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TSAPS Fall 2003

**Results for Positron-Acetylene Scattering
Using the Distributed Positron Model**

**Thomas L. Gibson — Texas Tech University
and**

Patrick J. Nichols — University of New Orleans

Special thanks to **Mr. Lee Burnside** for cluster construction and maintenance.

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Abstract for TSAPS 2003

Results for Positron-Acetylene Scattering Using the Distributed Positron Model,
THOMAS L. GIBSON, Texas Tech University, and PATRICK J. NICHOLS, University of New Orleans — The Distributed Positron Model (DPM) has been developed as a means to include the correlation/polarization interaction between a slow positron and an atomic or molecular target. This technique is based on a quantum chemistry approach that approximates the effect of virtual positronium formation near the target and automatically reduces to the correct long-range form of the polarization potential far from the target. Further, the codes that implement our DPM method have been written to run on low-cost concurrent computers known as GNU/Linux Beowulf clusters. In this talk we will report integrated and differential cross sections for low-energy positron-acetylene collisions.

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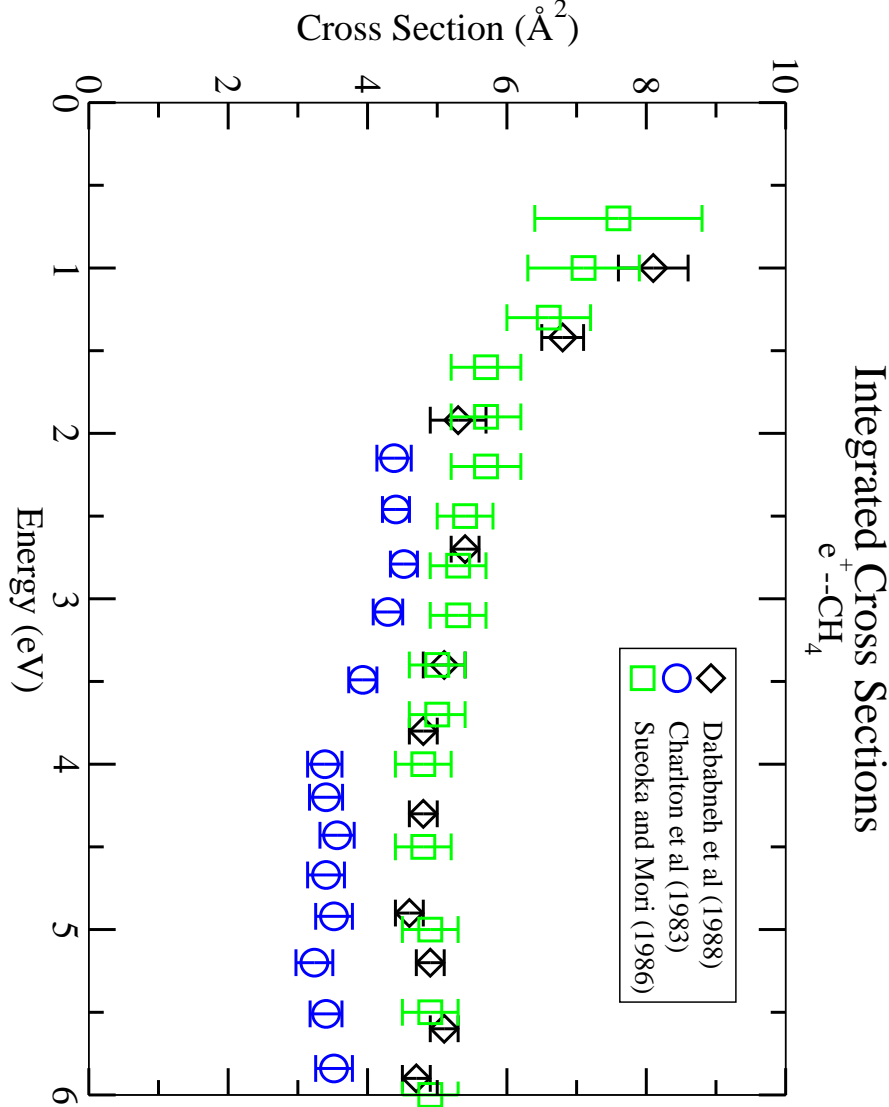
Motivation

Why calculate low-energy positron-molecule collisions?

- Very few measured results available
 - Intense low-energy beams are hard to achieve
- Existing experiments don't agree
- Most theories too parameterized or intractable for larger systems

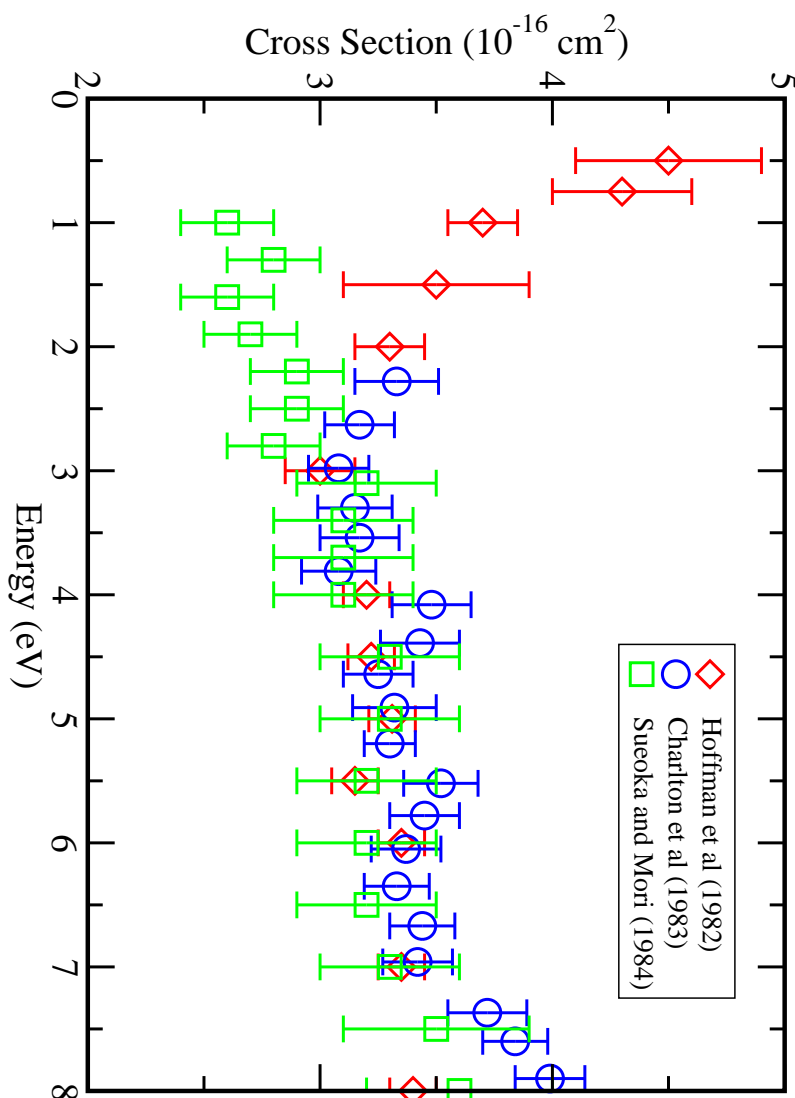
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Positron-CH₄



