

# High-Nitrogen and High-Oxygen Chemistry



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# Report Documentation Page

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# Program Objectives



***Discover, synthesize, characterize, and scale-up novel, highly energetic compounds***

## Technical Approach:

- Exploit synergism between theory and synthesis
- Use calculations to identify the most promising candidates and predict their properties
- Employ experimental expertise to design synthetic approaches, prepare novel compounds, and characterize products



- Polynitrogen Chemistry

$N_3^+$  Chemistry

$N_3NOF^+$  and  $N_7O^+$

New NMR Method

- Polyazide Chemistry

- Synthesis and Characterization of  $FN(NO_2)_2$

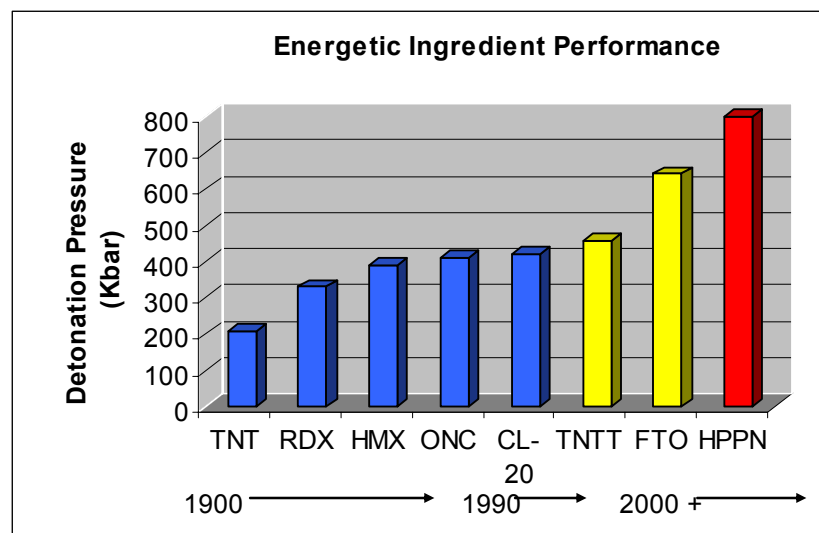
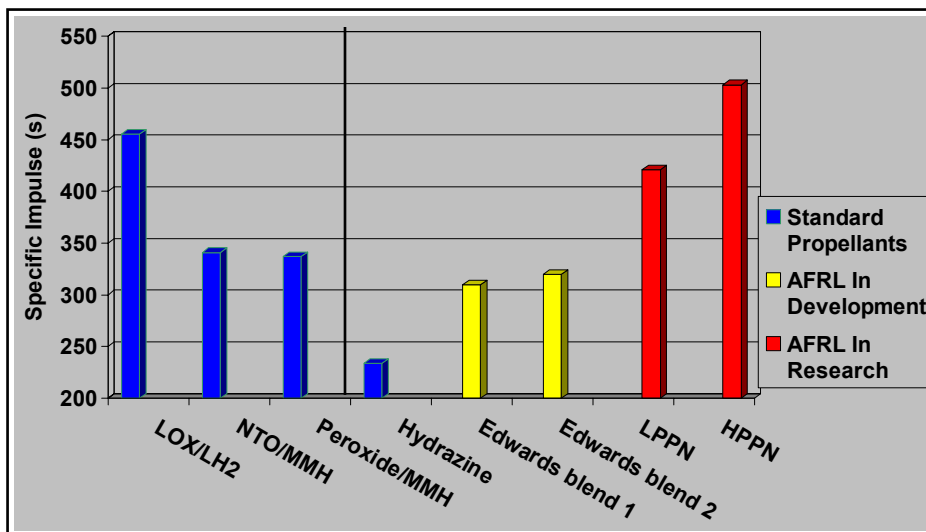
- Stable Difluorammine Sources

- High-Oxygen Carriers and Oxidizer Balanced Ionic Liquids

# Why are we interested in Polynitrogens?



**The performance of polynitrogens as monopropellants would dwarf that of hydrazine, would greatly exceed even many bipropellants, and result in reduced signature**



LPPN = Low performing polyN ( $N_5^+N_3^-$ );

HPPN = High performing polyN (cubic  $N_8$ )

**Polynitrogens would also have great potential as high-performance explosives**

# Bulk Synthesis of $N_3^+$

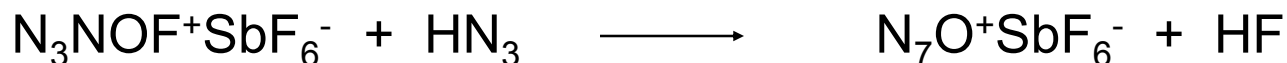
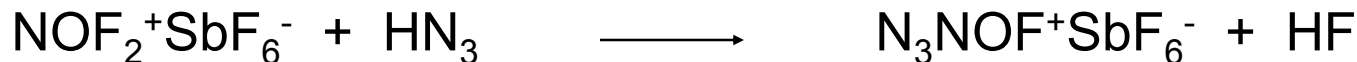


- $N_3^+$  is a very promising candidate (is derived from a vibrationally stable radical and has high decomposition barrier)
- Studied  $F^-$  abstraction from  $FN_3$  by strong Lewis acids
- Solved problem of synthesis and safe handling of  $FN_3$  on a preparative scale, but found that  $N_\alpha$  is a better donor than  $F$
- HF addition to the  $FN_3$ - $SbF_5$  adduct, followed by  $N_2$  elimination, results in the formation of  $NH_2F_2^+$  salts
- Are working on synthesis of  $XeN_3^+$  and its decomposition and on controlled photolytic decomposition of  $N_5^+$  as potential methods for the production of  $N_3^+$

