INT Structure of light nuclei, Oct. 2012, Seattle, WA

# Ab inito calculations of Be isotopes with JISP16

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#### PetaApps award

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INCITE award – Computational Nuclear Structure lead PI: James P Vary (ISU)

NERSC CPU time









## Ab initio nuclear physics – Quantum many-body problem

Given a Hamiltonian operator

$$\hat{\mathbf{H}} = \sum_{i < j} \frac{(\vec{p}_i - \vec{p}_j)^2}{2 \, m \, A} + \sum_{i < j} V_{ij} + \sum_{i < j < k} V_{ijk} + \dots$$

solve the eigenvalue problem for wave function of A nucleons

$$\mathbf{\hat{H}} \Psi(r_1, \dots, r_A) = \lambda \Psi(r_1, \dots, r_A)$$

Carbon 12 Proton Densities

 $\checkmark$  eigenvalues  $\lambda$  discrete (quantized) energy levels

• eigenvectors:  $|\Psi(r_1, \ldots, r_A)|^2$  probability density for finding nucleons 1, ..., A at  $r_1, \ldots, r_A$ 



## Ab initio nuclear physics – Computational challenges

- Self-bound quantum many-body problem, with 3A degrees of freedom in coordinate (or momentum) space
- Not only 2-body interactions, but also intrinsic 3-body interactions and possibly 4- and higher N-body interactions
- Strong interactions, with both short-range and long-range pieces
- Uncertainty quantification for calculations needed
  - for comparisons with experiments
  - for comparisons between different methods
- Sources of numerical uncertainty
  - statistical and round-off errors
  - systematical errors inherent to the calculational method
    - CI methods: finite basis space
    - Monte Carlo methods: sensitivity to the trial wave function
    - Lattice calculations: finite volume and lattice spacing
  - uncertainty of the nuclear potential

## Nuclear interaction

Nuclear potential not well-known ...

though in principle calculable from Quantum Chromo Dynamics

$$\hat{\mathbf{H}} = \hat{\mathbf{T}}_{\mathsf{rel}} + \sum_{i < j} V_{ij} + \sum_{i < j < k} V_{ijk} + \dots$$

In practice, alphabet of realistic potentials

- Argonne potentials: AV8', AV18
  - plus Urbana 3NF (UIX)
  - plus Illinois 3NF (IL7)
- Bonn potentials
- **Chiral NN interactions** 
  - plus chiral 3NF, ideally to the same order







## **Phenomeological** NN interaction: JISP16

J-matrix Inverse Scattering Potential tuned up to <sup>16</sup>O

- Constructed to reproduce np scattering data
- Finite rank seperable potential in H.O. representation
- Nonlocal NN-only potential
- Use Phase-Equivalent Transformations (PET) to tune off-shell interaction to
  - binding energy of <sup>3</sup>H and <sup>4</sup>He
  - Iow-lying states of <sup>6</sup>Li (JISP6, precursor to JISP16)
  - binding energy of <sup>16</sup>O





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Physics Letters B 644 (2007) 33-37

Realistic nuclear Hamiltonian: Ab exitu approach

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## J-matrix Inverse Scattering Potentials

- Constructed as matrix in H.O. basis
  - $2n + l \le 8$  for even partial waves, limited to  $J \le 4$
  - $2n + l \le 9$  for odd partial waves, limited to  $J \le 4$
  - $\hbar\omega = 40 \text{ MeV}$
- $\checkmark$   $\chi^2$ /datum of 1.05 for the 1999 np data base (3058 data)
- No charge symmetry breaking
- Use PET to improve
  - deuteron quadrupole moment
  - <sup>3</sup>H and <sup>4</sup>He binding energies
  - binding energies low-lying states of <sup>6</sup>Li: JISP6

Shirokov, Vary, Mazur, Zaystev, Weber, PLB 621, 96 (2005)

binding energy of <sup>16</sup>O: JISP16

Shirokov, Vary, Mazur, Weber, PLB 644, 33 (2007)

additional tuning, more accurate calculations: JISP16<sub>2010</sub> reproduces <sup>16</sup>O within numerical error estimates of 3% Shirokov, Kulikov, Maris, Mazur, Mazur, Vary, arXiv:0912.2967

