

GFMC Calculations

Steven C. Pieper

Work with

Ralph Butler (Middle Tennessee State U.)

Joseph Carlson (Los Alamos)

E. L. (Rusty) Lusk (Argonne)

Ivan Brida (Argonne)

Kenneth M. Nollett (Argonne)

Muslema Pervin (Argonne)

Robert B. Wiringa (Argonne)

- Making GFMC work on 65,536 processors
- GFMC ^{12}C results
- Benchmark results for NN potentials
- Neutron drops
- GFMC for scattering
- Deliverables and plans



Physics Division

Work supported by U.S. Department
of Energy, Office of Nuclear Physics,
and Grant DE-FC02-07ER41457

MAKING GFMC WORK ON 65,536 PROCESSORS

- Automatic Dynamic Load Balancing for sharing work between nodes
 - A general-purpose library to help application codes dynamically share work
 - Being developed by Rusty Lusk and Ralph Butler
 - Working very well on 8,192 nodes
 - Still trying to get good efficiency on 16,384 nodes
 - Rusty will talk about its structure
- OpenMP for work on one node
 - Individual nodes have multiple cores sharing a common memory (4 cores on BG/P)
 - Each core could be a separate MPI “node” but then each gets only 1/4 of memory
This is too small on BG/P (512 Mbytes) for ^{12}C
 - OpenMP allows one copy of the Fortran program to use all 4 cores – (2 Gbyte available)

CHANGES FOR OPENMP

Generally these were minor

- Need to identify loops that can be run in parallel on the 4 cores
- Insert directive to tell OpenMP to run loop in parallel

```
!$OMP PARALLEL DO PRIVATE( l, rcm, ... )  
  
...  
!$OMP END PARALLEL DO
```

- PRIVATE(l, rcm, ...) states that separate copies of these variables must be maintained for each thread
- Sometimes more extensive changes are needed

Original source

```
M = 0
do i = 1, ...
  do j = 1, function(i)
    M = M + 1
    x(M) = ...
  enddo
enddo
```

Modified for OMP

```
M = 0
do i = 1, ...
  MBASE(i) = M
  do j = 1, function(i)
    M = M + 1
  enddo
enddo

...
!$OMP PARALLEL DO PRIVATE( i, M, j, ...
do i = 1, ...
  M = MBASE(i)
  do j = 1, function(i)
    M = M + 1
  enddo
enddo
!$OMP END PARALLEL DO
```

RESULTS SO FAR

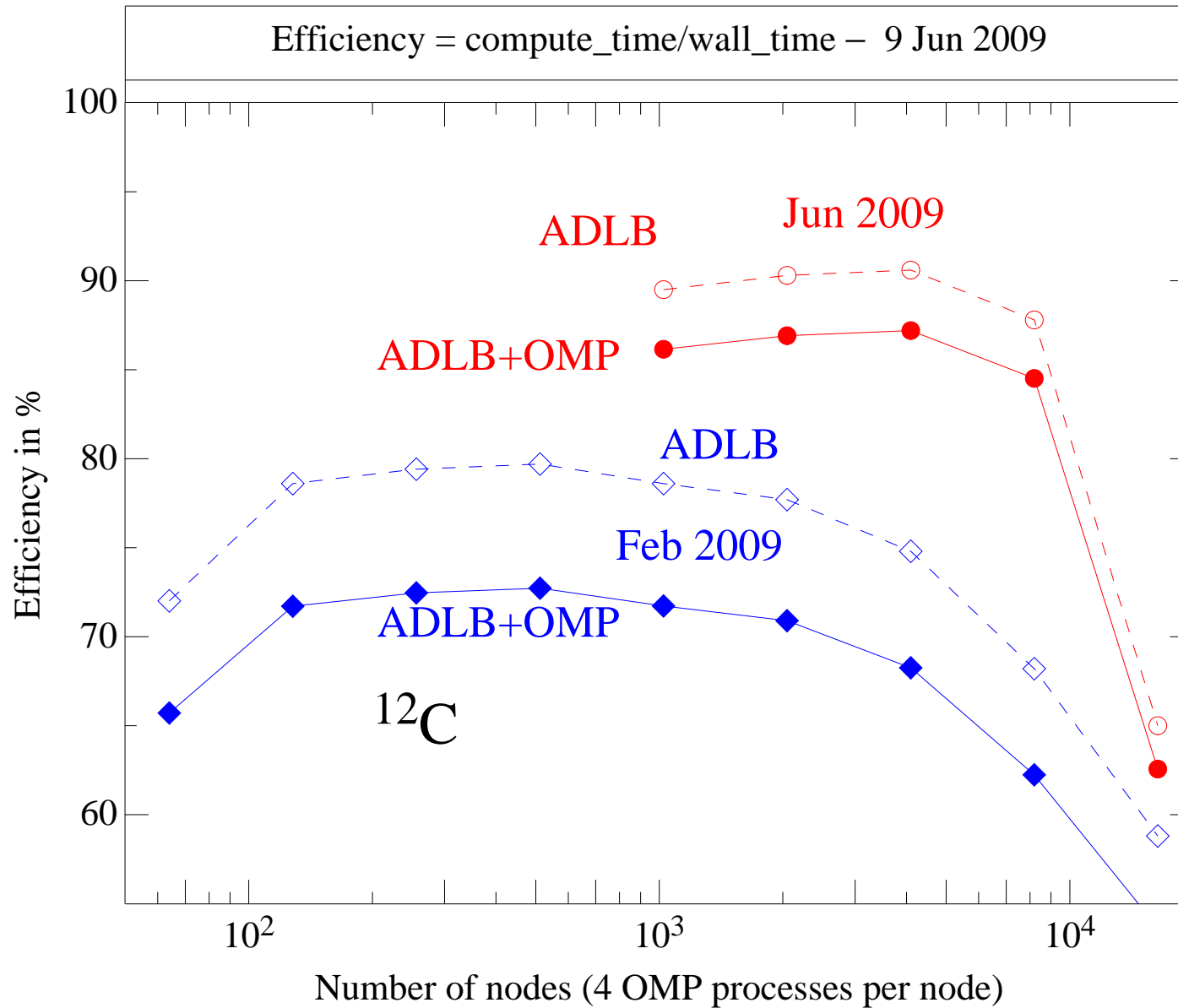
OpenMP is very successful, but the BG/P speeds are disappointing

¹²C Speeds for two key subroutines

Subroutine	MFLOPS		speed up
	1 thread	4 threads	
Wave function	239	1022	4.0
V_{ijk}	319	1233	3.9

RESULTS SO FAR

ADLB performance is very good up to 8192 nodes (32,768 cores)



^{12}C RESULTS

In Dec. 2008 & Jan. 2009, the first ADLB+GFMC calculation of the $^{12}\text{C}(\text{gs})$ was made.

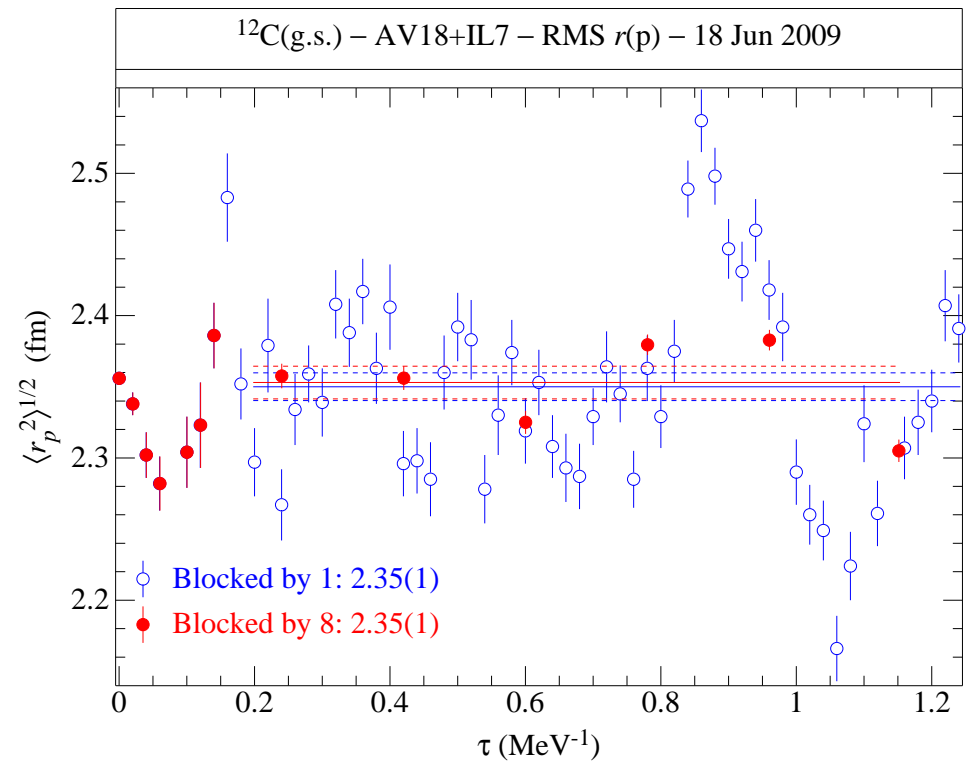
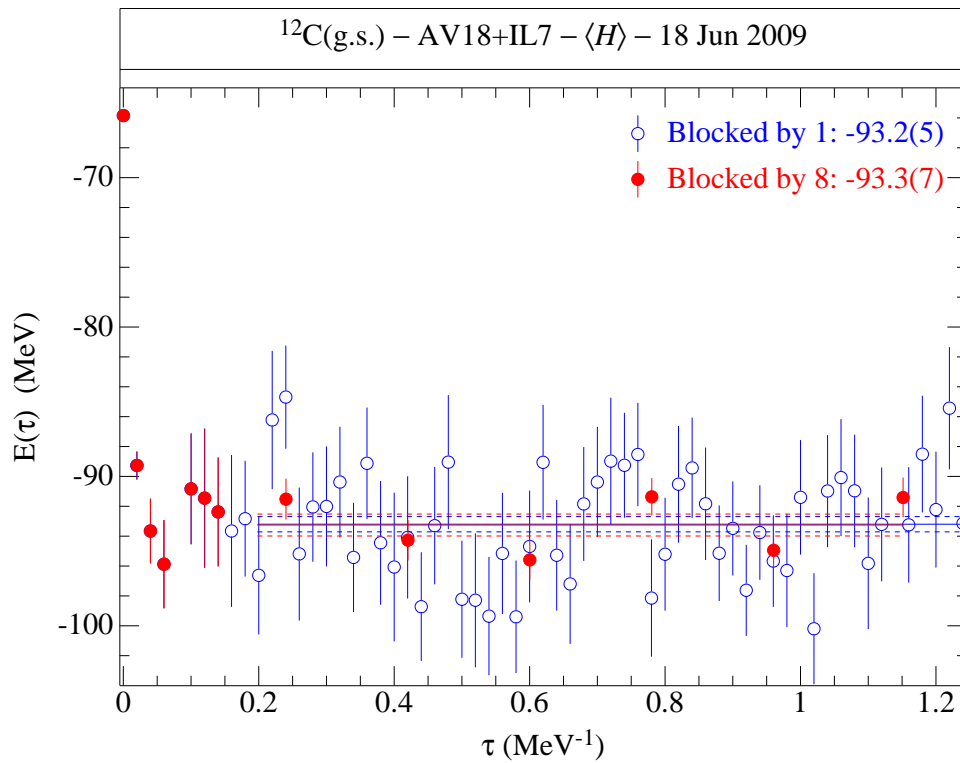
- AV18+IL7 Hamiltonian
- Improved (and slower) Ψ_T than in previous calculation
- GFMC result changed only a little
- 16,000 configurations propagated to $\tau = 1.24 \text{ MeV}^{-1}$ (2480 steps)
- 40 unconstrained time steps used before energy evaluations
- Used 8,192 nodes (32,768 cores) of BG/P
- 14 runs for total of 93 hours (first few very short)
- Speed of Ψ_T calculation significantly improved since
- Convergence is very good and shows that
 - smaller maximum τ can be used
 - fewer unconstrained time steps, and hence fewer configurations, can be used

