liquid which readily dissolves in the polymer to give a transparent mixture. AcAz is a white powder which disperses in the Telagen CT to give an opaque gumstock. The fact that FeOct gives faster reaction rates, especially during initial stages of the conversion, can be attributed to the presence of diluent (mineral spirits) in the catalyst which would contribute to the solution and diffusion of reactants and catalyst. A recurring phenomenon is the high degree of conversion observed for the 50°C CrOct catalyzed system. It has been noted before and still remains the subject of speculation.

Study of the effect of propellant components and contaminants on the catalyzed cure of the aziridine binders began with an examination of the AcAz system. Figure 33 displays the effect of AP,  $Fe_2O_3$ , and  $H_2O$  on the CrOct promoted reaction. The presence of AP has shown no effect on rate and stoichiometry at either 25° or 50°C. The presence of  $Fe_2O_3$  retards the the conversion at 25°C, possibly by aggravating the solubility situation.

Figure 34 shows the effect of AP and Fe<sub>2</sub>O<sub>3</sub> on the cure system without CrOct. AcAz and Telagen CT at 25°C will react to approximately 15% conversion in the time period shown. Hence, the reactivity displayed in the Fe<sub>2</sub>O<sub>3</sub> control would be due to the cure components themselves and not the extra ingredient. This suggests that Fe<sub>2</sub>O<sub>3</sub> is interfering (rate retardation of ~ 10%) with the catalytic process, perhaps by making it more difficult for CrOct to diffuse through the medium. A 5% loading of AP appears to catalyze the reaction slightly in the absence of CrOct (25% conversion in 10 days versus 15% for the cure functionalities alone).

In Figure 34, the effect of water in an amount equimolar to CrOct is shown; the rate pattern is identical within experimental error to that of the noncontaminated catalyzed system.

Hence, both FeOct and CrOct are effective cure catalysts for the acylaziridine-Telagen CT system. Rate differences between BITA and AcAz appear due to differences in solubility. AP and  $H_2O$  do not alter rate behavior in the AcAz cure reaction, though Fe2O3 does.

Supplemental aziridine evaluation data are reported in Table XXXIII.



Figure 32 - The Effect of Catalysis and Temperature on the Cure of Telagen CT with AcAz, a Comparison of the Catalytic Effectiveness of CrOct and FeOct







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### TABLE XXXIII

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# SUPPLEMENTAL AZIRIDINE CURE CATALYSIS DATA

System	<u>Catalyst Level</u>	Extra-Ingredient	<u>Temperature</u>	Time (hr.)	<u>%Rxn</u>
Telagen/BITA	0.5% CrOct	0	R.T.	2 10 23 48 3 days 7 days 10 days	9.7 34.3 42.2 50.0 53.4 57.5 58.5 Cured
Telagen/BITA	0.5% CrOct	5 <b>%</b> Ap	R.T.	10 23 48 3 days 7 days 10 days	28.8 39.4 47.7 51.1 55.7 56.4 Cured
Telagen/AcAz	0.5X CrOct	5% AP		2 10 23 48 3 days 7 days 10 days	0 1.0 4.3 10.4 15.6 22.6 26.6
Telagen/BITA	1% CrOct	17 Fe <sub>2</sub> 0 <sub>3</sub>		5 18.5 48 3 days 7 days 10 days	23.2 37.1 44.3 47.9 45.4 48.2
Telagen/BITA	1% Fe0ct	17 Fe <sub>2</sub> 03		5 18.5 48 3 days 7 days 10 days	23.7 38.5 47.0 49.5 54.1 57.6
Telagen/AcAz	l% FeOct	17 Fe <sub>2</sub> 0 <sub>3</sub>		5 18.5 48 3 days 7 days 10 days	0.0 10.0 13.7 18.4 28.1 28.4

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As reported previously, AcAz reacts at a significantly slower rate than the other aziridine curatives (BITA, MAPO, MAPS). The 0.5% CrOct is too low a catalyst level to bring about the cure of Telagen with AcAz in a reasonable time period. The presence of AP slows the cure for both AcAz and BITA by 5-10%. Iron oxide slows the cure rate of the CrOct catalyzed system (Telagen/BITA). A similar retardation effect by Fe<sub>2</sub>O<sub>3</sub> is noted in the FeOct catalyzed BITA/Telagen binder though it is not as pronounced as the chromium system.

### E. Ammonyx T Effect in Aziridine System

Figures 35 and 36 show the effect of 4% by weight of Anmonyx T on the aziridine cure of Telagen CT. In the BITA system the presence of the Anmonyx alone promotes the reaction during early stages of cure but not as efficiently as CrOct. When AP and CrOct are present the emulsifier diminishes the efficiency of the Cr catalyst and the ultimate extent of reaction is 10% less than in the absence of Ammonyx T. In the AcAz system, the presence of the emulsifier has no significant effect on the cure rate.





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### VII.

### POSTCURE SURVEILLANCE

It has been demonstrated that chromium salts will catalyze the reaction of carboxylic acid and epoxide at room temperature rapidly and stoichiometrically. This catalyst is also effective in controlling the cure reaction of CT polymers with aziridines. The occurrence of postcure reactions will be a function of prevailing conditions in the propellant, that is, temperature, amount and relative proportions of unconverted cure functionality, etc.

### A. Background - Epoxide Based Cure

The epoxide ring is a polar, strained structure susceptible to attack by nucleophilic species. The nature of the reaction is, of course, dependent on the type of attacking group, reaction conditions, and the type of catalysis, if any. King opening may proceed by way of acidic or basic m chanisms. In the basic system, attack occurs at the electron deficient carbon, leading to ring opening and the formation of an anionic product.

$$B \stackrel{\bigcirc}{\to} + c \stackrel{\bigcirc}{-c} c \stackrel{\downarrow}{\longrightarrow} B \stackrel{\bigcirc}{-c} c \stackrel{\downarrow}{\longrightarrow} B \stackrel{\bigcirc}{-c} c \stackrel{\downarrow}{-c}$$

In the acidic situation, the epoxide oxygen is protonated and ring opening may occur in a number of ways, that is, in a concerted fashion, or an open ion two-step mechanism, or through a four-center associated intermediate.

concerted



open ion

four-center  $\searrow \bigoplus_{\delta} \bigoplus_{k} \bigoplus_$ 

The reaction of specific interest here is the addition esterification of a carboxylic acid with an epoxide.

$$R = C = OH + C = C = R = C = O = C = OH$$
(1)

Several competing reactions are possible depending on conditions and reactant concentrations.





Reaction (2) is favored by high temperature (200°C), excess alcohol, and water removal. (Under actual polymerization or curing conditions, equilibrium conditions prevail and this reaction is essentially suppressed since the only route for water removal is evaporation and/or reaction with epoxide to form glycols.)

Etherification, reaction (3), is favored by low temporature (less than 200°C), excess epoxide, and acid catalysis. This route will lead to a polyether polymer of the following type  $R - \overset{H}{C} - O \leftarrow \overset{L}{c} - \overset{L}{c} - O \rightarrow_n H$ .

In the presence of a basic catalyst the reaction is specific to addition esterification (No. 1); only after complete consumption of acid will excess epoxide react with the product alcohol via the etherification route.

Hence, mild temperature, base catalysis, and equivalent concentrations of reactant acid and epoxide are conditions specific to reaction 1. Acid catalysis and low temperature favor etherification; excess epoxide will optimize this route. In the absence of acid or base catalysis, temperature becomes the controlling reaction parameter, with etherification being increasingly suppressed at higher temperatures.

In addition, several other interactions are possible; oxidative crosslinking of olefin groups present in the binder promoted by the cure catalyst; degradation (e.g., dehydration, decarboxylation) of the ester product through the influence of the metal.

### B. <u>Surveillance Studies Undertaken</u>

The intent of these studies was to determine the role which the cure catalyst may play in propellant aging. Specific interactions of interest (due to the presence of catalyst) include: reversal of the cure linkage, catalytic decomposition of the product ester, other reactions within the product (e.g., oxidative cross-linking of olefin groups), and promotion of reactions due to other propellant components. To separate the variables in the complex propellant, a series of surveillance studies is under way from simple model studies (to examine potential fundamental chemical interactions) to 40-lb. propellant blocks. The aging studies include the following:

1. <u>Model reaction systems with and without additional propellant in-</u><u>gredients</u>: These were intended to determine situations of cure reversal and/ or reaction of the product with the added ingredient attributable to the presence of the catalyst.

2. <u>Cured binders</u>: The intent is the same as above.

3. Excess epoxide studies: Both, model and polymer, were examined to assess the importance of etherification reactions as described above.

4. <u>Excess acid studies</u>: Both model and polymer, to determine the importance of postcure esterification reactions.

5. <u>Whole propellant surveillance</u>: Mechanical property changes with time on large samples.

### C. Binder Surveillance - Results

Cured binder systems under ambient storage for extended time periods were examined by infrared for evidence of postcure reactions. An IR spectrum was taken of each sample (cured and stored between salt plates); the ester content was measured and compared to the 30-day reading (a time at which cure has usually occurred) for the sample to determine whether cure continued (increase in ester) or whether cure degradation occurred (decrease in ester content). In addition, the spectrum was examined for the formation of or changes in other peaks, a situation which would indicate degradation of other groups within the polymer, or postcure reaction. The results from the stored samples are shown in Table XXXIV. Without exception, no significant infrared detectable chemical change has occurred in the epoxide cured systems after extended periods of ambient storage. This holds true for even those samples containing the various propellant solids. Only the aziridine curative AcAz shows evidence of a slow continuing ester formation reaction.

It can be safely concluded that the presence of the chromium salts (octoate, oleate) in a cured binder does not adversely affect the stability of the cure or the binder itself. In fact, the ambient cured and stored acid/epoxide based binders are chemically stable for periods of up to 22 months even in the presence of propellant solids.

The results for aziridine systems show a dependence upon the specific curative used, as expected. However, the only aging effect attributable to the presence of the Cr catalyst is continued ester formation with the slow reacting AcAz compound.

### D. Model System Surveillance - Results

Products of model reactions run at  $25\,^{\circ}$ C were stored at ambient temperature and at  $50\,^{\circ}$ C. These were analyzed after extended storage and the results compared to data obtained within about 2 weeks after initiation of the reactions. The results of these analyses are shown in Tables XXXV and XXXVI. A note of explanation is needed concerning this table. The short lines represent the relative concentrations of the side products and their position in the gas chroatogram, relative to the height and major product (long line). In the figure, where the major product is shown off-scale, the total amounts of the minor components never exceed 1% of the total theoretical yield.

In general, no change has occurred in the product composition of most of the CrOct catalyzed reactions. It is difficult to determine whether the minor products noted in the samples containing extra ingredients are due to solution of that ingredient or actual postcure reactions. Regardless, the presence of the cure catalyst does not cause reversal nor does it promote postcure reactions to any appreciable degree.