## JISP16 results for few-body systems

deuteron properties						
	Е	(MeV)	$r_p$ (fm)	${\cal Q}$ (e fm $^2$ )	$\mathcal{A}_s$ (fm $^{-rac{1}{2}}$ )	$\mathcal{A}_d/\mathcal{A}_s$
expt.	-2.224575		1.971(6)	0.2859(3)	0.8846(9)	0.0256(4)
JISP16	-2	2.224575	1.964	0.2886	0.8629	0.0252
AV18	-2.224575		1.967	0.270	0.8850	0.0250
selected $A = 3$ and 4 results						
		$E_b(^3H$	) $\mu(^{3}H)$	$\mu$ ( $^3$ He)	$E_b(^4He)$	
expt.		8.482	2.979	-2.128	28.296	
JISP16		8.369(2)	2.667	-1.819	28.299	
AV18		7.61(1)			24.07(4)	
AV18+IL2		8.43(1)	2.568(1)	-1.762(1)	28.37(3)	

Pieper, Wiringa, Annu. Rev. Nucl. Part. Sci. 51, 53 (2001)

## Many-Body systems

**Configuration Interaction methods** 

- Expand wave function in basis states  $|\Psi\rangle = \sum a_i |\Phi_i\rangle$
- Express Hamiltonian in basis  $\langle \Phi_j | \hat{\mathbf{H}} | \Phi_i \rangle = H_{ij}$
- Diagonalize Hamiltonian matrix  $H_{ij}$
- Complete basis  $\longrightarrow$  exact result
  - caveat: complete basis is infinite dimensional
- In practice
  - truncate basis
  - study behavior of observables as function of truncation
- Computational challenge
  - construct large ( $10^{10} \times 10^{10}$ ) sparse symmetric real matrix  $H_{ij}$
  - use Lanczos algorithm to obtain lowest eigenvalues & eigenvectors

## **Basis space expansion**

- Expand wave function in basis states  $|\Psi\rangle = \sum a_i |\Phi_i\rangle$
- Many-Body basis states  $|\Phi_i\rangle$  Slater Determinants of Single-Particle states  $|\phi\rangle$   $|\phi_i(r_1) \phi_i(r_2)| = \phi_i(r_1)$

$$\Phi_{i}(r_{1},...,r_{A}) = \frac{1}{\sqrt{(A!)}} \begin{vmatrix} \phi_{i1}(r_{1}) & \phi_{i2}(r_{1}) & \dots & \phi_{iA}(r_{1}) \\ \phi_{i1}(r_{2}) & \phi_{i2}(r_{2}) & \dots & \phi_{iA}(r_{2}) \\ \vdots & \vdots & & \vdots \\ \phi_{i1}(r_{A}) & \phi_{i2}(r_{A}) & \dots & \phi_{iA}(r_{A}) \end{vmatrix}$$

- Single-Particle basis states
  - eigenstates of  $\hat{\mathbf{L}}^2$ ,  $\hat{\mathbf{S}}^2$ ,  $\hat{\mathbf{J}}^2$ , and  $\hat{\mathbf{J}}_{\mathbf{z}}$ labelled by quantum numbers  $|n, l, s, j, m\rangle$
  - radial wavefunctions
    - Harmonic Oscillator
    - Wood–Saxon basis
    - Coulomb–Sturmian
    - Berggren

**\_** . . .

Negoita, PhD thesis 2010 Caprio, Maris, Vary, PRC86, 034312 (2012) Rotureau, last week

#### **Truncation scheme**

M-scheme: Many-Body basis states eigenstates of  $\hat{J}_z$ 

$$\hat{\mathbf{J}}_{\mathbf{z}}|\Phi_i\rangle = M|\Phi_i\rangle = \sum_{k=1}^A m_{ik}|\Phi_i\rangle$$

- single run gives spectrum
- alternatives: LS scheme, Coupled-J scheme, Symplectic basis, ...
- *N*max truncation: Many-Body basis states satisfy

$$\sum_{k=1}^{A} \left( 2 n_{ik} + l_{ik} \right) \leq N_0 + N_{\max}$$

- exact factorization of Center-of-Mass motion
- alternatives:

No-Core Monte-Carlo Shell Model, Importance Truncation, FCI (truncation on single-particle basis only), ...