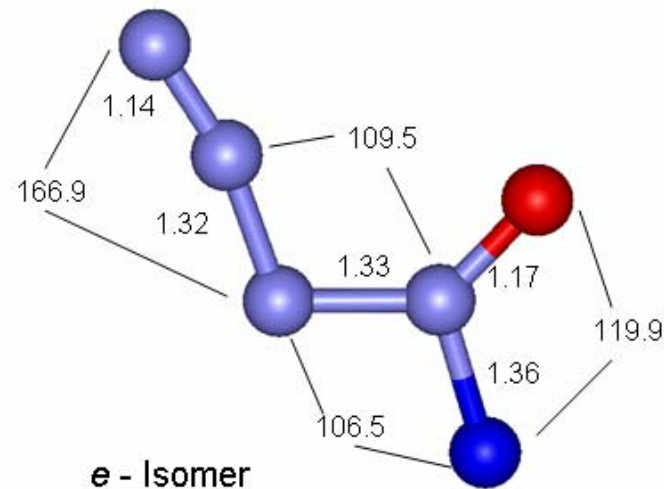
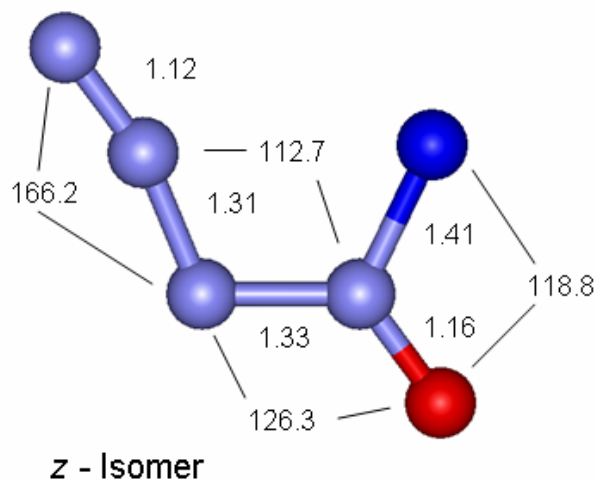
# Syntheses of $N_3NOF^+$ and $N_7O^+$



- The reactions



were studied in HF at  $-78^\circ\text{C}$ .  $N_3NOF^+SbF_6^-$  was isolated and is stable up to  $\sim 20^\circ\text{C}$ .  $N_3NOF^+$  exists as both a *z*- and an *e*-isomer.





- Nitrogen NMR spectroscopy is crucial for the identification of novel polynitrogen compounds. Unfortunately,  $^{14}\text{N}$  has a large quadrupole moment which broadens most signals to a point where they become unobservable, and the natural abundance of  $^{15}\text{N}$  (0.36%) is too low and its relaxation time too long to allow the observation of  $^{15}\text{N}$  spectra without  $^{15}\text{N}$  enrichment. Prof. Taylor has developed a **new NMR signal processing technique for the detection and enhancement of very weak NMR signals.**
- The power of this new method was demonstrated for the natural abundance  $^{15}\text{N}$  spectrum of  $\text{N}_5^+$  which allowed the observation of the complete spectrum with an excellent signal to noise ratio. We were also able to observe the  $^{14}\text{N}$ - $^{15}\text{N}$  and  $^{15}\text{N}$ - $^{15}\text{N}$  couplings in a partially  $^{15}\text{N}$  enriched spectrum.

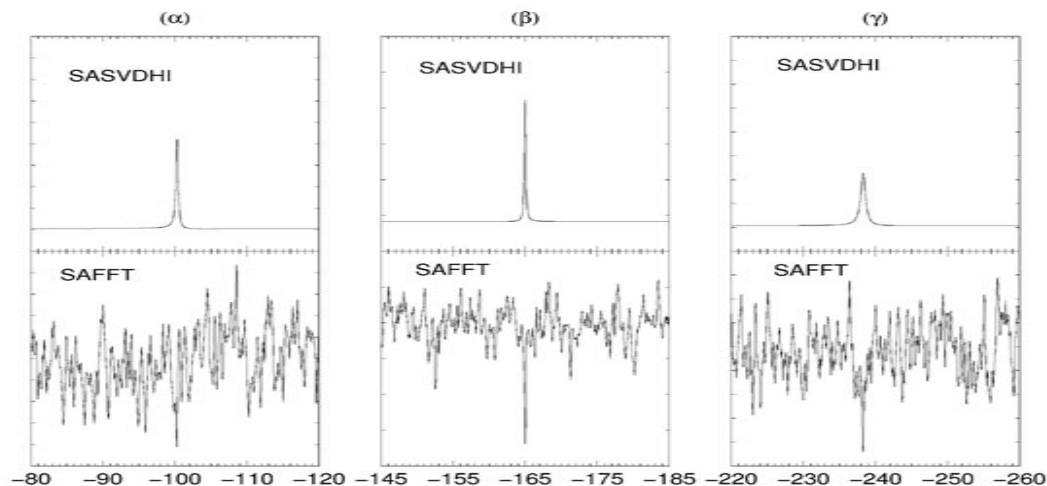


- Same raw natural abundance  $^{15}\text{N}$  data processed by the new and the conventional FT NMR methods for  $\text{N}_5^+$ .

