

# REPORT DOCUMENTATION PAGE

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1. REPORT DATE (DD-MM-YYYY)		2. REPORT TYPE Technical Papers		3. DATES COVERED (From - To)	
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				5c. PROGRAM ELEMENT NUMBER	
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<b>20030130 152</b>					
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2303 M2 CP

MEMORANDUM FOR PRS (In-House/Contractor Publication)

FROM: PROI (STINFO)

19 Apr 2001

SUBJECT: Authorization for Release of Technical Information, Control Number: **AFRL-PR-ED-AB-2001-103**  
Jerry Boatz; Don Thompson and Dan Sorescu (Oklahoma State Univ.), "Bond Dissociation Energies of Energetic Compounds: A Comparison of Theoretical Methods"

**AFOSR Contractors Meeting**  
**(Irvine, CA, 21-23 May 2001) (Deadline: 18 May 01)**

**(Statement A)**

1. This request has been reviewed by the Foreign Disclosure Office for: a.) appropriateness of distribution statement, b.) military/national critical technology, c.) export controls or distribution restrictions, d.) appropriateness for release to a foreign nation, and e.) technical sensitivity and/or economic sensitivity.

Comments: \_\_\_\_\_  
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Signature \_\_\_\_\_ Date \_\_\_\_\_

2. This request has been reviewed by the Public Affairs Office for: a.) appropriateness for public release and/or b) possible higher headquarters review.

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3. This request has been reviewed by the STINFO for: a.) changes if approved as amended, b) appropriateness of references, if applicable; and c.) format and completion of meeting clearance form if required

Comments: \_\_\_\_\_  
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Comments: \_\_\_\_\_  
\_\_\_\_\_

APPROVED/APPROVED AS AMENDED/DISAPPROVED

\_\_\_\_\_  
PHILIP A. KESSEL Date  
Technical Advisor  
Space and Missile Propulsion Division



# **Bond Dissociation Energies of Energetic Compounds: A Comparison of Theoretical Methods**

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Stillwater, OK 74078**

**AFOSR Molecular Dynamics/Theoretical Chemistry Contractors Conference  
May 21-23, 2001  
Irvine, CA**



# Outline

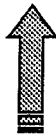
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- I. Overview of AFRL - Propulsion Directorate
  - High Energy Density Materials (HEDM) for rocket propulsion
  - Specific impulse as an assessment of energy density
  
- II. C-N bond energies of 1,1-diamino-2,2-dinitroethylene ("FOX-7") and prototypes
  
- III. Results
  
- IV. Summary and Conclusions



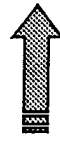
# Why is the Air Force interested in HEDM?

The performance limits of current propellants have been reached

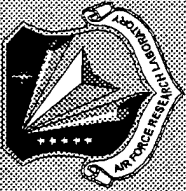


- The constituents of current propellants have been known for decades
- New missions require higher-performing propulsion systems

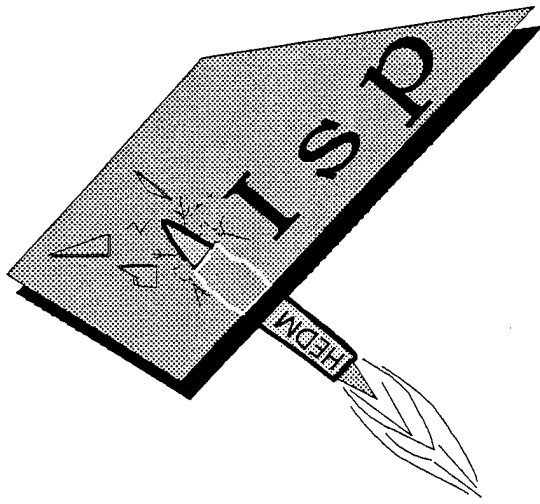
A revolutionary propulsion source would substantially improve our ability to access and exploit space



- Fusion, antimatter, and beamed energy are tantalizing but distant prospects
- Chemical propulsion will remain the method of choice for many applications
- Novel chemical propellants offer great potential for near-term improvements



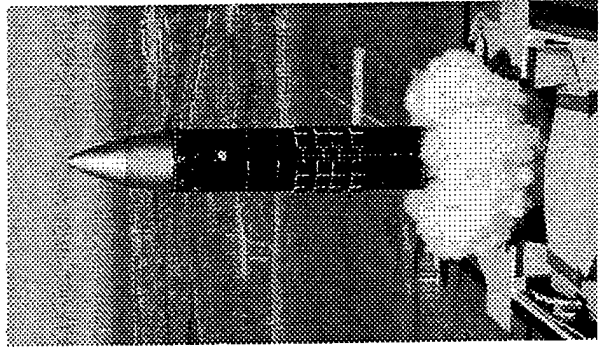
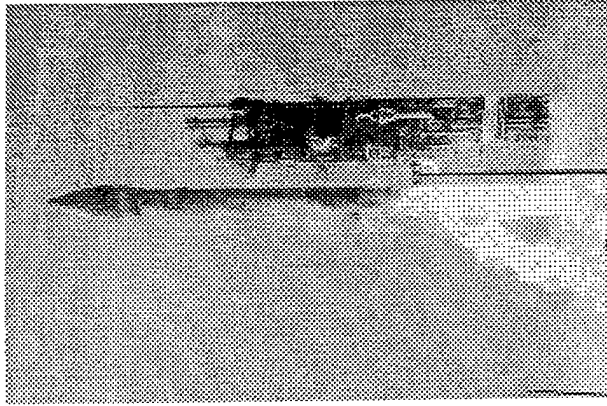
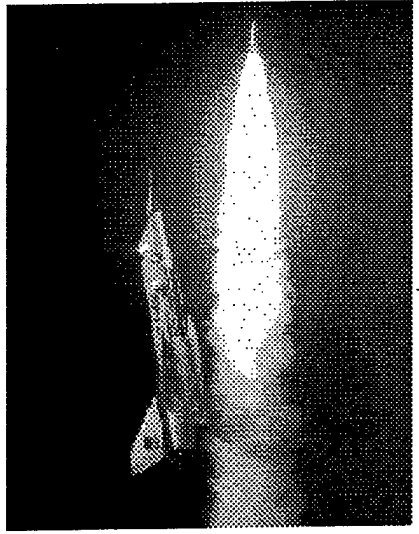
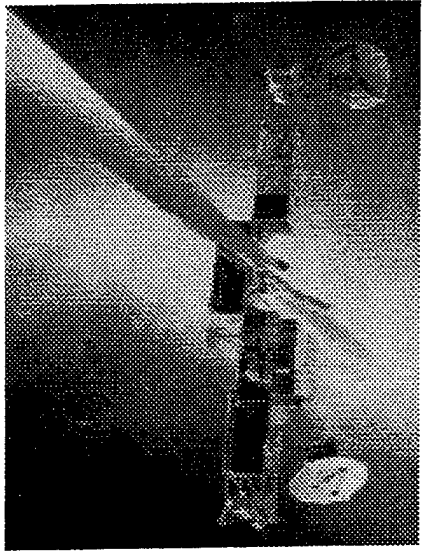
# HEDM Program Objective