#### Intermezzo: FCI vs. Nmax truncation





- exact factorization of Center-of-Mass motion
- converges much more rapidly than FCI truncation with basis space dimension
- Infinite basis space limit: No-Core Full Configuration (NCFC)

## Intermezzo: Center-of-Mass excitations

- Use single-particle coordinates, not relative (Jacobi) coordinates
  - straightforward to extend to many particles
  - have to seperate Center-of-Mass motion from internal motion
- Center-of-Mass wave function factorizes for H.O. basis functions in combination with N<sub>max</sub> truncation

$$\begin{aligned} |\Psi_{\mathsf{total}}\rangle &= |\phi_1\rangle \otimes \ldots \otimes |\phi_A\rangle \\ &= |\Phi_{\mathsf{Center-of-Mass}}\rangle \otimes |\Psi_{\mathsf{int}}\rangle \end{aligned}$$

where

$$\mathbf{\hat{H}}_{\mathsf{rel}} | \Psi_{\mathsf{j, int}} \rangle = E_{\mathsf{j}} | \Psi_{\mathsf{j, int}} \rangle$$

Add Lagrange multiplier to Hamiltonian (Lawson term)

$$\hat{\mathbf{H}}_{\mathsf{rel}} \longrightarrow \hat{\mathbf{H}}_{\mathsf{rel}} + \Lambda_{CM} \left( \hat{\mathbf{H}}_{CM}^{H.O.} - \frac{3}{2} \left( \sum_{i} m_{i} \right) \omega \right)$$

with  $\hat{\mathbf{H}}_{rel} = T_{rel} + V_{rel}$  the relative Hamiltonian

seperates CM excitations from CM ground state  $|\Phi_{CM}\rangle$ 

## **Configuration Interaction Methods**

- Expand wave function in basis states  $|\Psi\rangle = \sum a_i |\psi_i\rangle$
- Express Hamiltonian in basis  $\langle \psi_j | \mathbf{\hat{H}} | \psi_i \rangle = H_{ij}$
- Diagonalize Hamiltonian matrix  $H_{ij}$
- Variational: for any finite truncation of the basis space, eigenvalue is an upper bound for the ground state energy
- Smooth approach to asymptotic value with increasing basis space: No-Core Full Configuration calculation
- Convergence: independence of  $N_{\text{max}}$  and H.O. basis  $\hbar\omega$ 
  - different methods (NCFC, CC, GFMC, ...) using the same interaction should give same results within (statistical plus systematic) numerical uncertainties



#### No-Core CI calculations – main challenge



- $\blacksquare$  Increase of basis space dimension with increasing A and  $N_{max}$
- More relevant measure for computational needs
  - number of nonzero matrix elements

Challenge: achieve numerical convergence for no-core Full Configuation calculations using finite model space calculations

- $\blacksquare$  Renormalize interaction  $\longrightarrow$  effective interaction  $V_{\text{eff}}$ 
  - can improve quality of results in small model spaces
- Caveats
  - induces many-body forces
    - induced 3-body forces are often neglected
    - induced 4-, 5-, ..., A-body forces are always neglected
  - variational principle applicable to renormalized Hamiltonian not to original (bare) Hamiltonian
  - often complicates extrapolation to asymptotic values
  - need to renormalize operators as well
- Commonly used renormalization procedures
  - Lee–Suzuki effective interaction
  - Similarity Renormalization Group (in particular in combination with chiral interactions)

## **Results with Lee–Suzuki renormalization for JISP16**



Ground state energy of <sup>16</sup>O expected to be between variational upper bound without renormalization and lower bound (?) from Lee–Suzuki renormalized interaction

Used in tuning of JISP16 Shirokov, Vary, Mazur, Weber, PLB 644, 33 (2007)

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## Convergence Lee–Suzuki renormalization not monotonic