## Avoiding Enrichment

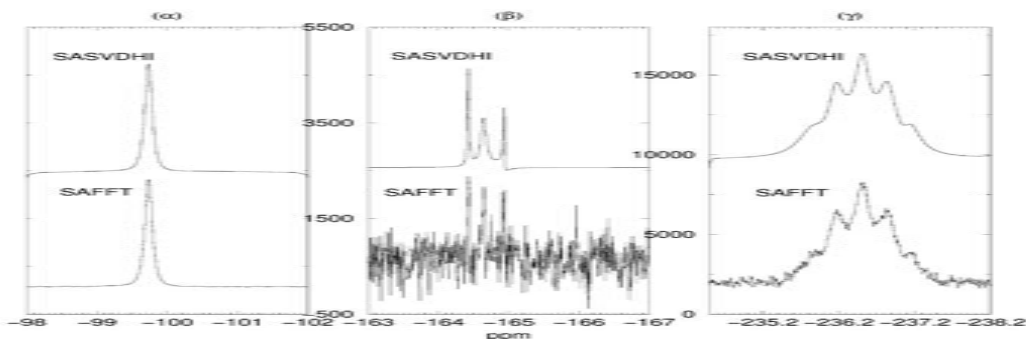
New Method

Conventional  
NMR



New Method

Conventional  
FT NMR



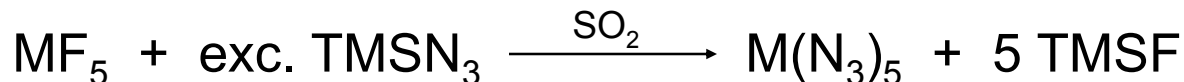


- One azido group contributes ~ 80 kcal/mol of endothermicity to a compound.
- Polyazides are highly energetic, sensitive materials which can be used for primary explosives. Typical example:  $\text{Pb}(\text{N}_3)_4$ .
- Have recently prepared and characterized numerous polyazides of S, Te, P, As, Sb, Ti, Mo and W, including spectacular compounds, such as  $\text{N}_5^+[\text{P}(\text{N}_3)_6]^-$  and  $\text{N}_5^+[\text{B}(\text{N}_3)_4]^-$ .
- Results were published in a series of papers in *Angewandte Chemie and Chemistry*, A European Journal.

# Synthesis of First Group 5 Binary Azides, $Nb(N_3)_5$ , $Ta(N_3)_5$ , $Nb(N_3)_5 \cdot CH_3CN$ and $[Nb(N_3)_6]^-$

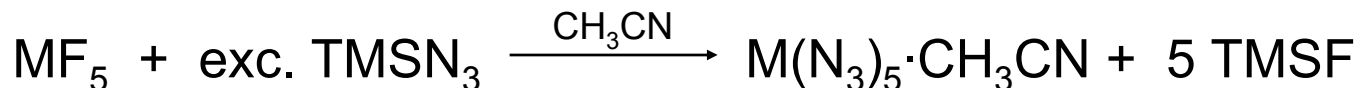


- Synthesized  $M(N_3)_5$  ( $M = Nb, Ta$ ) in  $SO_2$  according to

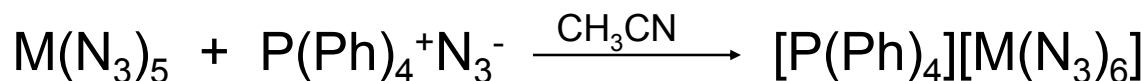


Both compounds are very sensitive and unstable. Raman spectra were recorded.

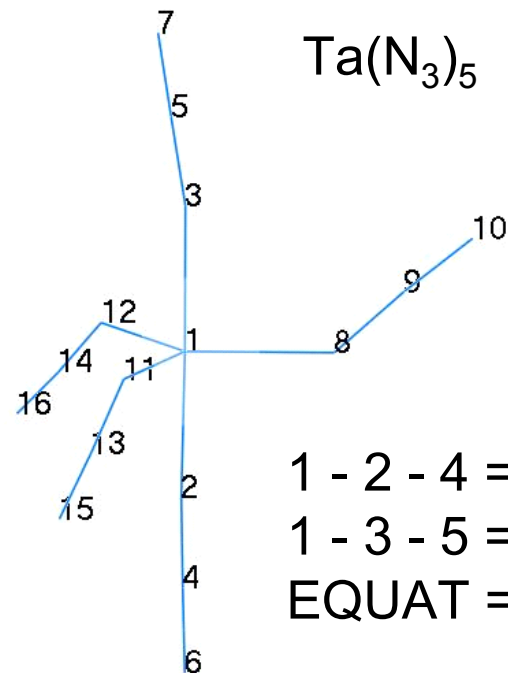
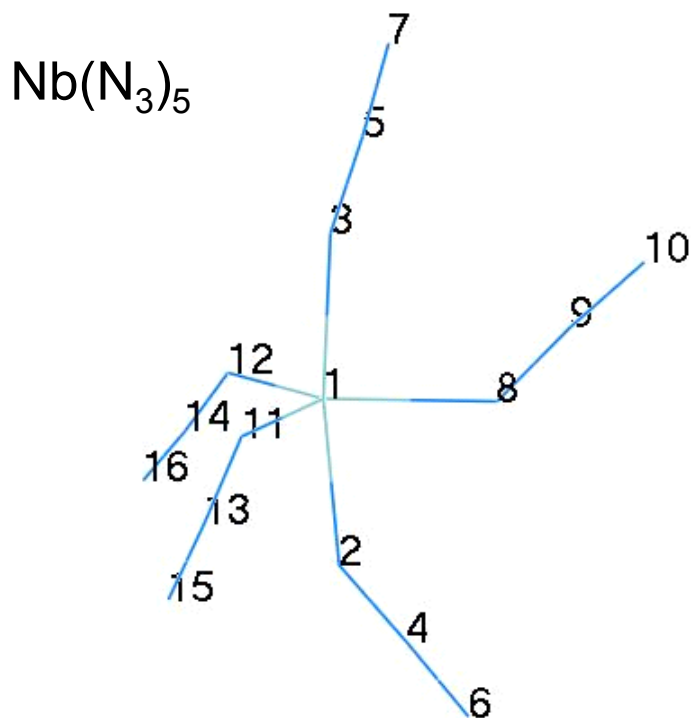
- Synthesized the adducts  $M(N_3)_5 \cdot CH_3CN$  ( $M = Nb, Ta$ ) according to