An ADLB+GFMC calculation using the benchmark modified SSCC v'_8 NN potential was also made; this is much faster

- No L^2 terms in NN potential
- No NNN potential

^{12}C CONVERGENCE STUDIES

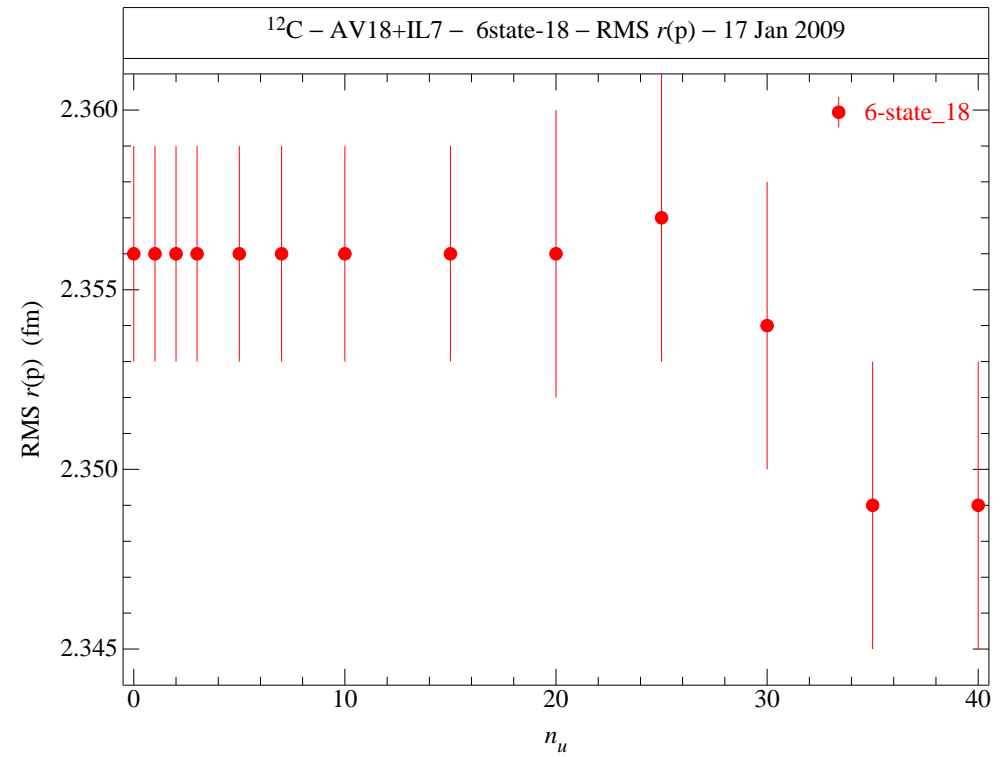
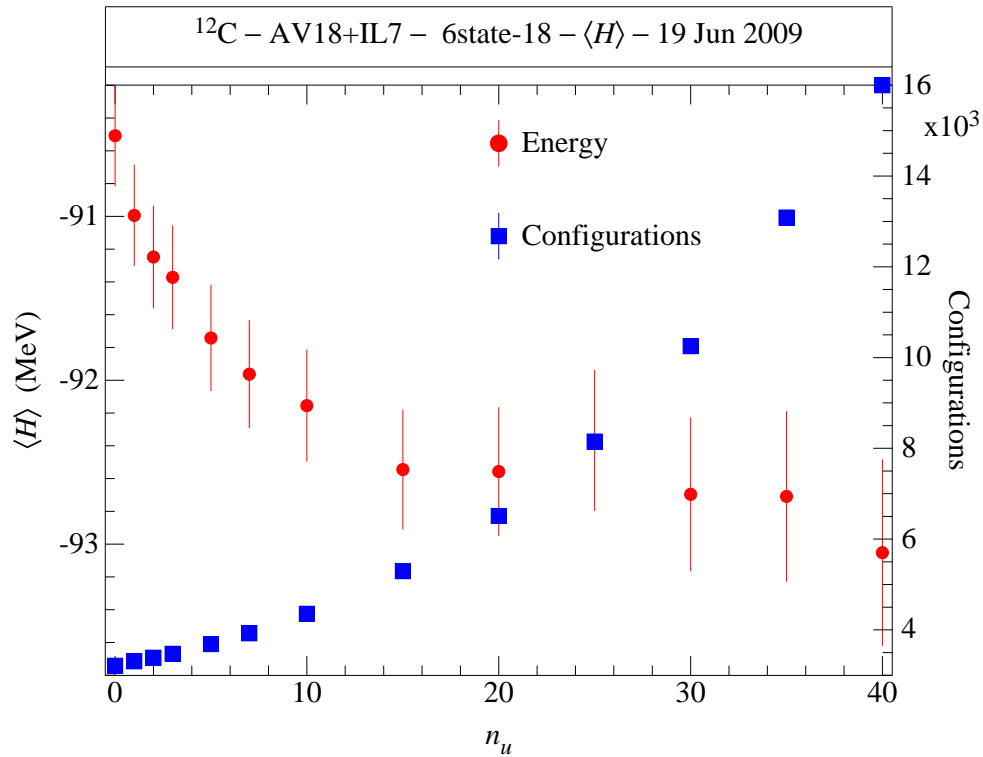
As a function of imaginary time (τ)



Modified SSCC v'_8 convergence with τ is similar

^{12}C CONVERGENCE STUDIES

As a function of number of unconstrained steps (n_u)



$n_u = 20$ is converged, needs only half as many configurations

Modified SSCC v'_8 convergence with n_u is better

^{12}C RESULTS – ENERGIES & RADII

	Energy			RMS radius		
	VMC	GFMC	Expt.	VMC	GFMC	Expt.
AV18+IL7	-65.8(2)	-93.2(6)	-92.16	2.36	2.35	2.33
Modified SSCC v'_8	-74.9(2)	-94.0(5)		2.21	2.24	

Modified SSCC v'_8 gives reasonable energies at least up to $A = 12$

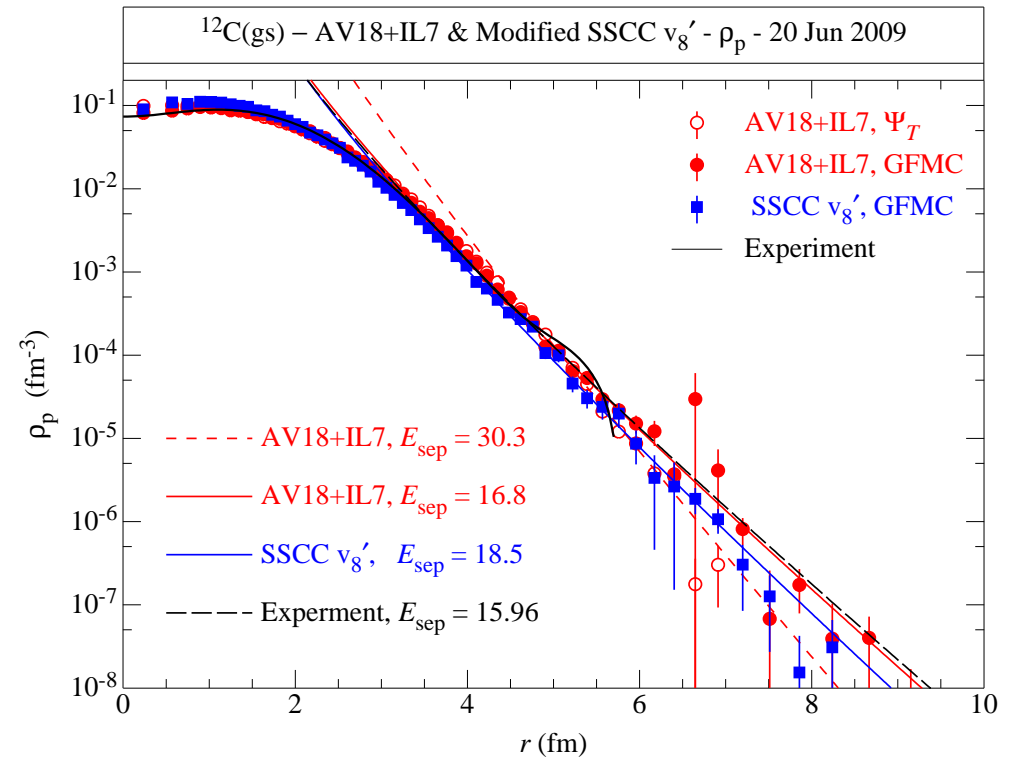
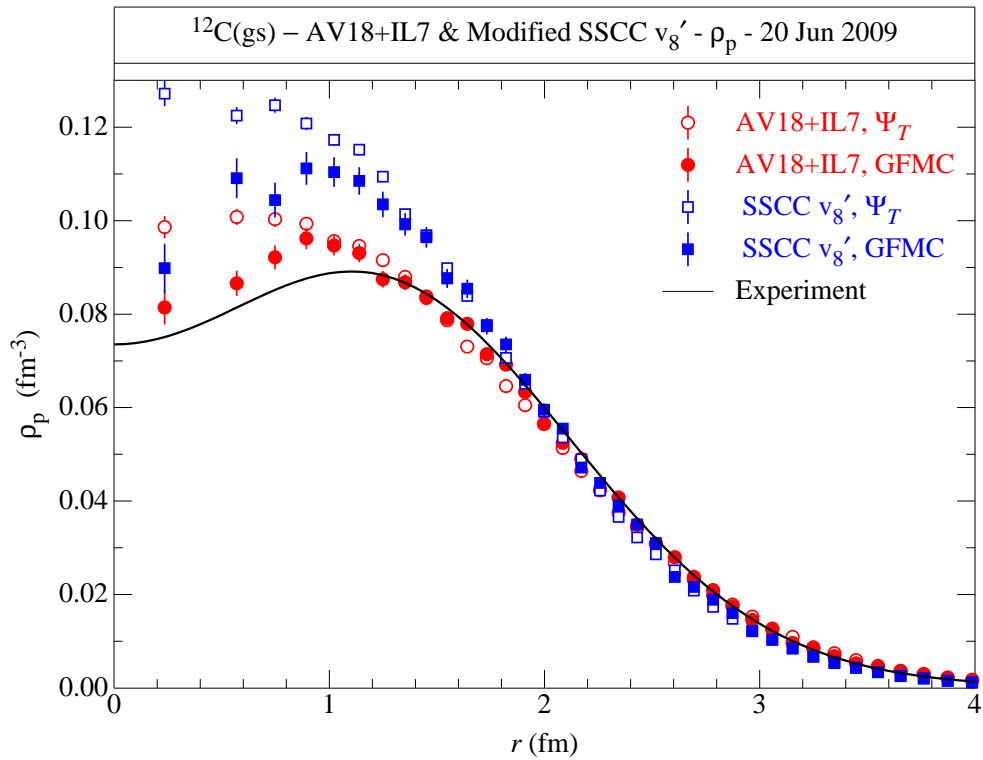
Remember that it does not accurately reproduce NN P -phase shifts

