Form Approved REPORT DOCUMENTATION PAGE OMB No. 0704-0188 Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources gathering and maintaining the data needed, and completing and reviewing this collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden to Department of Defense, Washington Headquarters Services, Directorate for Information Operations and Reports (0704-0188), 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302. Respondents should be aware that notwithstanding any other provision of law, no person shall be subject to any penalty for failing to comply with a collection of information if it does not display a currently valid OMB control number. PLEASE DO NOT RETURN YOUR FORM TO THE ABOVE ADDRESS. 1. REPORT DATE (DD-MM-YYYY) 2. REPORT TYPE 3. DATES COVERED (From - To) July 2015 **Briefing Charts** July 2015-August 2015 4. TITLE AND SUBTITLE 5a. CONTRACT NUMBER Isocyanate-Free Elastomers as Replacements for Isocyanate-cured In-House Polyurethanes (Briefing Charts) 5b. GRANT NUMBER 5c. PROGRAM ELEMENT NUMBER 6. AUTHOR(S) 5d. PROJECT NUMBER Josiah T. Reams, Andrew J. Guenthner, Jacob C. Marcischak, Michael D. Ford, Timothy S. Haddad, Gregory R. Yandek, Joseph M. Mabry 5e. TASK NUMBER **5f. WORK UNIT NUMBER** O0BG 7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES) 8. PERFORMING ORGANIZATION REPORT NO. Air Force Research Laboratory (AFMC) AFRL/RQRP 10 E. Saturn Blvd. Edwards AFB, CA 93524-7680 9. SPONSORING / MONITORING AGENCY NAME(S) AND ADDRESS(ES) 10. SPONSOR/MONITOR'S ACRONYM(S) Air Force Research Laboratory (AFMC) AFRL/RQR 11. SPONSOR/MONITOR'S REPORT 5 Pollux Drive NUMBER(S) Edwards AFB CA 93524-7048 AFRL-RQ-ED-VG-2015-301 12. DISTRIBUTION / AVAILABILITY STATEMENT Distribution A: Approved for Public Release; Distribution Unlimited. 13. SUPPLEMENTARY NOTES Briefing Charts presented at 250th ACS National Meeting; Boston, MA; 16-20 August 2015. PA#15447. 14. ABSTRACT **Briefing Charts**

15. SUBJECT TERMS

16. SECURITY CLASSIFICATION OF:			17. LIMITATION OF ABSTRACT	18. NUMBER OF PAGES	19a. NAME OF RESPONSIBLE PERSON Joseph Mabry
a. REPORT	b. ABSTRACT	c. THIS PAGE		24	19b. TELEPHONE NO
Unclassified	Unclassified	Unclassified	SAR		(include area code) 661-275-5857



Isocyanate-Free Elastomers as Replacements for Isocyanate-cured Polyurethanes

Josiah T. Reams, Andrew J. Guenthner, Jacob C. Marcischak, Michael D. Ford, Timothy S. Haddad, Gregory R. Yandek, Joseph M. Mabry



ACS National meeting 20 August 2015

Josiah Reams, Ph.D. ERC Inc. / Air Force Research Laboratory josiah.reams.ctr@us.af.mil (661) 275-5664

DISTRIBUTION A: Approved for public release; distribution is unlimited.



Outline



Goal

 Investigate polymeric cure systems with crosslinking chemistries that are unlikely to pose health concerns now and in the future

Objective

- Synthesis of new isocyante-free candidate binder formulations
- Determine impact physical and mechanical properties
- Demonstrate the use of new polymeric binders in binder formulation

Technical Challenges

- Maintain energy content and I_{sp}
- New system must conform to current binder processing operations
- Retention of physical and mechanical properties of HTPB / isocyanate systems
 - Low T_g
 - High elongation to break
 - Compatibility with high solids loading

Approach

- Nucleobase binding
- Thiolene "click" chemistry

Acknowledgements:

- Strategic Environmental Research and Development Program (SERDP)
- Air Force Office of Scientific Research, Air Force Research Laboratory – Program Support; PWG team members (AFRL/RQRP)