### TABLE XXXIV

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### CUREO BINOER SURVEILLANCE

					Continued	
		Time	New	Cure	Cure	
<u>cure System</u>	Temperaturea/	(month)	Peaksb/	<u>Reversal</u>	_(%) <u>d</u> /	<u>Othere</u> /
Tel/AcAz/0.5% CrOct	R.T.	11	No	No	54	No
Tel/AcAz/0.5% CrOct						
5% AP	R.T.	11	No	No	55	No
Tel/BITA/1% CrOct						
1% Fe2O3	R.T.	11	No	No	No	No
Tel/BITA/1% FeOct						
1% Fe203	R.T.	11	No	No	8	No
Tel/AcAz/1% FeOct						
1% Fe203	R.T.	11	No	No	50	No
Tel/AcAz/Control	R.T.	10	No	No	50	No
Tel/AcAz/5% AP	R.T.	10	No	No	35	No
Tel/BITA/1% Fe20a	R.T.	10	No	No	No	No
Tel/BITA/1% CrOct						
5% AP	R.T.	10	No	No	2	No
Tel/BITA/1% CrOct						
2 ul. H <sub>2</sub> O	R.T.	10	No	No	5	No
Tel/AcAz/4% Ammonyx T	R.T.	10	No	No	40	No
Tel/AcAz/1% CrOct		_				
4% Ammonyx T	R.T.	10	No	No	40	No
Tel/AcAz/1% CrOct				-		
4% Ammonyx T/3% AP	R.T.	10	No	No	35	No
Tel/BITA/4% Ammonyx T	R.T.	10	No	No	15	No
Tel/BITA/4% Ammonyx T					-	
1% CrOct	R.T.	10	No	No	10	No
Tel/BITA/47 Anmonys T				••••		
1% CrOct/4% AP	R.T.	10	No	No	No	No
lel/AcAz/5% CrOIPS	R.T.	10	No	No	30	No
Tel/BITA/5% CrOIPS	R.T.	10	No	No	No	No
Tel/ERLA/5% CrOIPS	R.T.	10	No	No	No	No
Tel/ERLA/5% CrDIPS						
32. AP	R.T.	10	No	No	No	No
Tel/ERLA/5% CrDIPS						
4% Astronyx T	R.T.	9	No	No	No	No
Tel/ERLA/5% CrD1PS						
2% AP/4% Amenonyx T	R.T.	9	No	No	No	No
Tel/ERLA/1% CrOct						
17. AP/47. Amonyx T	R.T.	9	No	No	No	No
[e1/ERLA/0.57. CrOct						
1% AF/4% Ammonyx T	R.T.	9	No	No	No	No
Tel/ERLA/0.52 CrOct						
37. AP/47. Ammonyx T	R.T.	9	No	No	No	No
Tel/ERLA/0.5% CrOct						
3% AP/1% Ammonyx T	R.T.	9	No	No	No	No
Tel/ERLA/17, CrOct	R.T.	24	No	No	No	No
"el/ERLA/1". CrOleate	R.T.	24	No	No	No	No
Tel/ERLA, 6% FeOct	R.T.	23	No	No	No	No
Tel/17 CrOct	R.T.	22	No	No	No	No
Tel/(23%) ERLA/1%						•••
CrOctf/	<b>R.</b> T.	22	No	No	No	No
(e1/(e6%) ERLA/1"						
(10:11/	R.T.	22	No	No	No	No
Tel (12) ERLA/12			•			•••
:00:1/	R.T.	22	No	No	No	No

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### TABLE XXXIV (Concluded)

					Continued	
<u>Cure System</u>	<u>Temperatura</u> <sup>2</sup> /	Time (month)	New Peeksb/	Cure <u>Reversel</u> C/	Cure ( <u>7)</u> <u>d</u> /	Othere/
Tel/(97%) ERLA/1%						
CrOct <u>f</u> /	R.T.	22	No	No	No	No
Tel/(120%) ERLA/1%						
CrOct <u>f</u> /	R.T.	22	No	No	No	No
Tel/(144%) ERLA/1%						
CrOctf/	R.T.	22	No	No	No	No
Tel/(166%) ERLA/1%						
CrOct <u>f</u> /	R.T.	22	No	No	No	No
Tel/ERLA/0.2% CrOct	R.T.	18	No	No	No	No
Tel/MAPS/1% FeOct	R.T.	18	No	No	No	No
Tel/AcAz/1% FeOct	R.T.	16	No	No	20	No
Tel/ERLA/1% TOPA						
0.4% CrOct	R.J.	16	No	No	No	No
Tel/AcAz/0.5% CrOct	R.T.	15	No	No	5	No
Tel/AcAz/1% CrOct	R.T.	15	No	No	20	No
Tel/BITA/1% CrOct	R.T.	12	No	No	No	No
Tel/AcAz/1% CrOct	R.T.	12	No	Nc	5	No
Tel/AcAz/1% FeOct	R.T.	12	No	No	30	No
Tel/BITA/17 FeOct	R.T.	12	No	No	No	No
Tel/AcAz/1% CrOct						
2 µ1. H <sub>2</sub> 0	R.T.	12	No	No	10	No
Tel/AcAz/1% CrOct						
5% AP	R.T.	12	No	No	10	No
Tel/AcAz/1% CrOct						
17. Fe203	R.T.	12	No	No	15	No
Tel/ERLA/5% CrOIPS	R.T.	11	No	No	No	No
XS Tel/ERLA/1% CrOct	R.T.	11	No	No	No	<b>N</b> (8)
Tel/ERLA/1% CrOct						
4% Ammonyx T	R.T.	11	No	Yes	No	No
Tel/ERLA/0.5% CrOct						
47. Ammonyx T	R.T.	11	40	No	No	No
Tel/ERLA/0.5% CrOct						
5% AP/4% Ammonyx T	R.T.	11	No	No	No	No
XS Tel/ERLA/0.5% CrOct	R.T.	11	No	No	No	No
Tel/XS ERLA/0.5% CrOct	R.T.	11	No	No	No	No
Tel/BITA/0.5% CrOct	R.T.	11	No	No	No	No
Tel/BITA/0.5% CrOct						
5% AP	RT.	11	No	No	No	No

<u>e</u>/ Tempereture of storege.

b/ The epperance of new pasks in the spectrum would indicate the formation of a new chemical antity during storage.

 $\underline{c}$  / Cure reversel would be indicated by loss of ester relative to previous reading.

d/ Continued cure would be indicated by en increase in ester absorbance accompanied by a decrease in the remaining acid content.

e/ This cetegory would include observations not felling under the other three classifications.

<u>f</u>/ The numbers in perentheces indicate the percent of the etoichiometric amount of epoxide used in the experiment. These mixes were used for the esteblishment of e celibration curve.

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TABLE XXXV

# SURVEILLANCE OF MODEL REACTION SAMPLES

4

AGE	COMPOSITION	REACTION PRODUCTS
+0 11 2	15 Croct/5\$AP/Age/BE	
• <b>01</b>	0.1% Croct/5%AP/Age/Ena	
• • • •	15 CrOct/PA/Age	
् म <b>्</b>	2≸ Crost/PA/Age	
• 0 <b>9</b>	15 Croct/0A/Age (sheken)	
- <b>0</b>	25 Croct/0A/Age	
• <b>08</b> ∑∑	15 Croct/941H20/Age/Eha	
جر <b>no.</b>	15 CrOct/18w1H <sub>2</sub> 0/Age/Ena	
•о <b>ц</b> 22	15 Croct/90u1H20/Age/Ena	
• <b>्य</b> ्र	15 Croct/0.55Fe202/Age/Ena	
जन्म र	15 Croct/15Fe203/Age/Eha	
-0 - ₽	10% CrOct/Age (Homopolymerization)	
E	65 Croct/Age/Eha	

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AGE	CONTOSITION	REACT AT PRODUCES
e E	1\$ Croct/1\$ A1/Age/Eha	
O EL	15 CrOct/0.55Roccal Age/Eha	
0 11	14 CrOct/0.55 Pherc1-B-N Age/Eha	
100	15 CrOct/160xm1du/Agc/ 12 n	
	0.5%CrOct/0.5%FeOct/Age/	*
• 0 <b>2</b>	1≸ CrOct(ROC)/Age/Ehs	
•0 <b>f</b>	14 Croct/PA/Age/Ena	
01		
	15 CrOct/OA/Age/Eha	
- 100-	0.4\$ CrOct/OA/Age/Ena	*
• 014	15 Crost/15 IDPA/Age/Era	
• <u>•</u> • • • •	14 Crost/N TDPA//gs/Ehe	
* *	20% CrOct/Age/Ehu	
3		

TABLE XXXV (Continued)

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AGE	COMPOSITI ON	REACT OF PRODUCTS	
+ 0 1 + 1	45 CrOct/Age/Ehn		
د <sup>ر</sup> <b>в</b> 0.	15 Croct/xs Age/OA		
16 <b>20</b>	15 Croet/Ester Product/Age		*
14 100.	15 Croct/18103/Age/Eha		<b>.</b>
14 Jao -	No Croct/HNOS/Age/Ehn		
14 mo.	15 Croct/NH4,02/Age/Eha		
14 20	No Croct/NH <sub>4</sub> Oil/Age/Eha		
14 100.	1\$ Croct/ROBC2/Age/Ease		*
14 100.	No Croct/HOHO2/Age/Ena		

- n-propyluturate
  Research Organic Chemicals, Inc.
  desfred ester product
  Phenyl-B-napt ylamine
  antioxidant viosiphenylmine
  ocianoic acid
  propi wie acid NONO (NOC) • TEPPA TEPPA PA

TABLE XXXVI

Contraction of

# SURVEILLANCE OF MODEL REACTION SAMPLES (50°C STORAGE)

ÅEe	Composition	Reaction Products
, <b>180</b> .	25 Cr-neo/Age/Bha	
2 6 <b>mo.</b>	15 CrOct/Age/Eha/55 AP	
. S. BO.	15 Croct/Age/Eha/18 ul H20	
• • • • •	15 CrOct/Age/Bha/9 µl H20	
23 ( <b>30.</b>	1\$ CrOct/Age/Eha/90 ul H20	
23 mo.	Product/Tol/9 ul H <sub>2</sub> 0	
7.5 ( <b>mo.</b>	15 Cr-neo/Age/Eha/90 Ll H20	
23 580.	1\$ CrOlea/Age/Eha/9 μl H20	
-30 <b>-00</b> -	15 Croles /Age/Eha/90 µl H20	
• <b>08</b> • 2 • 2	0.5\$ Fe203 /Age /Eta	
23   <b>BO.</b>	1% CrOct/0.5% Pe203/Age/Eha	
. <u>(</u> 2) <b>BO.</b>	1\$ CrOct/1\$ Fe20g/Age/Bha	
<i>े</i> टे <b>ह</b> 0.	1\$ Fe205/Age/Eta	
-1 mc.	1≸ ∰Age/Ens	

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TABLE XXXVI (Continued)

<u>^6</u>	Composition	Reaction Products
0	2% Fe(Oct)2/Age/Eha	
, 12 ( <b>BO</b> .	25 Pe(Oct) <sub>2</sub> /18 µ1 H2O/Age/Bha	
2. BO.	0.5\$ CrOct/0.5\$ Pe(Oct)2/Age/Eha	
20 <b>mo.</b>	1\$ TRU/Age/Bha	
20 mo.	15 DFLDC/Age/Eha	
50 <b>mo.</b>	15 TFL /Age /Zha	
20 20.	15 TEL/Age/Bha	9.1
20 000	15 Croct(HOC)/Age/Eha	*
20 20.	15 FeOct(ROC)/Age/Eha	
20 <b>BO.</b>	15 PbOct(ROC)/Age/Bha	
20 00.	1\$ CrFormate(ROC)/Age/Eha	
20 50.	1\$ TPLC/Age/Eha	
19 <b>BO</b> .	1\$ CrOct/PA/Aze/Bha	
19 <b>mo</b> .	C.45 CrUct/PA/Age/Eha	

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TABLE XXXVI (Continued)

4

Age	Composition	Reaction Products
9 5 1	1% CrOct/ON/Age/Eha	
10 20.	0.4% CrOct/OM/Age/Bha	
	15 CrHep/Age/Ena	
13 50.	1\$ CrHex/Age/Eha	
19 50.	0.5% TDFA/Age/Eha	
19 NO.	1\$ TDPA/Age/Eha	
19 mo.	24 TDFA/Ace/Eha	
<b>0</b> 1	1% Croct/1\$ TDPA/Age/Eha	
200 200 200 200	0.5\$ TDFA/0.5\$ Fe203/Age/Eha	
le mo.	9 μ1 HClO <sub>4</sub> /Age/Sha	
15 <b>m0</b> .	0.5% TDPA/0.5% Fe203/0.5% CrOct/Age/Ehm	
• <b>01</b> 51	1% CrOct/10 µ1 HCIC <sub>4</sub> /Age/Eha	
10 mo.	15 CrOcc/100 H1 HCLO4/Age/Eha	
1º BO.	15 CrOct/5 μl HCl04/Age/Eha	