**Identify and develop advanced chemical propellants for rocket propulsion applications**

- Hydrocarbons for liquid boosters
- Liquid & solid oxidizers for boost and upper stages
- Monopropellants for upper stages and satellites
- Cryogenic propellants for upper stages

*Breaking the performance barrier*





## Assessment of energy density: Specific Impulse ( $I_{sp}$ )

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$I_{sp}$  = pounds of delivered thrust / (pounds of propellant burned/second); similar to mpg.

$$I_{sp} \propto \sqrt{\frac{\Delta H_{comb}}{m_{products}}}$$

=> highly exothermic reactions AND combustion products with small masses are required  
(  $\approx$  50% of liquid  $H_2$  in SSME is not burned!)



# Specific impulse values of currently used propellants

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Solid propellant: Ammonium perchlorate(AP)/Al powder/hydroxy-terminated polybutadiene (HTPB)

$$I_{sp} = 267 \text{ sec}$$

Liquid propellant: RP-1/LOX

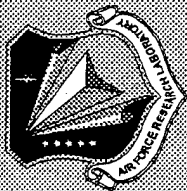
$$I_{sp} = 300 \text{ sec}$$

Monopropellant: Hydrazine ( $N_2H_4$ )

$$I_{sp} = 230 \text{ sec}$$

Cryogenic propellants:  $LH_2/LOX$

$$I_{sp} = 390 \text{ sec}$$



# HEDM Propellant Payoffs

- Larger payloads, smaller vehicles, and lower launch costs
- Greater capability to access and exploit space

Vehicle Type	Baseline Vehicle	Propellant	Takeoff Mass (lb)	Payload Mass (lb)	Payload Mass (lb) With 10% Isp Increase
Two-stage ELV	Atlas II // Centaur D-1A	RP-1/LOX (Isp = 295 s) // LH2/LOX (Isp = 455 s)	360,000	12,500	15,600 (+25%)
SSTO RLV	Lockheed SSTO	LH2/LOX (Isp = 455 s)	1,900,000	40,000	68,000 (+70%)
Missile Defense Interceptor	Boost-Phase Interceptor	HTPB/Al/HMX (Isp = 270 s)	1,847	74	110 (+49%)

**Our research is aimed at increasing propellant Isp by 5 to 50%**



## **Computational chemistry plays an important role in identification and characterization of HEDM**

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**Experimental synthesis and characterization is difficult**

- Little or no intuition to guide synthesis of new molecules.
- Energetic compounds are thermodynamically (and often kinetically) unstable.
- Synthesis is time-consuming, expensive, high-risk.



## **Energetic compounds present several challenges to theory**

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### **Exotic electronic structures**

- Large non-dynamical correlation effects => single configuration methods may not be applicable.
- Multiple potential energy surfaces (e.g., low-lying triplet state(s) often intersect the lowest singlet state.)
- Nonadiabatic interactions (e.g., spin-orbit coupling, radiationless transitions)



**Which theoretical method(s) give reliable predictions?**

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### **Density Functional Theory**

- Widely used due to its efficiency and accuracy (generally comparable to MP2.)

### **Single-configuration correlated methods**

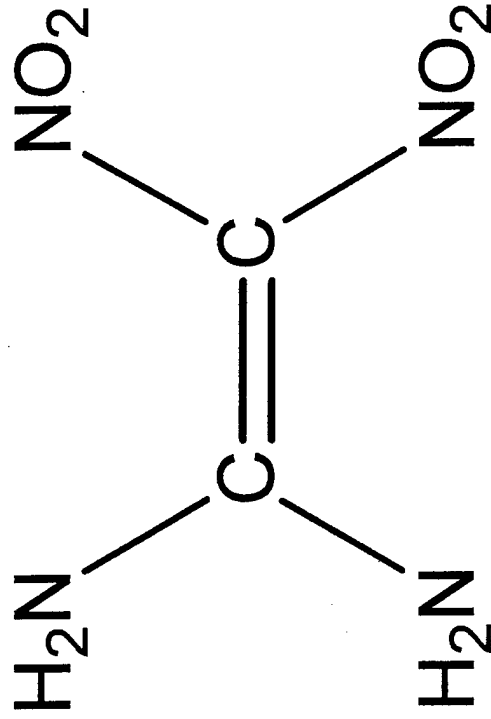
- MPn, CC, QCI, G2

### **Multiconfigurational methods**

- MCSCF, MCQDPT, MRCI, MRCC



# FOX-7: A prototypical energetic compound



## Advantages:

- Chemically balanced wrt decomposition products ( $2\text{CO} + 2\text{H}_2\text{O} + 2\text{N}_2$ )
- Lower impact/shock sensitivity than other  $\text{C}_n\text{H}_{2n}\text{O}_{2n}\text{N}_{2n}$  compounds (e.g., RDX and HMX).