	K.E.	v'_{1-4}	$v'_{t,t\tau}$	$V_{ijk'}$
AV18+IL7	427.	-216.	-282.	-23.
Modified SSCC v'_8	350.	-220.	-223.	0.

K.E. of SSCC v'_8 is not so small, despite much smaller central repulsion

Could be associated with the still large $v'_{t,t\tau}$

^{12}C RESULTS – ONE-BODY DENSITY



The Ψ_T density is significantly improved by GFMC

- Central dip is generated
- AV18+IL7 tail falls at rate dictated by $E_{\text{sep}} = E(^{11}\text{B}) - E(^{12}\text{C})$ instead of twice as fast
- SSCC v'_8 E_{sep} is also good

BENCHMARKS RESULTS FOR MODIFIED SSCC v'_8

	SSCC v'_8	Experiment
${}^4\text{He}(0^+)$	-25.36(1)	-28.30(0)
${}^6\text{He}(0^+)$	-26.48(1)	-29.27(0)
${}^6\text{Li}(1^+)$	-28.98(2)	-31.99(0)
${}^6\text{Li}(3^+)$	-26.22(2)	-29.800(2)
${}^7\text{Li}(\frac{3}{2}^-)$	-35.93(4)	-39.24(0)
${}^7\text{Li}(\frac{1}{2}^-)$	-35.78(5)	-38.760(0)
${}^8\text{He}(0^+)$	-28.60(3)	-31.41(1)
${}^9\text{Be}(\frac{3}{2}^-)$	-53.49(13)	-58.16(0)
${}^{10}\text{Be}(0^+)$	-61.34(16)	-64.98(0)
${}^{10}\text{Be}(2^+)$	-58.85(22)	-61.61(0)
${}^{10}\text{Be}(2^+, 2^{nd})$	-57.13(17)	-59.02(0)
${}^{12}\text{C}(0^+)$	-94.00(47)	-92.16(0)

NEUTRON DROPS

- Collection of neutrons interacting via standard NN and NNN Hamiltonian with added artificial external well
- Well can be adjusted to change density or surface thickness
- Well could be non-spherical
- If NN and NNN H is realistic, can provide input to EDF's
- GFMC can compute up to 16 neutrons (part-way through $S - D$ shell).

- Ψ_T has BCS one-body part with pairs of $0S, 0P, 1S, 0D$ neutrons
- $0S, 0P$ shells basically full; $1S, 0D$ occupations are variational parameters
- Ψ_T are pure Jastrow or Jastrow+ f_6 -pair correlations

NEUTRON DROPS

- Last year we computed a series of drops confined in Woods-Saxon external wells
- A few small improvements to these results were made this year
- Most effort has been on drops confined in oscillator wells, $\hbar\omega = 5\&10$ MeV
- Comparisons with AFDMC results obtained by Stefano Gandolfi are generally good
 - Energies agree to $\leq 1\%$
 - For $\hbar\omega = 5$ MeV external well agreement gets worse for larger drops; may be due to insufficient pairing in AFDMC.

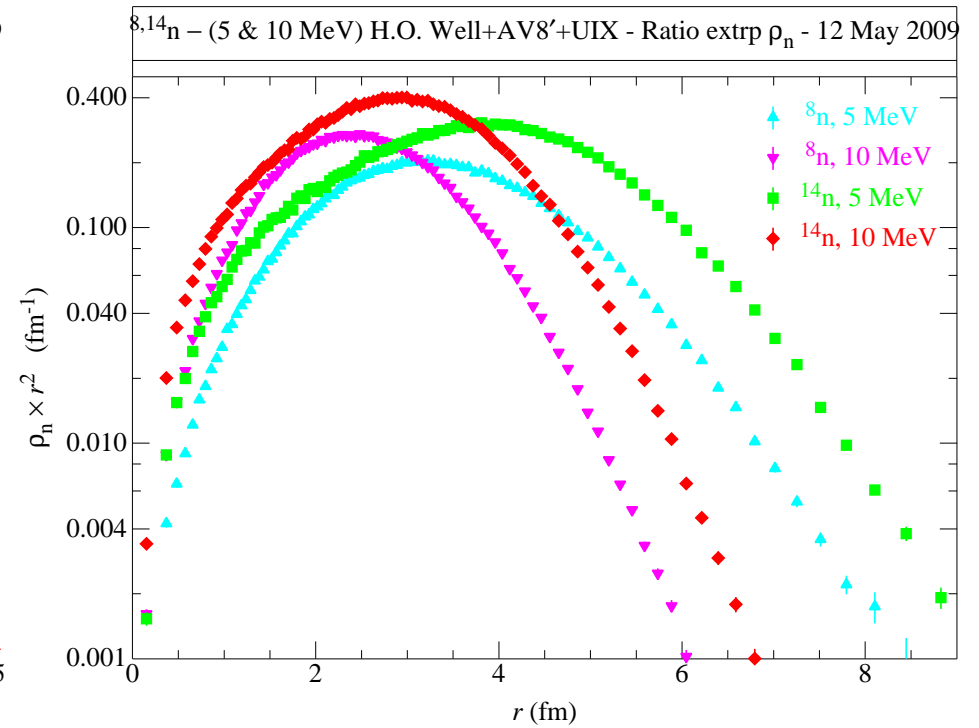
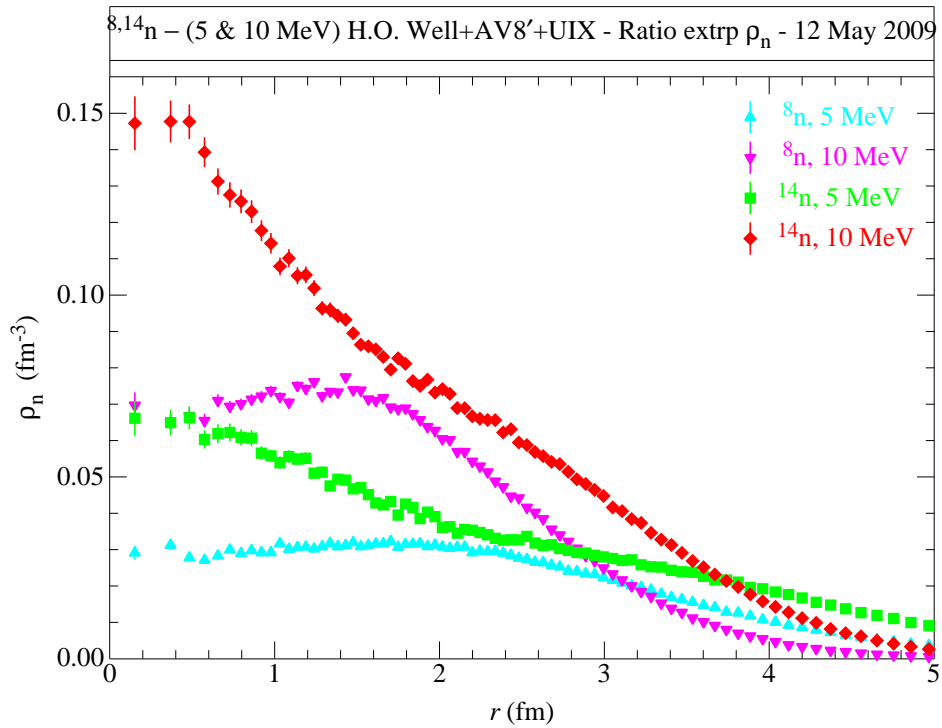
NEUTRON DROPS – ENERGIES