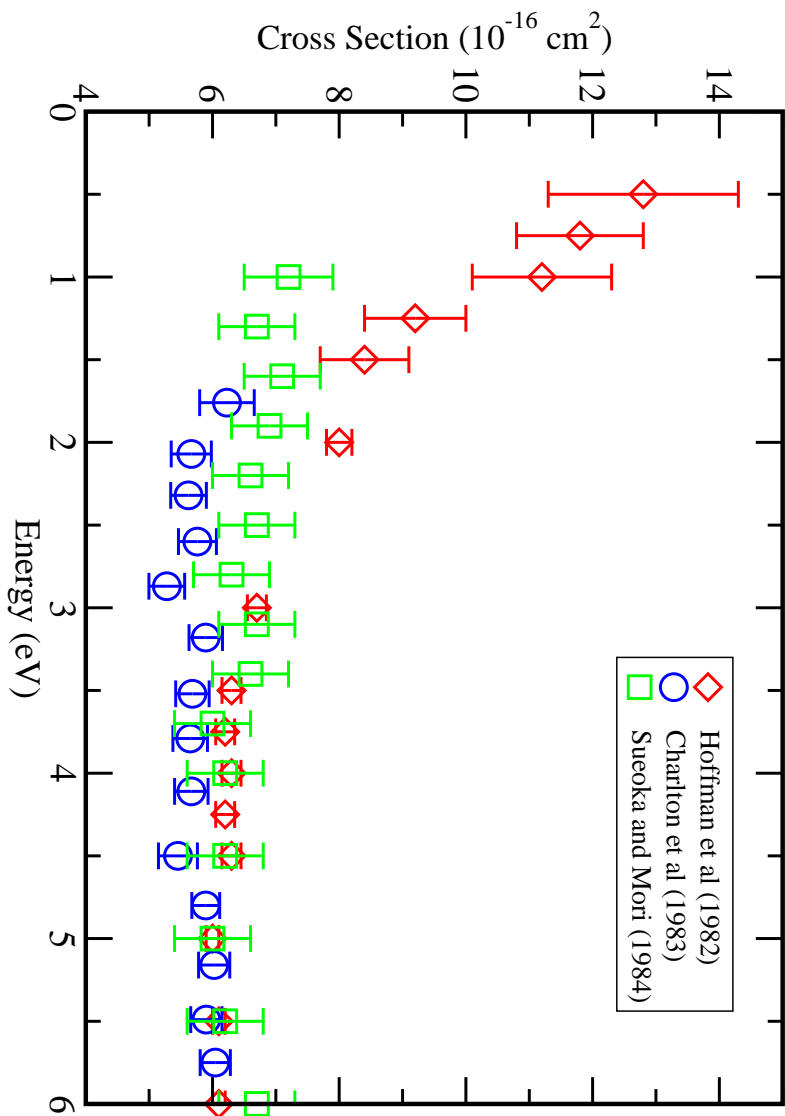
Positron- N_2

Integrated Cross Sections
 $e^+ - \text{N}_2$



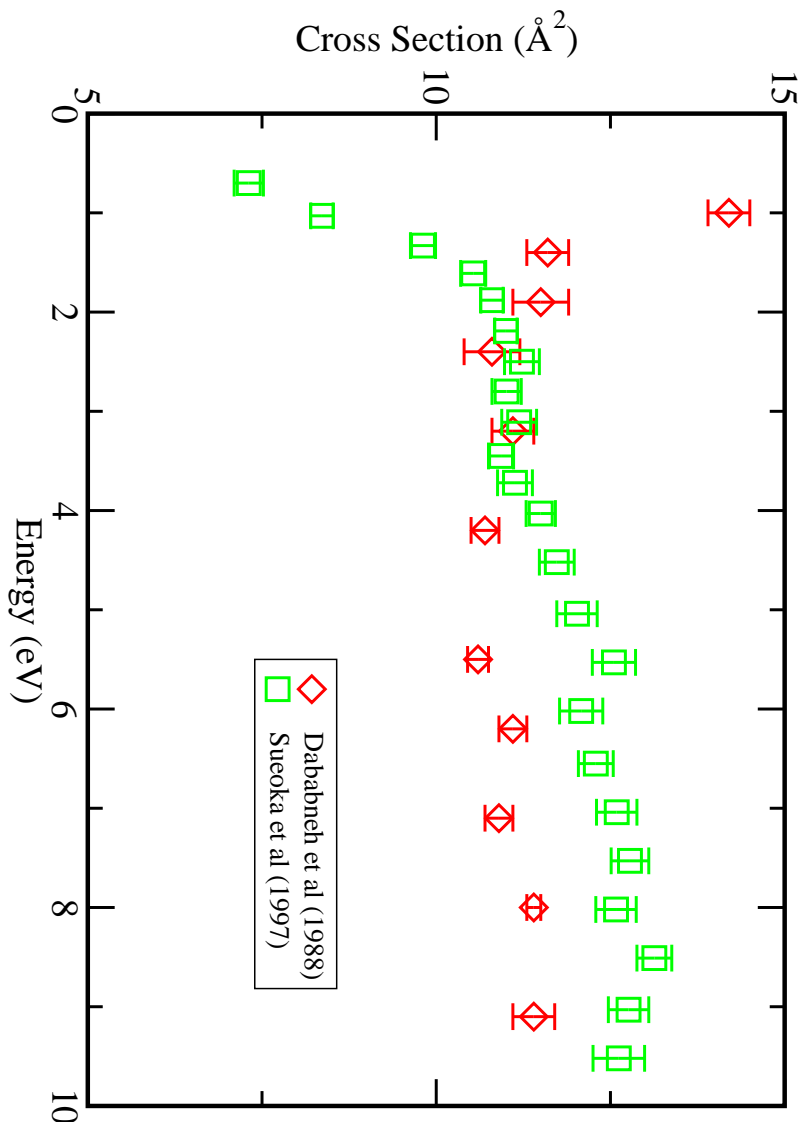
Positron-CO₂

Integrated e⁺ Cross Sections e⁺-CO₂



Positron-SF₆

Integrated Cross Section
 $e^+ - SF_6$



Our Theoretical Treatment

- Isolated collision between one target (atom or molecule) and one positron of known kinetic energy
- Only elastic scattering—stay below the energy threshold for electronic excitation of the target or for **real** positronium formation
- Local, energy-independent interaction potential based on physically-motivated approximations

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Reduction of Scattering Formulation

We “reduce” the problem by expanding the (N+1)-particle ket in a complete set of target and scattering states, viz.

$$|\Psi\rangle = \sum_{\gamma} F_{\gamma} \Phi_{\gamma}.$$

Using this expansion in the T.I.S.E. and operating on the left with $\langle \Phi_0 |$ eventually leaves the one-particle T.I.S.E. for elastic scattering of the positron

$$\hat{T}_p |F_0\rangle - \frac{k^2}{2} |F_0\rangle = -\hat{V}_{sp} |F_0\rangle .$$

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Interaction Potential

A local approximation to the interaction potential between a colliding positron and the molecular target can be represented as

$$V_{sp}(\vec{r}_p) = V_{st}(\vec{r}_p) + V_{pol}(\vec{r}_p),$$

where the static potential is given by,

$$V_{st}(\vec{r}_p) = \langle \Phi_0(\vec{r}_i) | \hat{V}_{int} | \Phi_0(\vec{r}_i) \rangle,$$

with \hat{V}_{int} representing the Coulomb interactions between the positron and the target and where $V_{pol}(\vec{r}_p)$ is the polarization potential.

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Polarization Potential

The polarization potential is defined as the energy difference

$$V_{pol}(\vec{r}_p) = E^R(\vec{r}_p) - E^U(\vec{r}_p),$$