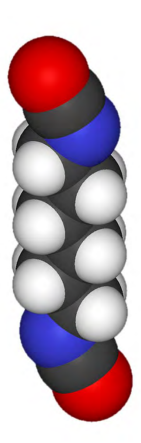




Background: Isocyanates



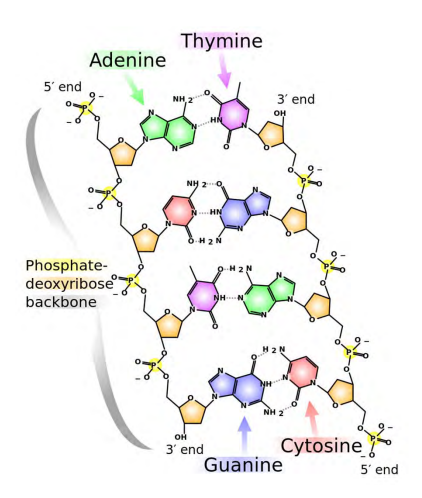
- Isocyanates are prized for their high reactivity; they are particularly useful for production of energetic materials where high temperatures must be avoided
- Isocyanates are a respiratory irritant and a significant cause of occupational asthma
- Sensitization can occur at very low levels of exposure, with effects continuing for many years afterward
- Cross-sensitization (e.g. dermal exposure leading to sensitization of respiratory tract) has also been reported.
- Isocyanates are not found in nature; their high reactivity without specificity makes them unsuitable for complex biological systems
- Increased regulation will result in increased cost and eventual unavailability in the U.S.





Nucleobase Binding





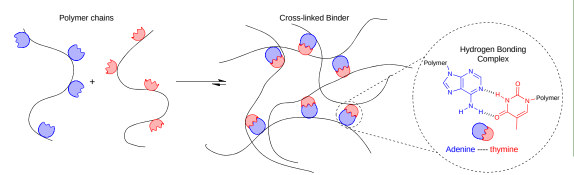
- DNA is biologically ubiquitous therefore no current environmental regulation and likely no regulation in the future
- Potential advantages
 - Dynamic covalent chemistry is a hot topic
 - Potential for self-healing properties
 - Reversible crosslinks have the potential for easy demilitarization
 - Insensitivity
- Potential risks of nucleobase crosslinked polymer systems
 - Polymers containing nucleobase monomers are rare and therefore are not commercially available
 - Properties not well understood



Incorporation of Nucleotide Bases in Polybutadiene



Copolymerization of either adenine or thymine methacrylate with butadiene gives an "A" and "B" complimentary copolymers Cheng, S; Zhang, M.; Dixit, N.; Moore, R. B.; Long, T. E. Macromolecules, 2012 45, 805-812.

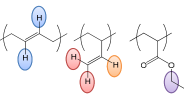


- When in the presence of each other complimentary copolymers associate and form a crosslinked system consisting of hydrogen bonds
- Both components exist as low viscosity liquids and do not self-associate
- When mixed together, the two part system is initially a low viscosity liquid that forms a crosslinked network as complimentary nucleobases associate

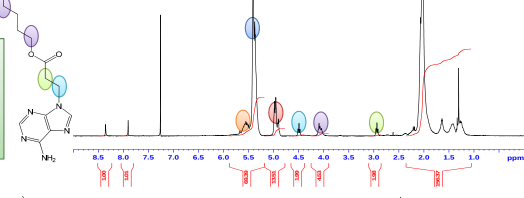


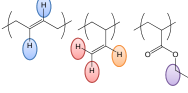
Adenine and Thymine Acrylate / Butadiene Copolymerization



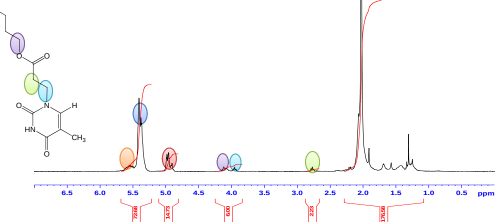


- Copolymerization of adenine acrylate with butadiene was performed in DMSO with AIBN as the initiator
- Adenine acrylate incorporation of 2.5 mol% was obtained from a 4 mol% feed ratio





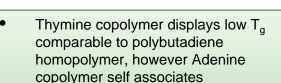
- Thymine acrylate incorporation of 2.7 mol% was obtained from a 4 mol% feed ratio
- Adenine acrylate and thymine acrylate appear to have near identical reactivity ratios as would be expected
- Overall yield of ~20% for adenine and thymine copolymerizations was consistent with butyl methacrylate copolymerization with butadiene