	Energy	Kinetic	V_{Extern}	v_{ij}	V_{ijk}	RMS r
<hr/> $\hbar\omega = 5 \text{ MeV}$ external well + AV18 + UIX <hr/>						
${}^8\text{n}(0^+)$	67.09(2)	76.2(4)	32.00(3)	−45.79(37)	0.26(1)	3.642(2)
${}^9\text{n}(\frac{1}{2}^+)$	80.95(4)	86.2(2)	39.06(3)	−53.64(38)	0.43(1)	3.793(2)
${}^9\text{n}(\frac{5}{2}^+)$	81.24(4)	86.2(2)	38.94(3)	−52.67(35)	0.36(1)	3.787(2)
${}^{10}\text{n}(0^+)$	92.22(8)	102.0(4)	45.44(5)	−65.41(61)	0.49(2)	3.881(2)
${}^{12}\text{n}(0^+)$	118.26(15)	128.0(5)	58.68(7)	−82.26(80)	0.79(4)	4.026(2)
${}^{13}\text{n}(\frac{1}{2}^+)$	130.88(13)	141.(1)	65.31(12)	−90.74(89)	0.75(4)	4.081(4)
${}^{13}\text{n}(\frac{5}{2}^+)$	131.63(13)	138.(1)	65.65(13)	−87.55(89)	0.90(4)	4.092(4)
${}^{14}\text{n}(0^+)$	142.33(16)	157.(1)	71.07(12)	−102.85(98)	1.08(4)	4.102(4)
<hr/> $\hbar\omega = 10 \text{ MeV}$ external well + AV18 + UIX <hr/>						
${}^8\text{n}(0^+)$	136.04(5)	136.0(5)	71.41(9)	−76.93(47)	1.53(3)	2.720(2)
${}^9\text{n}(\frac{1}{2}^+)$	163.84(9)	154.0(4)	85.8(1)	−90.75(57)	2.34(4)	2.811(2)
${}^9\text{n}(\frac{5}{2}^+)$	163.35(8)	154.0(3)	84.92(10)	−89.66(55)	1.93(3)	2.797(2)
${}^{12}\text{n}(0^+)$	242.12(58)	224(1)	131.69(32)	−138.24(2.10)	4.03(21)	3.016(4)
${}^{13}\text{n}(\frac{1}{2}^+)$	268.44(54)	248(2)	146.83(57)	−153.4(2.2)	4.11(22)	3.06(1)
${}^{13}\text{n}(\frac{5}{2}^+)$	267.94(60)	249(2)	147.18(53)	−156.34(2.26)	4.32(24)	3.063(6)
${}^{14}\text{n}(0^+)$	292.51(22)	275(1)	159.97(42)	−175.1(1.4)	5.28(15)	3.077(4)

NEUTRON DROPS – SINGLE-NEUTRON DENSITY DISTRIBUTIONS

Oscillator well + AV18 + UIX

$\hbar\omega = 5\&10\text{ MeV}$



GFMC FOR SCATTERING STATES

GFMC treats nucleus as particle-stable system

– Should be good for narrow resonances

Many cases should be done as scattering states

– Wide resonances: ${}^5,7\text{He}$, ${}^6\text{Li}(2^+)$, ${}^8\text{Be}(2^+,4^+)$, ...

– Will get widths of resonances

– Capture reactions: ${}^4\text{He}(d,\gamma){}^6\text{Li}$, ${}^7\text{Be}(p,\gamma){}^8\text{B}$, ...

1987 – early 1990's:

– Carlson *et al.* do ${}^5\text{He}$ states by VMC scattering

– Carlson also does preliminary ${}^5\text{He}$ GFMC scattering

Present:

– Joe Carlson doing ${}^5\text{He}$ for parity violation studies

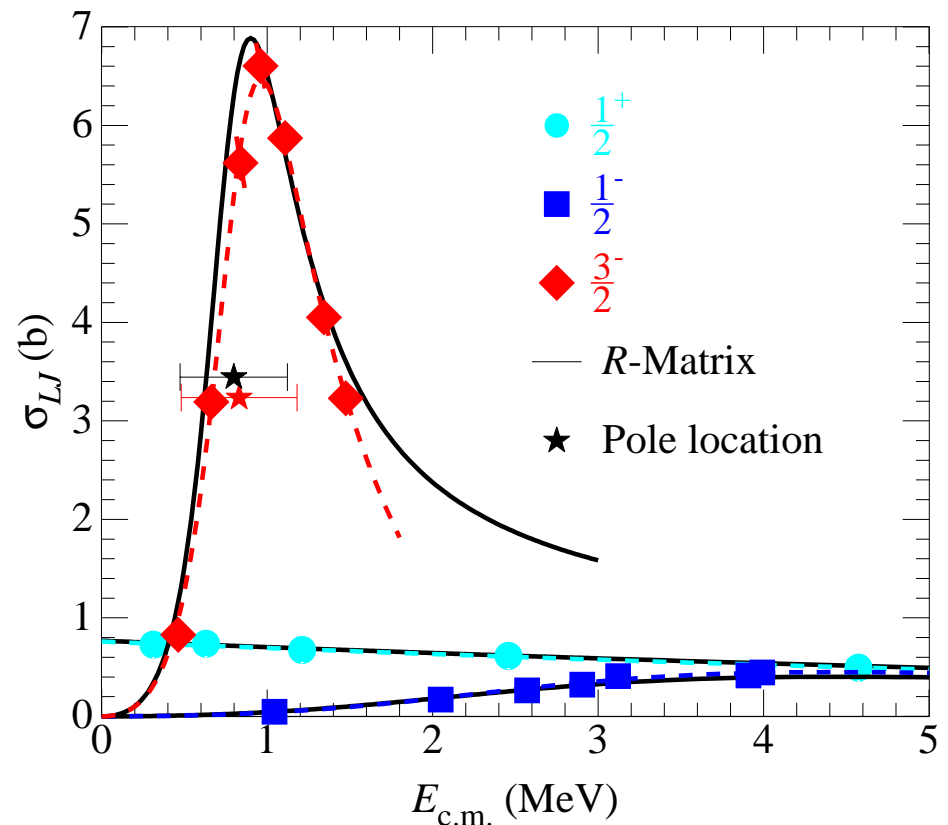
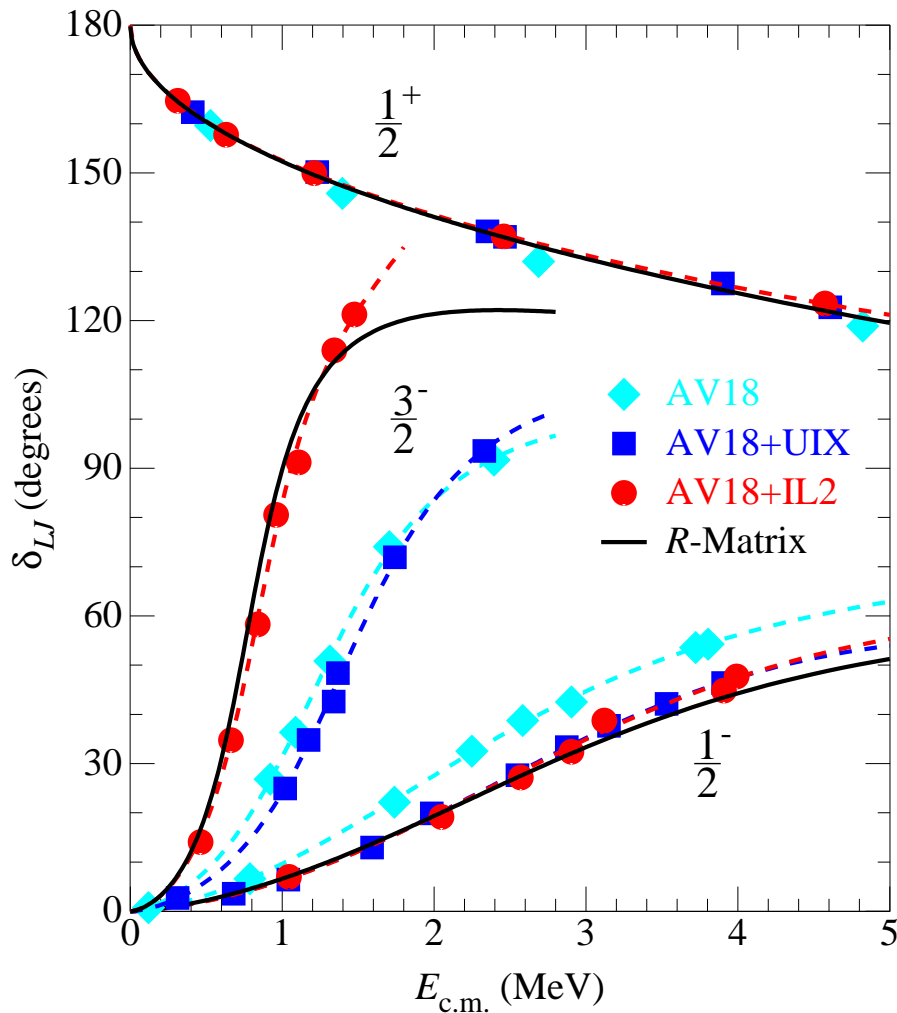
– Ken Nollett has modified Argonne GFMC program for scattering and done ${}^5\text{He}$