- Synthesized the anions  $[M(N_3)_6]^-$  ( $M = Nb, Ta$ ) according to



# Structures of $Nb(N_3)_5$ and $Ta(N_3)_5$



$$1 - 2 - 4 = 178^\circ$$

$$1 - 3 - 5 = 171^\circ$$

$$\text{EQUAT} = 137^\circ$$

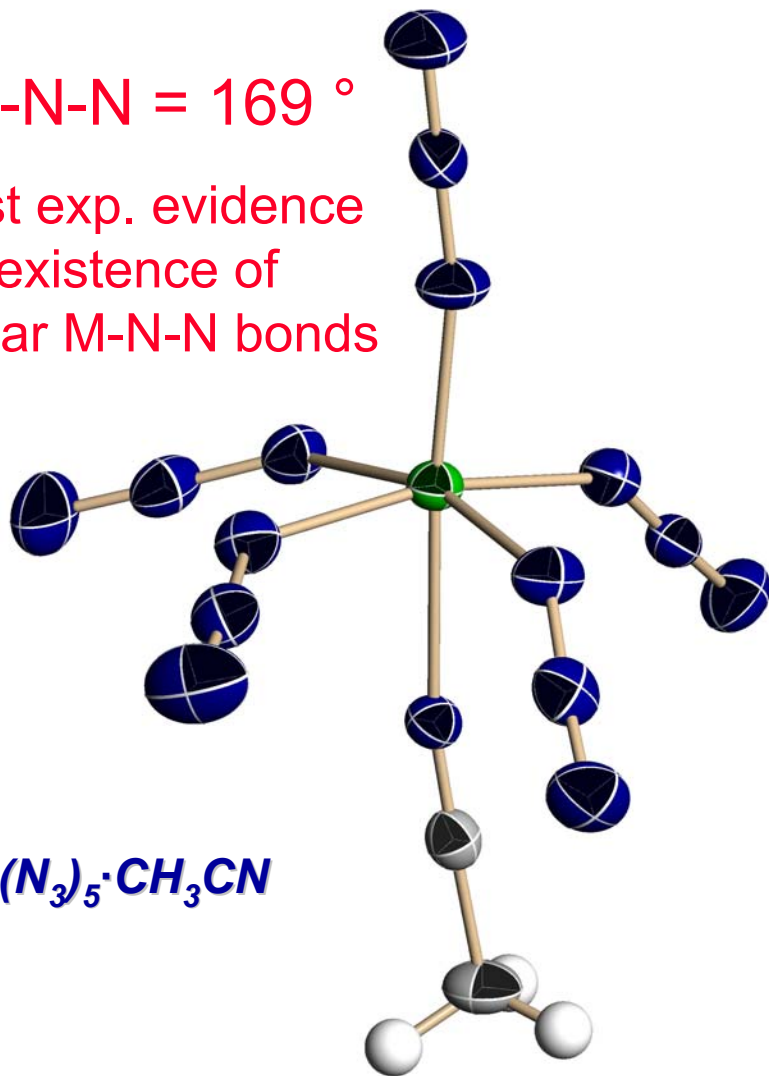
- Contain both linear and bent M-N-N bonds
- Calculated structures are supported by observed Raman spectra

# Crystal Structures of $Nb(N_3)_5 \cdot CH_3CN$ and $[Nb(N_3)_6]^-$

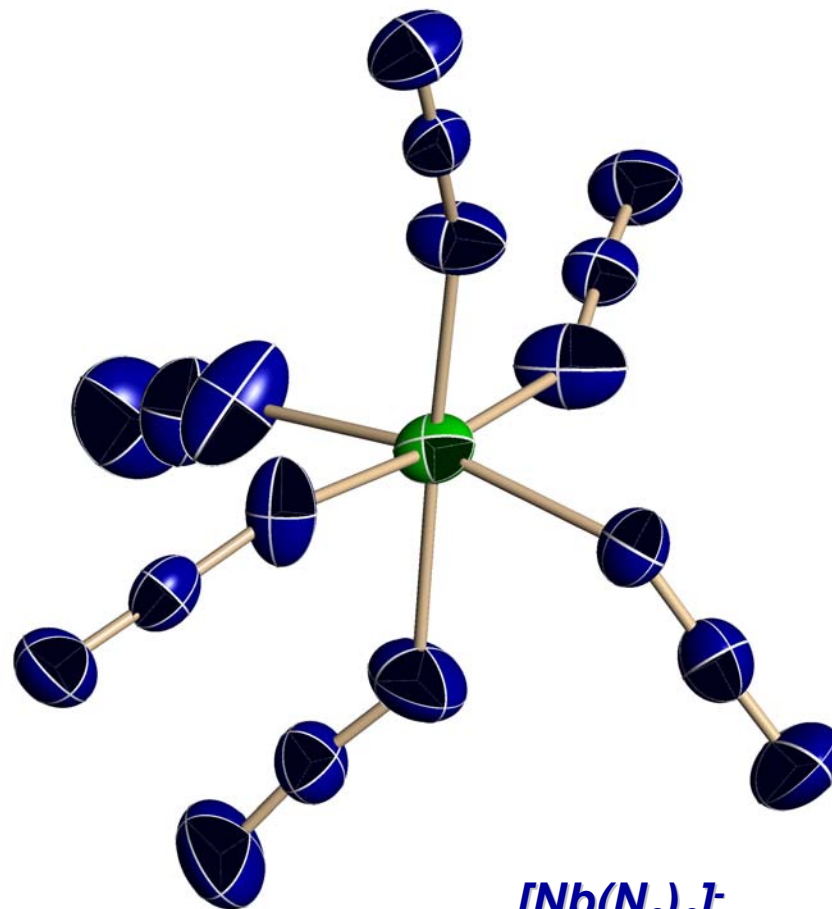


$Nb-N-N = 169^\circ$

First exp. evidence  
for existence of  
linear M-N-N bonds



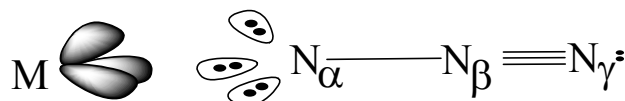
$Nb(N_3)_5 \cdot CH_3CN$



$[Nb(N_3)_6]^-$

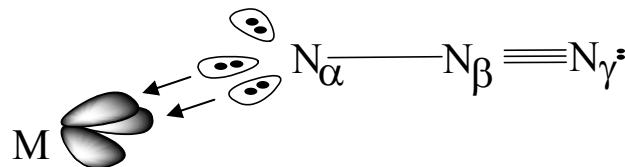
Bonding schemes for transition metal azides (from top to bottom): (i) ionic azide, showing for didactic reasons the azide ion in one of its asymmetric resonance structures and only some of the empty  $s_2d_{10}$  orbitals on M; (ii) strongly bent two-center/monodative bond; (iii) moderately bent two-center/bidative bond; (iv) linear two-center/tridative bond.