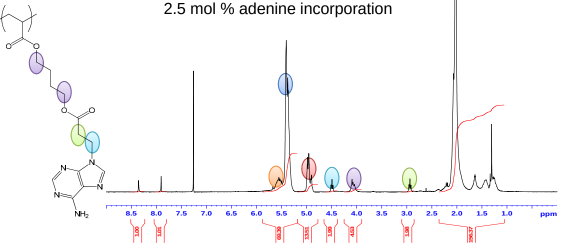


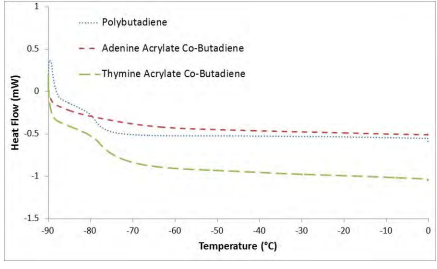
Adenine Acrylate Co-Butadiene Polymerization Variability

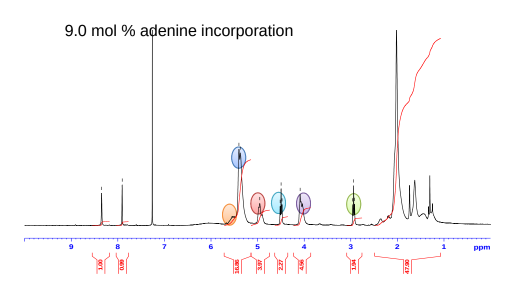




 Variability in Adenine copolymerization was observed with respect to nucleobase incorporation



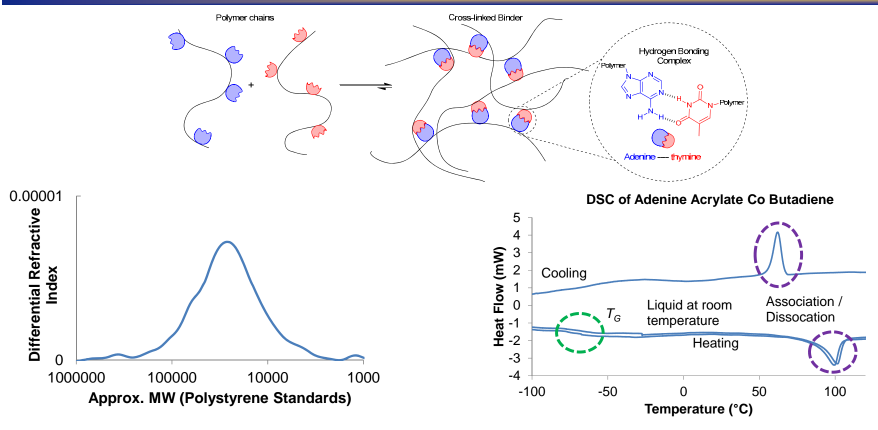






DNA Nucleobase Polymer Properties



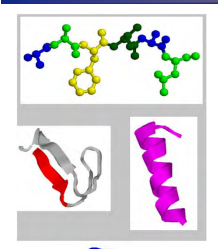


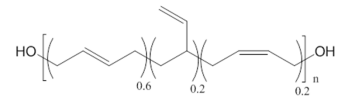
- In polar media adenine units do not self associate, however non-polar media promotes self-association
- Self association of individual "part A" or "part B' components makes mixing of two components difficult
- Too many unknown characteristics of self- and cross-association for near-term development of reliable cross-linking substitute



Carbon-Sulfur Bonds as Crosslinks







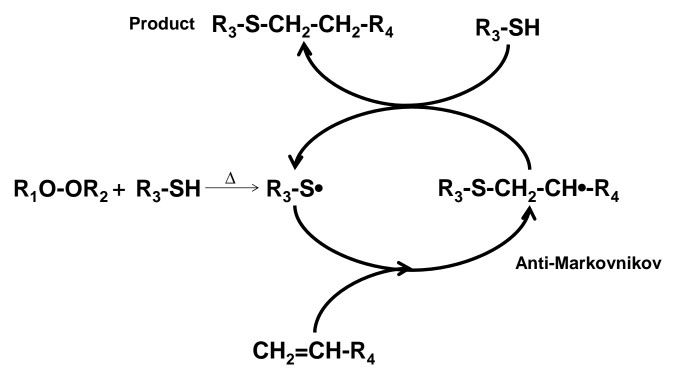
HSCH₂(CH₂)₇CH₂SH

- Many sulfur containing compounds exist in nature including proteins
- Thiol-ene chemistry not biologically ubiquitous like nucleobase binding
- Thiol-ene chemistry similar to free radical in terms of reactivity but with more control and selectivity
- Utilization of existing vinyl double bonds in commercial polybutadiene



Thiol-Ene Chemistry





- Thiol-ene chemistry is a hybrid of free radical and condensation chemistry.
 It combines the speed and ease of initiation of free radical chemistry with the product specificity of condensation chemistry.
- As a type of "click" chemistry, it generates no volatiles and minimal side products.