$I_{sp} = 254 \text{ sec}$  (calculated)  
 $\Delta H_f = -9.5 \text{ kcal/mol}$  (G2(MP2))



## Recent Studies of FOX-7

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### Experimental X-ray structure

U. Bemm and H. Östmark, Acta Cryst. C54, 1997(1998).

### Structures and C-N bond energies (B3P86/6-31+G(d,p))

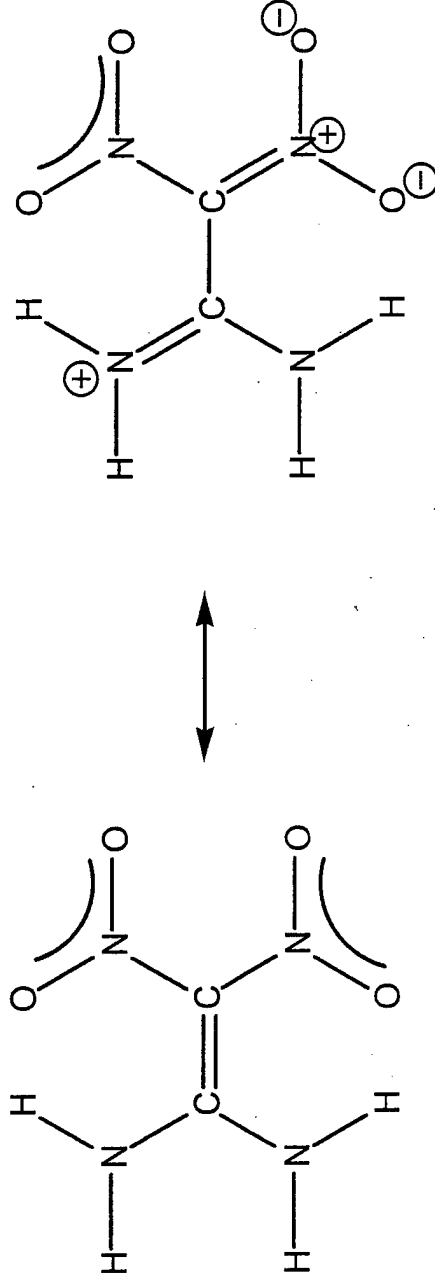
P.Politzer, M.C.Concha, M.E.Grice, J.S.Murray, P.Lane, and  
D.Habibollazadeh Theochem, 452, 75(1998)

### Decomposition mechanisms (B3P86/6-31+G(d,p), B3LYP/6-31+G(d,p))

A.Gindulyte, L.Massa, L.Huang, and J.Karl, J. Phys. Chem. A, 103,  
11045(1999)



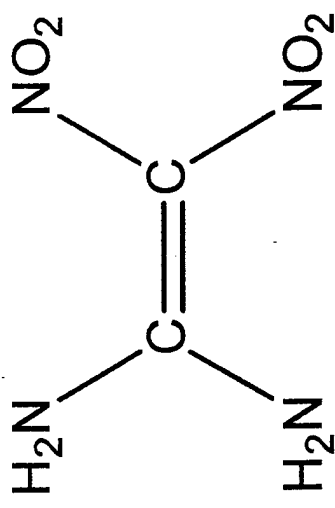
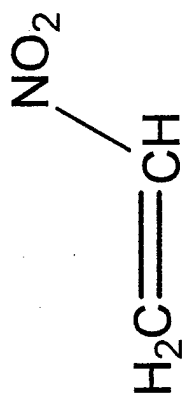
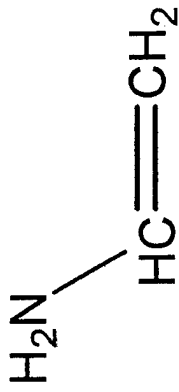
# FOX-7: A "Push-Pull" Ethylene



**Multiple Lewis structures suggest that FOX-7 may have high degree of multiconfigurational character => good testbed for SOTA single-reference methods and DFT.**

# DFT Geometries

C=C	C-NH <sub>2</sub>	C-NO <sub>2</sub>
1.331*		
1.327		
1.340*	1.386*	
1.336	1.392	
1.326*		1.459*
1.322		1.473
1.426*	1.339*	1.424*
1.428**	1.345**	1.432**
1.421	1.345	1.438
(1.456)	(1.319, 1.325)	(1.399, 1.426)



\* P. Politzer, M.C. Concha, M.E. Grice, J.S. Murray, P. Lane, and D. Habibollahzadeh Theochem, 452, 75(1998)

\*\* A. Gindulyte, L. Massa, L. Huang, and J. Karl, J. Phys. Chem. A, 103, 11045(1999)

() U. Bemm and H. Östmark, Acta Cryst. C54, 1997(1998).



# Approach

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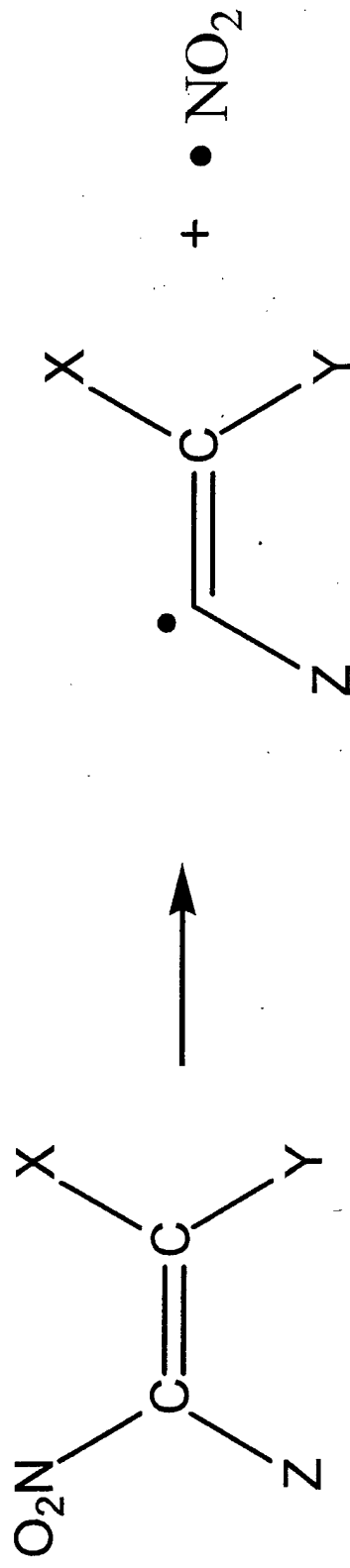
- 1. Compute C-NH<sub>2</sub> and C-NO<sub>2</sub> bond dissociation energies of FOX-7 and simpler prototypes using DFT (B3LYP), single-reference methods (MP2, G2(MP2), CCSDT//MP2), and a multireference method (MCQDPT(2)//CASCF). 6-311G(d,p) used throughout.**
- 2. Assess degree of multiconfigurational character via calculation of natural orbital occupation numbers (MP2, CCSD(T), MCSCF).**
  - MP2 and CCSD(T) “non-physical” occupation numbers indicative of significant degree of multiconfigurational character**

M.S. Gordon, M.W. Schmidt, G.M. Chaban, K. R. Glaesemann, W.J. Stevens, and C. Gonzalez, J. Chem. Phys., 110, 4199 (1999).