NCSM and CC are also computing resonance states

A benchmark comparison (${}^5\text{He}$ with SSCC v'_8 ?) would be useful

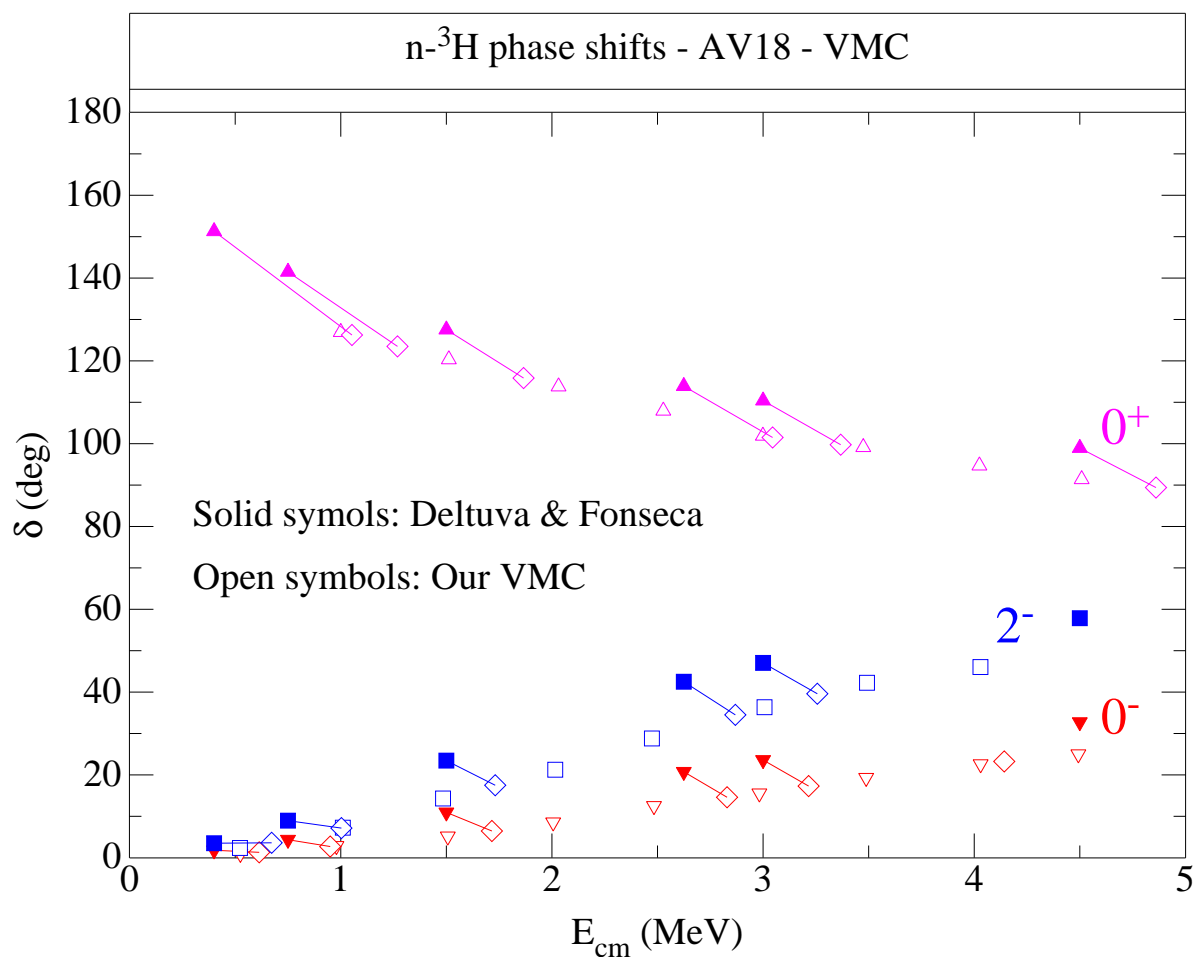
GFMC FOR ${}^5\text{He}$ AS $n+{}^4\text{He}$ SCATTERING STATES

- Black curves: Hale phase shifts from R -matrix analysis up to $J = \frac{9}{2}$ of data
- AV18 with no V_{ijk} underbinds ${}^5\text{He}(\frac{3}{2}^-)$; overbinds ${}^5\text{He}(\frac{1}{2}^-)$
- AV18+IL2 was not fit to ${}^5\text{He}$, reproduces locations and widths of both P -wave resonances
 - Spin-orbit splitting well reproduced by AV18+IL2



^4H AS $n-^3\text{H}$ SCATTERING

- We are starting to do $n-^3\text{H}$ and $p-^3\text{He}$ scattering
- Comparison with “exact” results of Deltuva & Fonseca will be a good test of our results
- So far have VMC results for several $n-^3\text{H}$ partial waves



STATUS OF DELIVERABLES FOR THIS YEAR

- Ab-initio calculations for neutron drops and asymmetric nuclear matter, including the response to external potentials
 - Done for spherical drops in several harmonic oscillator wells
- Calculate ab initio one-body densities for spherical and deformed nuclei and use them to inform DFT
 - Done: Densities of spherical nuclei
- Asynchronous Dynamic Load-Balancing improvements (microparallelization, debugging) for ^{12}C in GFMC
 - OpenMP fully implemented, much progress on ADLB
 - production AV18+IL7 ^{12}C calculation done on 32,768 processors
 - Benchmark SSCC v'_8 also done for ^{12}C
- Investigate reactions in light nuclei using ab initio methods: NCSM with RGM, and GFMC. Benchmark $n-^7\text{Li}$, $n-^8\text{He}$, and move towards nuclear projectiles
 - $n-^3\text{H}$ scattering being done in VMC

Following should have been in the list

- Improve ^{12}C VMC wave function
 - Structure of Ψ_T significantly improved since full GFMC calculation above. Still in progress

PLANS

Remainder of this year

- Continue ADLB work with aim of good efficiency on 65,536 processors
 - Non synchronous control of population growth or decay (presently uses a barrier)
 - Not allowing too great a dispersion in time steps being processed
- Further improvements to $^{12}\text{C } \Psi_T$
 - $J = 0^+$ basis states that allow Hoyle state to be represented
 - More than just one $J = 2^+$ basis state
- Deformed neutron-drop calculations if desired
- More $^{12}\text{C}(\text{gs})$ calculations using ADLB version of GFMC (Benchmark NN potentials?)
- GFMC nucleon-nucleus scattering and comparison with other methods

Next year: Not all of the following will be possible in one year!

- Continuing ADLB work in GFMC – finer-grain parallelization for BG/Q
- Many ^{12}C calculations
 - Full H for several states (Hoyle state?)
 - Transitions and transition densities
- VMC (GFMC?) computation of density matrix if interesting
- Neutron drops with new Illinois potential
- Real nuclei in external wells
- Non-spherical external wells
- GFMC nucleon-nucleus scattering and comparison with other methods

PLANS

Year 5: Some of these could be in year 4 with and some year 4 ones in year 5

- More changes of GFMC/ADLB for exascale class computers (e.g. BG/Q)
- More ^{12}C calculations, specifically Hoyle state.
- other $A = 11,12$ nuclei
- $A > 12$ nuclei?
- ^{11}Li and densities in $^9\text{Li-n-n}$ space

ADDITIONAL MATERIAL FROM PREVIOUS TALKS

GFMC needed to be redone for leadership class computers

- Old program did several Monte Carlo samples per processor
- Branching can kill samples – need enough not to fluctuate to zero
- ^{12}C will have 10,000 Monte Carlo samples
- Leadership class computers have 10,000's processors
- Need to split one sample over many processors

AUTOMATIC DYNAMIC LOAD BALANCING – CURRENT GFMC IMPLEMENTATION

Old GFMC

Each slave gets several configurations

Slave

propagates configurations

(few w.f. evaluations)

replicates or kills configs (branching)

→ periodic global redistribution

computes energies

(many w.f. evaluations)

Need ~ 10 configs per slave

^{12}C will have only $\sim 10,000$ configs.

Can't do on more than 2000 processors

Configurations cannot be unit of parallelization

With ADLB

A few “boss” slaves manage the propagation:

- Generate propagation work packages
 - Answers used to make 0,1,2, ... new propagation packages (branching)
 - Number of prop. packages fluctuates
 - Global redistribution may be avoided
- Generate energy packages – No answers

When propagation done, become worker slaves

Most slaves ask ADLB for work packages:

- Propagation package
 - Makes w.f. and $3N$ potential packages
- Energy package
 - Makes many w.f. packages
 - Makes $3N$ potential packages
 - Result sent to Master for averaging
- Wave Function or $3N$ potential package
 - Result sent to requester

Wave function is parallelization unit

Can have many more processors than configs

AUTOMATIC DYNAMIC LOAD BALANCING – CALCULATIONS

Development is still continuing but VMC and GFMC calculations using ADLB are being made.

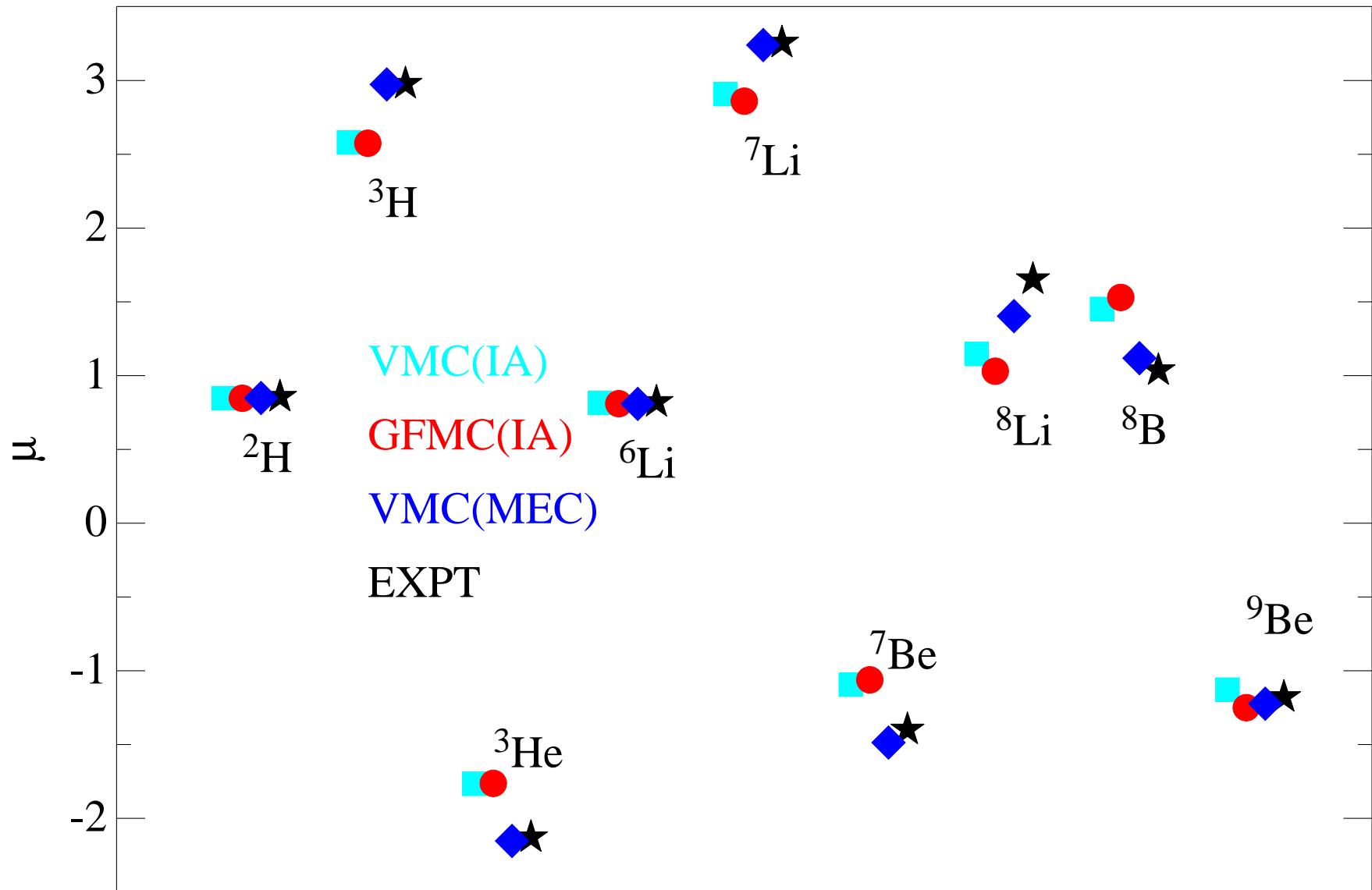
Calculations made principally to demonstrate ADLB

- 5,400-processor VMC for ${}^7\text{Li}$ – 95% efficiency
- 16,384-processor GFMC for 14-neutron drop – 83% efficiency

Calculations made for physics interest

- 2,000- to 8,192-processors: many 14-neutron drop with various H , Ψ_T , GFMC parameters
- 8,192-processor ${}^9\text{Be}$ with SSCC v'_8
- 4,096 – 8,192-processor ${}^{10}\text{Be}$ with SSCC v'_8
- up to 4,800 processors: nuclei up to ${}^{10}\text{B}$ with new Illinois V_{ijk}

Magnetic Moments



No effective charges or effective nucleon magnetic moments!