**IONIC**  
 $M^+ N_3^-$



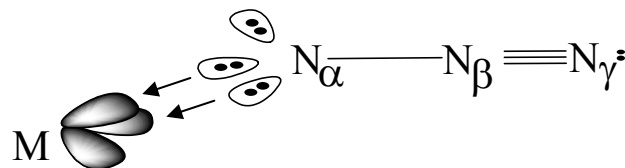
**2c-1d**

$(M-N_\alpha-N_\beta) = 109.5^\circ$



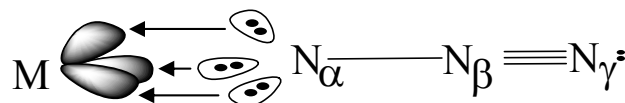
**2c-2d**

$(M-N_\alpha-N_\beta) = 125.5^\circ$



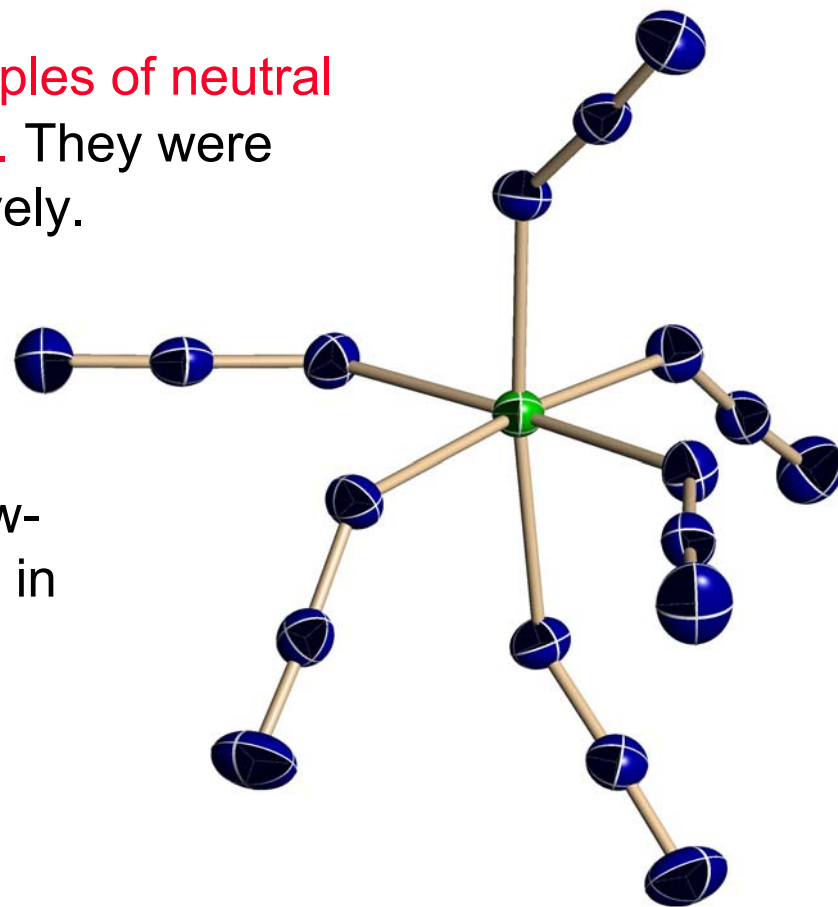
**2c-3d**

$(M-N_\alpha-N_\beta) = 180.0^\circ$



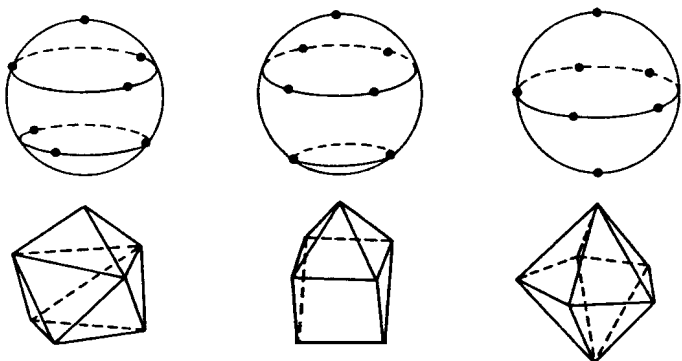
- $\text{Mo}(\text{N}_3)_6$  and  $\text{W}(\text{N}_3)__6$  are the **first examples of neutral hexaazides and binary Group VI azides**. They were prepared from  $\text{MoF}_6$  and  $\text{WF}_6$ , respectively.

- Both compounds are highly shock sensitive and were characterized by low-temperature Raman spectroscopy and, in the case of  $\text{W}(\text{N}_3)_6$ , also by its crystal structure.

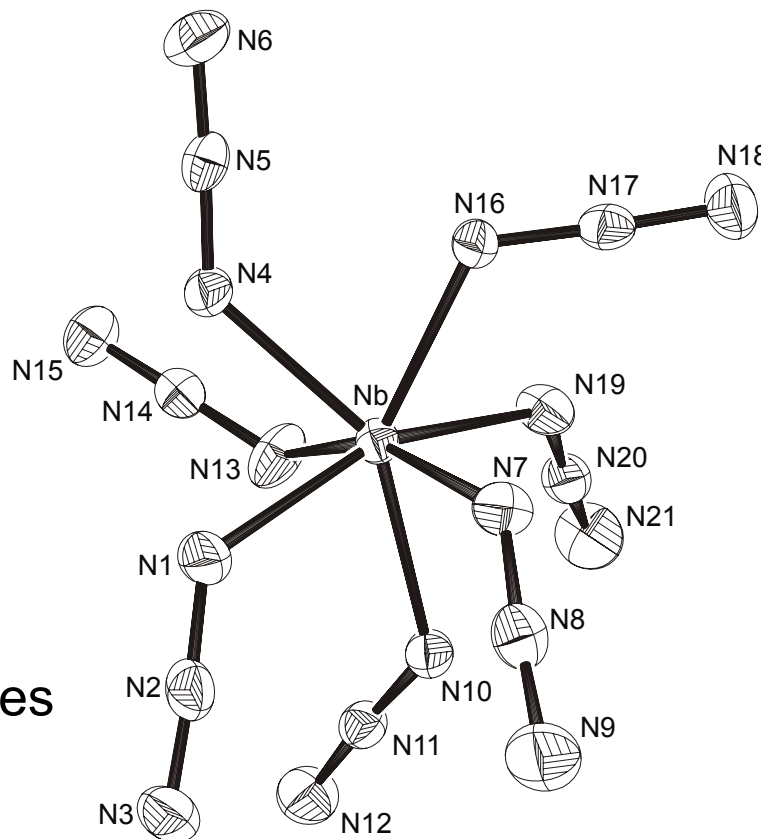


- Synthesized the  $[W(N_3)_7]^-$ ,  $[Mo(N_3)_7]^-$ ,  $[Nb(N_3)_7]^{2-}$ ,  $[Ta(N_3)_7]^{2-}$  anions