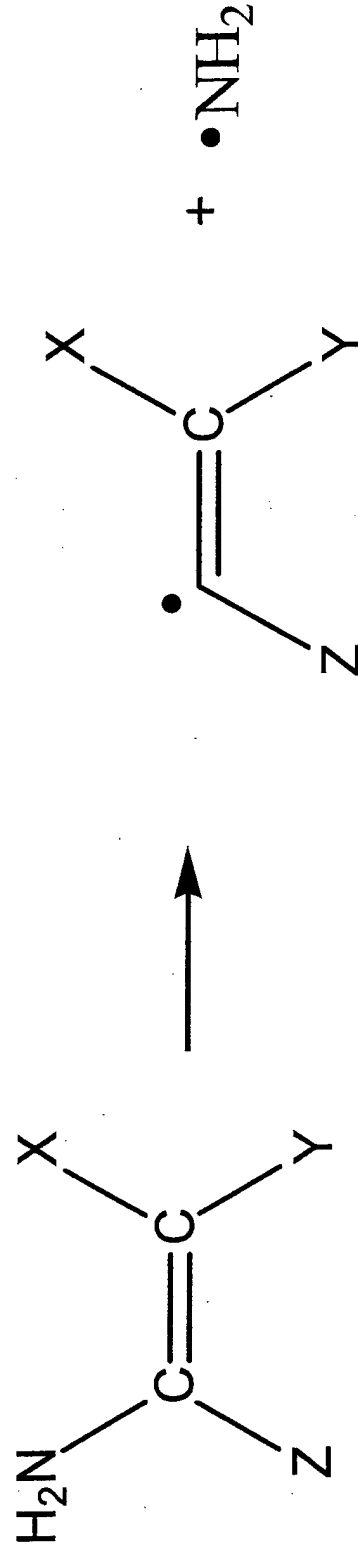
# Bond Dissociation Reactions

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**C-NO<sub>2</sub> BDEs:** X, Y = H, NH<sub>2</sub> ; Z = H, NO<sub>2</sub>



**C-NH<sub>2</sub> BDEs:** X, Y = H, NO<sub>2</sub> ; Z = H, NH<sub>2</sub>



# Choice of CASSCF Active Space

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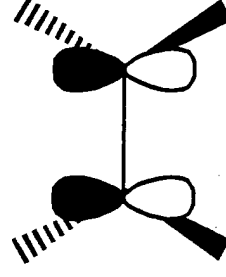
“Push-pull” Lewis structures suggest that delocalization of  $\pi$  electrons account for the most important non-dynamical correlation.



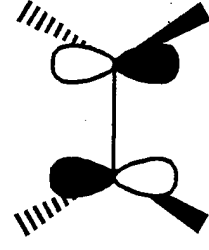
$\pi$

non-bonding

$\pi^*$



$\pi$



$\pi^*$



$\pi$

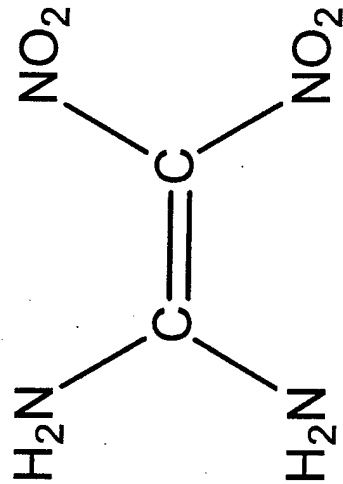


$\sigma$



$\sigma^*$

# Structures of FOX-7



\* P. Politzer, M.C. Concha, M.E. Grice, J.S. Murray, P. Lane, and D. Habibollazadeh *Theochem*, 452, 75(1998)

\*\* A. Gindulyte, L. Massa, L. Huang, and J. Karl, *J. Phys. Chem. A*, 103, 11045(1999)

( ) U. Bemm and H. Östmark, *Acta Cryst. C*54, 1997(1998).

## Level of theory

B3P86/6-31+G(d,p)

B3LYP/6-31+G(d,p)

B3LYP/6-311G(d,p)

MP2/6-311G(d,p)

CASSCF/6-311G(d,p)

## C=C

1.426\*

1.428\*\*

1.421

1.392

## C-NH<sub>2</sub>

1.339\*

1.345\*\*

1.345

1.359

## C-NO<sub>2</sub>

1.424\*

1.432\*\*

1.438

1.441

## X-ray structure

(1.456)

(1.319)

(1.325)

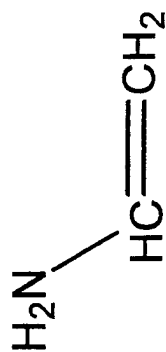
(1.399)

(1.426)

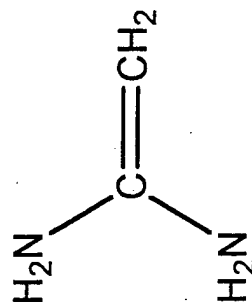
# Natural Orbital Occupation Numbers

(closed shell species)

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MP2: none  
CCSD(T): none



MP2: -0.001  
CCSD(T): none

<----- MCSCF (6e,5o) ----->

C=C	$\pi$	$\pi^*$
	1.927	0.079
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$
	1.980	0.020
NH <sub>2</sub> lp	1.994	

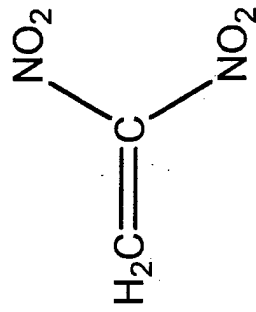
<----- MCSCF (8e,7o) ----->

C=C	$\pi$	$\pi^*$
	1.940	0.068
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$
	1.978	0.023
	1.981	0.019
NH <sub>2</sub> lp	1.995	

# Natural Orbital Occupation Numbers

(closed shell species)

---



MP2: 2.00001,  
-0.00009  
CCSD(T): none

MP2: -0.001  
CCSD(T): none

<----- MCSCF (8e,7o) ----->

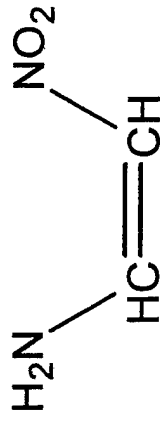
C=C	$\pi$	$\pi^*$	
	1.914	0.077	
NO <sub>2</sub>	$\pi$	n.b.	$\pi^*$
	1.983	1.897	0.130
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.977	0.023	