M1, E2, F, GT transitions

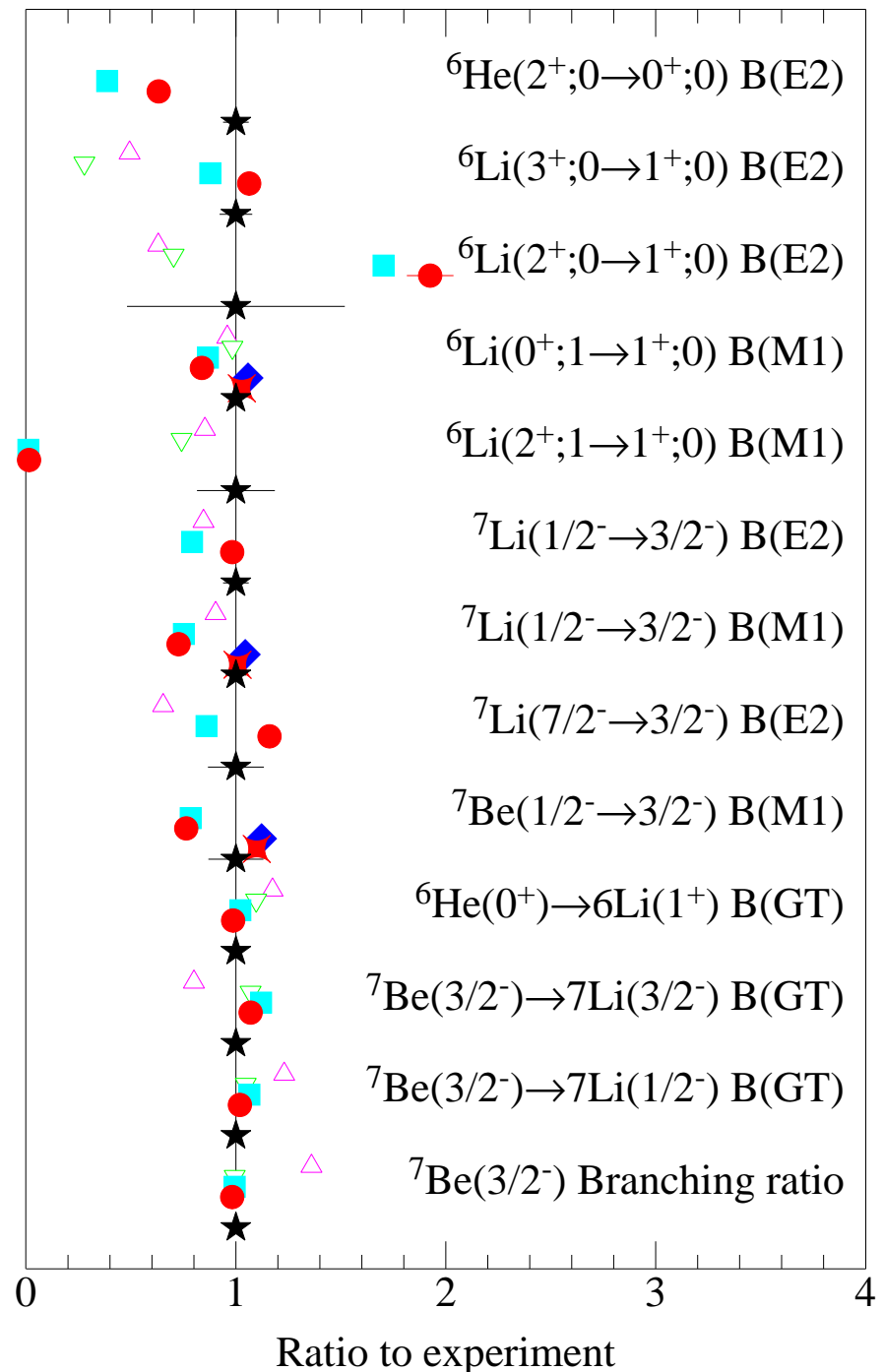
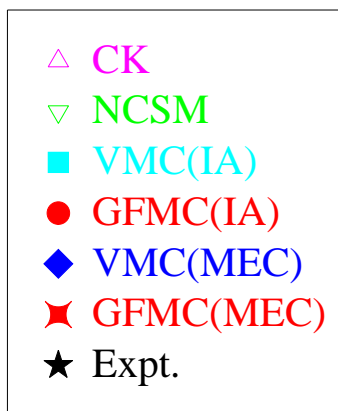
$$E2 = e \sum_k \frac{1}{2} [r_k^2 Y_2(\hat{r}_k)] (1 + \tau_{kz})$$

$$M1 = \mu_N \sum_k [(L_k + g_p S_k)(1 + \tau_{kz})/2 + g_n S_k (1 - \tau_{kz})/2]$$

$$F = \sum_k \tau_{k\pm} ; \text{GT} = \sum_k \sigma_k \tau_{k\pm}$$

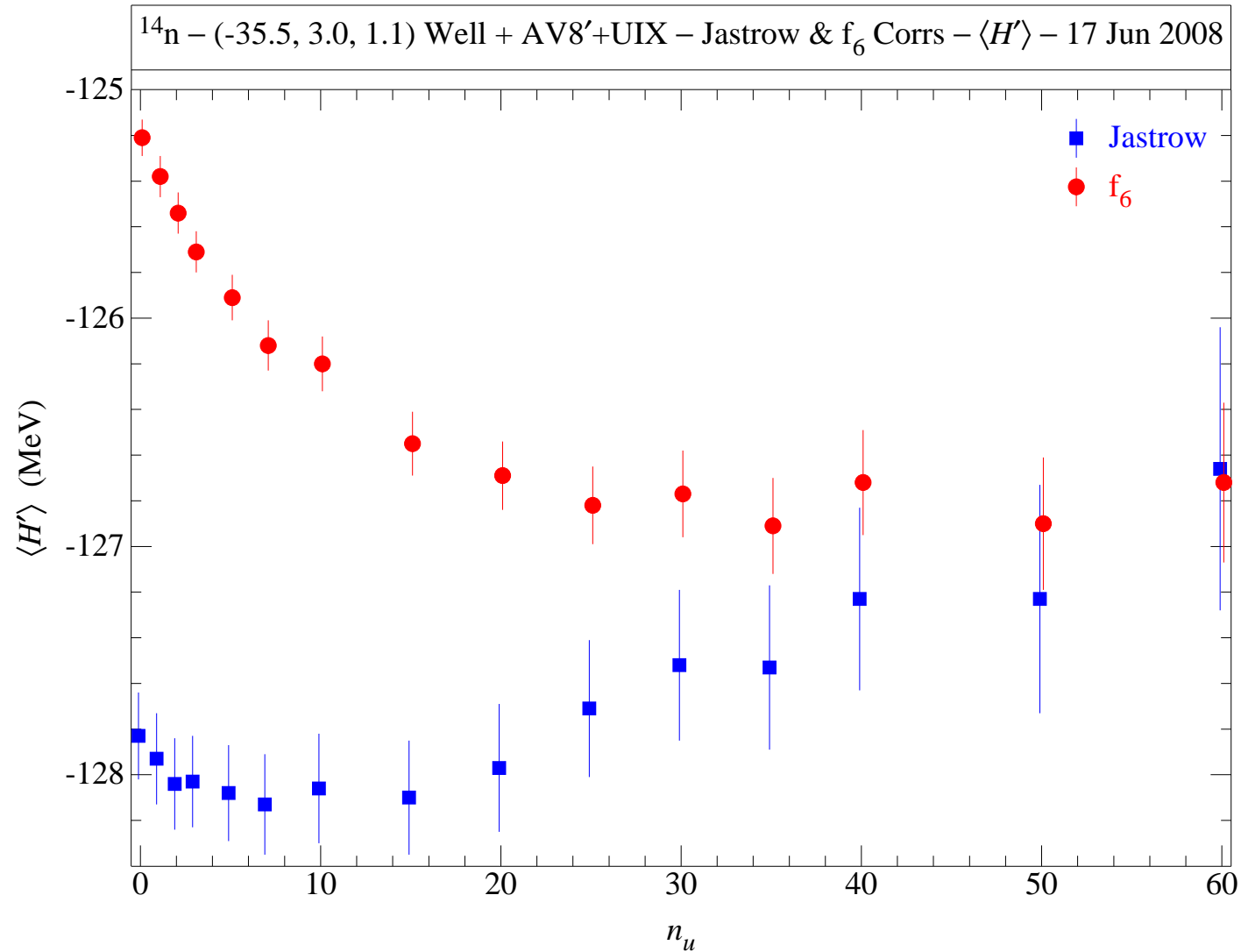
No effective charges or
effective nucleon magnetic moments!

