



Ab initio α - α scattering

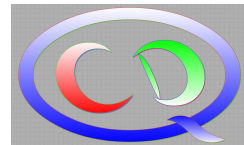
Ulf-G. Meißner, Univ. Bonn & FZ Jülich

Supported by DFG, SFB/TR-16 and by DFG, SFB/TR-110

and by CAS, PIFI

and by BMBF 05P15PCFN1

and by HGF VIQCD VH-VI-417



CONTENTS

- Short introduction
- Basics of nuclear lattice simulations
- Results from nuclear lattice simulations
- Ab initio calculation of α - α scattering
- Summary & outlook

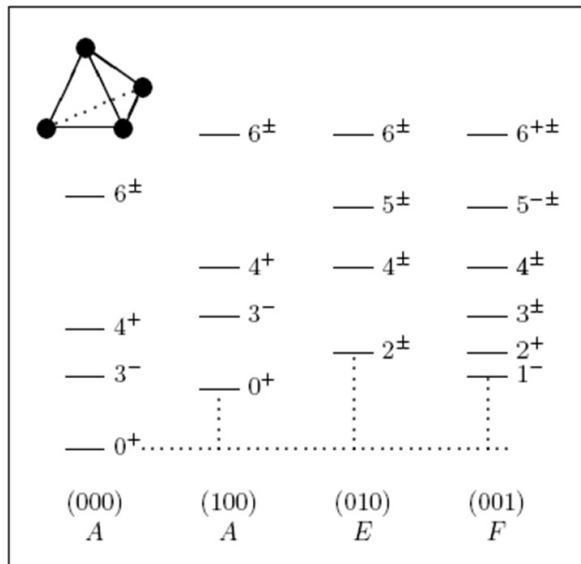
Short introduction

CLUSTERING in NUCLEI

- Introduced theoretically by Wheeler already in 1937:

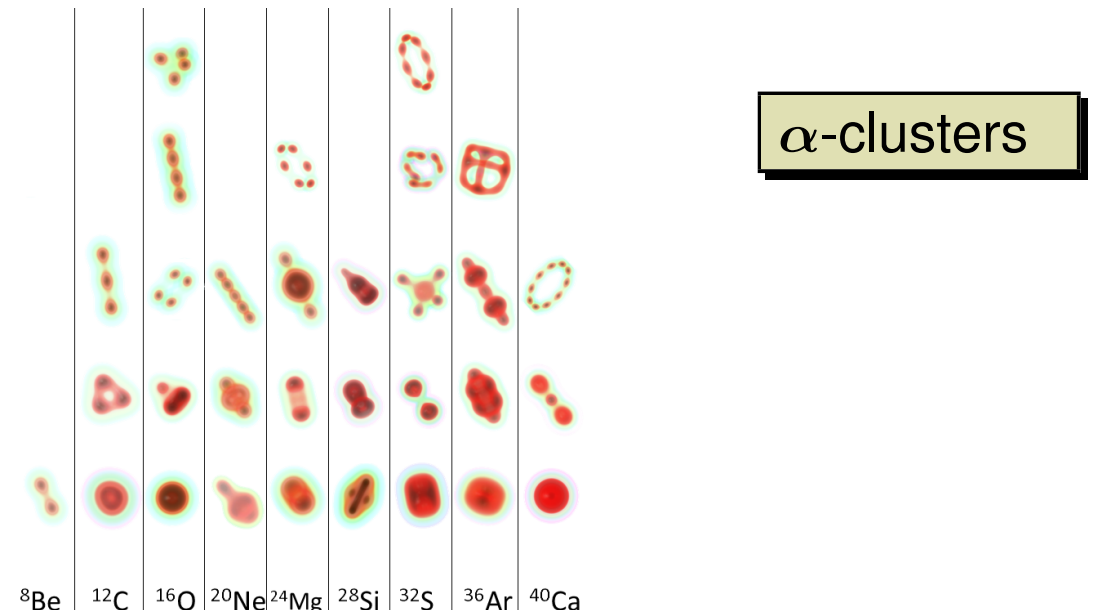
John Archibald Wheeler, “Molecular Viewpoints in Nuclear Structure,”
Physical Review **52** (1937) 1083

- many works since then...



Bijker, Iachello (2014)

Ikeda, Horiuchi, Freer, Schuck, Zhou, Khan, . . .



Ebran, Khan, Niksic, Vretenar (2014)

⇒ can we understand this phenomenon from *ab initio* calculations?

Basics of nuclear lattice simulations

for an easy intro, see: UGM, Nucl. Phys. News **24** (2014) 11

NUCLEAR LATTICE SIMULATIONS

Frank, Brockmann (1992), Koonin, Müller, Seki, van Kolck (2000) , Lee, Schäfer (2004), . . .
 Borasoy, Krebs, Lee, UGM, Nucl. Phys. **A768** (2006) 179; Borasoy, Epelbaum, Krebs, Lee, UGM, Eur. Phys. J. **A31** (2007) 105

- *new method* to tackle the nuclear many-body problem

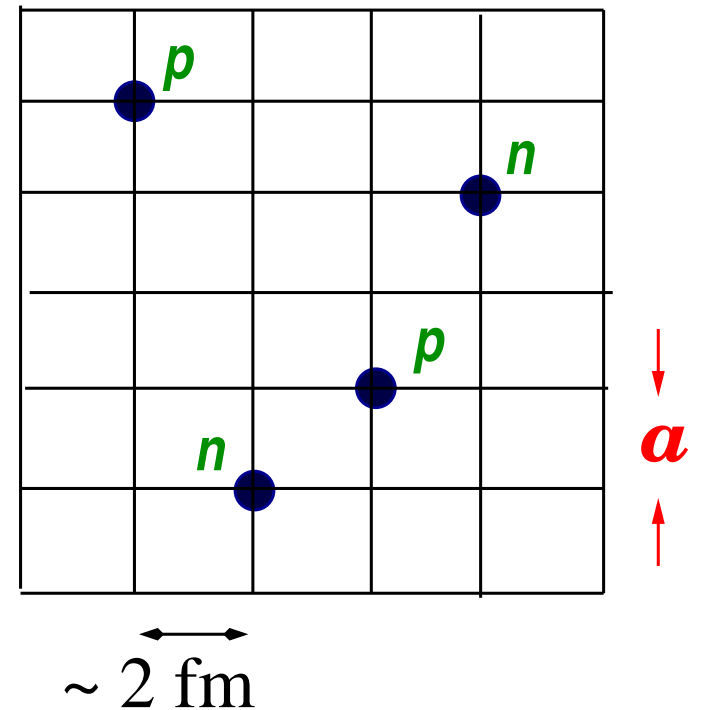
- discretize space-time $V = L_s \times L_s \times L_s \times L_t$:
 nucleons are point-like particles on the sites

- discretized chiral potential w/ pion exchanges
 and contact interactions + Coulomb

→ see Epelbaum, Hammer, UGM, Rev. Mod. Phys. 81 (2009) 1773

- typical lattice parameters

$$\Lambda = \frac{\pi}{a} \simeq 300 \text{ MeV} \text{ [UV cutoff]}$$



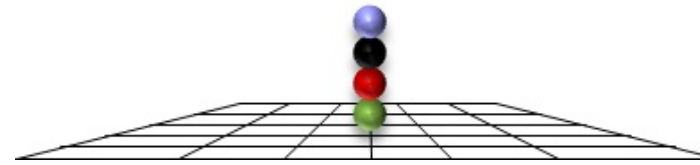
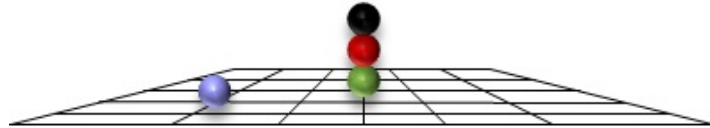
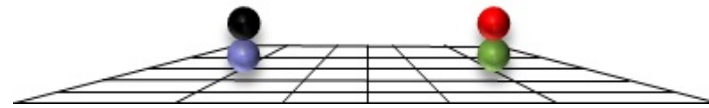
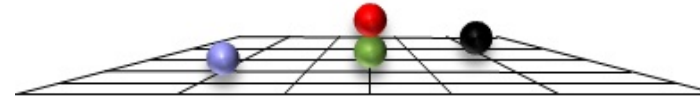
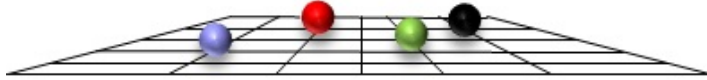
- strong suppression of sign oscillations due to approximate Wigner SU(4) symmetry

J. W. Chen, D. Lee and T. Schäfer, Phys. Rev. Lett. **93** (2004) 242302, T. Lähde et al., EPJA **51**: 92 (2015)

- hybrid Monte Carlo & transfer matrix (similar to LQCD)