<----- MCSCF (14e,12o) ----->

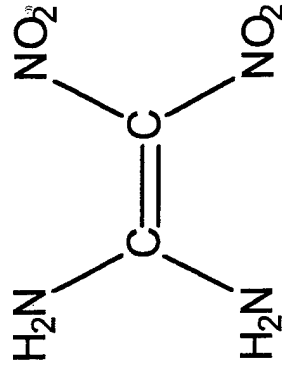
C=C	$\pi$	$\pi^*$	
	1.915	0.077	
NO <sub>2</sub>	$\pi$	n.b.	$\pi^*$
	1.984	1.889	0.132
	1.984	1.883	0.137
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.977	0.030	
	1.973	0.020	

# Natural Orbital Occupation Numbers

(closed shell species)



MP2: -0.00001(2)  
CCSD(T): none



MP2: 2.00001(3),  
-0.00016, -0.00009,  
-0.00004, -0.00001  
CCSD(T): none

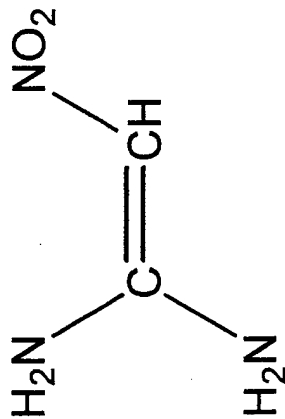
<----- MCSCF (12e,10o) ----->

C=C	π	π*	
	1.950	0.059	
NO <sub>2</sub>	π	n.b.	π*
	1.983	1.914	0.108
C-NH <sub>2</sub>	σ	σ*	
	1.981	0.019	
C-NO <sub>2</sub>	σ	σ*	
	1.978	0.023	
NH <sub>2</sub> lp	1.987		

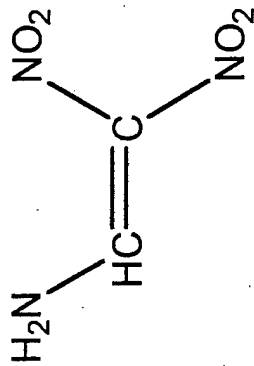
<----- MCSCF (18e,14o) ----->

C=C	π	π*	
	1.983	0.036	
NO <sub>2</sub>	π	n.b.	π*
	1.986	1.911	0.099
	1.989	1.906	0.113
C-NH <sub>2</sub>	σ	σ*	
	1.982	0.018	
	1.978	0.022	
C-NO <sub>2</sub>	σ	σ*	
	----- n/a -----		
NH <sub>2</sub> lp	1.988(2)		

# Natural Orbital Occupation Numbers (closed shell species)



MP2: -0.0001(2)  
CCSD(T): none



MP2: -0.0002(2),  
-0.00001  
CCSD(T): none

<----- MCSCF (16e,13o) ----->

C=C	$\pi$	$\pi^*$	
	1.962	0.050	
NO <sub>2</sub>	$\pi$	n.b.	$\pi^*$
	1.983	1.921	0.100
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.982	0.018	
	1.977	0.022	
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.979	0.024	
NH <sub>2</sub> lp	1.994		
	1.990		

<----- MCSCF (18e,15o) ----->

C=C	$\pi$	$\pi^*$	
		TBD	
NO <sub>2</sub>	$\pi$	n.b.	$\pi^*$
		TBD	
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$	
		TBD	
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$	
		TBD	
NH <sub>2</sub> lp		TBD	

# Natural Orbital Occupation Numbers

(open shell species)

---



MP2: none  
CCSD(T): none

<----- MCSCF (3e,2o) ----->  
NH<sub>2</sub> lp 2.000  
N rad. 1.000



MP2: 2.00001  
CCSD(T): none

<----- MCSCF (5e,4o) ----->  
NO<sub>2</sub> π n.b. π\*  
1.986 1.937 0.078  
N rad. 1.000

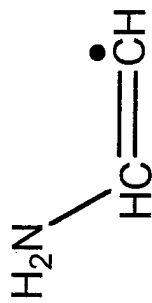


MP2: 2.07575,  
-0.08353

CCSD(T): none

<----- MCSCF ----->  
C=C π π\*  
1.899 0.101  
C rad. 1.000

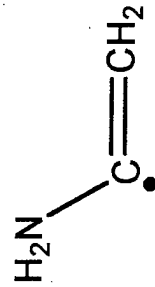
# Natural Orbital Occupation Numbers (open shell species)



MP2: 2.06629,

-0.08353

CCSD(T): none



MP2: TBD

CCSD(T): TBD

<----- MCSCF (7e,60) ----->

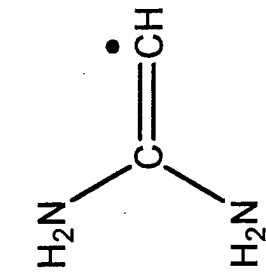
C=C	$\pi$	$\pi^*$	
	1.911	0.094	
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.980	0.021	
NH <sub>2</sub> lp	1.995		
C rad.	1.000		

<----- MCSCF (7e,60) ----->

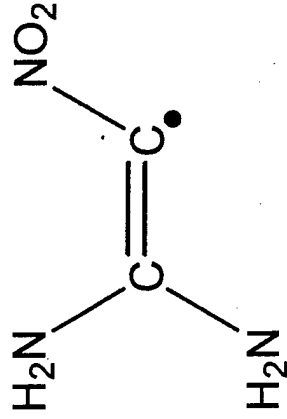
C=C	$\pi$	$\pi^*$	
	1.918	0.084	
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.981	0.019	
NH <sub>2</sub> lp	1.993		
C rad.	1.006		

# Natural Orbital Occupation Numbers

(open shell species)



MP2: 2.04703,  
2.00001, -0.06203,  
-0.00003  
CCSD(T): none



MP2: TBD  
CCSD(T): TBD

<----- MCSCF (11e,90) ----->

C=C	$\pi$	$\pi^*$	1.924	0.083
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$	1.977	0.023
			1.980	0.020
NH <sub>2</sub> lp			1.994	
			1.997	
C rad.			1.001	

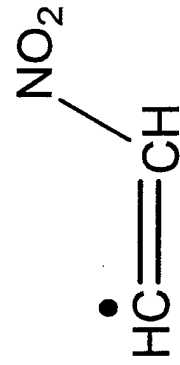
<----- MCSCF (13e,120) ----->

C=C	$\pi$	$\pi^*$	1.929	0.068
NO <sub>2</sub>	$\pi$	n.b.	1.9834	1.907
		$\pi^*$		0.114
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$	1.982	0.017
			1.980	0.020
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$	1.977	0.024
NH <sub>2</sub> lp			---	n/a