Peroxide Catalyzed Thiol-ene Click



Peroxides

Lauroyl Peroxide Luperox ® LP 64 °C

t-butyl peroxybenzoate Luperox ® P 104 °C

$$\begin{array}{c|c}
H_3C & CH_3 \\
N \equiv C & N & C \equiv N \\
H_3C & CH_3
\end{array}$$

AIBN

t-butylperoxy 2ethylhexyl carbonate, Luperox ® TBEC 100 °C

Benzoyl Peroxide 73 °C

Dicumyl Peroxide 117 °C (DCP)

Thiols

HSCH₂(CH₂)₇CH₂SH

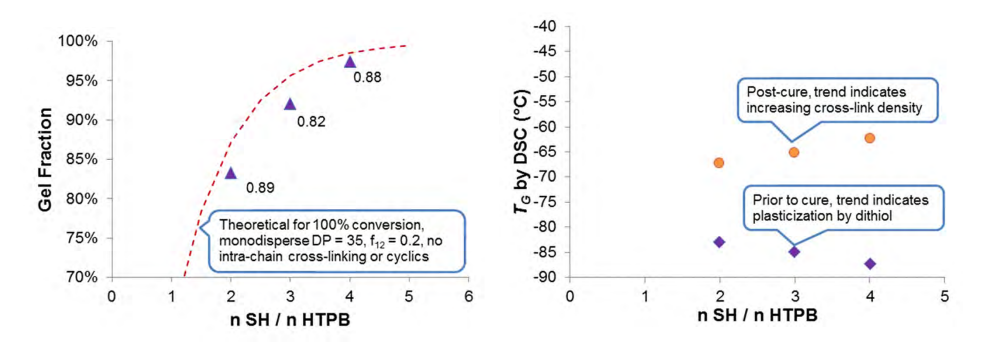
1,9-Nonanedithiol, 95%

- Many available peroxide initiators with different 10 hour half-life temperatures allow reactivity to be tuned to processing conditions
- Ratio of peroxide to thiol can be varied to tune rate of initiation
- Choice of thiol affects crosslink density and hydrolytic stability
- Peroxide-initiated nonanedithiol /HTPB cure was chosen to match current HTPB-isocyanate processing conditions at 60 °C



Thiol-ene Crosslinked HTPB



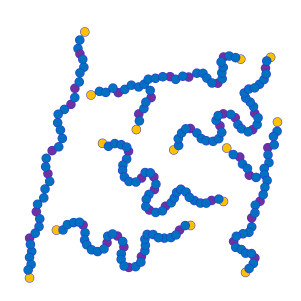


- HTPB cured with 1,9-nonanedithiol / TBEC (peroxide) at 5:1 SH / O•, 60 °C, for 8 days under N₂. Data point labels (left figure) indicate fraction of available –SH incorporated into gel, a measure of conversion.
- Glass transition temperatures remain at acceptable levels over a wide range of cure conditions



Importance of Network Architecture





Chain backbone (1,4 addition)
Vinyl goups (1,2 addition)
Hydroxyl end groups

Free radical polymerized HTPB

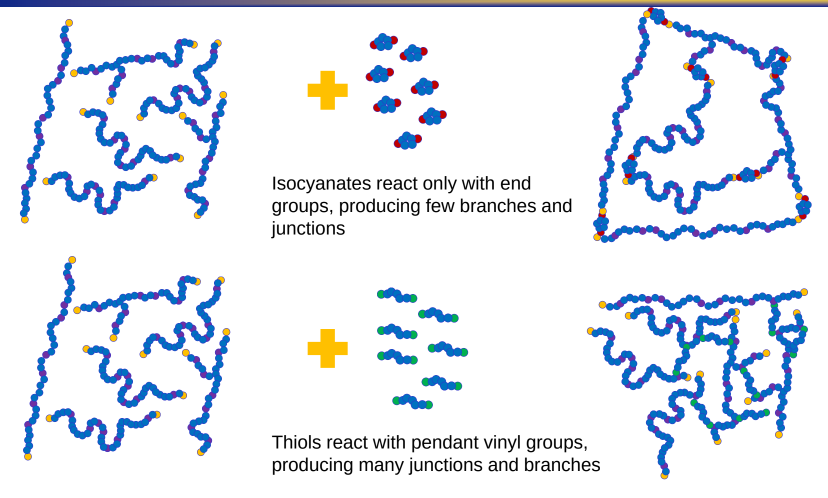
- Mixture of 1,4 and 1,2 addition
- Hydroxyl end-group functionality 2.4-2.6 / per chain
- T_α -75 °C
- Molecular Weight M_n = 2800 g/mol
- Polydispersity ~2.5