CONFIGURATIONS

7



⇒ all possible configurations are sampled
⇒ clustering emerges naturally

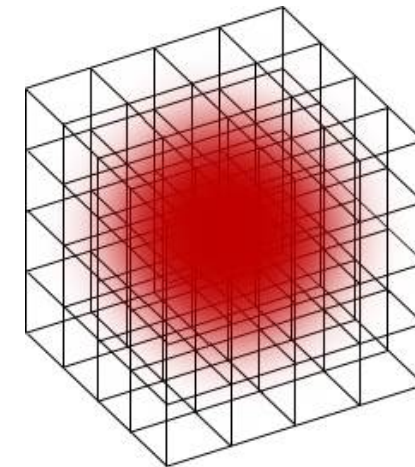
NUCLEAR WAVE FUNCTIONS

- General wave function:

$$\psi_j(\vec{n}) , \quad j = 1, \dots, A$$

- States with well-defined momentum (anti-symm.):

$$L^{-3/2} \sum_{\vec{m}} \psi_j(\vec{n} + \vec{m}) \exp(i \vec{P} \cdot \vec{m}) , \quad j = 1, \dots, A$$



- Insert clusters of nucleons at initial/final states (spread over some time interval)
 - allows for all type of wave functions (shell model, clusters, ...)
 - removes directional bias

shell-model type

$$\psi_j(\vec{n}) = \exp[-c\vec{n}^2]$$

$$\psi'_j(\vec{n}) = n_x \exp[-c\vec{n}^2]$$

$$\psi''_j(\vec{n}) = n_y \exp[-c\vec{n}^2]$$

$$\psi'''_j(\vec{n}) = n_z \exp[-c\vec{n}^2]$$

cluster type

$$\psi_j(\vec{n}) = \exp[-c(\vec{n} - \vec{m})^2]$$

$$\psi'_j(\vec{n}) = \exp[-c(\vec{n} - \vec{m}')^2]$$

$$\psi''_j(\vec{n}) = \exp[-c(\vec{n} - \vec{m}'')^2]$$

$$\psi'''_j(\vec{n}) = \exp[-c(\vec{n} - \vec{m}''')^2]$$

- shell-model w.f.s do not have enough 4N correlations $\sim \langle (N^\dagger N)^2 \rangle$

EXTRACTING PHASE SHIFTS on the LATTICE

9

- Lüscher's method:

Two-body energy levels below the inelastic threshold on a periodic lattice are related to the phase shifts in the continuum

Lüscher, Comm. Math. Phys 105 (1986) 153

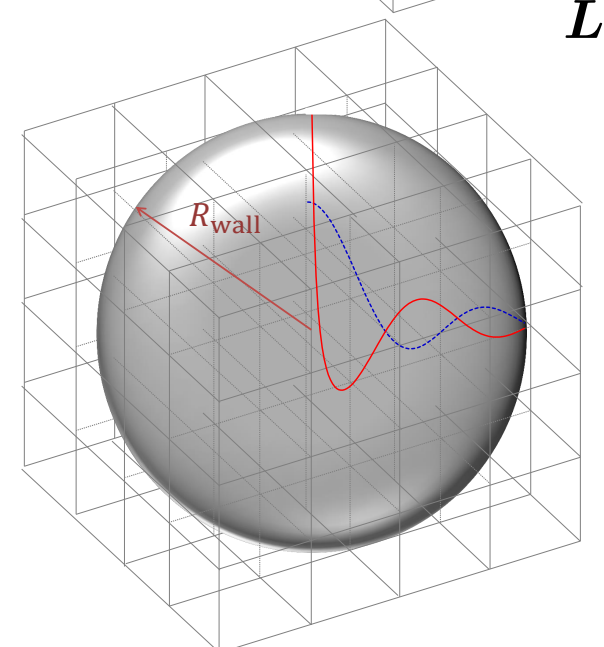
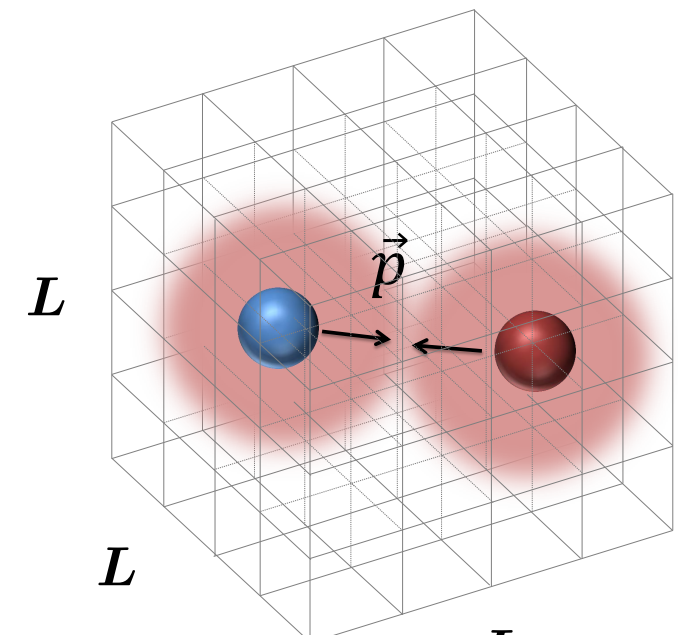
Lüscher, Nucl. Phys 354 (1991) 531

- Spherical wall method:

Impose a hard wall on the lattice and use the fact that the wave function vanishes for $r = R_{\text{wall}}$:

$$\psi_\ell(r) \sim [\cos \delta_\ell(p) F_\ell(pr) + \sin \delta_\ell(p) G_\ell(pr)]$$

Borasoy, Epelbaum, Krebs, Lee, UGM,
EPJA 34 (2007) 185



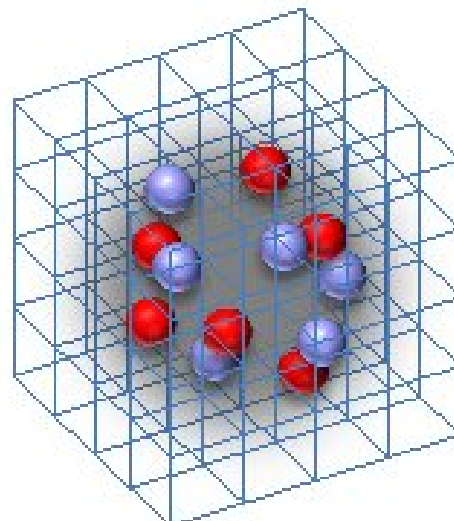
COMPUTATIONAL EQUIPMENT

- Present = JUQUEEN (BlueGene/Q)



6 Pflops

Lattice: some results



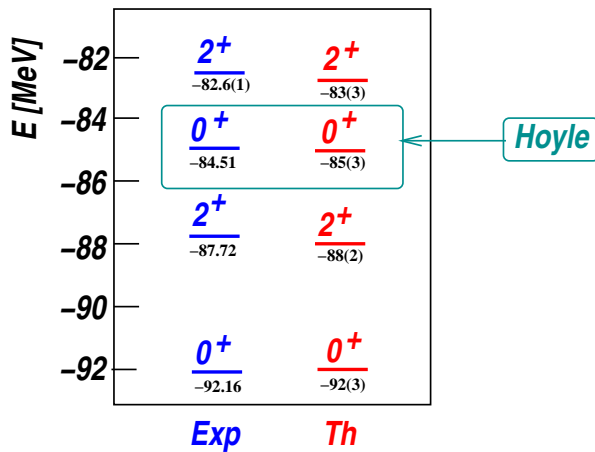
NLEFT

Epelbaum, Krebs, Lähde, Lee, Luu, UGM, Rupak + post-docs + students

RESULTS from LATTICE NUCLEAR EFT

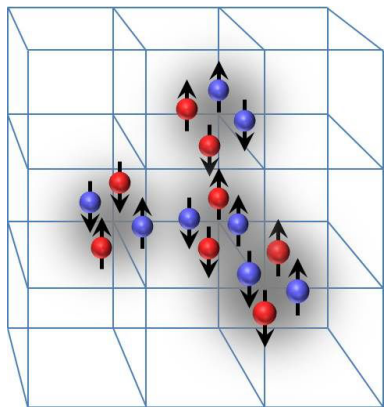
- Hoyle state in ^{12}C

PRL 106 (2011)



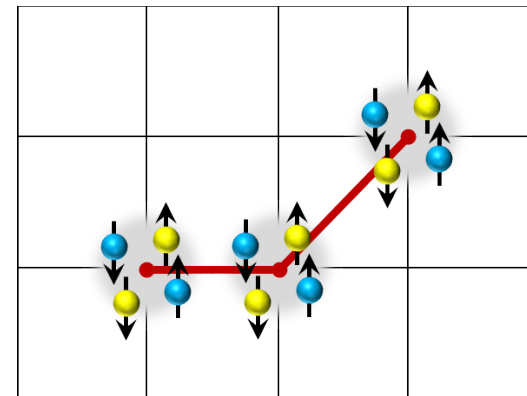
- Spectrum of ^{16}O

PRL 112 (2014)



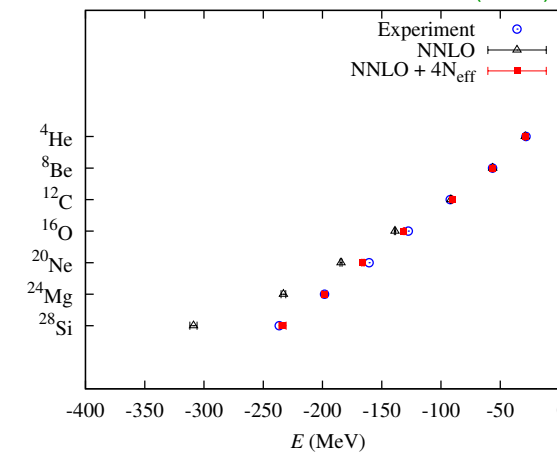
- Structure of the Hoyle state

PRL 109 (2012)