Possible arrangements for CN 7



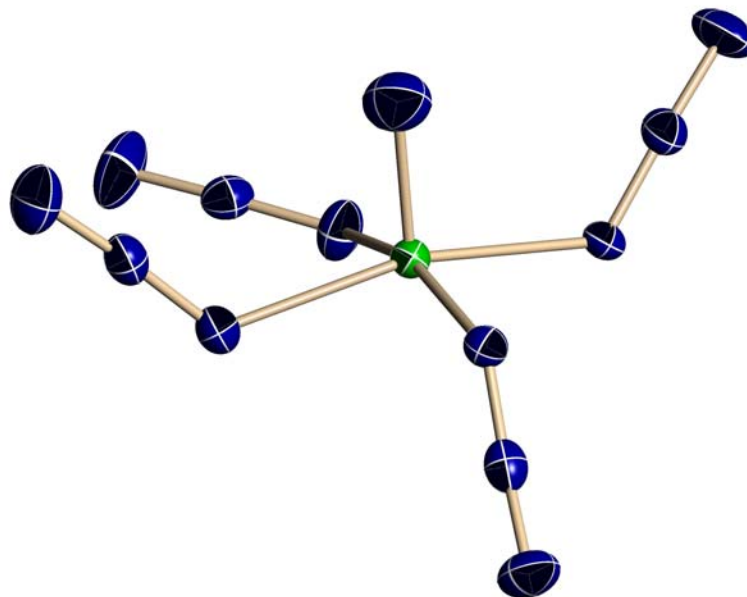
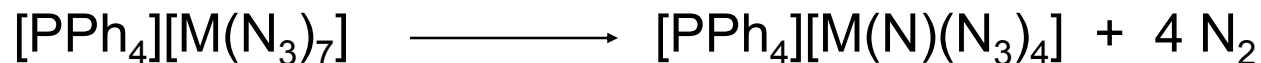
$M(N_3)_7$  anions have 2 : 4 : 1 structures