Importance of Network Architecture





Reproduction of correct polymer architecture is necessary to obtain correct mechanical properties



Thiol-ene cured HTPB Modulus



Luperox® P

Luperox® TBEC

Peroxide	Thiol	SH:HTPB	Theoretical Modulus (psi)	Measured Modulus* (psi)
Luperox P	1,9 Dithiol	3:1	215	141
Luperox P	1,9 Dithiol	4:1	360	331
TBEC	1,9 Dithiol	3:1	492	419
TBEC	1,9 Dithiol	4:1	851	502

^{*}Compressive modulus at 298K measured by thermomechanical analysis (TMA)

- Mechanical property data suggests thiol-ene cured HTPB has a lower molecular weight between crosslinks and isocyanate cured HTPB
- Reaction at chain ends rather than at pendant vinyl groups is necessary to achieve correct architecture and mechanical properties



Strategies for Desirable Architecture



- HTPB without vinyl side groups
- Polybutadiene (PB) with low vinyl incorporation
 - Polybutadiene with 1% vinyl
 - Difficult to tell the position of vinyl groups on polymer chain
 - Initial investigation with nonanedithiol cure produces elastomeric material
- Thiol-terminated polyethylene glycol (PEG) system
 - Co-cured with di and tri-functional vinyl compounds
 - Demonstration of principle in progress
- Thiol-terminated polybutadiene
 - Co-cure with di and tri-functional vinyl groups
 - Presence of main chain vinyl groups will result in intramolecular reaction / cyclization
 - Synthesis of polybutadiene with di-anionic initiator to produce cis-1,4 substituted polybutadiene



Thiol-Terminated Polybutadiene (from HTPB)



$$R = OH + CI =$$

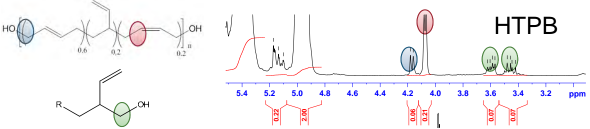
Thiol-terminated polybutadiene

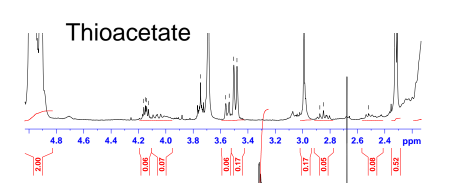
- Presence of main chain vinyl groups will result in intramolecular reaction / cyclization
- Co-cure with di and tri-functional vinyl groups

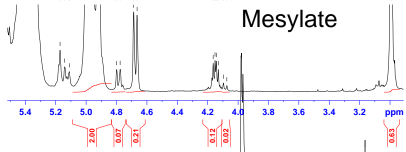


Thiol-terminated HTPB Synthesis

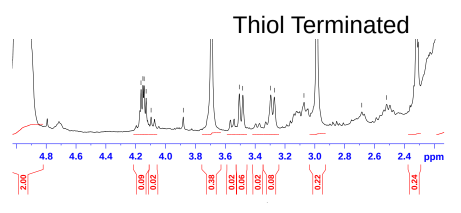








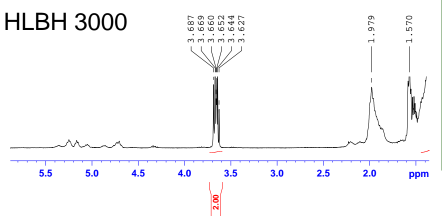
- Reaction conditions optimized for hydroxyl protection with mesylate. Side reactions were observed in the presence of triethylamine and methanol
- Trans mesylate groups were surprisingly unreactive
- Formation of thiol terminated HTPB was accompanied by concomitant thiol-ene formation
- Propensity of thiol groups to undergo initiator-free thiolene click chemistry makes use of HTPB backbone impractical for real world use