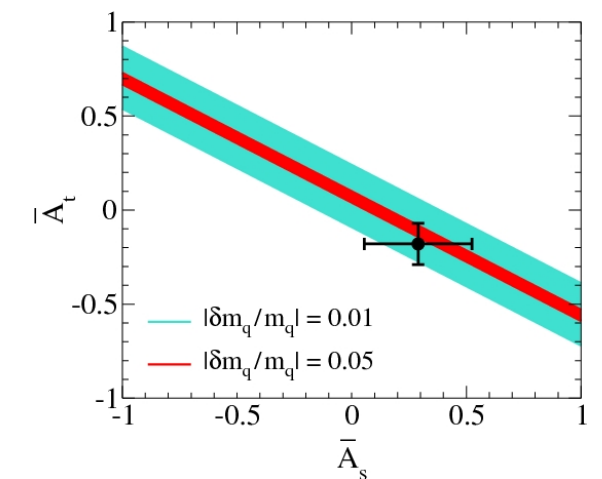
- Going up the α -chain

PLB 732 (2014)



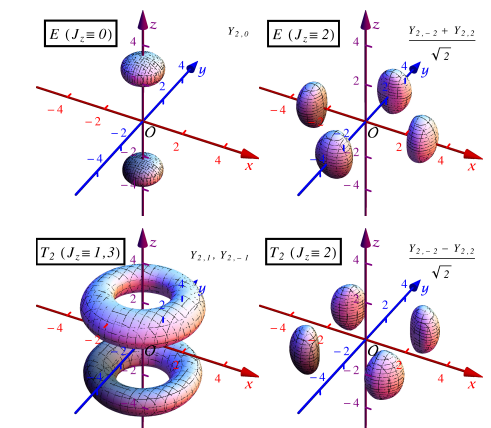
- Fate of carbon-based life

PRL 110 (2013), EPJ A49 (2013)



- Rot. symmetry breaking

PRD 90 (2014), PRD 92 (2015)



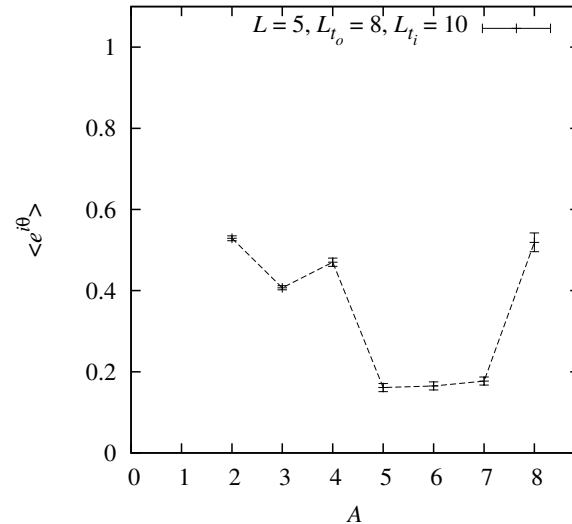
SYMMETRY-SIGN EXTRAPOLATION METHOD

Epelbaum, Krebs, Lähde, Lee, Luu, UGM, Rupak, EPJA 51: 92 (2015)

- so far: nuclei with $N = Z$, and $A = 4 \times \text{int}$
as these have the least sign problem
due to the approximate SU(4) symmetry

$$\langle \text{sign} \rangle = \langle \exp(i\theta) \rangle = \frac{\det M(t_o, t_i, \dots)}{|\det M(t_o, t_i, \dots)|}$$

$M(t_o, t_i, \dots)$ is the transition matrix



Borasoy et al. (2007)

- Symmetry-sign extrapolation (SSE) method: control the sign oscillations

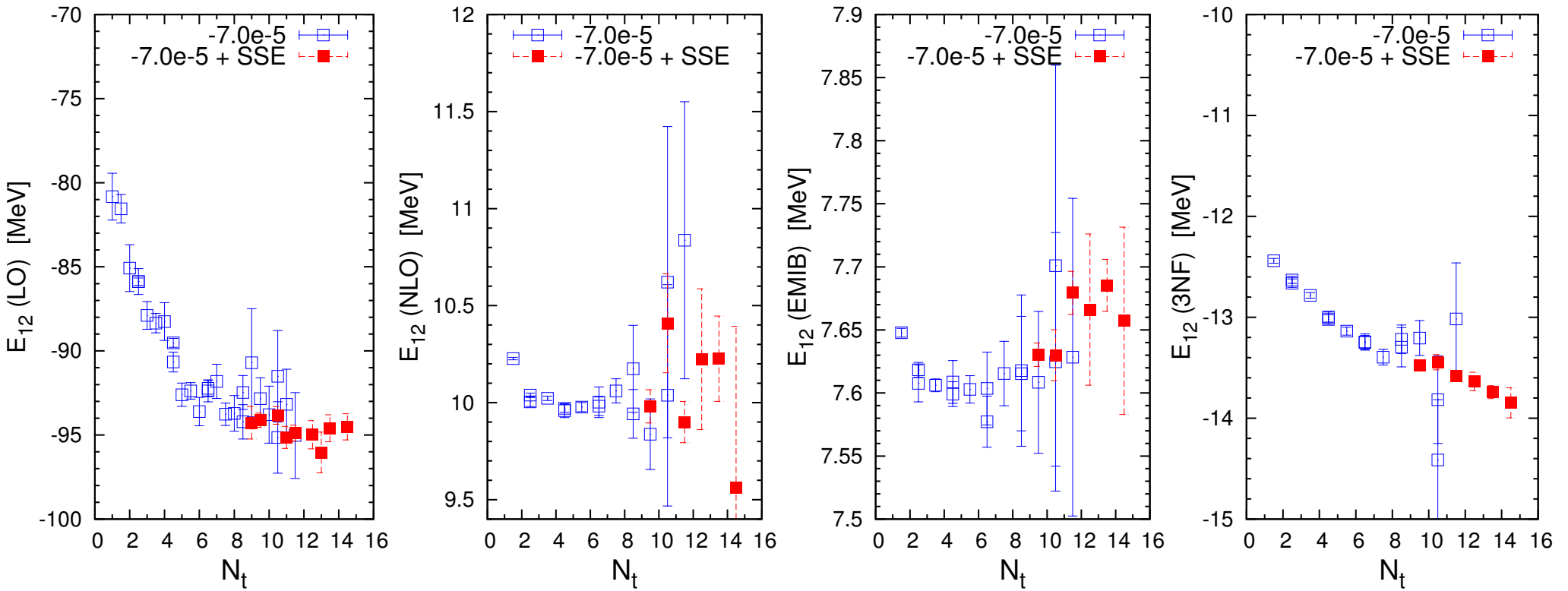
$$H_{d_h} = d_h \cdot H_{\text{phys}} + (1 - d_h) \cdot H_{\text{SU}(4)}$$

$$H_{\text{SU}(4)} = \frac{1}{2} C_{\text{SU}(4)} (N^\dagger N)^2$$

→ family of solutions for different SU(4) couplings $C_{\text{SU}(4)}$
that converge on the physical value for $d_h = 1$

RESULTS for ^{12}C

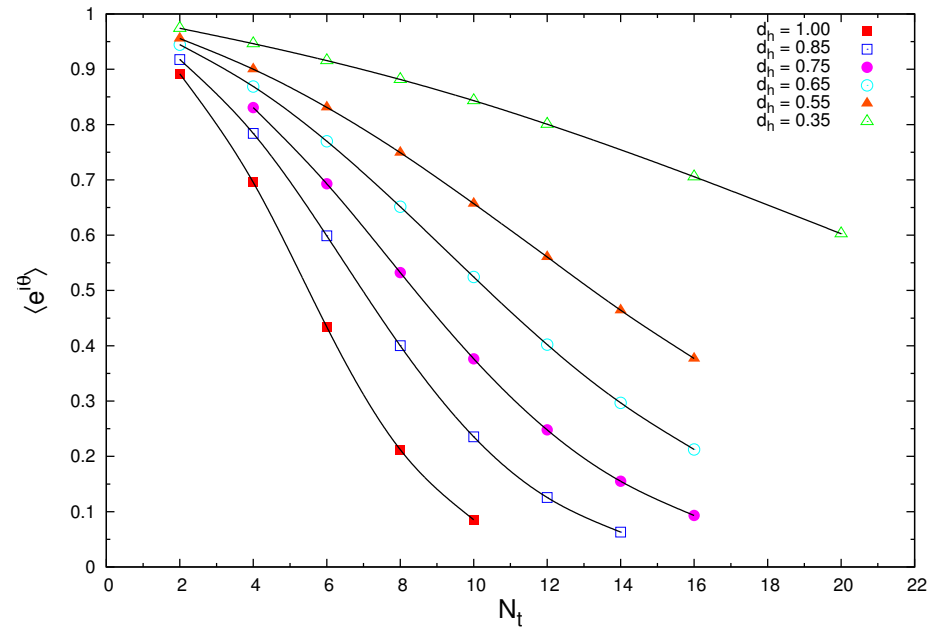
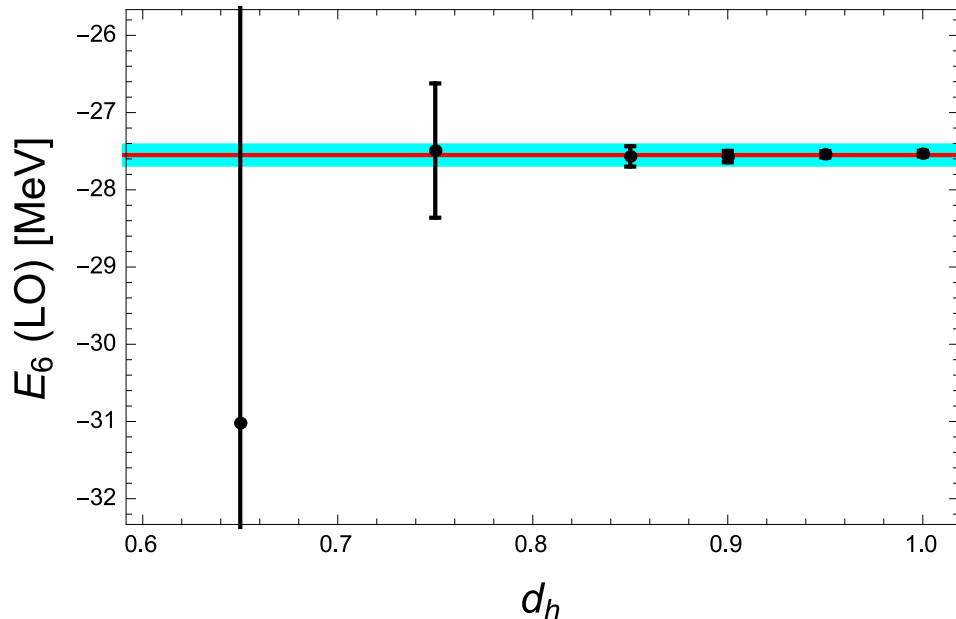
- generate a few more MC data at large N_t using SSE