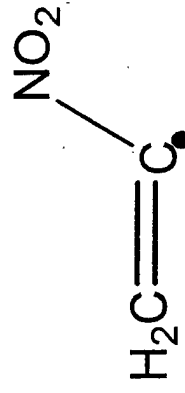
# Natural Orbital Occupation Numbers

(open shell species)

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**MP2:** 2.08805,  
 2.00001, -0.00001,  
 -0.00006  
**CCSD(T):** none



**MP2:** 2.09302,  
 2.00001, -0.00008,  
 -0.10792  
**CCSD(T):** TBD

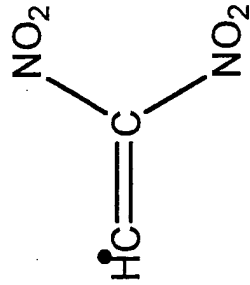
<----- MCSCF (9e,8o) ----->

C=C	$\pi$	$\pi^*$		
	1.904	0.083		
NO <sub>2</sub>	$\pi$	n.b.	$\pi^*$	
	1.983	1.900	0.134	
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$		
	1.975	0.025		
C rad.	1.000			

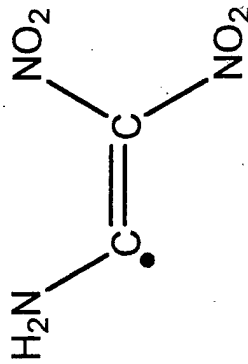
<----- MCSCF (9e,8o) ----->

C=C	$\pi$	$\pi^*$		
	1.896	0.090		
NO <sub>2</sub>	$\pi$	n.b.	$\pi^*$	
	1.983	1.891	0.141	
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$		
	1.977	0.023		
C rad.	1.000			

# Natural Orbital Occupation Numbers (open shell species)



MP2: 2.08603,  
2.00001(2),  
-0.10207, -0.00003  
CCSD(T): TBD



MP2: 2.00001(2),  
-0.0013, -0.00055  
CCSD(T): none

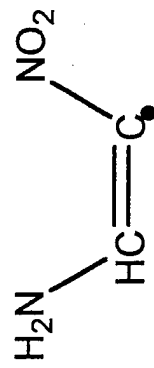
<----- MCSCF (15e,13o) ----->

C=C	$\pi$	$\pi^*$	
	1.904	0.084	
NO <sub>2</sub>	$\pi$	n.b.	$\pi^*$
	1.984	1.882	0.142
	1.984	1.888	0.133
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.972	0.031	
	1.976	0.021	
C rad.		1.000	

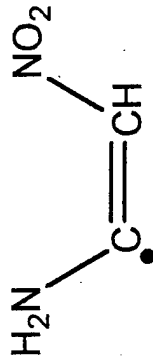
<----- MCSCF (15e,12o) ----->

C=C	$\pi$	$\pi^*$	
	1.976	0.043	
NO <sub>2</sub>	$\pi$	n.b.	$\pi^*$
	1.989	1.905	0.114
	1.987	1.909	0.103
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$	
	-----	n/a	-----
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.981	0.020	
NH <sub>2</sub> lp		1.974	
C rad.		1.000	

# Natural Orbital Occupation Numbers (open shell species)



MP2: 2.05145,  
2.00001, -0.00014,  
-0.06755  
CCSD(T): TBD



MP2: 2.00001,  
-0.00002, -0.00015,  
-0.01124  
CCSD(T): TBD

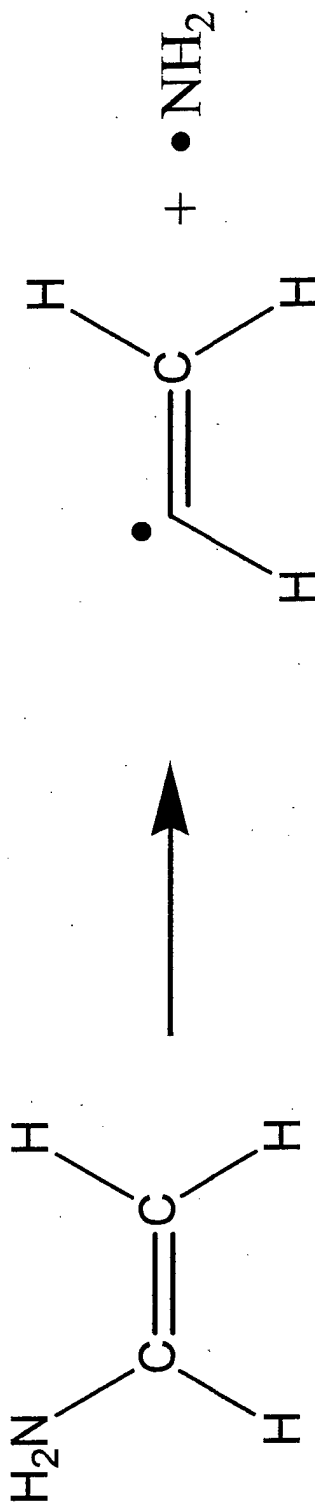
<----- MCSCF (13e,11o) ----->

C=C	$\pi$	$\pi^*$	
	1.923	0.079	
NO <sub>2</sub>	$\pi$	n.b.	$\pi^*$
	1.983	1.906	0.119
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.977	0.024	
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.979	0.019	
NH <sub>2</sub> lp		1.990	
C rad.		1.001	

<----- MCSCF (13e,11o) ----->

C=C	$\pi$	$\pi^*$	
	1.938	0.064	
NO <sub>2</sub>	$\pi$	n.b.	$\pi^*$
	1.982	1.911	0.112
C-NO <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.976	0.025	
C-NH <sub>2</sub>	$\sigma$	$\sigma^*$	
	1.980	0.019	
NH <sub>2</sub> lp		1.987	
C rad.		1.006	

# C-NH<sub>2</sub> Bond Dissociation Energies (kcal/mol)



DFT	105.2
	109.8 (103.3)*
MBPT(2)	115.6
G2(MP2)	110.8 (102.4)
CCSD(T)//MBPT(2)	104.7
MCQDPT(2)//MCSCF	101.9
experiment <sup>†</sup>	102