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TABLE XXXVI (Continued)

Acc	Composition	Reaction Products
17 0.	1\$ CrOct/Ot/Age	
17 20.	25 CrOct/ON/Age	
1	0.5\$ CrHep/OA/Age	
17 160.	1\$ CrHep/OA/Age	
16 <b>mo.</b>	1\$ CrOct/xs Age/Bha	
16 📷 .	1\$ CROct/xs Age/CM	
16 20.	1\$ CrOct/x. Age/Product	
16 <b>BO.</b>	1\$ Pyridine/Age/OA	
16 📷 .	1\$ TEA/Age/OA	
15 80.	1\$ Croct/HRO <sub>5</sub> /Age/Eha	
15 mo.	No CrOct/HNO <sub>3</sub> /Age/Eha	
1. Tho.	1≸ CrOct/RiteCH/Age/Eha	
li J <b>oo.</b>	No CrOct/NH <sub>4</sub> OH/Age/Sha	
1: 	1\$ CrOct/RONO2/Age/Eha	
11 <b>eo.</b>	No CrOct/RONO2/Age/Eha	

TABLE XXXVI (Concluded)

Age	Composition	Reaction Products
13 BC.	15 CrOct/15 AmyxT/Age/Bha	*
13 mo.	15 CrOct/45 AmyxT/Age/Eha	
13 🔟 .	No CrOct/1\$ AmyxT/Age/Eha	
13 mo.	No CrOct/45 AmyxT/Age/Eha	
13 <b>mo.</b>	No CrOct/HF/Age/Eha	
13 mo.	1\$ CrOct/HF/Age/Eha	
13 mo.	1\$ CrOct/N20/Age/Eha	

<ul> <li>n-propynitrate</li> </ul>	- Research Chemicals, Inc.	<ul> <li>desired ester product</li> </ul>	<ul> <li>Chromium-neo-decanoate</li> </ul>	- Chromium Oleate	B*trichloro-N-trimethylboratine	<ul> <li>Triphenyl germanium</li> </ul>	<ul> <li>Diphenyllead dichloride</li> </ul>	Triphenyllead chloride	<ul> <li>Tetraphenyllead</li> </ul>	<ul> <li>Tetracthyllead</li> </ul>
RONO2	(ROC)	*	Cr-neo	Croles	TCMB	TR	DPLDC	TFLC	1 1 1 1	ISI

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Chromium Heptano	Chromium Hexanos	antioxidant thio	ester product fr
9	I	H	•
CrHep	CrHex	TDPA	Product

- Chromium Heptanoate
  Chromium Hexanoate
  antioxidant thiodiphenylamine
  ester product from large batch of catalyzed Age and Eha
  Anmonyx T
  - AmyocT

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### VIII.

### PROPELLANT STUDIES

### A. Objective of Propellant Studies

The overall objective of this portion of the program was the demonstration of the ambient-temperature-cure catalyst system which Midwest Research Institute (MRI) studies indicated was the best for CTPB propellants.

### B. Approach

The following six carboxyl binder systems were examined: the carboxyterminated prepolymers, Telagen and HC-434, and the terpolymer polybutadiene acrylic acid acryloritrile (PBAN) cured with either epoxides or aziridines. This portion of the program was divided into two phases. The first part, "Catalyst Behavior with Composite Propellant Ingredients" (Phase III of the overall program), consisted of basic studies of various catalysts which were suggested by the results obtained at MRI. The basic studies were performed on the six binder systems and involved: (1) determining relative effectiveness of various catalysts, (2) effects of normal propellant solid ingredients on catalyzed ambient cure, and (3) selection of catalyst concentrations to be tested in actual composite propellant formulations. The second part of the ABL program, "Scale-Up Demonstration of Cure Catalysis" (Phase IV of the overall program), consisted of: (1) selection of baseline propellant formulations which would have the optimum cure catalyst needed to yield good slurry pot life, reasonably short ambient cure time and good cured mechanical properties, (2) determination of the effect of various other propellant ingredients and likely contaminants on catalyzed ambient cure, and (3) evaluation of the aging characteristics of solid propellants formulated with these cure catalyst systems.

### CATALYSIS BEHAVIOR WITH COMPOSITE PROPELLANT INGREDIENTS

### A. Summary

The Telagen prepolymer was found, in gumstock studies, to cure at ambient temperature with the triepoxide ERLA-0510 in 1 week with 0.40%, and in 2 weeks with 0.37% chrominum octoate (CrOct) cure catalyst. Ordinary propellant solids such as ammonimum perchlorate (AP), aluminum, and  $Fe_2O_3$  do not affect the cure significantly. The HC-434 prepolymer behaves in a similar fashion in gumstock studies. The potential cure catalysts, chromium oleate and chromium 2,6-diisopropyl salicylate were found to be less effective than CrOct at equal Cr concentrations.

Ambient cure of the two CTFB preprlymers with the aziridine HX-868 required about twice as much CrOct catalyst as did the expoxide-cured systems.

Normal propellant solids interfered with the CrOct ambient cure of binders based on the PBAN prepolymer. The PBAN-aziridine system was very slow to cure, even with high CrOct concentrations.

In each of the six binder systems, catalyst levels were selected to be used for the first 1-lb. propellant mixes to be made in Phase IV.

### B. Discussion of Results

1. <u>Procedure</u>: Model compound studies performed at Midwest Research Institute indicated that the chromium coordination compounds were the most effective catalysts for carboxy binder systems. Therefore, ABL examined three such compounds; chromium 2-ethyl hexanoate (generally called chromium octoate or CrOct), chromium oleate and chromium 2,6diisopropyl salicylate (CrDIPS). Testing consisted entirely of gumstock studies during this phase.

Binder studies were conducted to determine the concentration of cure catalyst which would provide cure at ambient temperature  $(25 \pm 1^{\circ}C)$ in a reasonable time period (1 to 2 weeks). The binders consisted of the prepolymer, curing agents, IDP or DOA plasticizer and cure catalyst. Some also contained various solid fillers. The degree of cure was followed by periodically measuring mechanical properties, gel content, and swelling. Experimentally the cure study procedure was to prepare 30-g mixes of the binder consisting of 70% polymer, the specified quantity of cure catalyst and solids, and the remainder was plasticizer. The binder was evacuated and cast into four 8-in. long by 1/4-in. diameter Teflon tubes. After curing for the specified time period, the binder was removed from the Teflon tube and specimens were prepared. Duplicate mechanical property specimens (1/4-in. by 1-in. cyclinders) were prepared by epoxy bonding to wooden end tabs. The samples were pulled on an Instron tester at a strain rate of 1 in/in/min until 35% elongation. The secant modulus at 30% elongation was then calculated. Samples were not tested till break because, frequently, the binder was stronger than the bond to the wooden end tab which resulted in specimens breaking at the bond rather than within the binder phase. The bond was always strong enough to permit testing the specimen to 30% elongation.

Gel content was determined on a l-g. binder sample by a 48-hr. methylene chloride extraction which was followed by a 4-day acctone extraction if AP was present. Gel samples had a 1/16-in. web thickness. Linear swelling was determined on duplicate 1-in. long specimens in contact with benzene for 3 days. The benzene was replaced with fresh solvent on each of the first 2 days.

### 2. <u>Chromium Octoate</u>:

a. <u>Telagen gumstocks</u>: It was determined that nearly complete cure of Telagen will take place within a week with 0.40% CrOct present and within 2 weeks at the 0.37% catalyst level. Mechanical property data which form part of the basis for this conclusion are shown in Figures 37 and 38. The presence of AP filler seems to accelerate the binder cure at the higher catalyst level, while aluminum and Fe<sub>2</sub>O<sub>3</sub> (not shown) do not affect the cure. At the 0.2 or 0.3% CrOct levels, cure was not complete after 2 weeks. In both of these studies, and in others performed with the Telagen prepolymer, mechanical preperties gel content, or swell, all change slightly after 2 or 3 weeks, indicating that cure is still progressing slowly.

An experiment was performed to measure the viscosity of Telagen binder with 0.40% chromium octoate as a function of time, to determine if this system had a reasonable pot life at 26°C. The results, given in Figure 39, show that the viscosity of the binder triples in about 4 hr. This is an acceptable increase. To meet the goals of the program, the propellant must have at least a 4-hr. pot life ( $\leq$  35 K<sub>p</sub>) and a reasonable ambient temperature cure time of 2 weeks or less.

Binder gumstock studies were performed with Telagen and the aziridine HX-868 as a preliminary to propellant manufacturing. In the first study it was learned that the cure reaction was slower than the




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Telagen-epoxide reaction at equal CrOct levels. It was also found that the presence of solids had no effect on the rate at which binders cured and that using equivalent amounts of aziridine and carboxyl (A/C - 1.0)would give a binder which had too high a gel content for use in propellant. One possible explanation for the slow cure is the fact that HX-868 is an acyl aziridine and is subject to rearrangement to form oxazolines. The oxazoline will react with carboxyls but at a slower rate than do the aziridines.

A second set of binders was made in which the CrOct level was varied from 0.4 to 1.0% and the A/C ratio was lowered to 0.8. Figure 40 shows the cure curves for three of the catalyst levels. Cure is not quite complete in 3 weeks even at the 1% CrOct level. Postcure of CTPB propellants cured with HX-868 is well known, for example, the Aerojet Minuteman propellant ANB-3066 (Ref. 7). The differences in strengths between the three binders shown is attributed to a loss of curatives to side reactions at the lower catalyst levels. In order that the cured propellant have as high an amount of the desired crosslinks as possible the catalyst level to be used should be the maximum concentration which gives an acceptable pot life.

A binder viscosity study was carried out which indicated that 1% Cruct should be close to the maximum allowable level due to pot life considerations.

The first Telagen-aziridine propellant which was prepared had an A/C of 0.70 since the second binder gumstock study also gave binders whose gel contents were too high at the high catalyst level.

b. <u>HC-434 gumstocks</u>: Figure 41 is a plot of percent gel content vs. cure time for various HC-434 binder compositions at the two CrOct levels, 0.4 and 0.2%. At the 0.4% catalyst level, the maximum gel content was generally reached within 7 days, while at the lower level cure was not complete at 7 days. At an intermediate cure satalyst level of 0.3% which is not shown on the figure, gel content was still increasing in the 7- to 10-day period. The presence of the various propellant fillers (AP, oxamide,  $Fe_2O_3$ , and A1) when 0.4% CrOct was present did not materially alter the gel content achieved by the binder.

An HC-434-aziridine gumstock study was performed prior to propellant manufacturing. Since it was determined in the Telgen case that more CrOct cure catalyst was required to cure the CTPB with aziridine (HX-868) than with epoxides, CrOct levels from 0.9 to 1.5% were tested here. The study was conducted for a period of 93 days and the axiridineto-carboxyl ratio was also varied in order to determine its effect on cure time.





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Figure 42 presents cure data in terms of secant modulus at 30% elongation vs. ambient cure time. It can be seen that the higher A/C, as well as higher CrOct levels, give the binder a higher modulus and also that the modulus of all four systems was increasing even beyond the 46-day point. Figure 43, which is a plot of percent of final secant modulus vs. cure time, more clearly shows the effect of catalyst level at the two stoichiometries. From Figure 43 it can be seen that cure was faster at the lower stoichiometry, i.e., when carboxyl was in excess, even though the catalyst level was also lower. Therefore, for the fastest possible ambient cure the CrOct level should be the highest value permitted by pot life considerations and the aziridine-to-carboxyl ratio should be the lowest value permitted by the ultimate mechanical properties desired of the propellant.