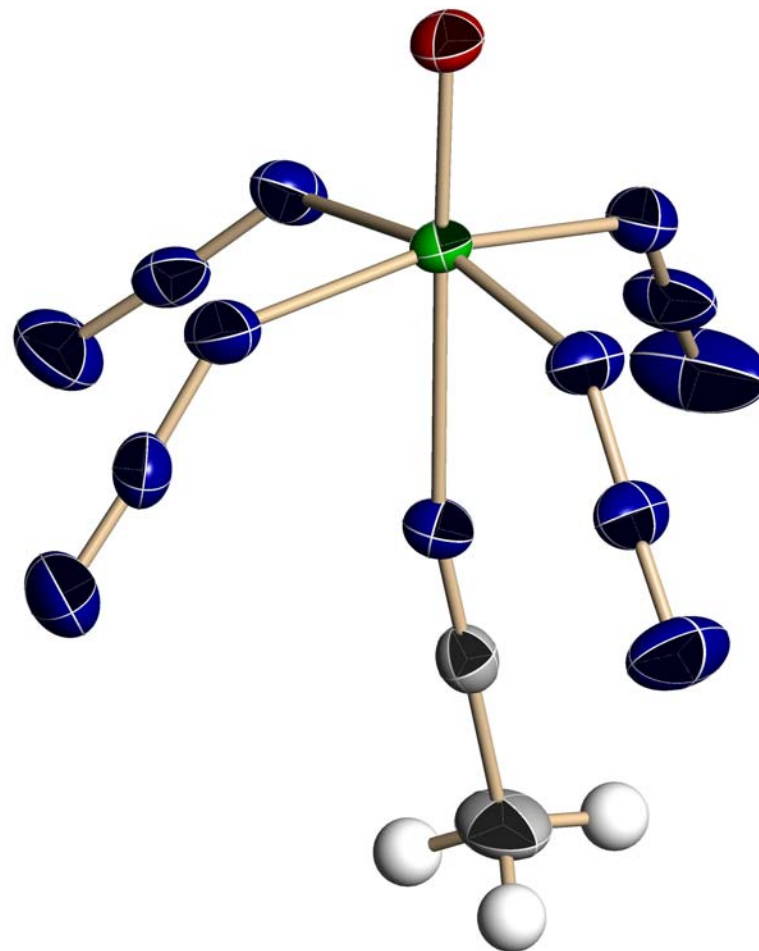
# Synthesis of Nitrido-Azides



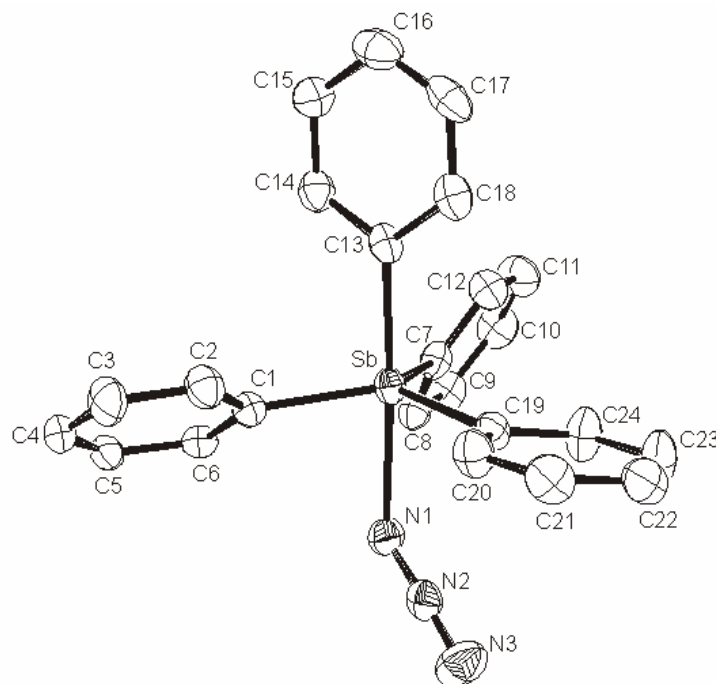
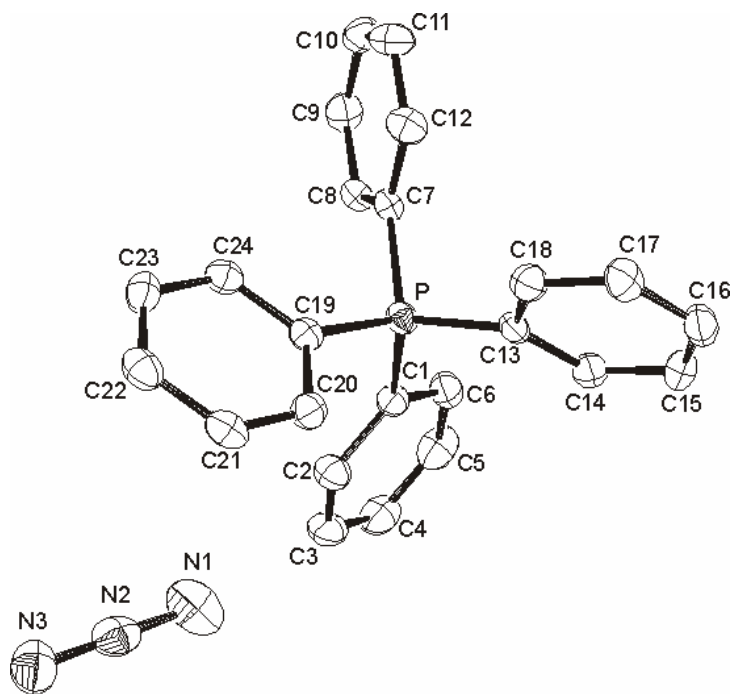
•Solid  $[\text{PPh}_4][\text{W}(\text{N}_3)_7]$  and  $[\text{PPh}_4][\text{Mo}(\text{N}_3)_7]$  are very sensitive compounds and decompose explosively. However, in solution at  $-30^\circ\text{C}$ , partial  $\text{N}_2$  evolution occurs with formation of nitrido-azides.



- $\text{WO}(\text{N}_3)_4$ ,  $[\text{cis-WO}_2(\text{N}_3)_4]^{2-}$ ,  $[\text{trans-WO}_2(\text{N}_3)_4]^{2-}$ ,  $[\text{MoO}(\text{N}_3)_5]^{2-}$ , and  $[\text{NbO}(\text{N}_3)_5]^{2-}$ , **the first examples of transition metal oxoazides**, were prepared and characterized by their crystal structures and vibrational spectroscopy
- $[\text{trans-UO}_2(\text{N}_3)_4]^{2-}$ , **the first example of an actinide oxoazide**, was also prepared and characterized by its crystal structure



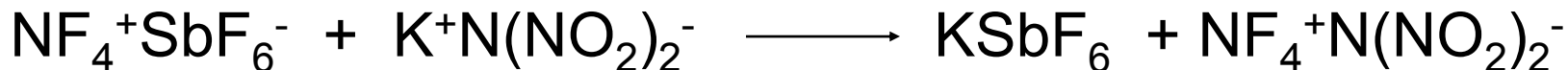
- Obtained crystal structures of  $M(\text{Ph})_4\text{N}_3$  where  $M = \text{P}, \text{As}, \text{Sb}$
- $\text{P}(\text{Ph})_4\text{N}_3$  and  $\text{As}(\text{Ph})_4\text{N}_3$  are ionic, while  $\text{Sb}(\text{Ph})_4\text{N}_3$  is covalent



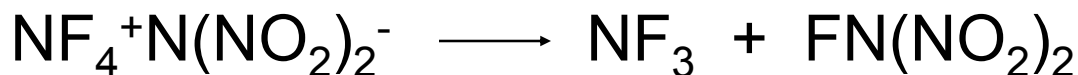
# Synthesis and Characterization of $\text{FN}(\text{NO}_2)_2$



- Prepared  $\text{NF}_4^+\text{N}(\text{NO}_2)_2^-$  at low temp in  $\text{SO}_2$  solution



and decomposed it according to:

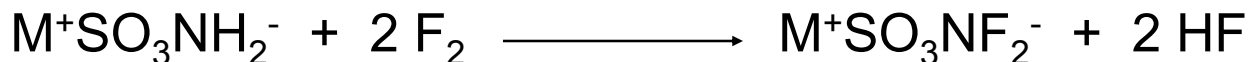


- $\text{FN}(\text{NO}_2)_2$  was characterized by low-temperature multinuclear NMR spectroscopy and is unstable at room temperature
- Efforts are in progress to isolate it in pure form