Pervin, Pieper & Wiringa, PRC 76, 064319 (2007)

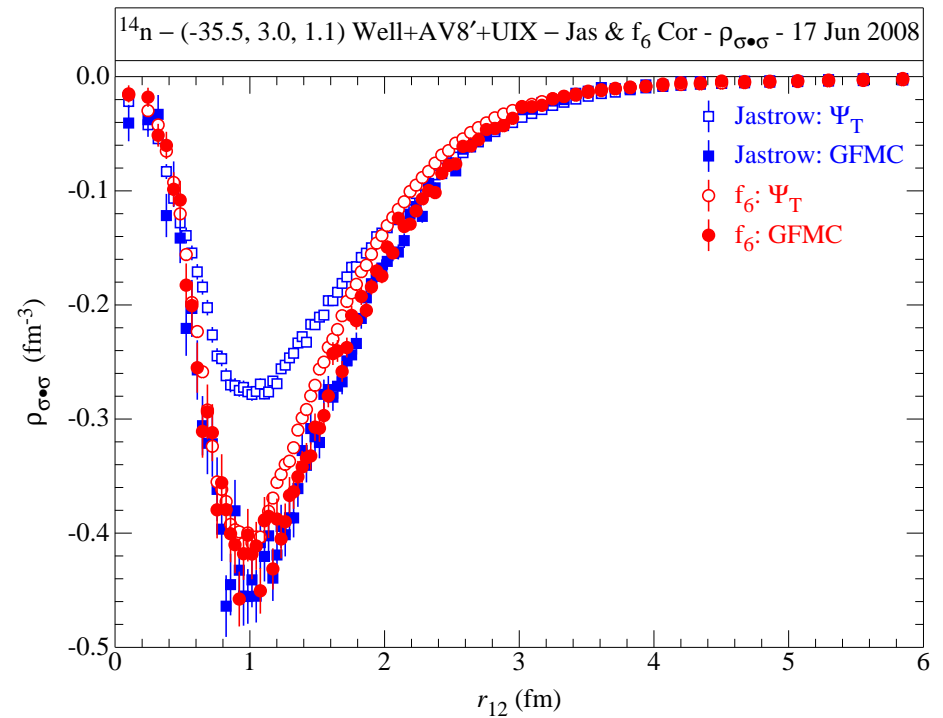
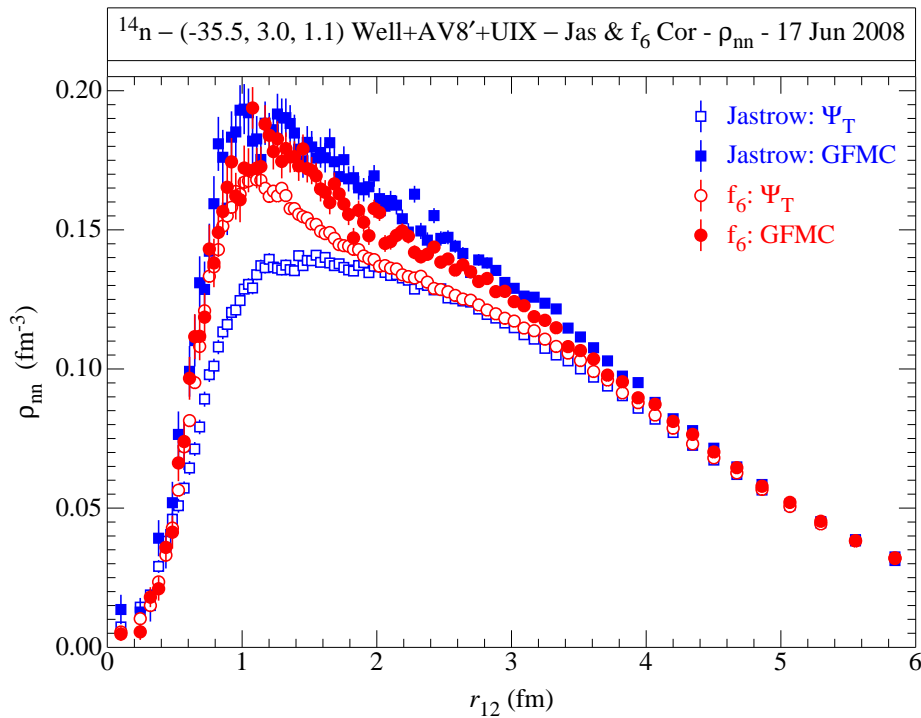


NEUTRON DROPS - GFMC CONSTRAINED-PATH CONVERGENCE

- We use a number of unconstrained GFMC steps before computing energies
- Usually 10–20 unconstrained steps are adequate
- ^8He and neutron drops require more.

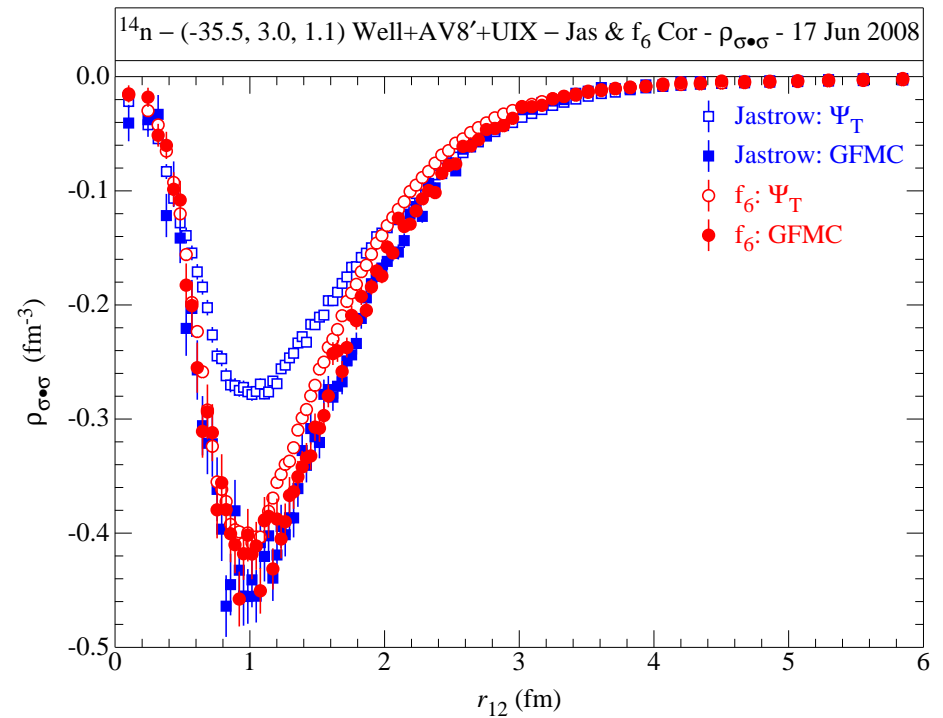
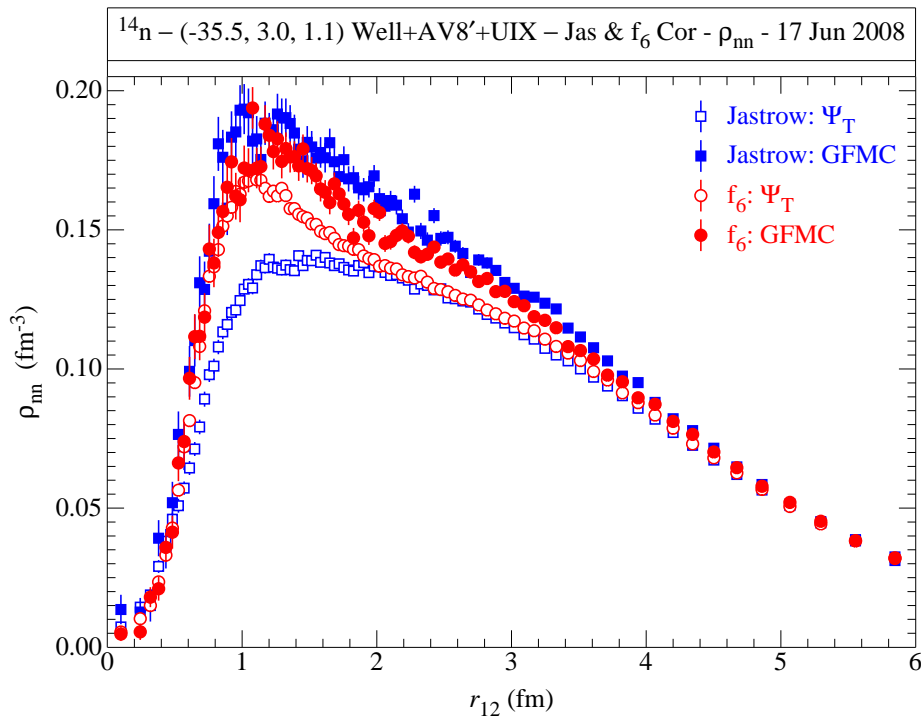