- promising results → no more exponential deterioration of the MC data
- results w/ small uncertainties for $d_h \geq 0.8$

RESULTS for $A = 6$

- Simulations for ${}^6\text{He}$ and ${}^6\text{Be}$



⇒ methods works for nuclei with $A \neq Z$

⇒ neutron-rich nuclei can now be systematically explored (larger volumes)

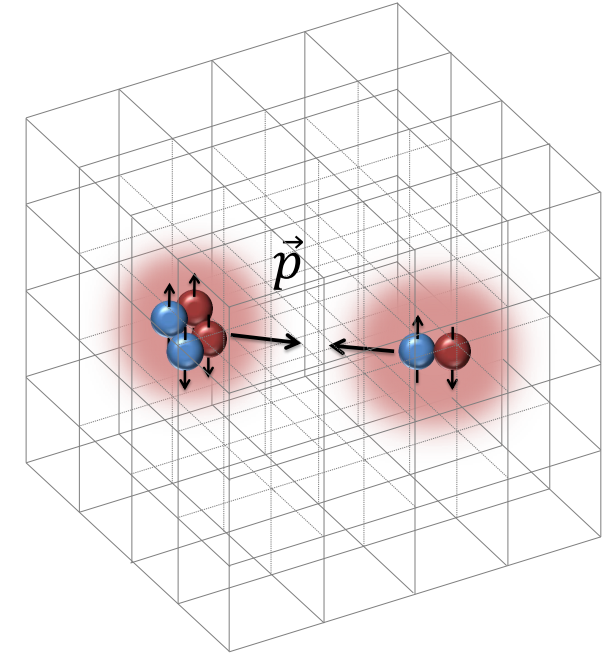
⇒ stay tuned!

Ab initio calculation of α - α scattering

Elhatisari, Lee, Rupak, Epelbaum, Krebs, Lähde, Luu, UGM,
Nature **528** (2015) 111

TWO-BODY SCATTERING on the LATTICE

- Processes involving α -particles and α -type nuclei comprise a major part of stellar nucleosynthesis, and control the production of certain elements in stars
- Ab initio calculations of scattering and reactions suffer from computational scaling with the number of nucleons in the clusters



Lattice EFT computational scaling $\Rightarrow (A_1 + A_2)^2$

Rupak, Lee, Phys. Rev. Lett. 111 (2013) 032502
 Pine, Lee, Rupak, Eur. Phys. J. A49 (2013) 151
 Elhatisari, Lee, Phys. Rev. C90 (2014) 064001
 Rokash et al., Phys. Rev. C92 (2015) 054612

ADIABATIC PROJECTION METHOD

- Basic idea to treat scattering and inelastic reactions:
split the problem into two parts

First part:

use Euclidean time projection to construct an *ab initio* low-energy cluster Hamiltonian, called the **adiabatic Hamiltonian**

Second part:

compute the two-cluster scattering phase shifts or reaction amplitudes using the adiabatic Hamiltonian

ADIABATIC PROJECTION METHOD II

- Construct a low-energy effective theory for clusters

- Use initial states parameterized by the relative separation between clusters

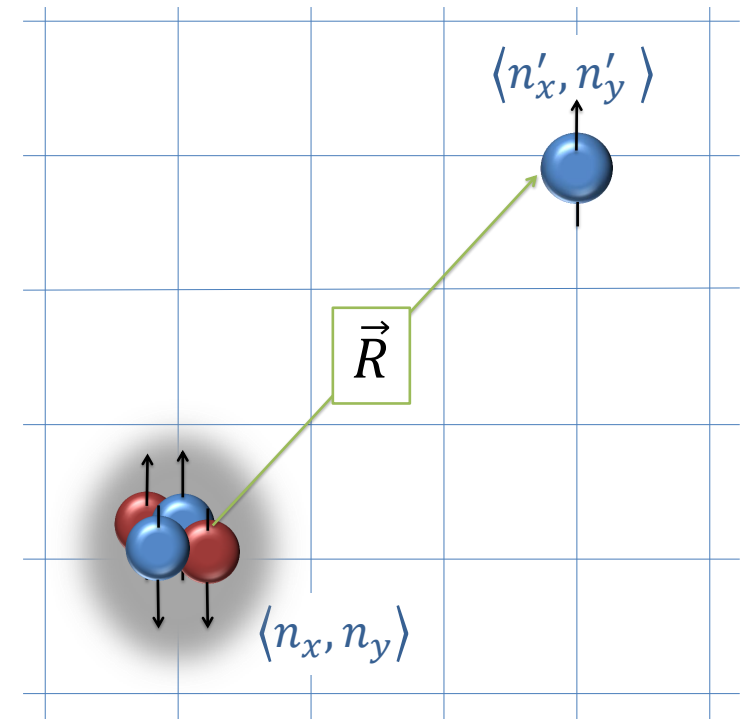
$$|\vec{R}\rangle = \sum_{\vec{r}} |\vec{r} + \vec{R}\rangle \otimes \vec{r}$$

- project them in Euclidean time with the chiral EFT Hamiltonian H

$$|\vec{R}\rangle_\tau = \exp(-H\tau)|\vec{R}\rangle$$

→ “dressed cluster states”

- The adiabatic projection in Euclidean times gives a systematically improvable description of the low-lying scattering states
- In the limit of large Euclidean time, the description becomes exact



ADIABATIC HAMILTONIAN

- Construct the adiabatic Hamiltonian from the dressed cluster states:

$$[H_\tau]_{\vec{R}\vec{R}'} = {}_\tau \langle \vec{R} | H | \vec{R}' \rangle_\tau$$

- States are i.g. not normalized, require *norm matrix*:

$$[N_\tau]_{\vec{R}\vec{R}'} = {}_\tau \langle \vec{R} | \vec{R}' \rangle_\tau$$

- construct the full adiabatic Hamiltonian:

$$[H_\tau^a]_{\vec{R}\vec{R}'} = \sum_{\vec{R}_n \vec{R}_m} [N_\tau^{-1/2}]_{\vec{R}\vec{R}_n} [H_\tau]_{\vec{R}_n \vec{R}_m} [N_\tau^{-1/2}]_{\vec{R}_m \vec{R}'}$$

→ The structure of the adiabatic Hamiltonian is similar to the Hamiltonian matrix used in recent ab initio NCSM/RGM calculations

Navratil, Quaglioni, Phys. Rev. C 83 (2011) 044609
 Navratil, Roth, Quaglioni, Phys. Lett. B 704 (2011) 379
 Navratil, Quaglioni, Phys. Rev. Lett. 108 (2012) 042503

TESTING the ADIABATIC HAMILTONIAN

- Consider fermion-dimer scattering:

Microscopic Hamiltonian

$$L^{3(A-1)} \times L^{3(A-1)}$$



Two-cluster adiabatic Hamiltlonian

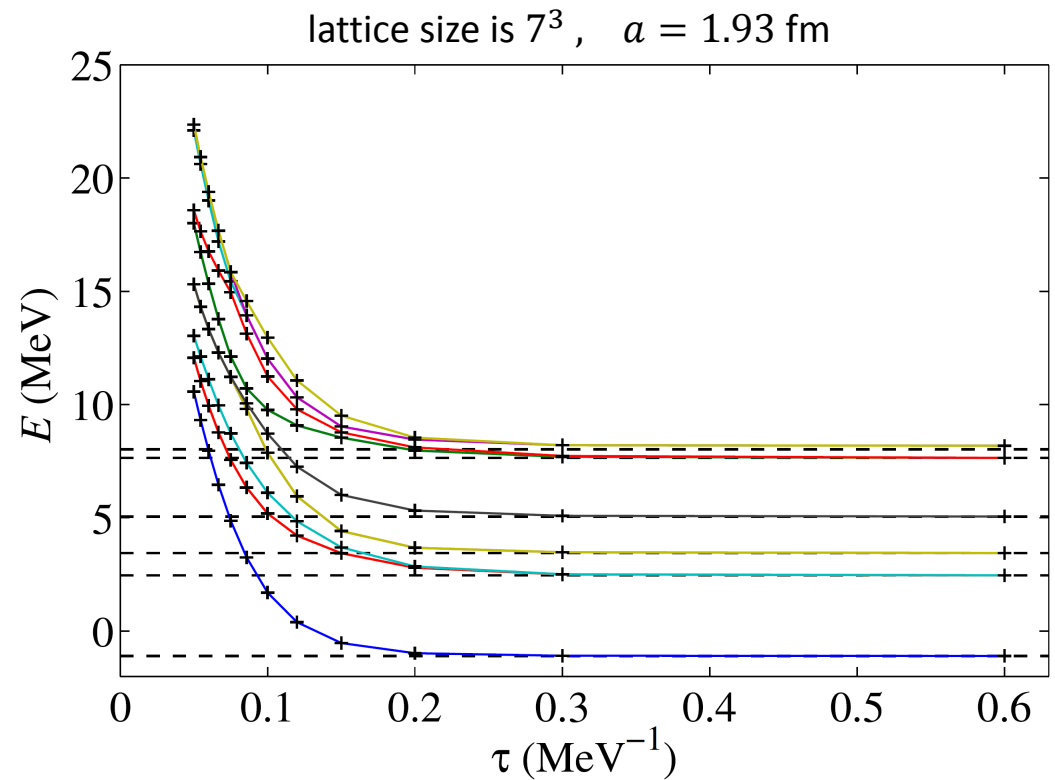
$$L^3 \times L^3$$

- calculation of a 7^3 lattice,
lattice spacing $a = 1.93$ fm

Pine, Lee, Rupak, EPJA 49 (2013) 151

exact Lanczos: black dashed lines

adiab. Ham.: solid colored lines



SCATTERING CLUSTER WAVE FUNCTIONS

- During Euclidean time interval τ_ϵ , each cluster undergoes spatial diffusion:

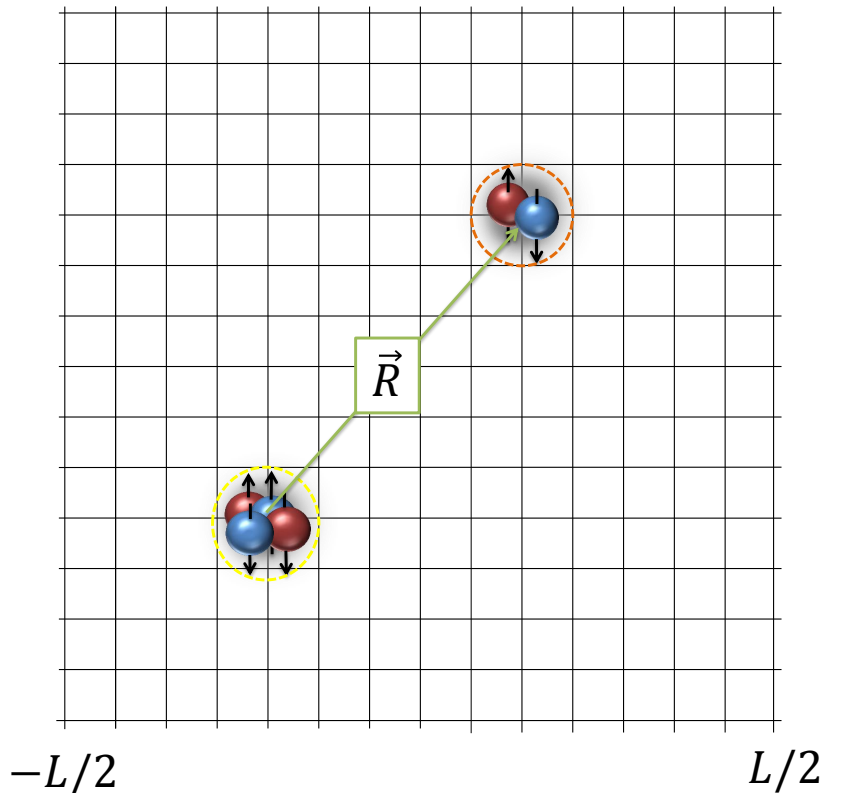
$$d_{\epsilon,i} = \sqrt{\tau_\epsilon/M_i}$$

- Only non-overlapping clusters if

$$|\vec{R}| \gg d_{\epsilon,i} \Rightarrow |\vec{R}\rangle_{\tau_\epsilon}$$

- Defines asymptotic region, where the amount of overlap between clusters is less than ϵ

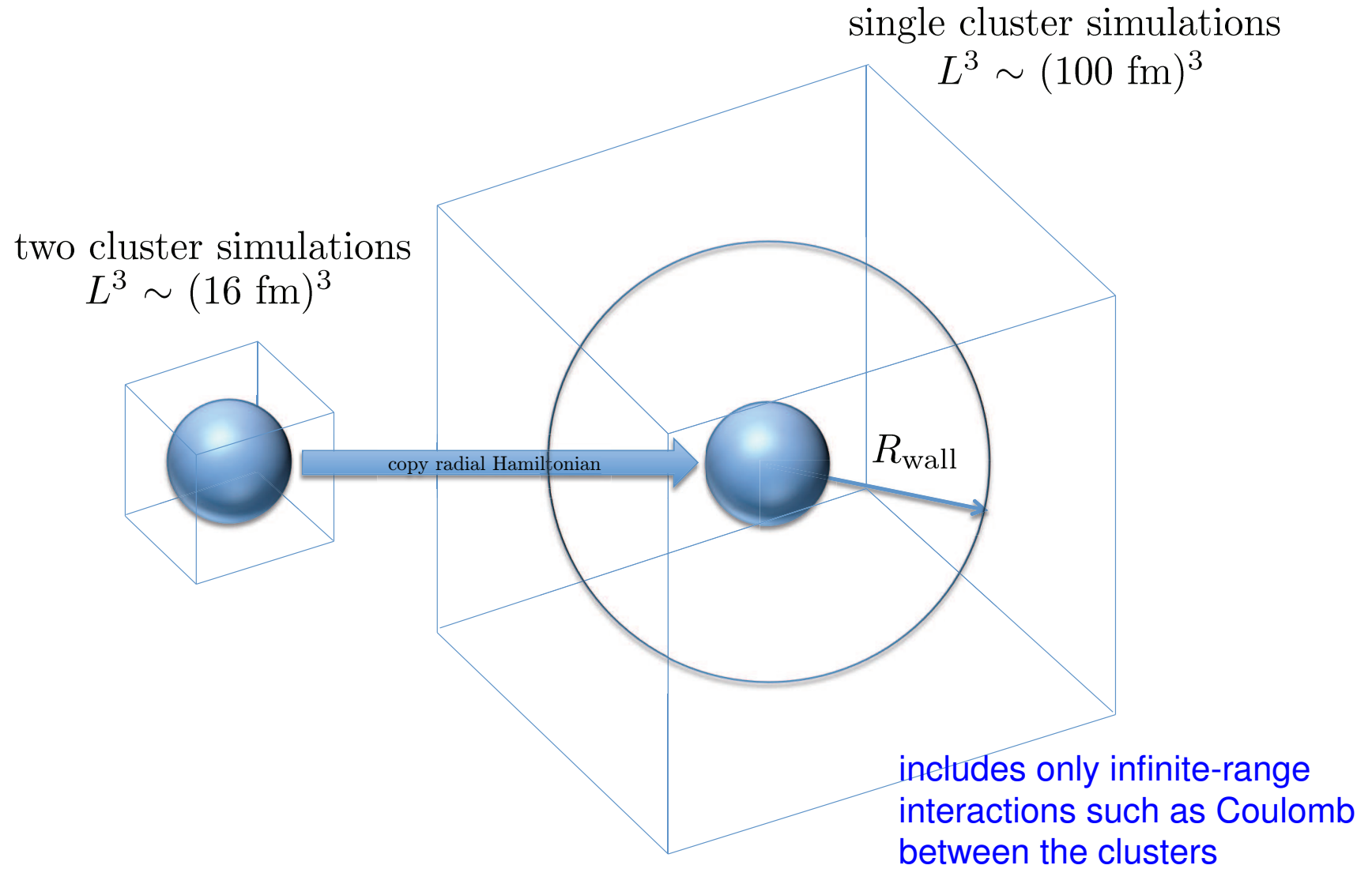
$$|\vec{R}| > R_\epsilon$$



In the asymptotic region we can describe the system in terms of an effective cluster Hamiltonian (the free lattice Hamiltonian for two clusters) plus infinite-range interactions (like the Coulomb int.)