# $SO_3NF_2^-$ , a *Stable Difluoramine Source*

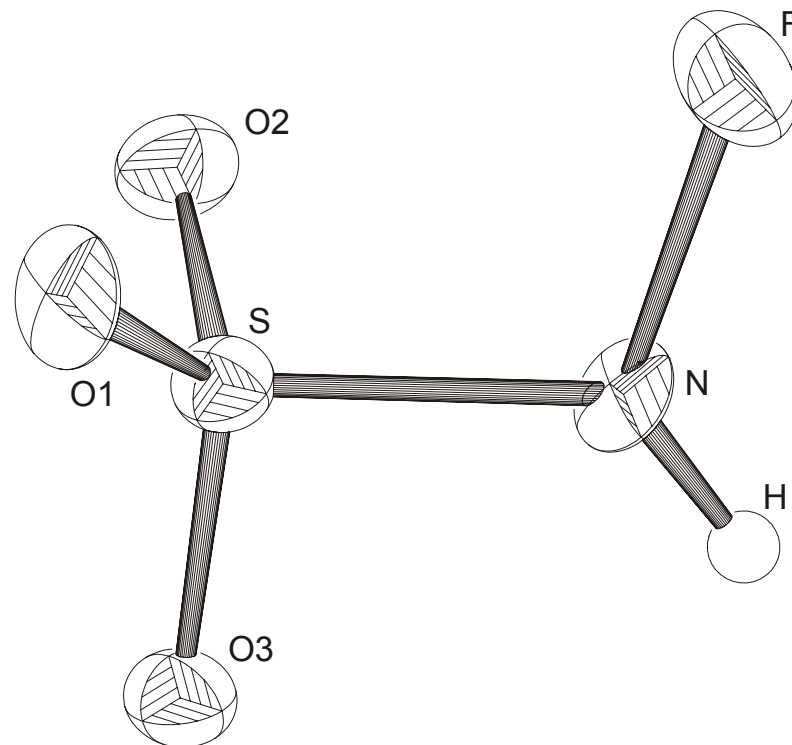
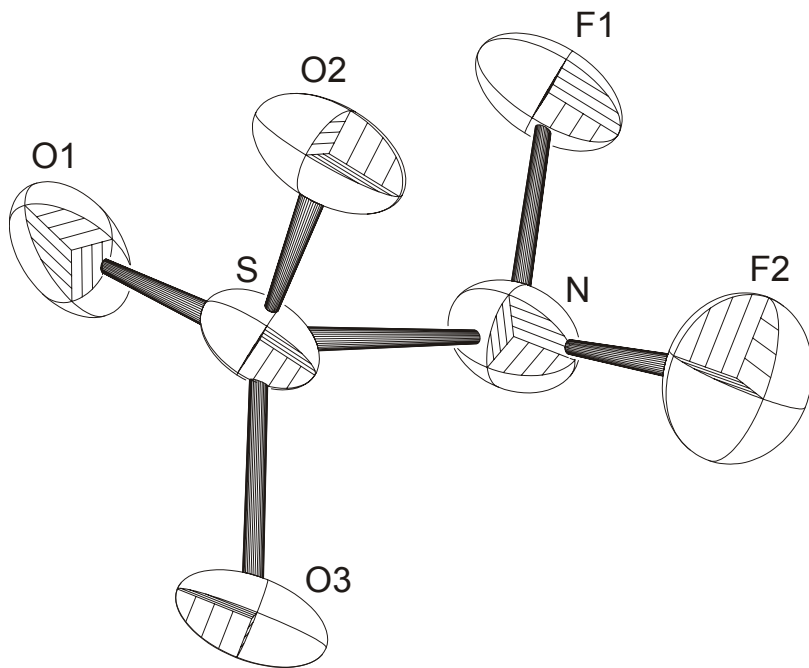


- Stable, safely storable sources for the generation of the highly explosive  $HNF_2$  are of great interest for the synthesis of *gem-bis-difluoramino compounds*.
- In previous work, we have demonstrated the potential of  $(C_6H_5)_3CNF_2$  for this application. However, the synthesis of  $(C_6H_5)_3CNF_2$  requires pressurizing  $N_2F_4$  in chlorobenzene solution and mercury and is cumbersome and dangerous.
- We have searched for a more convenient and accessible stable  $HNF_2$  source. We found that alkali metal salts of the previously unknown  $SO_3NF_2^-$  anion can be prepared by a simple, one-step, direct fluorination of the sulfamate anion in water



The  $SO_3NF_2^-$  and related  $SO_3NHF^-$  salts were characterized by their crystal structures, vibrational and NMR spectroscopy, and theoretical calculations, and *shown to be excellent stable reagents for converting carbonyl groups into gem-bis-NF<sub>2</sub> groups.*

# Crystal Structures of $\text{SO}_3\text{NF}_2^-$ and $\text{SO}_3\text{NH}^+$



# High-Oxygen Carriers and Ionic Liquid Propellants



- Ionic liquids have great potential for liquid propellants (Christe, Drake, pending US Patent).
- Because the bulky cations require large amounts of oxygen for complete combustion and high performance, there is a great need for high-oxygen carrying anions.
- We have investigated the usefulness of the tetranitrato-borate and tetranitrato-aluminate anions for these applications, and their ability to form room temperature ionic liquids.
- We have prepared and characterized numerous tetranitrato-borate salts. Although 1-ethyl-3-methyl imidazolium tetranitrato-borate forms a low freezing ( $-30\text{ }^{\circ}\text{C}$ ) ionic liquid, its thermal stability of  $60\text{ }^{\circ}\text{C}$  is insufficient for practical applications.

# High-Oxygen Carriers and Ionic Liquid Propellants



- Replacement of  $[B(NO_3)_4]^-$  by  $[Al(NO_3)_4]^-$  was highly successful.
- We prepared and characterized 1-ethyl-3-methyl imidazolium tetranitrato-aluminate:

Thermal stability (DSC):  $>200\text{ }^\circ\text{C}$

Freezing Point =  $-30\text{ }^\circ\text{C}$

$\rho = 1.508\text{ g/cm}^3$

Dropweight, negative at  $10\text{ kg}\cdot\text{cm}$

Friction,  $24\text{ kg}$

Calcd Isp =  $261\text{ sec}$

- We are presently preparing tetrazolium tetranitrato-aluminates with predicted Isp values of  $280\text{-}290\text{ sec}$ .

# Conclusions



- Excellent progress is being made in the area of polynitrogen chemistry ( $N_3^+$ ,  $N_3NOF^+$ ,  $N_7O^+$ ).
- A large number of novel polyazides, nitrido-azides and oxo-azides were prepared and characterized.
- A new NMR data processing method was developed and applied to  $N_5^+$  which allows the observation of  $^{15}N$  NMR spectra in natural abundance.
- The new  $FN(NO_2)_2$  molecule was synthesized and characterized.
- The novel  $M^+SO_3NF_2^-$  salts were synthesized and characterized, and their usefulness for preparing *gem*-bis-difluoramino compounds was demonstrated.
- High-oxygen carrying anions hold great promise for oxidizer-balanced, ionic liquid propellants.