----- Non-physical NOONS -----  
 MBPT(2) CCSD(T) none n/a  
 ----- MCSCF -----

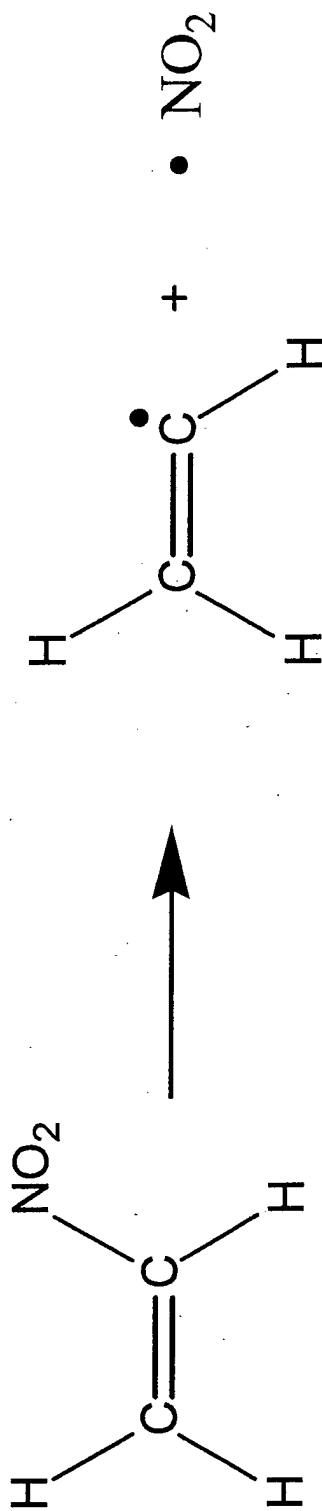
•CH=CH<sub>2</sub>  
 ----- Non-physical NOONS -----  
 MBPT(2) CCSD(T) 2.07575 none  
 -0.09409  
 ----- MCSCF(3e,3o) -----  
 C=C π 1.899 π\* 0.101  
 C rad. 1.000

NH<sub>2</sub>CH=CH<sub>2</sub>  
 ----- Non-physical NOONS -----  
 MBPT(2) CCSD(T) none none  
 ----- MCSCF(6e,5o) -----  
 C=C π 1.927 π\* 0.079  
 NH<sub>2</sub> Ip 1.994  
 C-NH<sub>2</sub> σ 1.980 σ\* 0.020

\* P. Politzer, M.C. Concha, M.E. Grice, J.S. Murray, P. Lane, and D. Habibollazadeh Theochem, 452, 75(1998). Values in () include ZPE corrections.

† S.G. Lias, J.E. Bartmess, J.F. Liebman, J.L. Holmes, R.D. Levin, W.G. Mallard, J. Phys. Chem. Ref. Data 17 (Suppl. 1), 1988.

# C-NO<sub>2</sub> Bond Dissociation Energies (kcal/mol)



DFT	68.8
	74.5*
MBPT(2)	77.7
G2(MP2) <sup>†</sup>	79.4
CCSD(T)//MBPT2	70.9
MCQDPT(2)//MCSCF	59.5

----- Non-physical NOONS -----	•NO <sub>2</sub>	----- MCSCF(5e,4o) -----
MBPT(2) CCSD(T)	NO <sub>2</sub> π n.b. π*	
2.00001 none	1.986 1.937 0.078	
	N rad. 1.000	

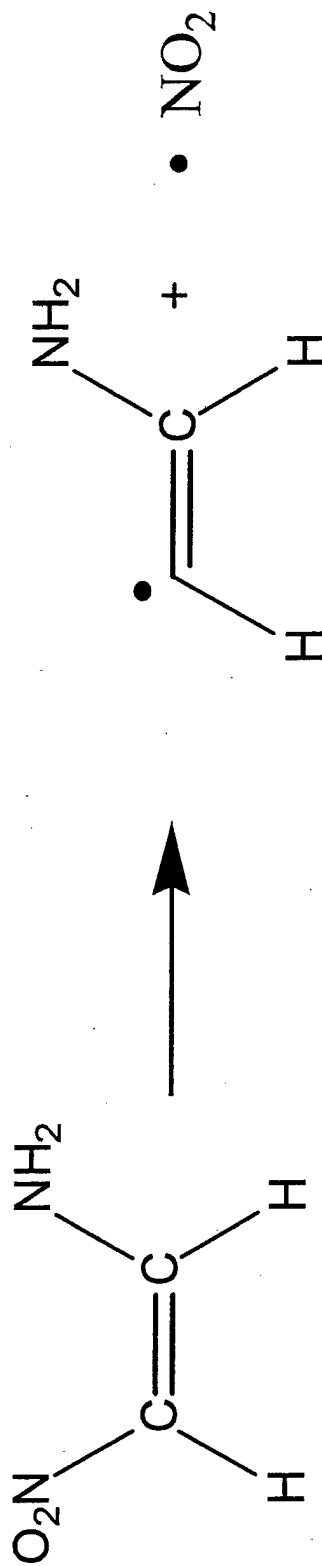
----- Non-physical NOONS -----	•CH=CH <sub>2</sub>	----- MCSCF(3e,3o) -----
MBPT(2) CCSD(T)	C=C π π*	
2.07575 none	1.899 0.101	
-0.09409	C rad. 1.000	

----- Non-physical NOONS -----	NO <sub>2</sub> CH=CH <sub>2</sub>	----- MCSCF(8e,7o) -----
MBPT(2) CCSD(T)	C=C π π*	
2.00001 none	1.914 0.077	
-0.00009	NO <sub>2</sub> π n.b. π*	
-0.00004	1.983 1.897 0.130	
	C-NO <sub>2</sub> σ σ*	
	1.977 0.023	

\* P. Politzer, M.C. Concha, M.E. Grice, J.S. Murray, P. Lane, and D. Habibollahzadeh Theochem, 452, 75(1998).

<sup>†</sup> Does not include ZPE corrections.