ADIABATIC HAMILTONIAN plus COULOMB

23



25

ALPHA-ALPHA SCATTERING

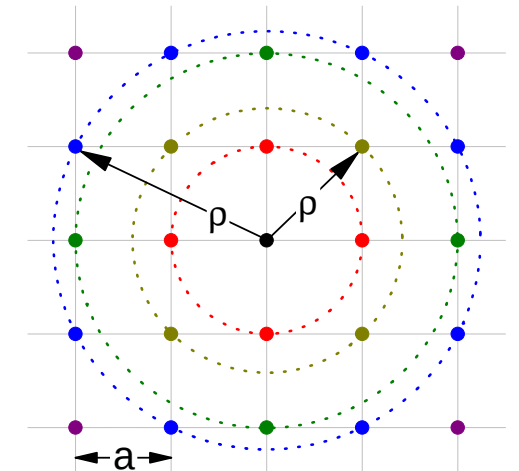
- same lattice action as for the Hoyle state in ^{12}C and the structure of ^{16}O
- 9 NN + 2 3N LECs, coarse lattice $a = 1.97 \text{ fm}$, $N = 8$
- new algorithm for Monte Carlo updates and alpha clusters
- adiabatic projection method to construct a two-alpha Hamiltonian
- spherical wall method to extract the phase shifts using radial Hamiltonian

$$|\vec{R}\rangle^{\ell,\ell_z} = \sum_{\vec{R}'} Y_{\ell,\ell_z}(\vec{R}') \delta_{\vec{R},|\vec{R}'|} |\vec{R}'\rangle$$

→ precise extraction of phase shifts & mixing angles

Lu, Lähde, Lee, UGM, arXiv:1506.05652

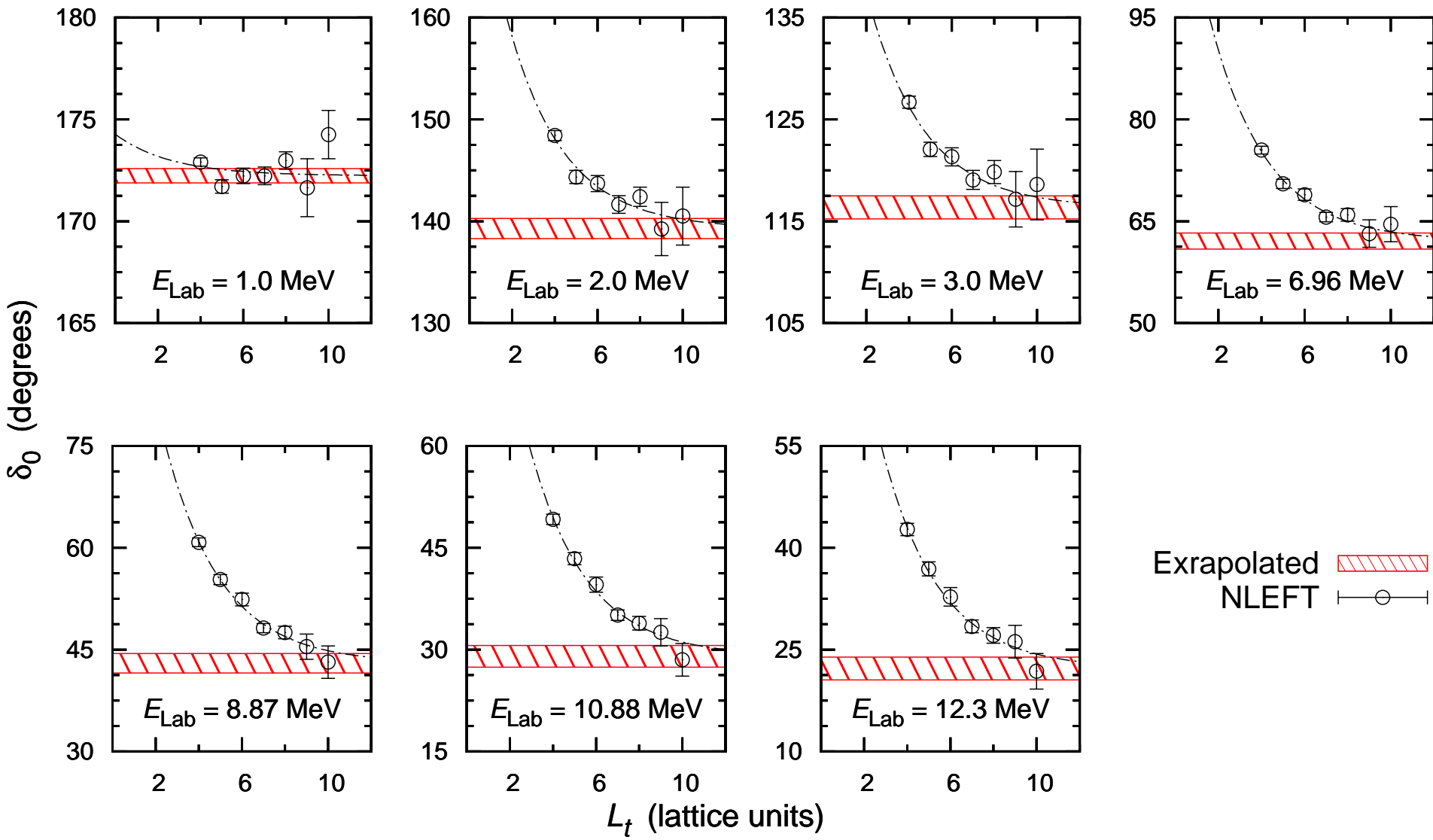
Moinard et al., work in progress



LATTICE DATA I

25

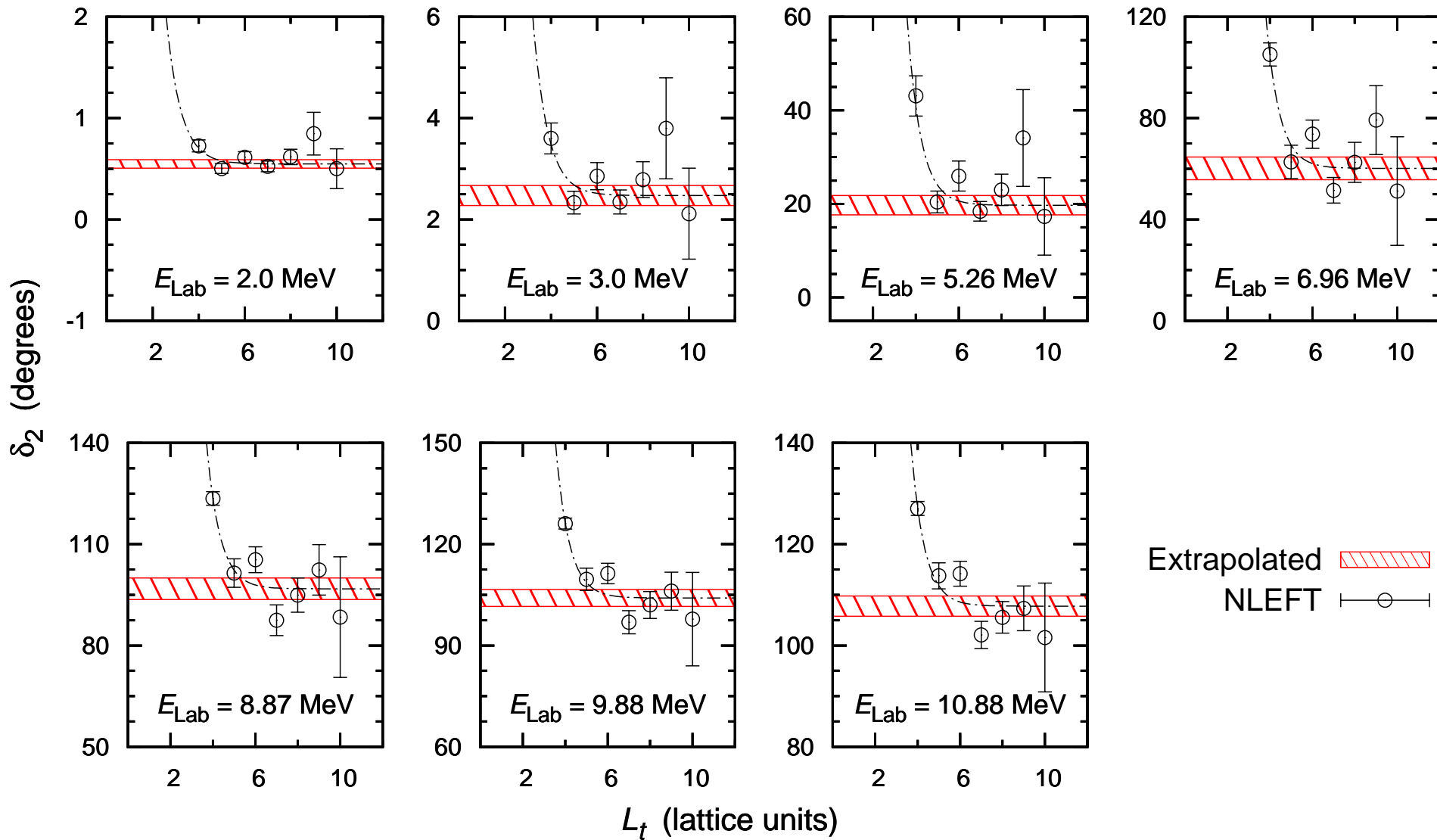
- Show data for the S-wave:



LATTICE DATA II

26

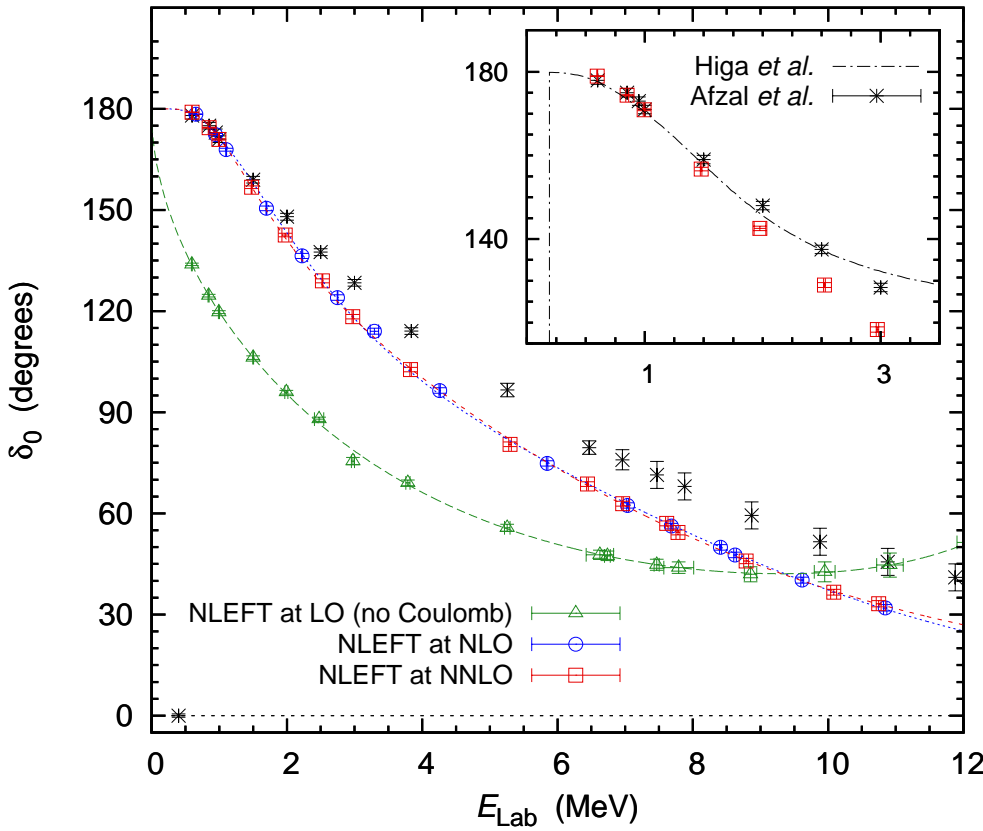
- Show data for the D-wave:



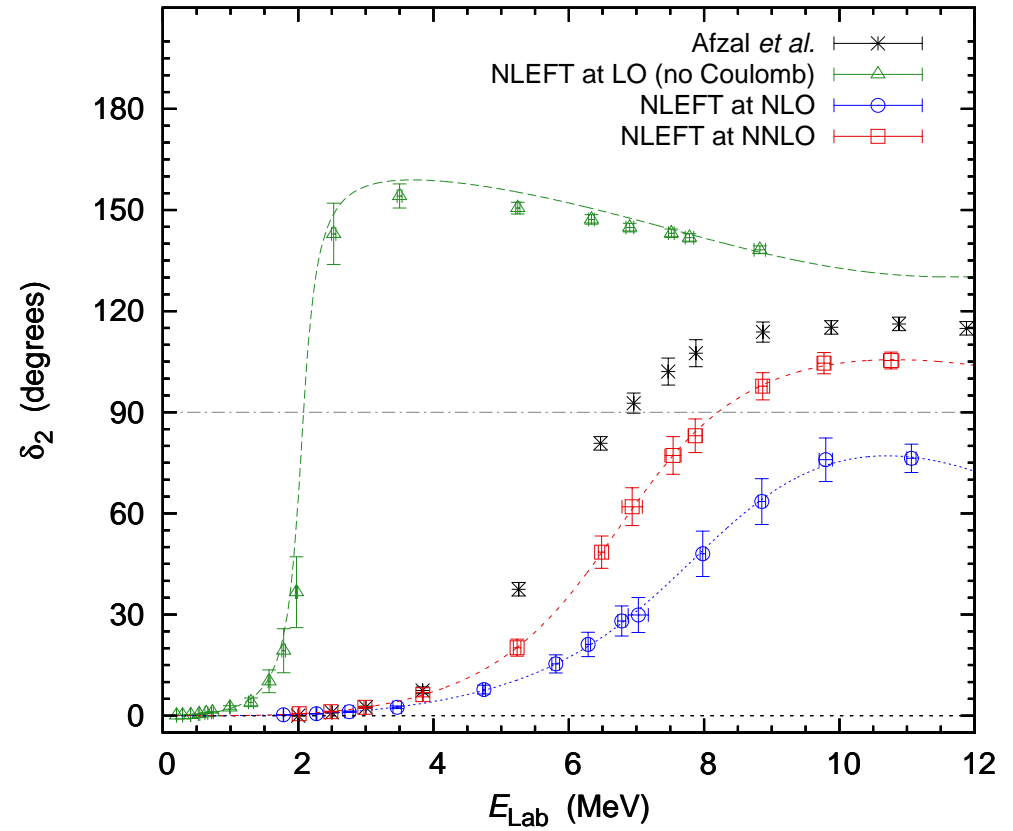
PHASE SHIFTS

27

- S-wave and D-wave phase shifts (LO has no Coulomb)



$$E_R^{\text{NNLO}} = -0.11(1) \text{ MeV} \quad [+0.09 \text{ MeV}]$$



$$E_R^{\text{NNLO}} = 3.27(12) \text{ MeV} \quad [2.92(18) \text{ MeV}]$$

$$\Gamma_R^{\text{NNLO}} = 2.09(16) \text{ MeV} \quad [1.35(50) \text{ MeV}]$$

Afzal et al., Rev. Mod. Phys. 41 (1969) 247 [data]; Higa et al., Nucl.Phys. A809 (2008) 171 [halo EFT]

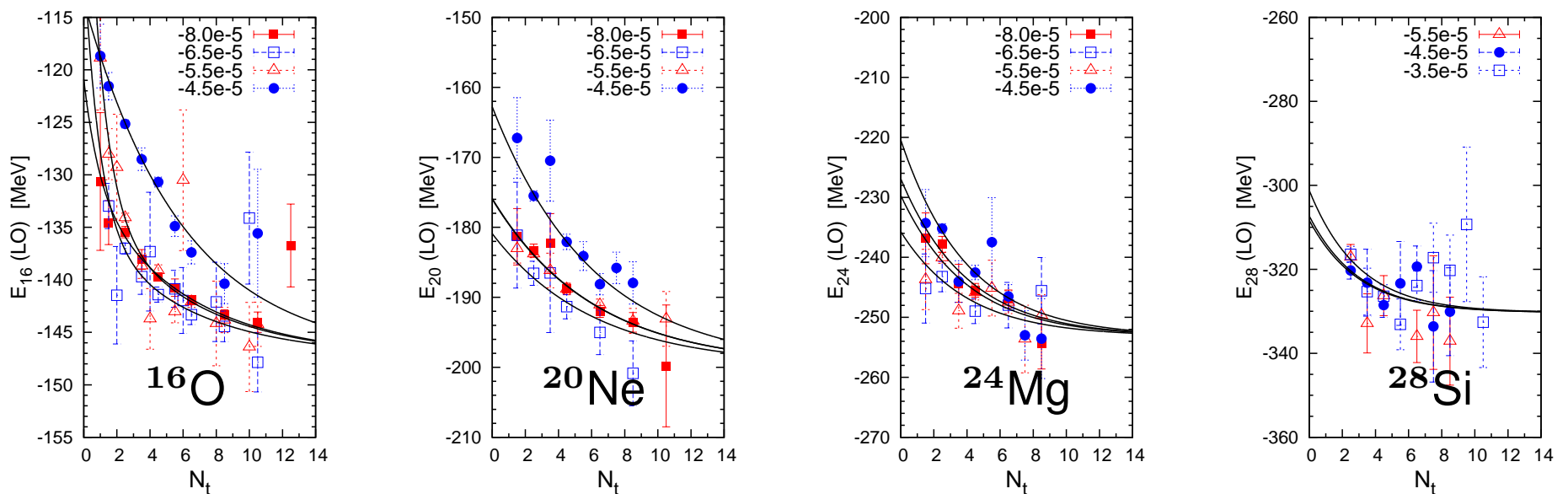
SUMMARY & OUTLOOK

- Nuclear lattice simulations as a new quantum many-body approach
 - based on the successful continuum nuclear chiral EFT
 - clustering emerges naturally, α -cluster nuclei
 - symmetry-sign extrapolation method allows to go to the drip lines
 - ab initio study of α - α scattering: promising results
 - holy grail of nuclear astrophysics ($\alpha + {}^{12}\text{C} \rightarrow {}^{16}\text{O} + \gamma$) in reach
- Some on-going activities:
 - improving the forces (N^3LO , sph. harmonics)
 - systematic studies of a -independence
 - Klein, Lee, Liu, UGM, PLB747 (2015) 511
 - finite size effects/averaging procedures
 - Lu, Lähde, Lee, UGM, Phys. Rev. D90 (2014) 034507 & D92 (2015) 014506
 - a quantum phase transition? ... and much more ...