where the energy from the relaxed (distorted) target orbitals is

$$E^R(\vec{r}_p) = \langle \Phi^R(\vec{r}_i; \vec{r}_p) | \hat{\mathcal{H}}_t + \hat{V}_{int} | \Phi^R(\vec{r}_i; \vec{r}_p) \rangle_{d\vec{r}_i},$$

and the energy from the unrelaxed (ground state) orbitals is

$$E^U(\vec{r}_p) = \langle \Phi_0^U(\vec{r}_i) | \hat{\mathcal{H}}_t + \hat{V}_{int} | \Phi_0^U(\vec{r}_i) \rangle_{d\vec{r}_i}.$$

In the adiabatic approximation, both $E^U(\vec{r}_p)$ and $E^R(\vec{r}_p)$ are calculated by treating the projectile as an additional “proton” (located at \vec{r}_p) and using an LCAO:MO—SCF Quantum Chemistry code to provide variational estimates of these energies.

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Nuclear Attraction Integrals

The distortion of the molecular orbitals is driven by the nuclear attraction integrals that involve the positron as an additional “nucleus” at \vec{r}_p .

$$I_{i,j}^{N\!A\!I} = \langle \alpha_i(\vec{r}_e) | V(\vec{r}_e; \rho_{pos}) | \beta_j(\vec{r}_e) \rangle ,$$

where the interaction V from a distribution of positive charge ρ_{pos} is given as

$$V(\vec{r}_e; \rho_{pos}) = \int d\vec{r} \rho_{pos}(\vec{r}) \frac{-1}{|\vec{r}_e - \vec{r}|}.$$

For the adiabatic approximation, $\rho_{pos}(\vec{r}) = \delta(\vec{r} - \vec{r}_p)$, which is appropriate for the positive charge distribution in a virtual H atom.

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Virtual Positronium vs Virtual Hydrogen

- Adiabatic approximation is much too attractive in the near-target region
- Use ρ_{pos} more nearly correct for virtual Ps
 - Hence, the name **Distributed Positron Approximation** (DPM)

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Ps C.O.M. Figure

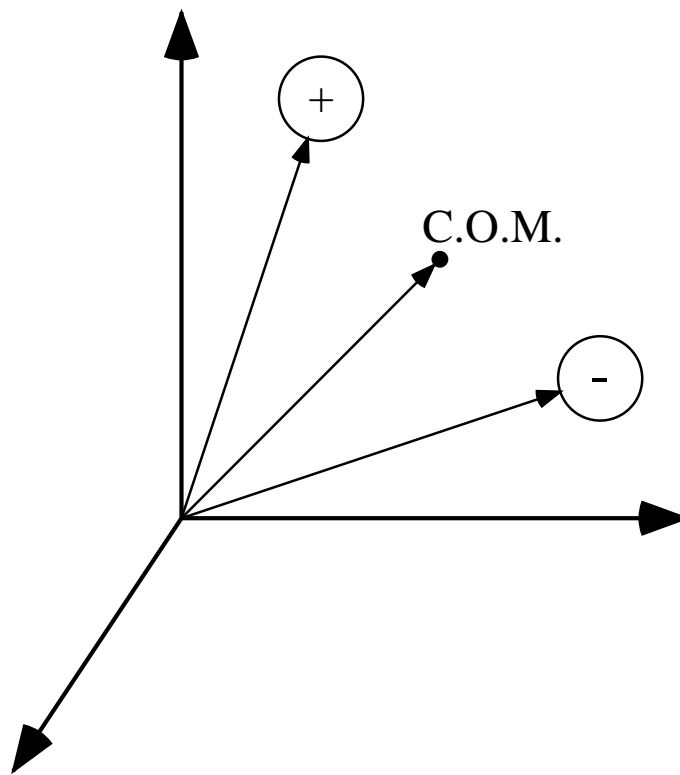


Figure 1: Unlike the H atom, the positive charge in the Ps atom is not localized at the C.O.M. of the atom.