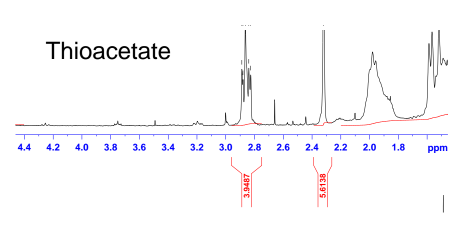


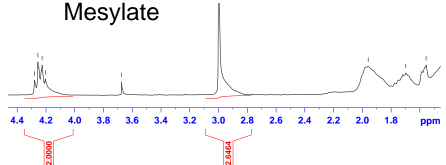
Thiol Terminated Hydrogenated Polybutadiene Synthesis

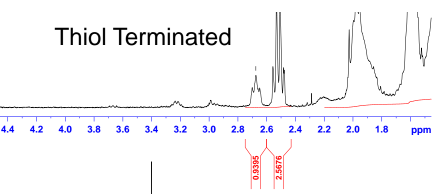




- Hydroxyl protection with mesylate groups proceeds without formation of side products as occurs with HTPB
- Formation of thiol terminated HLBH 3000 is obtained in good yield with high conversion, i.e. all hydroxyl groups present have equal reactivity
- Reduction of thioacetate to thiol is accompanied by disulfide formation (~25%)



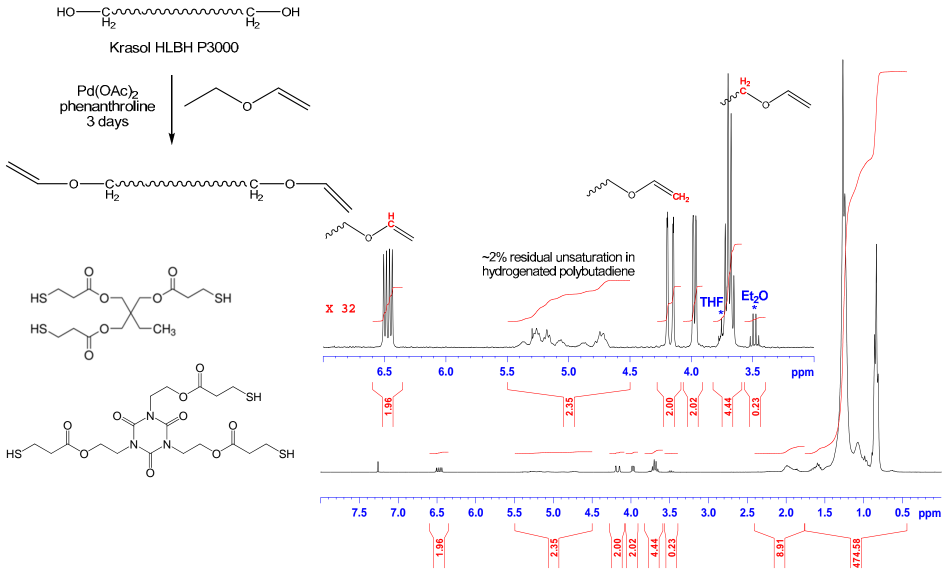






Vinyl Ether-Terminated Hydrogenated Polybutadiene







Impact of Results on Future DoD Operations



- Elimination of isocyanates alleviates a major occupational health and safety concern in the manufacture and use of solid rocket motor propellants.
- Elimination of isocyanates also mitigates issues related to moisture sensitivity during propellant mixing and casting, and may reduce some forms of degradation during long-term storage in humid environments.
- Replacement of isocyanates with chemical groups that are ubiquitous in the biosphere will greatly reduce the risk associated with future regulation / obsolescence throughout the manufacturing, use, and de-militarization life cycle.
- Alternative cure chemistries that are successful for solid rocket motors may be transitioned to other applications, such as paints, foams, sealants, and adhesives that also represent a significant source of occupational health risk for DoD and DOE.



Conclusions



- Increasing regulation of isocyanates in the workplace poses a significant, but not insurmountable, challenge to DoD/DOE operations
- DNA nucleobase technology represents a promising future path to isocyanate replacement, but is not yet mature enough for near-term development in solid rocket motors
- Thiol-ene chemistry represents a safer alternative to isocyanate chemistry that remains a promising candidate for near-term development in solid rocket motor propellants
- When paired with the proper polymer architecture, thiol-ene based propellants will offer simpler manufacturing, improved quality control, greatly reduced occupational health and safety concerns, and some potential decrease in humid ageing in solid rocket motor propellants.

QUESTIONS?



U.S. AIR FORCE

DISTRIBUTION A: Approved for public release; distribution is unlimited.