The cause for slower cure at higher aziridine concentration may be attributed to the aziridine rearrangement reaction provided that the rate of rearrangement is more dependent upon aziridine concentration than is the aziridine-carboxyl reaction. If this is true, more aziridine will rearrange to the oxazoline structure during cure at the higher A/C. Since oxazoline reacts with carboxyl at a lower rate than does the aziridine, a buildup of oxazoline in the propellant will result in very long cure time.

c. PBAN gumstocks: PBAN is a terpolymer of butadiene, acrylic acid and acrylonitrile manufactured by American Synthetic Rubber Corporation. This liquid rubber contains 4.8% of the emulsifying agent Ammonyx T, an ammonium salt. An initial PRAN binder gumstock study which contained just the triepoxide and 0.3-0.4% CrOct showed that unfilled binders were cured in 2 weeks at ambient with the higher CrOct level. However, when AP, Fe<sub>2</sub>O<sub>3</sub> and Cab-O-Sil, all at the 1% level, were present the binders were not cured enough for mechanical property testing until 20 days of ambient cure. Since these solids were not found to interfere with the cure of the other prepolymer gumstocks this behavior was tentatively attributed to the presence of the Ammonyx T. Amines do react with AP to liberate ammonia which could interfere with cure but ammonia was not detected when AP was added to a sample of Ammonyx T. Later in the program it was determined that the presence of NH3 does not affect CrOct-catalyzed ambient propellant cure. No satisfactory explanation for the interference of solids in PBAN cure was obtained. Model compound studies were performed at MRI using PBAN or Telagen and neat Ammonyz T, and evidence of a side reaction in the PBAN system was noted. The exact nature of the cure inhibition reaction is not defined.

A second binder gumstock was performed using both di- and triepoxide, filler and higher cure catalyst levels. The epoxide-to-carboxyl ratios were unity. The all-diepoxide bieders containing up to 2.0% CrOct





failed to cure but those with all-triepoxide cured at a rate only slightly slower than the unfilled binders (of 0.4% catalyst) when 1-2% CrOct was present. A binder viscosity study was performed which indicated that 1.5% CrOct was the maximum concentration of catalyst which would give good pot life. Therefore, the cure inhibition observed in the PBAN-epoxide system relative to the two CTPB prepolymers can be overcome simply by increasing the catalyst concentration.

Two binder gumstock studies were carried out on the PBAN-aziridine cure system in order to determine the best catalyst level and A/C ratio for ambient cure and good pot life. Propellant solids (1%) were included in all the mixes since it was found that the presence of these solids in the PBAN-epoxide system required a different CrOct cure catalyst concentration than did the unfilled binders. In the first study, the CrOct level was varied from 0.4 to 2.0% of the binder and both 0.8 and 1.0 A/C were used. None of these binders cured well enough in 1 week to obtain mechanical property specimens. In the second study 2%, 3%, 4% and 5% CrOct were used with an A/C of 1.0. The binder with 5% CrOct had a viscosity which indicated that a propellant could be manufactured with high concentration. All gumstocks were very slow to cure and no mechanical property data were obtained. Qualitatively, a cure time of several months at ambient was indicated.

d. <u>EA/AA gumstocks</u>: A fourth prepolymer which was to be studied in this program was the copolymer of ethylactylate and acrylic acid (EA/AA) synthesized by Celanese Corporation. The prepolymer, which had an acid content of 5.2%, was not very soluble in the plasticizers DOA or IDP but was miscible with either tri-n-butylphosphate (TBP) or triacetin (TA). The TA-prepolymer solution slowly cured with the triepoxide at ambient without any added catalyst. The EA/AA-TBP solution did not have this characteristic but both plasticizers were used in the subsequent binder gumstock cure studies.

In gumstock studies, plasticizer-to-polymer ratios of 1 to 5 were used with the epoxides ERLA-0510, DER-332, ERLA-4221 or UNOX-221, E/C ratios of 0.8 to 3.6, CrOct levels of 0.2-0.4 and AP filler to stimulate homopolymerization of the epoxide when excess epoxide was present. Cured gumstocks had very little strain at maximum stress and problems were anticipated when EA/AA was to be used in the 85.5% solids propellant of this program. A plasticizer-to-polymer ratio of 3 was necessary in order to obtain binders with TBP which had about the same initial viscosity as the CTPB binders at 25°C.

The best gumstock cures were obtained with TBP/EA/AA - DER-332, E/C = 3.6 and 0.4% CrOct. Two l-lb. propellant mixes were made with DER. In both cases the pot life was very short and end-of-mix viscosity exceeded 160 Kp. After 5 weeks of ambient cure, the propellant was still too soft to cut mechanical property specimens.

It was concluded that propellants based on the EA/AA copolymer were not processable at the normal binder and plasticizer levels used in 85 to 88% solids propellants. The intrinsic viscosity of EA/AA is such that it must be highly plasticized in order for it to be the basis of a processable propellant; yet, enough prepolymer must be present in the propellant so that a sufficient crosslink density is achieved and the grain will be consolidated after curing. The limit of processability of EA/AA propellants is about 20% binder when elevated mix temperature and high aluminum levels are used. Normally, 30% or more binder is used in highenergy EA/AA propellants. The high energy is attained by the use of an energetic plasticizer such as TVOPA. Since the use of explosive plasticizers is beyond the scope of this program, the remaining EA/AA studies were deleted from the program. The funds saved by this deletion were directed to an expanded inveatigation, through the 5-gal. mix of the other three prepolymers of this program.

3. <u>Chromium oleate</u>: A gumstock study of Telagen prepolymer cured with chromium oleate was performed to see if the oleate might offer any advantages over the octoate. The catalyst materials used were prepared by Shepherd Chemical Company and contained 4.7 and 6.0% Cr (theoretical is 6.16, while the MRI tested material was 4.0% Cr). Results, which are given in Figure 44, show that even at the high 0.80% catalyst level, cure is not complete in 15 days with the purer chromium oleate sample. This high catalyst level corresponds to a Cr content equivalent to 0.44% CrOct. Since the Cr oleate appeared to be inferior to CrOct no further work was performed with the oleate.

4. <u>Chrowium 2,6-diisopropyl salicylate</u>: Chromium 2,6-diisopropyl salicylate (CrDIPS) was a cure catalyst recommended and supplied by the Explosives Research and Development Establishment, Waltham Abbey, England. The matrix of Telagen binders was prepared and is shown on the following page. A plot of modulus versus ambient cure time for these binders is shown in Figure 45. The epoxide binders cured with CrDIPS were not strong enough to prepare mechanical property specimens, even when a higher Cr level than the CrOct baseline formulation (0.5% of the binder) was used. In the aziridine case, the CrDIPS sample was weak but tractable. From the figure, it can be seen that both catalysts cure the aziridine binder in 14 days at ambient, but the higher modulus of the CrOct sample indicates a higher degree of crosslinking. Since the low modulus indicates that undesirable side reactions took place (perhaps homopolymerization) CrDIPS would not be recommended as a replacement for CrOct in ambient-cured propellants on the basis of this study.



Figure 44 - Secant Modulus at 30% Elongation vs. Cure Time X196-79

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A 1-1b. propellant mix was prepared with CrDIPS which confirmed the gumstock results.

# AMBIENT CURE OF TELAGEN BINDERS

Curative	<u>% Catalyst</u>	<u>% Cr</u>	<u>Results (see Figure 45)</u>
Epoxide	0.5 CrOct	0.05%	Baseline formulation, full cure ~ 15 days
Epoxide	0.83 CrDIPS	0.054	30 day cure too soft to test
Epoxide	0.664 CrDIPS	0.045	30 day cure too soft to test
Epoxide	0.996 CrDIPS	0.065	30 day cure too soft to test
Aziridine	1.0 CrOct	0.108	Baseline formulation, full cure $\sim$ 15 days
Aziridine	1.67 CrDIPS	0.108	Full cure ~ 15 days

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# SCALE-UP DEMONSTRATION OF CURE CATALYSIS

# A. Summary

The CrOct catalyst concentration and curative stoichiometry were optimized for the six propellant binder systems. Calling the levels and cure times for the six baseline formulations are shown below:

#### Baseline Propellant Compositions

System	% CrOct in Binder	Cure time, days
Telagen-epoxide	0.50	14
HC-434-epoxide	0.45	14
PBAN-epoxide	1.75	28
Telagen-aziridine	1.00	21
HC-434-aziridine	1.20	21
PBAN-aziridine	5.00	> 150

CrDIPS was shown to be inferior to CrOct as a cure catalyst for these ambient-cured carboxyl propellants.

Water contamination was found to lower propellant tensile strength but did not affect ambient cure time. Water solutions of the contaminants NH<sub>3</sub>, HClO<sub>4</sub>, HNO<sub>3</sub> and HF were also tested in the Telagen-epoxide baseline propellant. Cure times were unchanged with the exception of the HF case in which the cure rate appeared to be increased.

Oxamide shortens the pot life of the Telagen-epoxide baseline propellant. The antioxidant, thiodiphenylamine, consumes curative in these amhient cured propellants to a higher degree than in 140°F-cured propellants. Cruct does act as a surfactant for these propellants.

A surveillance program conducted for 6 months at 75° and 160°F on propellants cured at both ambient temperature and 140°F showed that the ambient-cured propellants agad as well as the heat-cured propellants.

The Telagen-epoxide propellant had the best aging characteristics of the four binder systems examined. Although it was shown that a higher degree of desired ester crosslinks formed during ambient cure than during 140°F cure, there was no evidence that this resulted in improved aging characteristics.

# B. Discussion of Results

1. <u>Procedure</u>: Initially, 1-1b. propellant mixes were manufactured with cure catalyst concentrations selected on the basis of the binder gumstock studies. If necessary, further 1-1b. mixes were then made in order to optimize the catalyst level to achieve the best balance between short ambient cure time and adequate ( $\geq 4$  hr.) slurry pot life. Curative stoichiometry would also be varied, if necessary, to obtain good mechanical properties. The optimized propellant was then made on a 10-1b. scale to better define the actual cure characteristics and determine if there were any changes brought about due to scale-up. Having completed this work, a baseline composition was then chosen for each of the six binder systems. The effects of contaminants on cure and the aging characteristics of several of the baseline propellants were then determined.

All propellants had the same basic composition consisting of 14.5 wt. % (27.5 vol. %) binder, 80% AP, 5% aluminum and 0.5% Fe<sub>2</sub>0<sub>3</sub>. The binders were 70% polymer with catalyst and plasticizer taking up the remaining 30% of the binder phase. The 1-lb. and 10-lb. propellant mixes were manufactured in a l-quart and l-gal. Baker-Perkins double verticle planetary mixer, respectively. The 40-lb. mixes were prepared in a 5-gal. Day mixer. Mixer temperature was 25°C in all cases. The solids were added incrementally and the catalyst was added as a THF solution 40 min. before the end of the mix cycle. Viscosities were measured on a Haake rotoviscometer and the values reported here are at a shear rate of 0.41 sec.<sup>-1</sup>. Cure time was followed by either gel content analysis or mechanical property measurements. Propellant mechanical properties were measured on Type II JANAF tensile specimens using an Instron tester and a strain rate of 0.74 in/in/min. Since CrOct acts as a surfactant for these propellants, propellant processing was aided. Because of its superior cure catalyzing efficiency, CrOctwas choosen as the catalyst for all the baseline propellants.

#### 2. Selection of baseline propellants:

a. <u>Telagen-epoxide propellant</u>: The first propellants processed and cured at ambient temperature  $(75^{\circ} \pm 5^{\circ}F)$  in this program contained the Telagen-epoxide binder, cured with CrOct. All propellants were mixed, tast and post-evacuated without difficulty at ambient temperature. A listing of the Telagen-epoxide 1-lb. mixes is given in Table XXXVII. Initially, 0.37% CrOct in the binder was tested buy later it was determined that as much as 0.5% satalyst could be used and still maintain a good pot life.