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The Distributed Positron Model

We have investigated several simple choices of ρ_{pos} that reflect the distribution of positive charge in a Ps atom and lead to a polarization potential that reduces to the correct long-range behavior of the polarization potential, including:

- Uniform Spherical
 - $R_p = 1.5$ Bohr (Average radius of Ps atom)
 - $R_p = 1.0$ Bohr (Maximum radial probability of Ps atom)
- H1S distribution
- STO-3G distribution

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DPM Figure

For specificity, consider the following case:

- Diatomic target with nuclei at \vec{A} and \vec{B}
- Additional “nucleus” (positron) with charge distribution ρ_{pos} located at $\vec{C} \equiv \vec{r}_p$

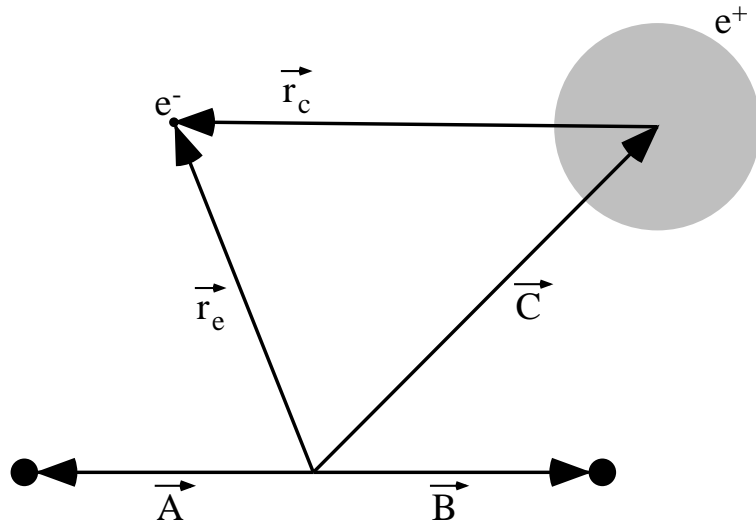


Figure 2: A distribution of positive charge simulates the effect of virtual Ps formation in the DPM.

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DPM Distortion Interaction Figure

DPM Distortion Interactions

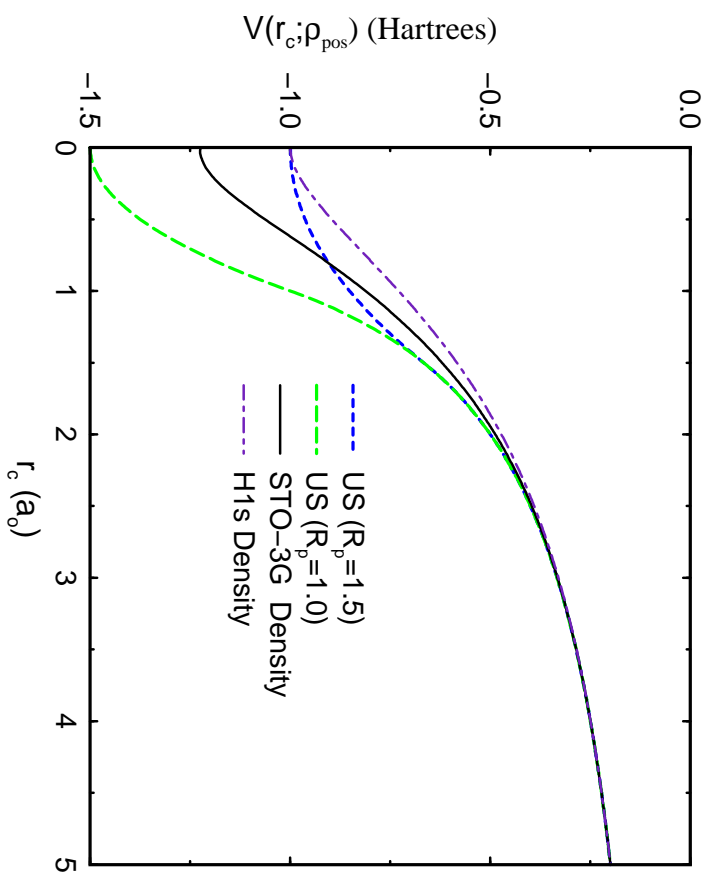


Figure 3: The distortion interaction $V(\vec{r}_c; \rho_{pos})$ for various choices of ρ_{pos} in the DPM.

Computing the Interaction Potential

To calculate the polarization component of the interaction potential, we fix the positron at a large number of points near the target and then compute a variational estimate of the system energy for each of these points.

- **Bigger Systems \Rightarrow More CPU Time**
 - Time to compute V_{pol} for Ar: 12 minutes (serial code)
 - Time to compute V_{pol} for SF₆: 125 days (estimated)
- **Bigger Systems \Rightarrow More Memory**
- **Bigger Systems \Rightarrow More I/O**

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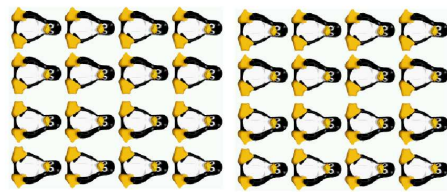
Solution

- **Beowulf Cluster**

- Performance: 100's to 1000's of Megaflops
- Memory: Gigabytes
- I/O: Distributed Across Multiple Nodes
- **Very Affordable Hardware**
- Standard Software Available for Free
- Local Control
 - * Configuration Optimized for Problem
 - * No Sharing Required