TWO-NEUTRON PAIR AND $\sigma \cdot \sigma$ DENSITY DISTRIBUTIONS



GFMC produces same final densities starting from very different Ψ_T densities

TWO-NEUTRON PAIR AND $\sigma \cdot \sigma$ DENSITY DISTRIBUTIONS



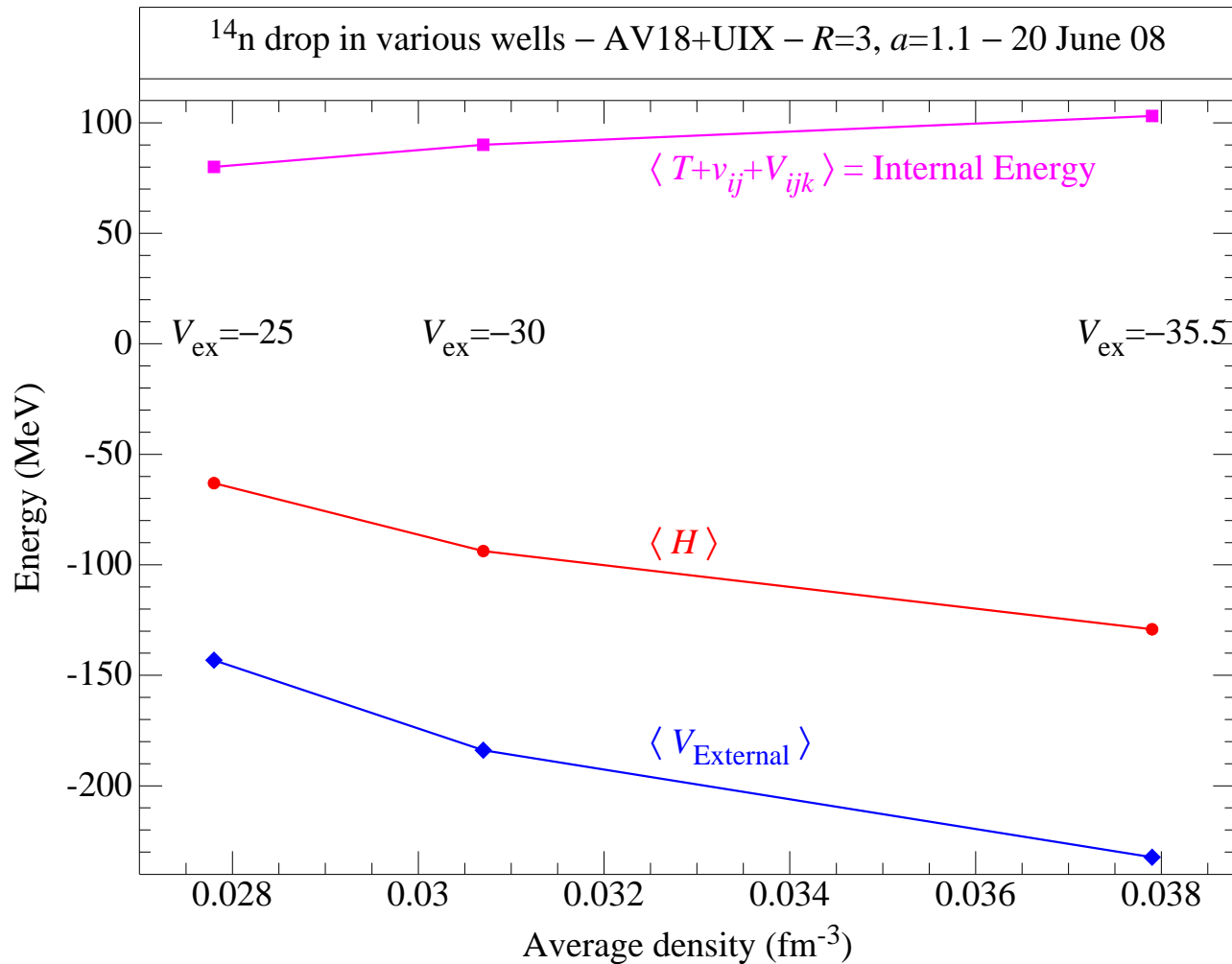
GFMC produces same final densities starting from very different Ψ_T densities

NEUTRON DROPS – DEPENDENCE ON EXTERNAL WELL DEPTH

Calculations of 14 neutrons with AV18+UIX and three external wells:

$R = 3.0$ fm; $a = 1.1$ fm; $V_{\text{ex}} = 25, 30, \& 35.5$ MeV

The computed energies have a (slightly) nonlinear dependence on the average density

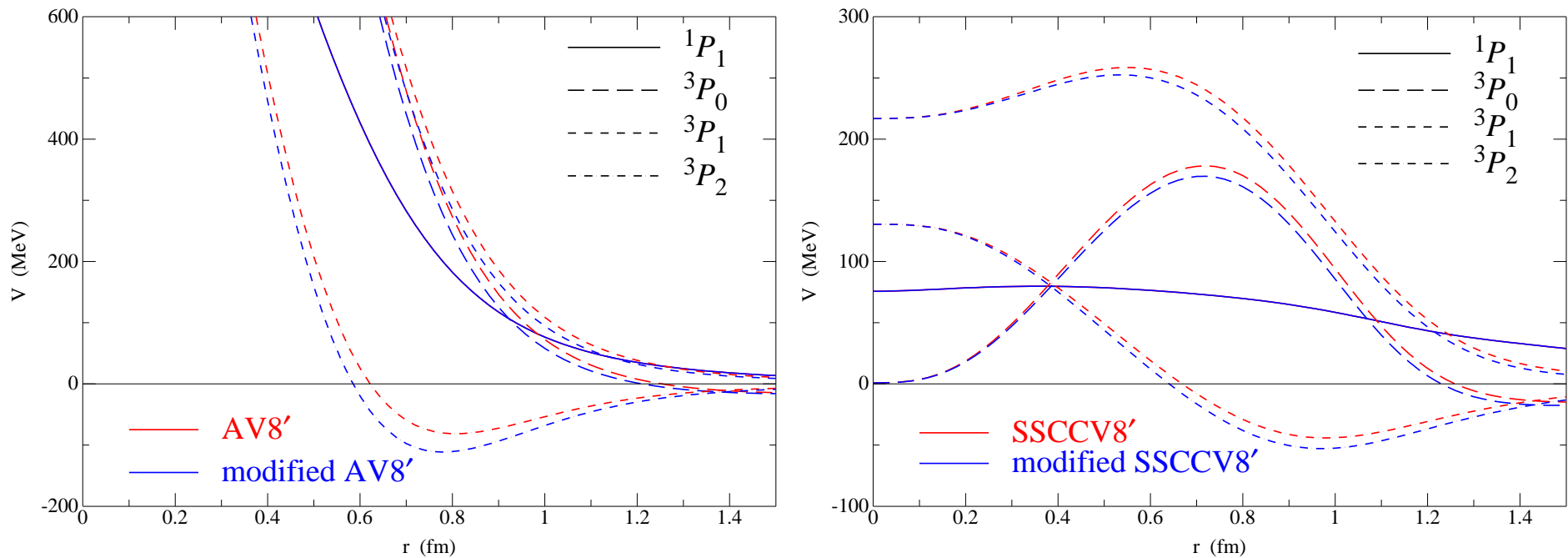


NN BENCHMARK POTENTIALS

The 2001 ${}^4\text{He}$ benchmark paper* used AV8' with no V_{ijk} as the Hamiltonian
Ultimately we want a ${}^{12}\text{C}$ benchmark using AV8' and AV8' with some V_{ijk} (UIX or TM').
However it was felt that

- 1) There is a need for intermediate benchmarks (perhaps ${}^6\text{Li}$, ${}^6\text{He}$, ${}^8\text{He}$).
- 2) A softer NN potential, such as SSCC, would also be desirable.

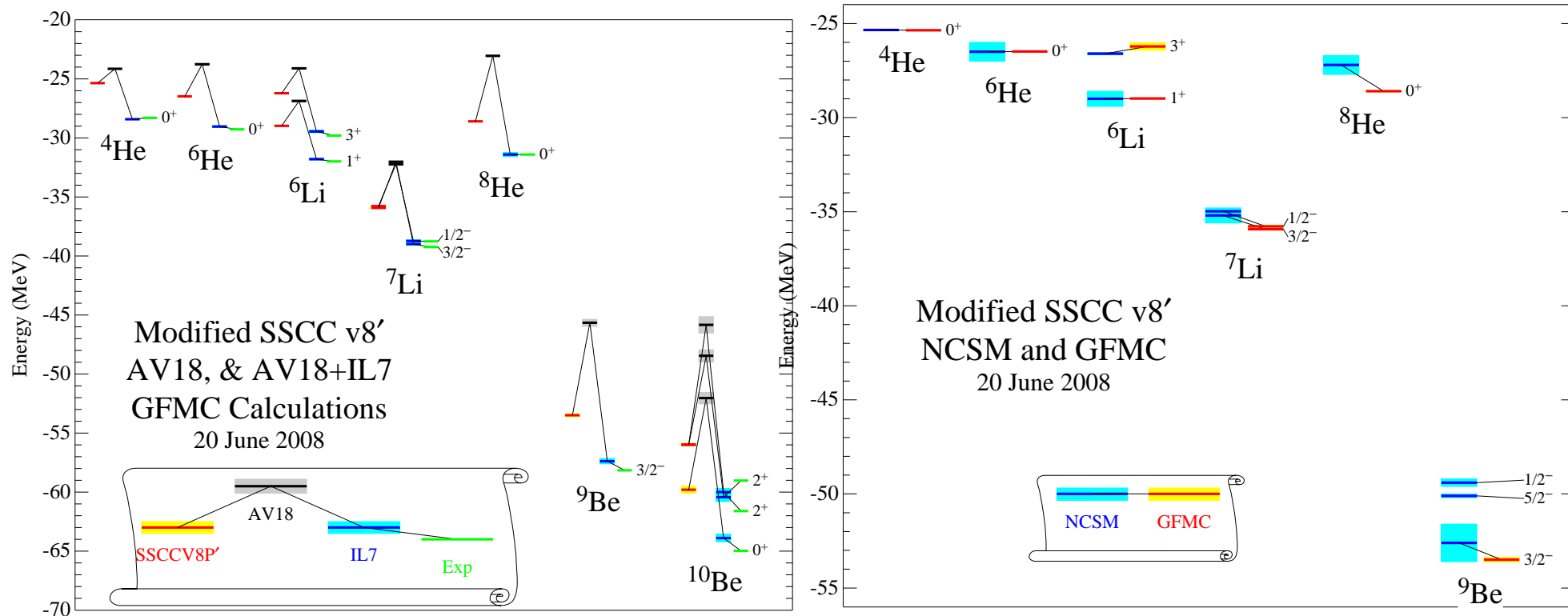
The v'_8 projection of the SSCC had to be modified to produce binding of $A = 6 - 8$ nuclei:



No changes in all even and singlet-odd partial waves

*H. Kamada *et al.*, Phys. Rev. C **64**, 044001 (2001)

NN BENCHMARK POTENTIALS



- The modified SSCC v_8' qualitatively reproduces experimental binding energies
 - Spin-orbit splittings are too small
- NCSM and GFMC energies are in general agreement within the quoted error bars.

OTHER NCSM AND GFMC RESULTS USING MODIFIED SSCC V8'

	$\langle r_p^2 \rangle^{1/2}$		Quadrupole Moment		Magnetic Moment*	
	NCSM	GFMC	NCSM	GFMC	NCSM	GFMC
^4He	1.51(1)	1.51(2)				
^6Li	2.33(5)	2.55(4)	0.00(5)	+0.1(1)		
^6He	1.88(5)	1.96(4)				
^7Li	2.24(5)	2.42(4)	≤ -2.85	-3.9(5)	+3.01(3)	+2.879(3)
^8He	1.85(5)	1.83(5)				
^9Be	2.32(5)	2.46(5)				

*Exchange currents not included