It was predicted by the gumstock study that cure of Mix No. X196-87 with 0.40% CrOct should be complete within 8 days. The modulus values for the propellant, which are shown in Figure 46, confirm this. The strain at maximum stress values and tensile data were less precise but indicated that cure was still progressing at a measurable rate beyond the 8-day period.

In subsequent propellant mixes it was determined that a mixture of di- and triepoxides should be used in order to improve the cured mechanical properties and that 0.5% CrOct in the binder was necessary to insure a 2-week ambient temperature cure. The X237-25 formulation was selected as the Telagen-epoxide baseline. A 10-lb. mix (X237-26) of the baseline was prepared which defined and confirmed the 2-week cure time (see Figure 47).

The purpose for using the 10-1b. grains in addition to the 1-1b. grains was to provide many more mechanical property specimens (1/4-in. thick by 5-in. long JANAF dogbones) for testing so that the cure cure-time curve could be unambiguously defined. The baseline X237-26 had a 2-week cure time, since the mechanical properties did not change beyond 2 weeks at ambient or after heating the propellant for 3 days at 60°C. The propellant mold was disassembled after 6 days of cure and the grain was quite strong when properties were first measured. The average mechanical properties of five specimens after 9 days of cure was 320 psi modulus, 102 psi tensile strength and 48% strain at maximum stress. Processing pot life was adequate with the 0.5% CrOct cure catalyst present in the binder phase. Therefore, no unusual problems are anticipated if motors were to be made of this propellant. Mechanical property vs. time results show that cores can be pulled in 1-2 weeks after casting.

Propellant X237-20 containing 0.4% CrOct (in the binder) was also prepared on a 10-lb. scale. The grain, which is pictured in Figure 48, was cured well enough after 8 days to be removed from the mold. Mechanical properties (see Figure 49) did not change after 20 days of cure at ambient, but cure was not complete in 20 days because exposing the propellant to 3 days cure of 60°C after 32 days at ambient gives a significantly stronger propellant grain.

### TABLE XXXVII

# TELAGEN-EPOXIDE PROPELLANTS#/

				Slurry	urry 77°F, 0.74 m			rties at <sup>a</sup> / hin <sup>-1</sup>		
<u>Mix No.</u>	% CrOct	Ероку С0 <sub>2</sub> н	di/tri <u>Epoxy</u>	Visc. Kp Time (Hr)	Cure Time <u>(Days)</u>	Modulus (psi)	Strength (psi)	Strain at Max. Stress (%)		
X196-70	0.37	1.00	tri	-	-	1,730	133	9.2		
X196-72	0.40	1.00	tri	11 at 5	-	1,800	133	9.4		
X196-74	0.40	0.80	tri	15 at 1 15.6 at 1.5	≥ 21	670	118	24.1		
X196-87	0.40	0.75	tri	12 at 1 19 at 3	8	510	112	29.6		
X222-2	0.37	0.86	0.50	20 at 1 26 at 4	14	860	116	19		
X222-12	0.43	0.75	tri	-	14	630	138	31		
X222-16	0.40	0.75	0. <b>50</b>	10 at 1 11 at 2	19	490	111	31		
X222-55	0.40	0.70	0.50	12 at 1 13 at 2	≤33	430	87	25		
X237-15	0.40	0.75	0.75	16 at 1 18 at 3	≤ 23	480	120	37		
x237-25 <u>b</u> /	0.50	0.75	0.82	11 at 1 12 at 4.5	14	420	110	37		
x237-26 <u>b</u> /	0.50	0.75	0.82	17 at 1 23 at 2.5	17	390	116	44		
x237-71 <sup>b/</sup>	0.50	0.75	0.82	9 at 1 12 at 4	14	390	113	43		

a/ Propellant composition is 32% 200 μ AP, 24% 130 μ AP, 24% 11 μ AP, 5% H-10 A1, 0.5% Fe<sub>2</sub>O<sub>3</sub>, and 14.5% binder. The binder is 70% polymer and 30% IDP plasticizer and cure catalyst.

b/ Baselina formulation on 1-1b. (X237-25), 10-1b. (X237-26) and 40-1b. (X237-71) scale.

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Figure 48 - Ten-Pound Telagen Grain Cured 8 Days at 75°F



The baseline propellant X237-26 (Figure 47) may have increased in strangth slightly after heating for 3 days at 60°C. However, if this was the case, a small degree of post-cure is no problem since the cure was complete enough after 1 week so that a propellant motor could be disassembled and shipped. The minor degree of cure which would take place in the propellant subsequent to this could easily be predicted and would not alter the reliability of the motor. The surveillance of the baseline propellants, discussed in the next section, has as its objective, the determination of post-cure aging characteristics. It was determined that the baseline ages in a similar fashion to 140°F-cured Telagen-epoxide propellant.

b. HC-434-Epoxide propellant: A propellant ambient cure study, similar to the Telagen study, was performed using the HC-434 prepolymer. A summary of the results obtained is tabulated in Table XXXVII. Early 1-1b. mixes indicated that CrOct was an efficient cure catalyst at the 0.40% level (based on binder) but a higher level might be more desirable. E/C of 0.75 coupled with a di/tri epoxide ratio of 0.50 gave good mechanical properties. Viscosity of HC-434 propellants is higher than Telagen propellants. The modulus vs. ambient cure time for propellant X237-21, a 10-lb. casting, is given in Figure 50. The results are quite similar to those obtained for the Telagen grain X237-20, Figure 49. Ambient cure appeared to be complete in 24 days but further cure took place when 31-day old propellant was subjected to 3 days of 60°C. The 10-1b. batch of X237-27, which contained 0.50% CrOct in the binder, was quite viscous and due, presumably to poor dispersion of the curatives, the propellant's homcgeniety was very poor. It appears that a 0.45% Cr0ct level, such as that used in the 1-lb. mix X237-10, would give a better balance between processability and minimum ambient cure time. Mechanical property data for propellant X237-10 are given in Figure 51.

A 10-1b. propellant mix (X237-49) was made with this intermediate concentration (0.45% of the binder) of CrOct cure catalyst. The cure was complete in 2 weeks at 75°F. Mechanical properties remained constant for the remainder of the 46-day period of the study (see Figure 52). However, when the ambient cured propellant was subjected to a week at 60°C, the modulus and strength dropped slightly. This composition was therefore selected as the HC-434-epoxide baseline formulation.

End-of-mix viscosities for the 10-1b. mix (X237-49) and later for the 40-1b. mix (X233-81, see below) were 15 and 16 Kp, respectively, compared to 21 Kp for the 1-1b. (X237-10) propellant mix. This lower mix viscosity for the larger mixes was found to he a general occurrence in this program. The 10-1b. mix was softer than the baseline made on either the 1- or 40-1b. scale; however, this was not a general occurrence and is considered to be due to a small formulation error rather than poor reproducibility of cure.

				End of Mix			Phys 7	ical Pro 7°F, 0.74	perties at 4 min <sup>-1</sup>	
Notebook No.	% CrOct	Epoxide Carboxyl	d1/tri Epoxide	Visc. at 26°C (Kp)	Visc. Kp/Time (Hr.)	Time Max. Mod. (Days at 26°C)	Modulus S (psi)	trength (psi)	Strain at Max. Stress (%)	Remarks
X196-93	0.40	0.80	tri	24	ı	14	920	119	17	
X222-15	0.40	0.75	0.50	16.5	24/5	12	470	131	43	
X222-33	0.40	0.75	0.50	29	26/3	> 22	270	69	41	0.5% TDP
X222-40	0.40	0.75	0.50	15	12/5	≥ 23	420	111	37	
X222-41	010	0.75	0.50	> 50	1	> 34	206	48	44	0.5% TDF
X227-61	0.40	0.75	0.50	15	1	< 19	450	120	39	
<b>X</b> 222 <b>-</b> 70	0.40	0.826	0.44	24	16/5	< 17	860	147	28	0.5% TDPA
<b>X</b> 222-95	0.45	0.75	0.50	26	30/4.5	< 15	640	154	36	
X237-10 <sup>a</sup> /	0.45	0.75	0.50	21	27/3.5	12	620	126	32	
X237-21	0.40	0.75	0.50	,	31/2.5	ı	460	104	33	
<b>X</b> 237-27	0.56	0.75	0.50	High	ı		7 00	150	35	
X237-49 <u>a</u> /	0.45	0.75	0.50	15	11/4	14	350	119	49	
<b>X</b> 23 7 -8 1 <sup>a</sup> /	0.45	0.75	0.50	16	15/3	14	690	146	41	
<u>a/</u> Baseli	ine fo	rmulation	on 1-1b.	(X237-10),	10-1b. (X	(237-49) and 40-1b	. (X237-81	) scale.		

TABLE XXXVIII

AMBIENT CURED HC-434 PROPELLANTS







c. <u>PBAN-epoxide propellant</u>: The terpolymer polybutadiene, acrylic acid and acrylonitrile (PBAN) was found in the Phase III gumstock studies to require considerably more CrOct cure catalyst to cure with epoxides than did the two CTPB prepolymers. This was true only when propellant solids were present; cure of unfilled binder was similar to that of the CTPB's. Presence of the emulsifying agent, Ammonyx T in the PBAN, was suspected of being the cause since this was the only known new constituent of the system. Ammonyx T, an amine salt, could interact with AP in the propellant environment to cause a reduction in curative concentration.

Several PBAN-epoxide propellants were prepared and are given in Table XXXIX. The first 1-1b. mix (X237-46) contained 1.50% CrOct in the binder and had an acceptable pot life (Figure 53) but long cure time (see Figure 54). When the CrOct concentration was increased to 1.75% of the binder (X237-79), the pot life was too short (viscosity 48 Kp in 4 hr.) but cure was complete in 12 days with properties of 500/81/24 (modulus/tensile strength/elongation). Since other propellants had lower mix viscosities when they were scaled-up, it was decided to use 1.75% CrOct in the binder for a 10-1b. PBAN-epoxide mix. This propellant (X237-94) cast well and had a viscosity of only 18 Kp 4 hr. after catalyst addition.

The 1-1b. mix cured faster than did the 10-1b. casting (see Figure 55). Although completion of cure for X237-94 was 4 weeks or longer, this composition was selected as the baseline for surveillance.

d. <u>Telagen-aziridine propellant</u>: The Telagen-aziridine binder gumstock study found that 1% CrOct cure catalyst was probably the maximum permissible level due to pot life considerations and that an A/C of 0.70 should give good mechanical properties.