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Gamera Mark II



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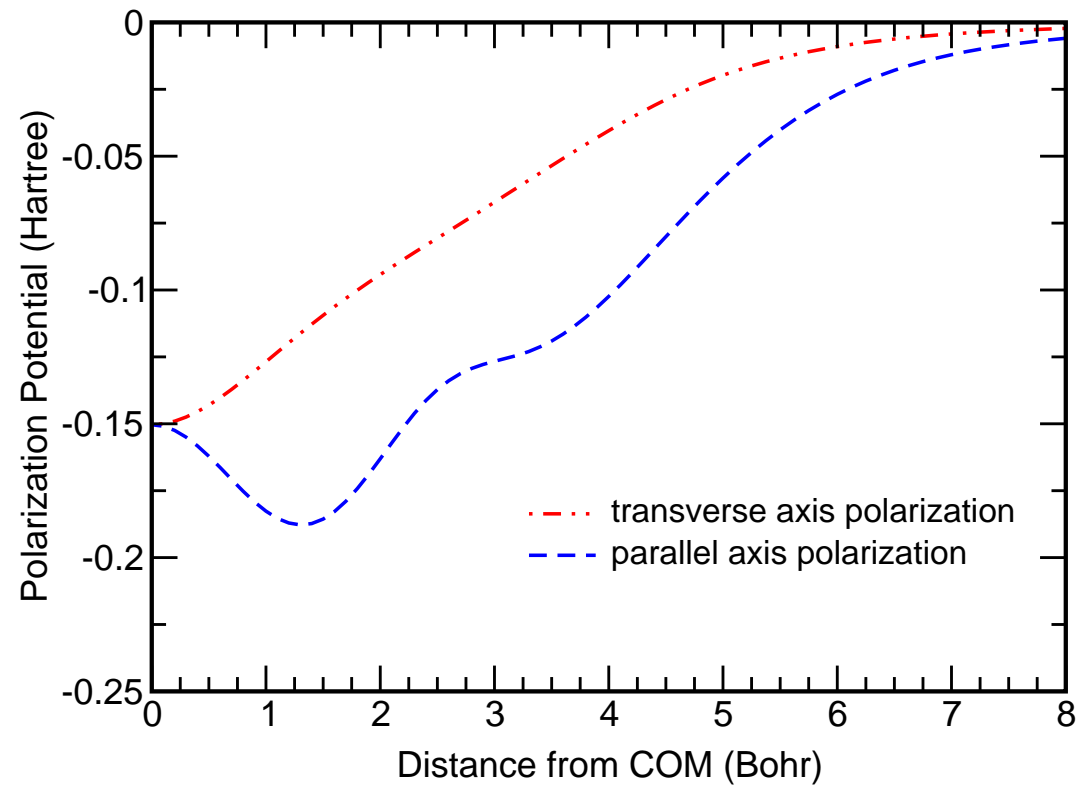
Gamera Mark II Hardware

- 32 1-GHz Athlon CPUs
- 16 Gigabytes aggregate ram
- 640 Gigabytes aggregate disk
- 2 100 base-T 24-port switches
- **Cost** ~ \$30,000



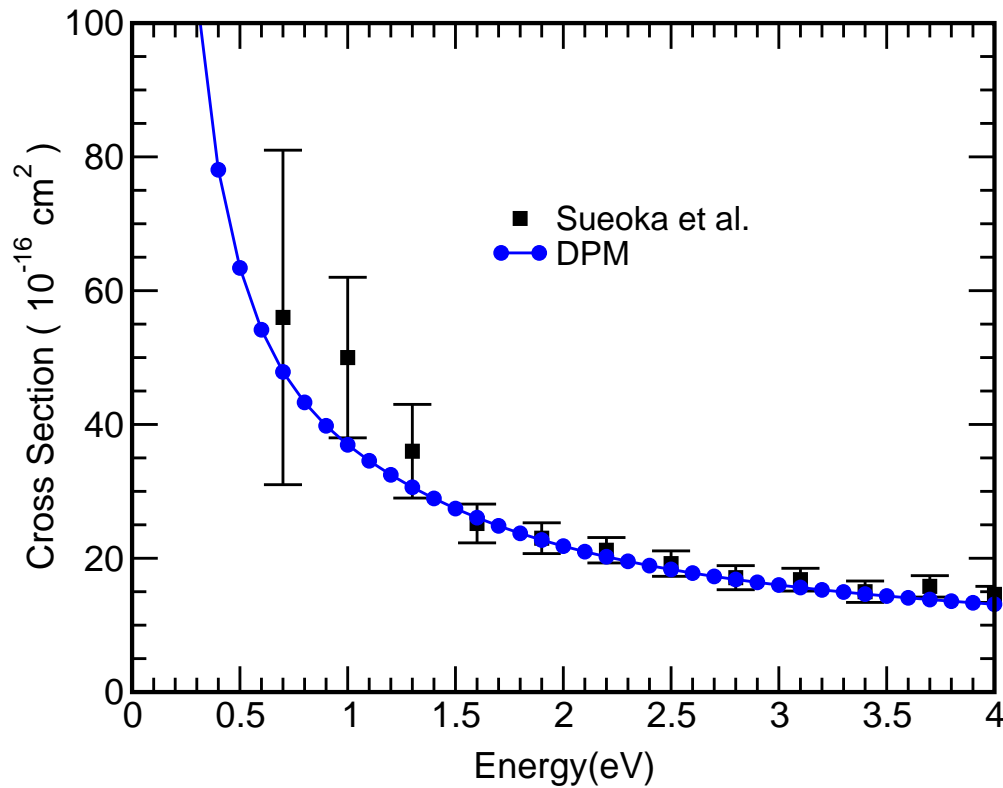
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Positron-C₂H₂ Polarization Potential



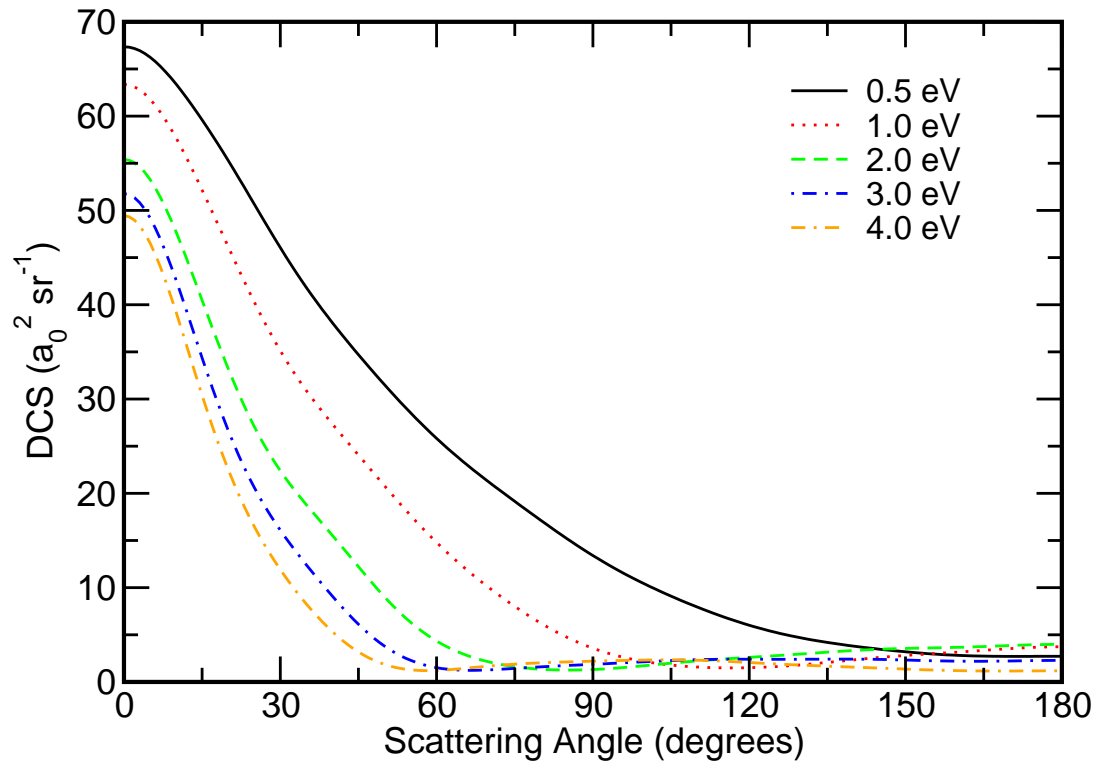
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Positron- C_2H_2 Integrated Cross Sections