# C-NO<sub>2</sub> Bond Dissociation Energies (kcal/mol)



DFT	81.4
	88.0*
MBPT(2)	86.2
G2(MP2) <sup>†</sup>	88.8
CCSD(T)//MBPT2	78.2
MCQDPT(2)//MCSCF	66.7

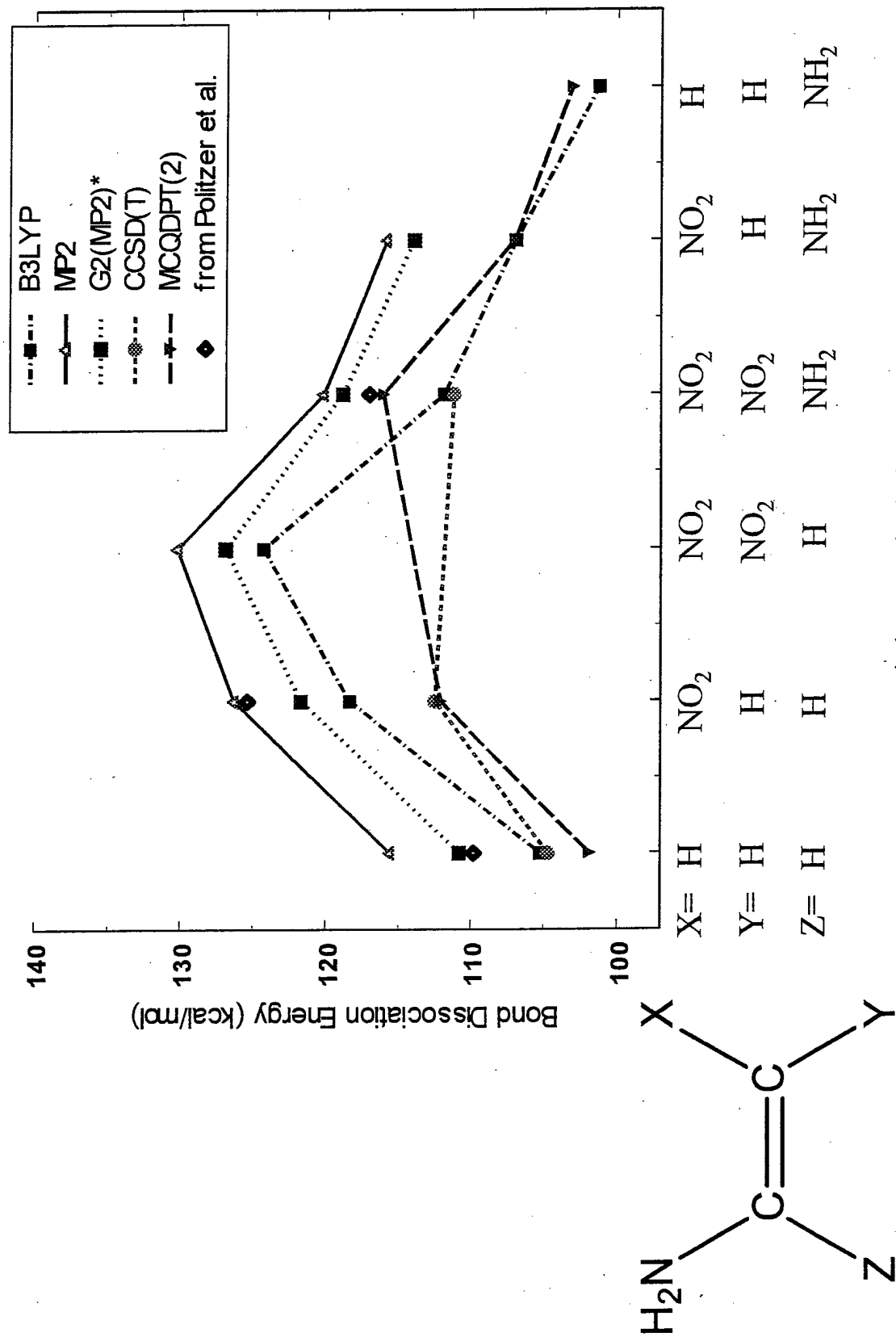
----- Non-physical NOONS -----	•NO <sub>2</sub>	----- MCSCF(5e,4o) -----
MBPT(2) CCSD(T)	NO <sub>2</sub> π	n.b. π*
2.00001	none	1.986 1.937 0.078
		N rad. 1.000
----- Non-physical NOONS -----	•CH=CH-NH <sub>2</sub>	----- MCSCF(7e,6o) -----
MBPT(2) CCSD(T)	C=C	π π*
2.06629	none	1.911 0.094
-0.08353		C rad. 1.000
-0.00001		NH <sub>2</sub> lp 1.995

\* P. Politzer, M.C. Concha, M.E. Grice, J.S. Murray, P. Lane, and D. Habibollahzadeh Theochem, 452, 75(1998).

<sup>†</sup> Does not include ZPE corrections.

----- Non-physical NOONS -----	NO <sub>2</sub> CH=CH-NH <sub>2</sub>	----- MCSCF(12e,10o) -----
MBPT(2) CCSD(T)	C=C	π π*
2.00001	TBD	1.950 0.059
-0.00009		NO <sub>2</sub> π n.b. π*
-0.00004		1.983 1.914 0.108
		NH <sub>2</sub> lp 1.987

# C-NH<sub>2</sub> Bond Dissociation Energies (kcal/mol)



\* does not include ZPE corrections



# Summary and Conclusions

1. The C-NH<sub>2</sub> and C-NO<sub>2</sub> bond dissociation energies of FOX-7 and simpler prototypes have been computed using DFT (B3LYP), single-reference (SR) methods (MP2, G2(MP2), CCSD(T)//MP2), and a multireference (MR) method (MCQDPT(2)//CASSCF).
2. With the exception of aminoethylene and amino radical, all of the closed shell and radical species considered in this study have non-physical NOONs at the MP2 level.
3. CCSD(T) is better able to "capture" non-dynamical correlation than MP2.
4. Species containing a nitro group generally have a higher degree of multiconfigurational character than those without NO<sub>2</sub>.
5. The DFT BDEs generally are in better agreement with the SR methods (MP2, G2(MP2), CCSD(T)) than with MCQDPT(2)), particularly for C-NO<sub>2</sub>.
6. The MCQDPT(2) BDEs are lower than those of the SR methods. The difference between the SR and MR predictions is greater for C-NO<sub>2</sub> than C-NH<sub>2</sub>.
7. The most stringent comparison of these methods will likely be the C-NO<sub>2</sub> BDE for (NH<sub>2</sub>)<sub>2</sub>C=C(NO<sub>2</sub>)<sub>2</sub>, for which the radical (NH<sub>2</sub>)<sub>2</sub>C=C-NO<sub>2</sub> is exceptionally MR in nature.



# Future Directions

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1. Include additional MR methods (MRCI, MRCC).
2. For the smaller systems, try larger basis sets (cc-pVTZ).
3. Replace UHF reference with ROHF for MP2, CCSD(T).

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AFOSR

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