Propellant X237-52 was prepared with 1.0% CrOct (in the binder) and the viscosity curve is given in Figure 53. The viscosity of 31 Kp at 4 hr. was within the program limit of 35 Kp. The curve time was not well defined by this 1-lb. mix but the catalyst level and A/C ratio appeared optimum. Therefore, 10-lb. mix X237-78 was prepared with the same composition and cure data as shown in Figure 56. It appeared that the propellant was fully cured in 4 weeks or less at ambient. Mechanical properties remained constant through 52 days at ambient but propellant which was then subjected to 7 days at 60°C did cure further. Properties at 52 days at ambient were 510/104/31 and after 59 days at ambient plus 7 days at 60°C were 650/128/28. This indicates that wither the cure reaction was not complete during the ambient phase of cure or side reactions are causing an increase in gel content. A comprehensive study of aging properties of this propellant were performed in the surveillance program. TABLE XXXIX

# PBAN-EPOXIDE PROPELLANTS

				End of Mix			Ч	ysical Pro 77°F, 0.7	perties at 4 min <sup>-1</sup>
		Enoxíde	di/tri	Visc. at 26°C	Visc. Kn/Time	Time to Max. Mod.	Modulus	Streneth	Strain at Max. Stress
M'X No.	Z CrOct	Carboxyl	Epoxide	(Kp)	(Hr.)	(Days at 26°C)	(psi)	(psi)	(%)
X237-46	1.50	1.0	tri	16	28/5	34	430	89	30
X237-79	1.75	1.0	trí	21	48/4	12	500	88	24
X237-94	1.75	1.0	trí	12	18/4	28	500	ះ <b>6</b>	31
X248-21	1.75	1.0	tri	I	ı	28	400	78	31

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Propellant X248-9, the 40-1b. casting of the baseline composition, had mechanical properties of 635/111/26 after 4 weeks cure at  $75^{\circ}F$ .

e. <u>HC-434 - aziridine propellant</u>: Since the Telagen-aziridine baseline propellant required twice the CrOct level as the epoxide-cured propellant, the first HC-434-aziridine propellant also used twice the epoxide cured baseline catalyst level, i.e., 0.9% CrOct in the binder (Table XL). Cure for this propellant, X237-64 (Figure 57), took about 4 weeks. Higher catalyst levels and lower A/C ratios were also tested since binder gumstock studies indicated these changes would bring about faster cure. A short pot life was obtained with 1.5 and 1.3% CrOct but the baseline propellant (X248-4) having 1.2% CrOct and an A/C of 0.75 had acceptable pot life. Cure for the baseline was nearly complete in 3 weeks although the propellant continued to harden slightly for 80 days, see Figure 58. No further work was scheduled or performed on HC-434azirdine propellants.

f. <u>PBAN - aziridine propellant</u>: Of the six binder systems examined in propellants, the PBAN-aziridine was the most difficult to process and cure at ambient temperature. The catalyst requirements were very high compared to the PBAN/epoxide or the CTPB systems.

	Relative Concentration of CrOct
Binder System	Needed for Ambient Cure of Propellant
CTPB/Epoxide	1
CTPB/Azir idine	2-1/2
PBAN/Epoxide	4
PBAN/Aziridine	10

Another factor in processing was the higher viscosity of the PBAN prepolymer compared to the CTPB's.

A PBAN-aziridine propellant with an A/C = 1.0 and 5.0% CrOct in the binder phase was very viscous and slow to cure. Mechanical properties after 4 months at  $75^{\circ} \pm 5^{\circ}F$  were 760/60/15. A mix was prepared (X248-82) im which the A/C was lowered to 0.9 which, from the HC-434-aziridine binder study mentioned above should give a faster cure. Cure data are shown in Figure 59. Mix viscosity was 55 Kp and the cure time was greater than 9 weeks. A 10-1b. mix of this composition was made which was processable, (again lower viscosity was observed on scale-up) but cure time was greater than 5 months. No further work on this binder system was performed. TABLE XL

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HC-434 AZIRIDINE PROPELLANTSA/

	Note	X237	X237	X237	X248	X248	
	ook	-60	-64	-84	-81	-4	
	CrOct	0 <b>6</b> - 0	06*0	1.50	1.30	1.20	
	<u>A/C</u>	0.6	0.8	0.8	0.7	0.75	
End of Mix	Visc. at 26°C (Kp)	26	27	34	30	30	
	Visc. Kp/Time (Hr.)	20/4	21/4	48/4	43/4	5/4	
	Modulus (psi)	ı	550	1,100	630	680	
Physical Prop at 77°F, 0.74	Strength (psi)	ı	134	156	134	128	
perties 4 min <sup>-1</sup>	Strain at Max. Stress (%)	ı	34	21	29	33	
	Cure Time (days)	> 35	35	> 21	28	21	

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 $\frac{\epsilon}{2}$  / Propellant composition is 32% 200  $\mu$  AP, 24\% 130  $\mu$  AP, 24% 10-12  $\mu$  AP, 5% H-10 A1, 0.5% Fe203 and 14 .5% The binder is 70% polymer and 30% IDP plasticizer and cure catalyst. binder.






### 3. Effects of Other Constituents on Propellant Cure

a. Other cure catalysts: A 1-lb. Telagen epoxide baseline formulation was made with CrDIPS (chromium 2,6-diisopropyl salicylate) replacing CrOct on a chromium for chromium basis. After 18 days cure, the properties were only 76/25/50 while the CrOct baseline cures fully in about 10 days to properties of 390/113/43. After a 7-week cure at 75°F the CrDIPS propellant properties were 87/32/53. This propellant confirmed the binder gumstock studies conclusion that CrDIPS was inferior to CrOct as a carboxyl-epoxide cure catalyst.

b. <u>Water contaminant</u>: Water is the most likely contaminant to be present in composite propellants and its influence on catalyzed ambient cure was therefore given the most consideration of any of the contaminants tested.

Water was first tested as an added contaminant in the Telagenepoxide baseline propellant. The quantity of water added was equal to the amount of CrOct present. The water was dissolved in THF and added to the propellant mix after all solids had been added. The mix viscosity was the same as for the baseline propellant 1-lb. mix. Cure was complete within 2 weeks at 75°F and final mechanical properties were 400/99/39. These properties comoare well with those of the baseline except for an 11 psi drop in tensile strength. See Table XLI.

This propellant was then made on a 10-lb. scale (X237-66). Cure time was confirmed to be 2 weeks or less (Figure 60) and again a drop in tensile strength was observed (400/96/43) compared to the baseline propellant.

The Telagen-aziridine baseline was made with 0.5% water added to the binder. Cure time was unchanged but the final tensi'e strength was lowered (see Table XLI). Similar results were obtained when water was added to the PBAN-epoxide propellant.

It is therefore concluded that water contamination does not change the cure time but does lower ambient-cured propellant tensile strength. The other contaminants (Section C) were added as water solutions and the effects of the combined contaminants were expected to be additive.

c. <u>NH3</u>, HC104, HN03 and HF contaminants: One-pound propellant mixes were prepared with the contaminants NH<sub>3</sub> and HC10<sub>4</sub> in the Telagenepoxide baseline formulation. The contaminants were added as water solutions. The contaminant concentration was 0.01% of the propellant while the water concentration was at the same level (0.07%) as was tested

TABLE XLI

## PROPELLANTS WITH CONTAMINANTS<sup>a</sup>/

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			Fine	al Properti	es
<u>Contaminant/% of Binder</u>	Baseline	Cure Time (Days)	E (psi)	σ (psi)	<mark>е</mark> п (%)
None	Telagen-Enoxide	14	420	011	37
		17	390	116	7 <del>7</del>
н <sub>2</sub> 0/0.5		14	400	66	39
		21	400	96	43
NH		·	320	101	42
,		21	370	83	40
HC104		21	310	78	33
		21	310	100	47
HNO3		t	250	74	39
1		28	260	80	43
51. 21		< 13	300	82	35
	<b>→</b>	8	350	66	40
None	Telagen-Aziridine	21	680	120	23
H20/0.5	Telagen-Aziridine	21	610	88	23
None	PBAN-Epoxide	28	500	95	31
H20/0.5	PBAN-Epoxide	≥ 28	500	80	24
$\underline{a}$ The first listing is the	e l-lb. míx whíle the sec	cond set of values	is from the 10	)-lb. casti	.ngs.

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previsouly in the baseline propellant. Mix viscosities were normal, being 13 Kp at the end of mix. By periodically measuring mechanical properties of these two propellants, it was determined that cure time ( $\approx 2$  weeks) at 75 ± 5°F was unaffected by the contaminants but that final properties, particularly in the case of HClO<sub>4</sub>, may have been affected. Final properties of the 1-1b. control propellant were (E, psi/ $\sigma_m$ , psi/ $\epsilon_m$ , %) 400/99/39 while those of the NH<sub>3</sub> and HClO<sub>4</sub> propellants were 320/101/42 and 310/78/33, respectively. The lower tensile strength of the HClO<sub>4</sub>-containing propellant is particularly at variance with the control propellant.

Propellants were made in the same fashion with  $HNO_3$  and HF contaminants. Mechanical properties after 24 days of ambient cure were 250/74/39 and 300/82/35, respectively. All four propellants had lower moduli than the control which probably indicate some loss of epoxide curative, perhaps due to homopolymerization catalyzed by the presence of acid.

Cure studies were then performed on a 10-1b. scale with each of the four contaminants in the Telagen-epoxide baseline formulation. Cure time and final properties are given in Table XLI. The only feature which stands out is the short cure time in the HF-contaminated propellant. This was in contrast to the model system experiments performed at MRI which indicated that HT only consumed epoxide.

d. <u>Oxamide</u>: One of the ingredients to be tested for its effect on ambient-cured propellants was oxamide. A propellant (X237-69) was prepared of the Telagen-epoxide baseline composition in which 4% of the fine AP was replaced with oxamide. The propellant cured normally; however, the mix unexpectedly had a short pot life. Two additional 1-1b. propellant mixes were made, which used a new lot (ABL No. 4) of oxamide. The pot life results are shown in Table XLII. Evidently oxamide does shorten the pot life of these ambient cured propellants with the "lab sample" being particularly active.

e. <u>Thiodiphenylamine</u>: Another additive to be tested was the antioxidant thiodiphenylamine (TDPA). This was added to Telagen propellant of the type X222-16 at a 1% level in the binder. The TDPA propellant (X222-34, Table XXXVII) cured more slowly and was weaker than the control as can be seen in Figure 61. Since this behavior could be due to a loss in epoxide, because of a reaction between TDPA and the epoxides, another composition (X222-85) was prepared in which the amount of triepoxide (ERLA-0510) was increased by a quantity equivalent to the TDPA added. This propellant cured at a good rate (see Figure 61) and was atronger. From this result it was concluded that, in the propellant environment, TDPA probably reacts with the epoxides but that the equivalent weight

### TABLE XLII

### EFFECT OF OXAMIDE ON SLURRY VISCOSITY OF TELAGEN-EPOXIDE PROPELLANT

	Oxamide		Viscosity <sup>a</sup> /
Mix Number	Lot Number	(%)	<u>(Kp)</u>
x-237-25	-	0	13
x-237-69	Lab Sample	4.0	42
<b>x -</b> 248 - 98	ABL No. 4	4.0	<b>3</b> 2
<b>x -</b> 248 - 44	ARL No. 4	4.0	28

a/ Measured at 77°F, 4 hr. after cure catalyst addition.

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of the TDPA is higher than the theoretical value. Loss of cure catalyst may also explain the observed behavior.

TDPA was again tested in the Telagen-epoxide baseline propellant. As in the previous Telagen case, 1% TDPA was added to the binder. This propellant (X237-54) cured on the normal time schedule but was softer than the baseline propellant. Confirming the previous finding that loss of curatives, when TDPA is present, is significant in ambient-cured propellants. This does not happen at 140°F cure temperature.

f. <u>CrOct as a slurry viscosity depressant</u>: It was reported above that CrOct was a surfactant in the CTPB system. This was verified by comparing the viscosities of Telagen-epoxide baseline formulations with and without CrOct at 75°F. The slurry made with 0.0725% CrOct had a viscosity of 16 Kp while the identical slurry made without CrOct had a viscosity of 50-80 Kp. A more exact determination of viscosity could not be made because, above 50 Kp, the slurry had a tendency to rotate with the bob instead of being sneared between cup and bob in the Rotovisco.

g. <u>Mix cycle variation</u>: In our normal mix cycle chromium octoate was added last. Since chromium octoate is a surfactant for these propellants, it was thought that processing might be improved if the cure catalyst could be added to the mix earlier. In Telagen-epoxide propellant X237-48 CrOct was present in the binder premix and the two epoxides were added last. The mix was, as expected, of lower viscosity when the epoxides were added last relative to the mix viscosity when CrOct was added last. After addition of the epoxides the viscosity visibly increased so that the end of mix viscosity was about normal for the 1-1b. baseline propellant. The propellant cured in the normal time period but the strength was quite low (220/66/42).

A repeat of propellant X237-48 was made in which CrOct was in the binder premix and a mixture of the two epoxides was added last. The repeat mix, X248-2, duplicated the properties of the original mix in every respect.