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Positron-C₂H₂ Differential Cross Sections



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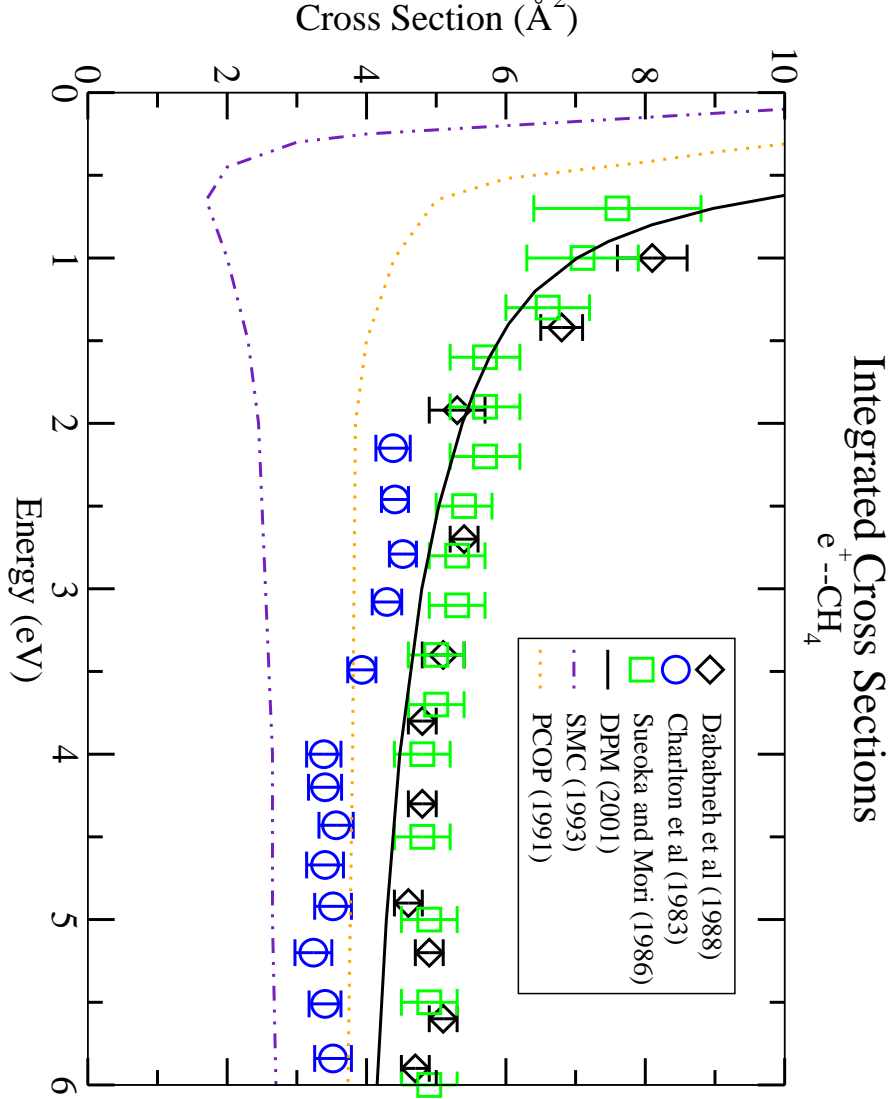
Conclusions

- High-Performance Computing \equiv Concurrent Computing
- Affordable Parallel Systems Are Available Now
- Open Source Software and Scientific Applications Are Compatible
- The DPM has been shown to yield consistently good agreement with measured low-energy positron-atom and positron-molecule collisions