16O JISP16 bare and Lee-Suzuki Veff truncated to 2-body clusters

- Lee—Suzuki result for ground state energy not a lower bound
- JISP16 overbinds <sup>16</sup>O by 10% to 15%

Maris, Vary, Shirokov, PRC79, 014308 (2009)

Challenge: achieve numerical convergence for no-core Full Configuation calculations using finite model space calculations

- Perform a series of calculations with increasing N<sub>max</sub> truncation
- Extrapolate to infinite model space  $\longrightarrow$  exact results
  - Empirical: binding energy exponential in Nmax

 $E_{\text{binding}}^{N} = E_{\text{binding}}^{\infty} + a_1 \exp(-a_2 N_{\text{max}})$ 

- use 3 or 4 consecutive  $N_{\text{max}}$  values to determine  $E_{\text{binding}}^{\infty}$
- use  $\hbar \omega$  and  $N_{max}$  dependence to estimate numerical error bars

Maris, Shirokov, Vary, PRC79, 014308 (2009)

- Recent studies of IR and UV behavior
  - exponentials in  $\sqrt{\hbar\omega/N}$  and  $\sqrt{\hbar\omega N}$  Coon *et al*, arXiv:1205.3230;

Furnstahl, Hagen, Papenbrock PRC86, 031301(R) (2012)

## **Extrapolating to complete basis – in practice**

- Perform a series of calculations with increasing N<sub>max</sub> truncation
- **J** Use empirical exponential in  $N_{max}$ :





• H.O. basis up to  $N_{max} = 16$ :  $E_b = -31.49(3)$  MeV

Cockrell, Maris, Vary, PRC86 034325 (2012)

Hyperspherical harmonics up to  $K_{max} = 14$ :  $E_b = -31.46(5)$  MeV
Vaintraub, Barnea, Gazit, PRC79 065501 (2009)

## Accelerating convergence – Coulomb-Sturmian basis

Asymtotic behavior

- H.O. basis  $\exp(-a r^2)$
- Coulomb–Sturmian basis  $\exp(-cr)$
- Disadvantage
  - no exact factorization of Center-of-Mass motion
  - in practice,
     approximate factorization
     Hagen, Papenbrock, Dean,
     PRL103, 062503 (2009)
  - can use Lagrange multiplier to remove spurious state

Caprio, Maris, Vary, PRC86, 034312 (2012)



### **Coulomb-Sturmian – binding energies**



- at N<sub>max</sub> = 4 further from convergence than H.O. basis
- extrapolate to the same results as H.O. basis
- dashed line: extrapolated result from N<sub>max</sub> = 16 calculations in H.O. basis

Cockrell, Maris, Vary, PRC86 034325 (2012)

## **Coulomb-Sturmian – radius**

Caprio, Maris, Vary, PRC86, 034312 (2012)



- exponential extrapolation does not work for radii in H.O. basis
- exponential extrapolation seems to work for radii in C.S. basis
- **best estimate based on**  $N_{max} = 16$  H.O. calculations: 2.3 fm

Cockrell, Maris, Vary, PRC86 034325 (2012)

experimental point-proton radius: 2.45 fm

#### Ground state energy Be-isotopes with JISP16



## 7Be – Ground state properties



- Binding energy converges monotonically, with optimal H.O. freugency around  $\hbar\omega = 20$  MeV to 25 MeV
- Ground state about 0.7 MeV underbound with JISP16
- Proton point radius does not converge monotonically
  - Coulomb–Sturmian basis likely to improve convergence

## 7Be – Proton density

Translationally-invariant density – center-of-mass motion taken out w. Cockrell, PhD thesis 2012



- Slow build up of asymptotic tail of wavefunction
- Proton density appears to converge more rapidly at  $\hbar \omega = 12.5$  MeV than at 20 MeV because long-range part of wavefunction is better represented with smaller H.O. parameter

## 7Be – Proton radius



• Calculation one-body observables  $\langle i|\mathcal{O}|j\rangle \sim \int \mathcal{O}(r) r^2 \rho_{ij}(r) dr$ 

- **P** RMS radius:  $\mathcal{O}(r) = r^2$
- Slow convergence of RMS radius due to slow build up of asymptotic tail
- Ground state RMS radius in agreement with data