We have observed that ambient-cured propellants and unfilled binders of identical stoichiometries can vary in strength, when cured, if the quantity of cure catalyst is varied. Propellants and binders with lower catalyst levels have lower strength. A possible explanation for obtaining softer propellant when the CrOct was added to the premix was that some of the CrOct prereacted somehow and the effective catalyst concentration was lowered. The cause for this apparent loss in CrOct to the system is not understood. Under this reverse addition, the CrOct has an opportunity to more completely react with the carboxyls on the CTPB before the epoxide was introduced, but this was not expected to effectively remove catalyst from the system.

### 4. <u>Surveillance</u>:

a. <u>Results</u>: Under Phase IV, Large-Scale Demonstration, the effects of catalyzed ambient temperature cure on the storage life of propellant was to be measured and compared to the storage life of 140°Fcured propellant. Ten propellant compositions were selected for the surveillance phase. These consisted of four 75°F-cured baseline propellants, two 75°F-cured baseline propellants with water contaminant added and four 140°F-cured control propellants. These propellants are listed in Table XLIII along with their initial mechanical properties. A 40-lb. batch of each propellant was made, then cut into ten 2 in. x 4 in. x 6 in. blocks and placed in controlled storage. The surveillance schedule was:

			Time -	Months		
Temperature	<u>1</u>	2	<u>3</u>	<u>4</u>	<u>5</u>	<u>6</u>
75 ± 5°F			х			х
160 ± 5°F		х		х		х

Some 12-month surveillance measurements were also made. Mechanical properties were measured on propellant in the interior (~ 1/2-in. from any surface) of each block at the specified time intervals. By testing interior propellant, mechanical property changes due to air oxidation were minimized.

The gel content of the propellant at the surface of these aged blocks was determined and compared with the gel content of the interior propellant to separate the effects of oxidative crosslinking from bulk propellant reactions. Two temperatures were chosen for the surveillance: vis., the 75°F cure temperature and an accelerated aging temperature of 160°F.

Mechanical property and gel content results of the 6-months surveillance are shown in Tables XLIV and XLV. Twelve-month aging data are given in Table XLVI. Figures 62 and 63 show the change in modulus with time and temperature for the Telagen baseline and heated-cured control propellants.

For the Telagen-epoxide propellants, the ambient-cured grain aved in a fashion similar to the 140°F-cured grain and the ambient baseline with water added aged as well as the baseline. There was only a small drop in modulus and tensile strength during ambient aging in each and a larger loss due to 160°F aging. Gel content was nearly constant of each case.

TABLE XLIII

### PROPELLANTS FOR SURVEILLANCE<sup>a</sup>/

		Cross-	di/tri	Z CrOct	Cure	Time <sup>c/</sup>		Tensile	Strain at
Type	Batch <u>Number</u>	linker/ Carboxyl	Crosslinker Functionality	in Binder	(Da) at 75°F	<u>s)</u> at 140°F	Modulua (psi) <u>d</u> /	Strength (psi) <u>d</u> /	Max. Stres $(7) \underline{d}/$
Telagen-Epoxide Baseline	X237-71	0.75	0.82	0.50	14	1	390	113	43
Telagen-Epoxide with water <sup>b/</sup>	X248-15	0.75	0.82	0.50	14	ı	380	67	41
Telagen-Epoxide heated cure	X237-72	0.92	0.82	0.12	ı	7	480	120	38
Telagen-Asividine Baseline	X248-9	0.70	tri	1.00	21	ı	635	111	26
Telegen-Aziridine with water ${\rm }^{\rm b'}$	X248-22	0.70	tri	1.00	21	ı	500	101	33
Telagen-Aziridine heated cure	X 248 - 35	1.00	tri	0.40	ı	7	470	130	45
HC-434-Epoxide Baseline	X237-81	0.75	0.50	0.45	14	ı	690	146	41
HC-434-Epoxide heated cure	X248-23	0.87	0.50	0.12	ı	٢	1,060	140	23
PBAN-Epoxide Baseline	X248-21	1.00	tri	1.75	26	ł	400	78	31
PBAN-Epoxide heated cure	X248-37	1.00	tri	0.40	ı	7	695	96	22
<pre>a/ Propellant composition - 3 The binder is 70% polyment</pre>	27 200 μ A	P, 24% 130 IDP plasti	μ AP, 247 10-1 cizer and cure	.3 μ AP, catalyst	5% H-10A1	, 0.5% Fe <sub>2</sub>	03 and 14	.5% binder	,

Amount of water added is equal to amount of CrOct present.

Cure time at 75°F was determined by testing 10-lb. mixes.

Mechanical properties were measured and surveillance begun 7 days after 75°F cure time to assure complete cure. 

TABLE XLIV

# SURVETILIANCE OF 40 POUND BATCHESA/ TELAGEN PROPELLANTS

				Mechanical	Properties		
		Modu	lus (psi)/Tens	ile Strength (	psi)/Strain at	Maximum Stre	88 (%)
		7	7*F, 0.74 in/i	n/min at Vario	us Aging Time	and Temperatu	res
			Aging at	75°P	Ÿ	ting at 160'F	
	Propellant	<u>lnitial</u>	3 Months	6 Months	2 Months	4 Months	6 Months
Ι.	Ambient-Cured Telagen-Epoxide	420/112/41	430/115/41	400/108/42	330/91/43	300/79/44	219/59/42
2.	140°F-Cured Telagen-Epoxide	480/120/38	480/110/35	449/99/34	410/105/39	340/105/49	285/98/54
е,	Ambient-Cured Telagen-Epoxide with H <sub>2</sub> 0 added	380/97/41	365/103/44	353/108/45	290/82/42	236/71/45	223/63/44
4.	Ambient-Cured Telagen-Aziridine	635/111/26	813/140/25	763/144/26	590/137/32	534/115/31	490/103/29
5.	140°F-Cured Telagen-Aziridine	470/130/45	577/140/36	530/145/43	390/117/45	260/86/51	263/81/50
<b>.</b>	Ambient-Cured Telagen-Aziridine with H <sub>2</sub> 0 added	500/101/33	522/113/34	512/111/35	380/89/34	293/65/37	213/52/39
			HC -	434 Propellant	5		
7.	Ambient-Cured HC-434-Epoxide	690/146/41	760/143/36	461/117/48	650/122/36	440/119/48	332/103/58
80	140°F-Cured HC-434-Epoxide	1060/140/23	998/129/21	870/131/25	770/130/30	650/130/35	540/138/43
			81	AN Propellants	ml		
6	Ambient-Cured PBAN-Epoxide	400/78/31	613/73/17	689/81/17	820/93/17	872/95/16	898/101/11
10.	140°F-Cured PBAN-Epoxide	695/96/22	869/109/18	945/117/20	835/109/19.4	946/116/19	980/121/19

a/ Cut into 2 in. x 4 in. x 6 in. blocks w.d wrapped in aluminum foil.

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### TABLE XLV

### BINDER CEL CONTENT AND MODULUS OF SURVEILLANCE PROPELLANTS

								Propellar	IC MODULUS
					Gel Content	t of Binder		6 Months	6 Months
			6 M	onth	2 Months	4 Months	6 Months	Ambient	160°F
	Propellant		Amb:	ient	160°F	160°F	160°F	(nsi)	(psi)
1.	Ambient-Cured	Interiors/	37.2	(38.3)	-	40.2	34.4	400	219
	Telagen-Epoxide	Surface <sup>b/</sup>	36.9	(38.1)	-	43.7	44.6	-	-
	5 1-			•		40.11			
2.	140°F Cured	Interior	50.3	(51.1)	-	50.4	47.5	449	285
	Telsgen-Epoxide	Surface	50.3	(51.0)	-	53.2	51.2	*	-
				• •		50.0			
3.	Ambient-Cured	Interior	-	(39.5)	42.8	54 . 3	41.0	353	223
	Telsgen-Epoxide	Surface	-	(41.7)	43.0	41.0	49.9	-	-
	with H <sub>2</sub> O sdded								
	2								
4.	Ambient-Cured	Interior	45.3	(45.2)	44.1	39.2	41.8	763	490
	Telagen-Aziridin	e Surface	44.5	(45.7)	45.0	44.2	49.3	•	-
	-								
5.	140°F-Cured	Interior		(45.7		38.8	41.2	530	263
	Telagen-Aziridin	e Surface		(43.4)		48.9	47.6	-	-
	U			•					
6.	Ambient-Cured	Interior	-	(41.3)	38.0	31.7	24.1	512	213
	Telagen-Aziridin	e Surface	-	(39.2)	46.1	49.5	47.6	-	-
	with H <sub>2</sub> 0 added			•					
	٤								
7.	Ambient-Cured	Interior	38.7	(41.0)	-	42.2	38.1	461	332
	HC-434-Epoxide	Surface	39.2	(41.5)	-	33.?	30.6	-	-
8.	140°F-Cured	Interior	67.6		-	57.0	59.9	870	540
	HC-434-Epoxide	Surface	67.2		-	54.9	53.7	-	-
9.	Ambient-Cured	Interior	•	(35.9)	40.6	39.4	38.4	689	898
	PBAN-Epoxide	Surface	-	(36.5)	38.4	37.5	37.2	-	-
10.	140°F-Cured	lnterior	+	(34.9)	-	34.9	35.4	945	980
	PBAN-Epoxide	Surface		(34.6)	-	34.8	34.3	-	-

 $\underline{a}$ / Gel content of propellant (expressed as percent of binder) from the interior of the block.

 $\vec{b}$ / Gel content of propellant from the surface of the block Note: ( ) "3" month at ambient samples that were tested for gel content after 6 months at ambient.

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### TABLE XLVI

### TWELVE-MONTH AGING PROPELLANT PROPERTIES

	Mechan 0.7	ical Properties 4 in/in/min_	at 77°F,
Propellant	Modulus (psi)	Strength (psi)	Strain at Max. Stress (%)
Ambient-Cured Telagen-Epoxide			
Aged 75°F	430	112	39
Aged 160°F	260	63	38
140°F-Cured Telagen-Epoxide			
Aged 75°F	470	112	37
Aged 160°F	290	100	51
Ambient-Cured HC-434-Epoxide			
Aged 75°F	430	112	45
Aged 160°F	330	92	49





Ambient cure with high GrOct level did not materially affect the aging characteristics of Telagen-epoxide propellant.

The three Telagen-aziridine propellants had small increases in modulus and tensile strength on ambient aging which abows that post-cure of the ambient cured propellants was the same as for the  $140^{\circ}$ F-cured propellant. When aged at  $160^{\circ}$ F the propellants eventually lost strength. In this case, the heated control and water-contaminated baseline had high losses in strength while the baseline did not. The two HC-434-epoxide propellants lost modulus and tensile strength on aging, hoth at 75° and  $160^{\circ}$ F. The modulus losses were similar.

The two PBAN-epoxide propellants post-cured both at ambient and 160°F to about equal degrees. Therefore, ambient-cured propellants age as well as the 140°F-cured control propellants. There was no evidence that the higher degree of deaired cure reaction, described below, improved the aging characteristics of ambient-cured propellants. It had been speculated that a higher degree of stable ester crosslinks in the binder of the ambient-cured propellants would provide better aging stability relative to the 140°F-cured propellants.

It was expected that, with aging, the exterior of a propellant block would increase in gel content relative to the interior due to airinduced oxidative crosslinking of the surface. All of the 6-month-at- $160^{\circ}F$  propellants, except the HC-434, had higher gel content at the surface than in the interior of the blocks. Both the ambient and  $140^{\circ}F$ -cured HC-434 propellants exhibited anomalously low gel contents at the surface after 6 months at  $160^{\circ}F$ .

One general conclusion that can be drawn is that there appears to be a general oxidative crosslinking effect on the surface of  $160^{\circ}$ Faged specimens made with Telagen.