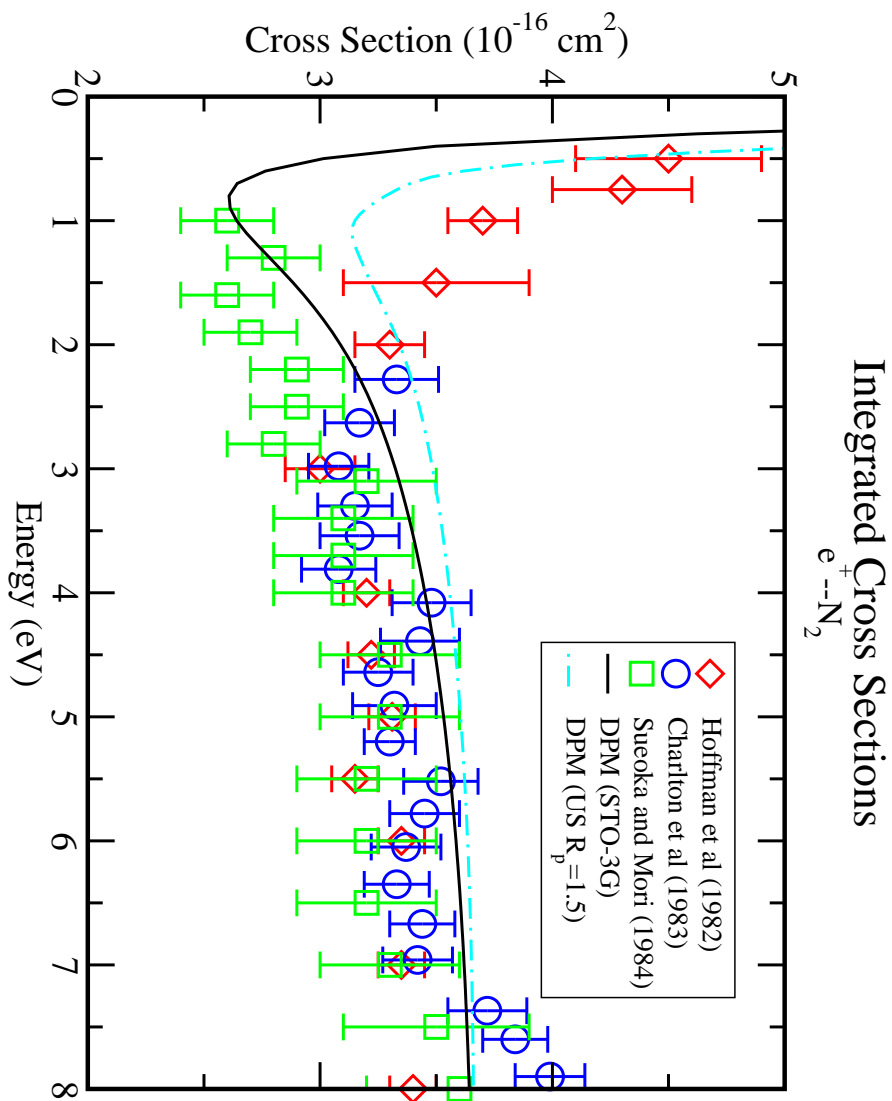
We are grateful for the collaborative efforts of Dr. Robert Lucchese and Dr. Franco Gianturco on the CH₄ and SF₆ systems, as well as our continuing work on C₈H₈.

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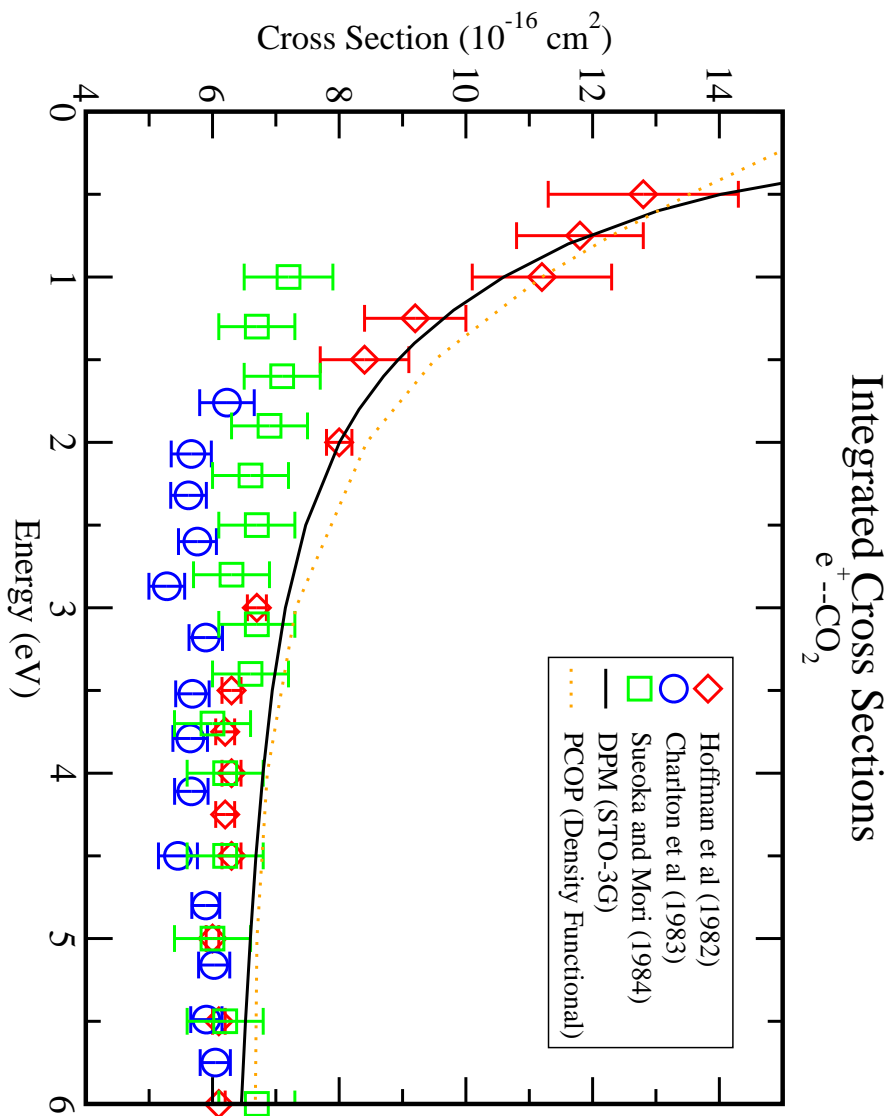
Positron-CH₄ Cross Sections