## 7Be – Quadrupole moment



- Ground state quadrupole moment in agreement with data
- Optimal basis space around  $\hbar\omega = 10$  MeV to 12 MeV
- Similar slow convergence for E2 transitions

## 7Be – Excited states



- Excitation energy of narrow states
  - converge rapidly
  - agree with experiments
- Broad resonances depend  $\hbar\omega$

- Magnetic moments well converged
  - 2-body currents needed for agreement with data (meson-exchange currents)

#### 7Be – Emergence of rotational band? in progress, w. M. Caprio

E2 observables suggest rotational structure for  $\frac{3}{2}$ ,  $\frac{1}{2}$ ,  $\frac{7}{2}$ ,  $\frac{5}{2}$  states



$$Q(J) = \frac{\frac{3}{4} - J(J+1)}{(J+1)(2J+3)} Q_0$$
$$B(E2; i \to f) = \frac{5}{16\pi} Q_0^2 \left( J_i, \frac{1}{2}; 2, 0 \middle| J_f, \frac{1}{2} \right)^2$$

## 7Be – Structure of $(\frac{5}{2}^-, \frac{1}{2})_1$ (broad) and $(\frac{5}{2}^-, \frac{1}{2})_2$ (narrow) states



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## 9Be – Ground state properties



- Convergence pattern natural and unnatural parity looks similar
- **9** Ground state about  $1.0 \pm 0.2$  MeV underbound with JISP16
- $\checkmark$  Lowest unnatural parity state underbound by about  $2.7 \pm 0.8$  MeV
  - need next basis space for unnatural parity
  - need improved interaction?

## Positive vs. negative parity states of Be-isotopes



- Unnatural parity states systematically underbound by about
   1 MeV to 2 MeV compared to lowest natural parity states
  - interaction JISP16 not good enough?
  - difference in convergence of pos. and neg. parity states?

### 9Be – Positive and negative spectrum



- **Solution** Excitation energy  $\frac{5}{2}^{-}$  at 3 MeV well converged (narrow)
- Excitation energy  $\frac{7}{2}^-$  reasonably converged
- Excitation energies broad neg. parity not well converged
- Excitation energies pos. parity well converged

#### 9Be – Emergence of rotational bands in progress, w. M. Caprio



- Solutional energy for states with axial symmetry  $E(J) \propto J(J+1)$
- Quadrupole moments for rotational band

$$Q(J) = \frac{3K^2 - J(J+1)}{(J+1)(2J+3)} Q_0$$

Quadrupole moments not converged, but ratio of quadrupole moments agree with rotational band structure

## **9Be – Structure: Density** $(\frac{3}{2}^{-}, \frac{1}{2})$ **ground state**

x (fm)



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#### Rotational bands odd Be isotopes in preparation, w. M. Caprio



Quadrupole moments not converged, but ratio of quadrupole moments agree with rotational band structure

#### Rotational bands odd Be isotopes in preparation, w. M. Caprio

Also for the unnatural parity states



Quadrupole moments not converged, but ratio of quadrupole moments agree with rotational band structure

#### Rotational bands even Be isotopes in preparation, w. M. Caprio





#### B(E2) transistions Be isotopes in preparation, w. M. Caprio



Ratio's B(E2)/ $Q^2$  in agreement with rotational structure as well

## **Conclusions**

- No-core Configuration Interaction nuclear structure calculations
  - Binding energy, spectrum
  - $\langle r^2 \rangle$ ,  $\mu$ , Q, transitions, wfns, one-body densities
- Main challenge: construction and diagonalization of extremely large (D > 1 billion) sparse matrices
- Need realistic basis function to improve convergence  $\langle r^2 \rangle$ , Q

#### JISP16

- Nonlocal phenomenological 2-body interaction
- Good description of a range of light nuclei
- Rapid convergence for binding energies
- Emergence of rotational bands and clustering in Be-isotopes
- Would not have been possible without collaboration with applied mathematicians and computer scientists Aktulga, Yang, Ng (LBNL); Çatalyürek, Saule (OSU); Sosonkina (ODU/AL)