It should be noted that this measure of gel in the system is only a gross measure of the crosslinks, not to be confused with the effective crosslink density defined by the Flory - Rohner (Ref. 9) or Mooney (Ref. 10) - Rivlin (Ref. 4) equations.

b. The ambient cure reaction: It was discovered that catalyzed ambient cure of these propellants required less crosslinking agent than did 140°F cure in order to obtain equivalent mechanical properties. This demonstrates that cure at ambient means that a higher degree of the desired crosslinking reaction is occurring than in 140°F cure, where loss of curative takes place due to side reactions.

This was noticed when the  $140^{\circ}$ F-cured control propellants for the surveillance program were being made. A Telagen-epoxide 1-1b. mix was made of the baseline formulation in which the only change was to reduce the CrOct concentration from 0.50 to 0.12% of the binder. Good pot life was obtained and cure was complete in 7 days but mechanical properties (modulus 130 psi, tensile strength 56 psi, strain at maximum stress 62%) indicated a very low gel content. Some loss in strength was expected, compared to the ambient-cured propellants, since undesired side reactions take place to a higher degree at  $140^{\circ}$ F. However, the magnitude of this loss in strength was not expected. It was necessary to increase the E/C from 0.75 to 0.92 to give a  $140^{\circ}$ F-cured propellant with good mechanical properties. Similar occurrences took place in the Telagen-aziridine and HC-434 epoxide cases (see Table XLIII).

However, there is still a loss of curative to side reactions taking place during ambient cure. This is evidenced by the fact that propellants with high CrOct levels have not only a shorter cure time but also yield stronger propellant; for example, propellants X196-87 and X222-12 in Table XXXVI, X222-61 and X222-95 or X237-21 and 27 in Table XXXVII. This indicates that when cure is slower and there is more time for side reactions to occur, that they will occur even at ambient.

### EXPERIMENTAL

XI.

A. Model System Studies

1. <u>Experimental considerations</u>: The analytical requirements for the kinetic study system were as follows:

a. The rate of the reaction must be followed quantitatively and accurately.

b. The stc.chiometry of the reaction must be followed, that is, the rates of disappearance of reactants should be followed simultaneously.

c. The rate of product formation should be readily determinable. This applies to the expected product plus any side products formed.

d. A means of indicating the products, especially side products should be available.

e. The entire technique should be as rapid and inexpensive as possible.

f. It would be desirable if all these requirements could be met in as few manipulations as possible.

The use of gas liquid chromatography comes close to meeting all the requirements enumerated above. Aliquots are withdrawn from the reaction vessel periodically and analyzed by GLC. The chromatograph indicates the total number of components present in the sample at that time. From retention time these components can be identified and relative concentration of components calculated.

2. Experimental procedures:

a. The internal standardization method was used to calibrate the gas chromatography method for quantitative analysis of the reaction mixtures. Known weight or mole ratios of the sample and a standard were prepared and chromatographed. The peak areas are measured. Area ratios are plotted against weight or mole ratios to obtain a calibration curve. An accurately known amount of the internal standard is then added to the unknown sample and the mixture chromatographed. Area ratios are measured and from the calibration graph the weight ratio of the unknown to the standard is known.

The advantages of this calibration method are that the quantitites injected need not be measured accurately and the detector response need not be known or remain constant since any change in response will not alter the area ratio.

An internal standard must:

- 1. Be well resolved from the other peaks;
- 2. Be eluted close to the peaks of interest;
- 3. Be approximately the same concentration as the unknown; and
- 4. Have a structural similarity to the unknown.

Peak height measurement is more rapid than peak area; however, plots of peak height vs. sample size have a more limited linear range than corresponding plots for peak area. Peak height and width are frequently dependent on sample size and sample feed volume while peak area is not.

After optimization of gas chromatographic conditions to give good separations (i.e., convenient retention and sharp peaks reproducibly), calibration curves were prepared using toluene as solvent and internal standard for AGE, epoxy butane, EHA, and octanoic acid and for the AGE-EHA combination. After establishment of the calibration plot its reproducibility was demonstrated on several test solutions. Variation in sample size between 1 and 2 µl. was shown not to affect the reproducibility of the peak height integration method.

b. Summary of GC conditions:

Column packing	20% SF-96 on Chromosorb P (60/80 mcsh), silanized
Column dimensions	6 ft. x $1/4$ in. aluminum
T <b>em</b> perature	100° to 275°C
lleating rate	10°C/min

Gas flow rate	80 cc/min (helium
Block temperature	275°C
Bridge current	175 ma.
Injection temperature	275°C

c. <u>Catalyst screening runs</u>: Stock solutions containing equimolar quantities of reactants are prepared in advance of a given series. A typical composition is given below:

Component	Moles	<u>Weight</u>	Weight (%)
Allyl glycidyl ether	0.204	23.256	22.342
2-Etnyl hexanoic acid	0.203	<b>29.23</b> 2	28.083
Toluene		51.600	49.573

Five milliliters of this solution are added to a vial, 1% by weight catalyst is added (0.0447 g.); the vial is sealed with a serum cap and subjected to the desired environmental conditions. At prescribed time intervals samples are removed by hypodermic syringe and subjected to GC analysis. Solution loss does not occur using this method.

### B. Infrared Analysis of Cure Systems

Absorbance changes in the infrared spectrum of binder mixtures provides a method for observing reaction rates in neat cure systems (Refs. 4 and 5).

Cure reaction formulations are prepared in small sample vials. A sample of known weight is transferred to a salt disc fitted with a Teflon spacer. A second salt disc is placed over the sample and spacer to complete the test specimen. The resulting cell is gently compressed in a nut and screw type ball joint clamp to expel air and to evenly distribute the cure formulation in the cavity between the salt discs. At temperatures as high as 100°C, the use of NaCl plates was shown not to have a catalytic effect on the reaction (Refs. 4 and 6).

The absorbance changes are noted at carbonyl bands related to COOH at 5.9  $\mu$  and to COOR at 5.8  $\mu$ . In the Telagen CT systems characteristic absorbance bands of the ethylenic group at 5.5  $\mu$  and 6.1  $\mu$  serve as convenient internal standard points for quantitative observations of the acid-to-ester conversion which takes place during the cure.

NEAR THE

A secondary group of bands, one at 2.9  $\mu$  related to development of CHOH and one at 3.2  $\mu$  due to COOH are also employed for noting changes during the progress of cures. The baseline absorbances at 2.7  $\mu$  and 4.7  $\mu$  are convenient reference points for quantitative estimation of changes in the two components as they occur.

Cell thicknesses for conducting the analyses were established to be 0.050 mm. for observing changes in the 5.8- $\mu$  and 5.9- $\mu$  bands and 0.20 mm. for noting changes in the 2.9- $\mu$  and 3.2- $\mu$  bands.

The baseline method was used to measure absorbances. A calibration curve relating absorbance ratios (5.8 and 5.9 to 6.1  $\mu$ ) to concentration (meq) of carboxylic acid and ester was established as follows: A series of CTPB samples containing different ratios of ester to acid were prepared by esterification of CTPB with less than equivalent amounts of epoxybutane. Residual acid content in each sample was determined by titration. The calibration curve was then established by relating the known acid and ester concentrations of the various samples to their respective infrared responses.

As the cure reaction proceeds, the intensity of the CGOH peak at 5.9  $\mu$  decreases continuously while the ester peaks at 5.8  $\mu$ g. During the latter stages of the conversion, the 5.9  $\mu$  peak is visible only as a shoulder in the spectrum and carboxylic acid content cannot be determined accurately by IR at this point. In this region, the extent of reaction was determined from the ester peak intensity-concentration relationship.

### C. Gumstock Studies

The method employed was to prepare 30-g. mixes of the binder consisting of 70% polymer, the specified quantity of cure catalyst and solids, and the remainder DOA. The binder was evacuated and cast into four 8-in. long by 1/4-in. diameter Teflon tubes. After cure for the specified time period, the binder was removed from the Teflon tube and specimens were prepared. Duplicate mechanical property specimens (1/4-in. by 1-in. cylinders) were prepared by epoxy bonding to wooden end tabs. The samples were pulled on an Instron tester at a strain rate of 1 in/in/min until 35% elongation. The secant modulus at 30% elongation was then calculated. Samples were not tested until break because with an E/C of 1 and the use of all tri-epoxide, frequently, the binder is stronger than the bond to the wooden end tab which results in specimens breaking at the bond rather than within the binder phase. The bond was always sirong enough to take the specimen to 30% elongation. Gel content was determined on a 1-g. binder sample by a 48-hr. methylene chloride extraction which was followed by a 4-day acetone extraction when AP was present. Gel samples had a 1/16-in. web thickness. Linear swelling was determined on duplicate 1-in. long specimens in contact with benzene for 3 days. The benzene was replaced with fresh solvent on each of the first 2 days.

### D. Propellant Mixes

The 1-lb. propellant mixes were manufactured in a 1-qt. Baker-Perkins double vertical planetary mixer. The solids were added incrementally and the CrOct was added as a THF solution 40 min. before the end of the mix cycle. Viscosities were measured on a Haake rotoviscometer at 26°C, and the values at a shear rate of 0.411 sec<sup>-1</sup> are reported in Table XVII. Propellant mechanical properties were measured on Type II JANAF tensile specimens using the Instron tester and a strain rate of 0.74 in/in/min.

### E. <u>Materials</u>

The acids, 2-ethyl hexanoic, octanoic, and propionic, and the epoxidesepoxy butane and allyl glycidyl ether--were all reagent grade and supplied by Matheson Chemical Company. The polymer cure studies employed liquid carboxyl-terminated polybutadiene (CTPS) with an equivalent weight of 2,990 supplied by General Tire and Rubber Company and the epoxides ERLA-0510 (97-101 equivalent weight) supplied by Union Carbide and DER-332 (1.2-174 equivalent weight) from Dow. Moat of the metal salts used in the acreening studies were obtained from Shepherd Chemical Company.

### CONCLUSIONS AND RECOMMENDATIONS

1. Propellants based on carboxylic acid-epoxide and carboxylic acidaziridine binder systems and containing up to 85.5% solids (AP, Al) can be mixed, cast, and cured completely and successfully at  $25^{\circ} \pm 1^{\circ}$ C. High solids loadings and ease of processing are possible because the chromium catalyst used (Cr 2-ethyl hexanoate) acts as a surfactant at the concentration level employed. The binder prepolymers to which this cure system has been applied include Telagen CT, PBAN, and HC-434.

2. For the ambient cure system, chromium salts are excellent catalysts for both epoxide and aziridine systems. Catalyst levels vary from 0.5 to 1.5 wt. % of the binder depending on the specific prepolymer.

3. Certain propellant ingredients and additives were shown to affect processing and/or cure. Minor adjustments in the formulation will compensate for the observed effects.

4. Surveillance studies of model, binder, and prope!"int samples have shown no detrimental aging effects due to the presence of the cure catalyst. The CrOct catalyzed propellants age as well as 140°F cured control propellants.

5. The Telagen-epoxide ambient-cured binder system is ready for scale-up beyond the 40-lb. batch scale size, 85.5% total solids propellant demonstrated in this program.

### XII.

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### GLOSSARY OF TERMS





BITA (HX-868)

AcAz



 $\stackrel{0}{\mathbb{P}} \left( \underset{CH_2}{\mathbb{N} \leq \binom{CH_3}{CH_2}} \right)_3$ 

$$\frac{S}{P} \left( -N \left( \frac{CH_3}{CH_2} \right) \right)$$

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UNCLASSIFIED Security Classification

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KEY WORDS LINK A LINK B	LINK C	
ROLE WT ROLE WT	ROLE	WT
Carboxy Terminated Binders		
Cure Catalysts		
Ambient Temperature Cure		
Epoxy Curatives		
Organometallics		
Solid Propellants		
Aziridine Curatives		
Organometallic Cure Catalysts		
Telagen CT		
PBAN		
HC 434		

Security Classification