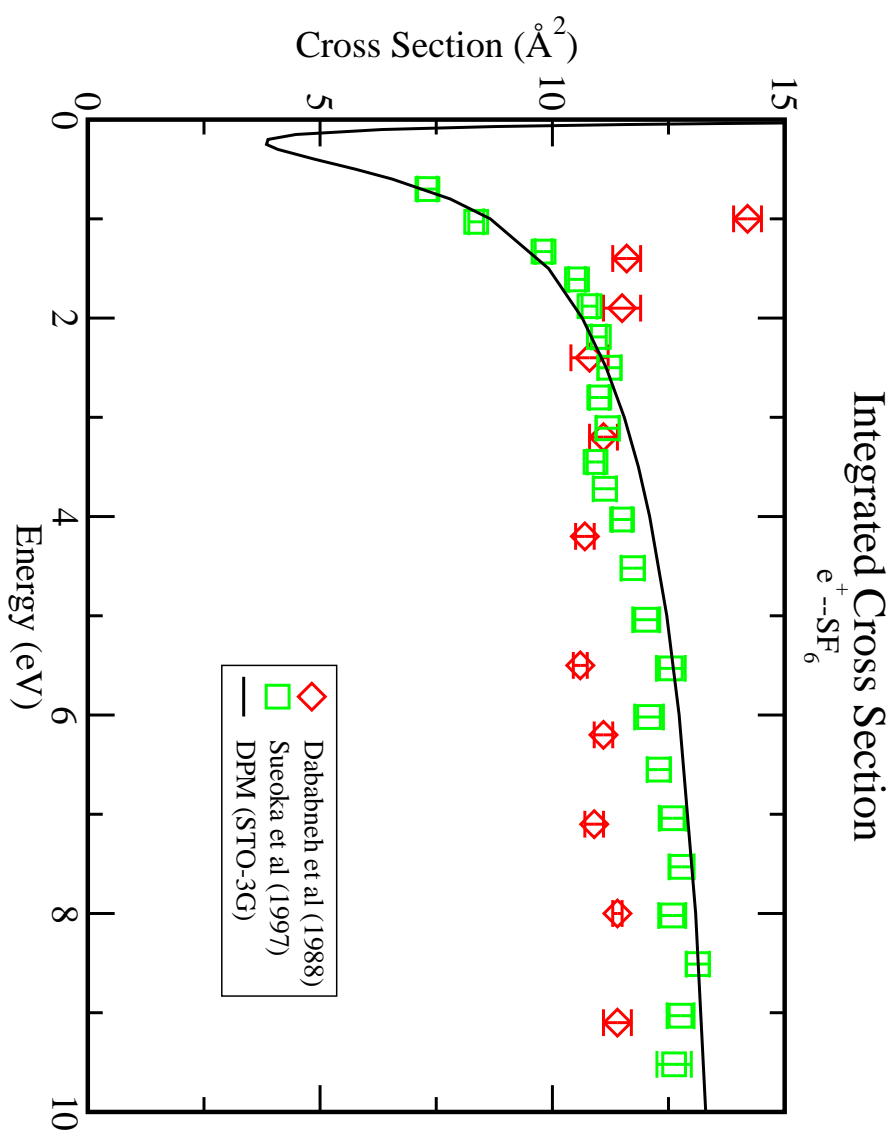
Positron- N_2 Cross Sections



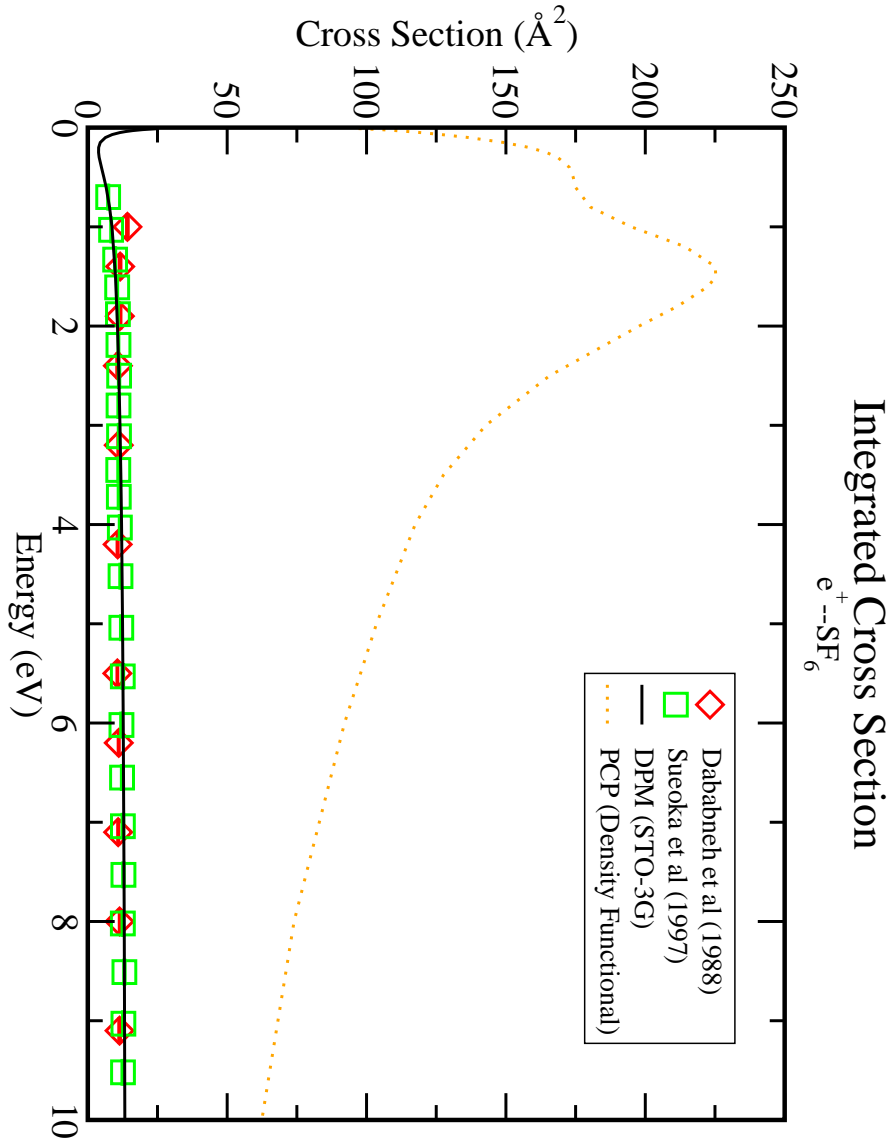
Positron- CO_2 Cross Sections



Positron-SF₆ Cross Sections - Close



Positron-SF₆ Cross Sections - Far



C₂H₂ Properties - 1

Table 1: Total Energies

Research Group	Energy (Hartree)
this work	-76.84885
Jameson and Fowler	-76.84857
Lindh and Liu	-76.85500

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C₂H₂ Properties - 2

Table 2: Quadrupole Moments

Group	θ (au)
this work	5.43
Jameson and Fowler	5.47
Amos	5.47
Lindh and Liu	5.24-5.44
experimental	2.23-6.24

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C₂H₂ Properties - 3

Table 3: Polarizabilities

Research Group	$\bar{\alpha}$	α_{zz}	α_{xx}
this work	21.22	32.95	15.35
Jameson and Fowler	23.11	31.34	19.34
Gough	22.68		
Gough	14.30	24.842	9.025
experimental	23.53	31.91	19.34

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