Spares

GOING up the ALPHA CHAIN

- Consider the α ladder ^{12}C , ^{16}O , ^{20}Ne , ^{24}Mg , ^{28}Si as $t_{\text{CPU}} \sim A^2$
- Improved “multi-state” technique to extract ground state energies
 - \Rightarrow higher A , better accuracy
 - \Rightarrow overbinding at LO beyond $A = 12$ persists up to NNLO



$$E = -131.3(5) \quad [-127.62]$$

$$E = -165.9(9) \quad [-160.64]$$

$$E = -232(2) \quad [-198.26]$$

$$E = -308(3) \quad [-236.54]$$

REMOVING the OVERBINDING

31

Lähde, Epelbaum, Krebs, Lee, UGM, Rupak, Phys. Lett. B 732 (2014) 110

- Overbinding is due to four α clusters in close proximity

⇒ remove this by an effective 4N operator [long term: N3LO]

$$V^{(4N_{\text{eff}})} = D^{(4N_{\text{eff}})} \sum_{1 \leq (\vec{n}_i - \vec{n}_j)^2 \leq 2} \rho(\vec{n}_1) \rho(\vec{n}_2) \rho(\vec{n}_3) \rho(\vec{n}_4)$$

- fix the coefficient $D^{(4N_{\text{eff}})}$ from the BE of ${}^{24}\text{Mg}$

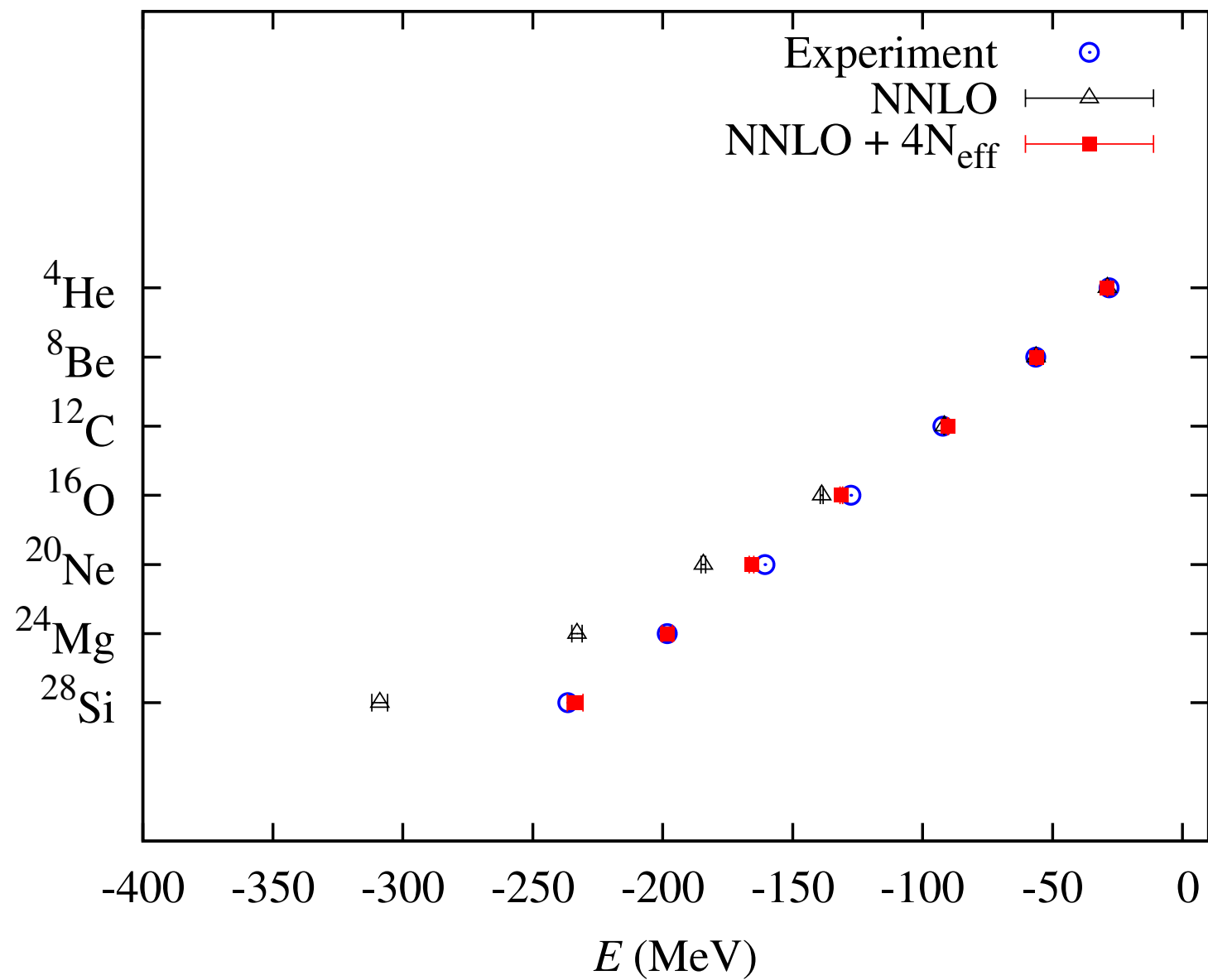
⇒ excellent description of the ground state energies

A	12	16	20	24	28
Th	-90.3(2)	-131.3(5)	-165.9(9)	-198(2)	-233(3)
Exp	-92.16	-127.62	-160.64	-198.26	-236.54

→ ultimately, reduce lattice spacing [interaction more repulsive] & N³LO

GROUND STATE ENERGIES

32



STRUCTURE of ^{16}O

33

- Mysterious nucleus, despite modern ab initio calcs

Hagen et al. (2010), Roth et al. (2011), Hergert et al. (2013)

- Alpha-cluster models since decades, some exp. evidence

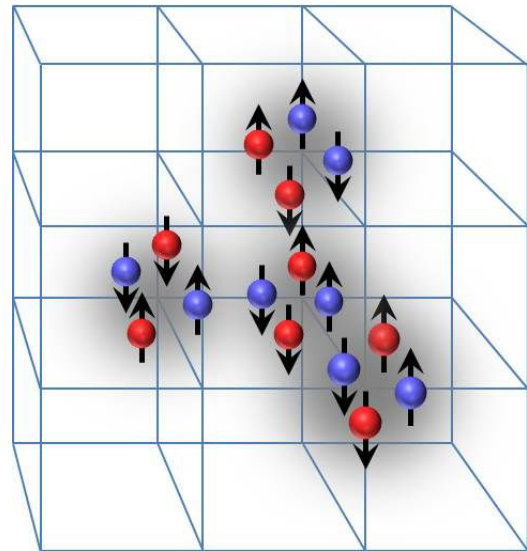
Wheeler (1937), Dennison (1954), Robson (1979), . . . , Freer et al. (2005)

- Spectrum very close to tetrahedral symmetry group

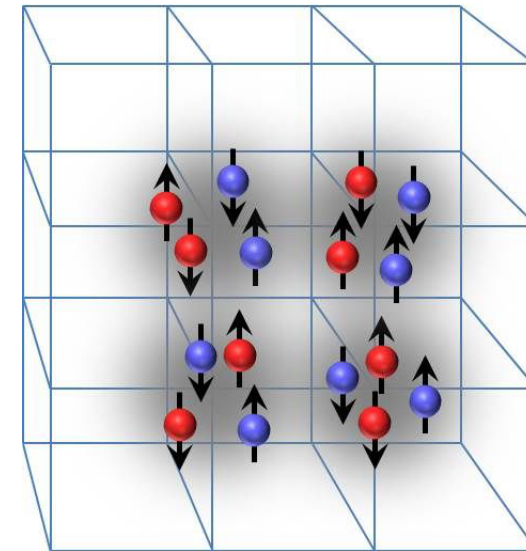
Bijker & Iachello (2014)

- Relevant configurations in lattice simulations:

Tetrahedron (A)



Square (narrow (B) and wide (C))



DECODING the STRUCTURE of ^{16}O

Epelbaum, Krebs, Lähde, Lee, UGM, Rupak, Phys. Rev. Lett. **112** (2014) 102501

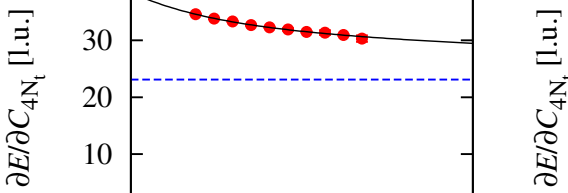
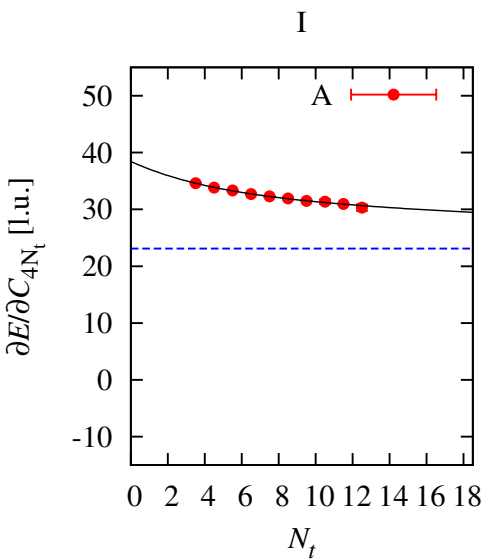
- measure the 4N density, where each of the nucleons is placed at adjacent points

$\Rightarrow 0_1^+$ ground state: mostly tetrahedral config

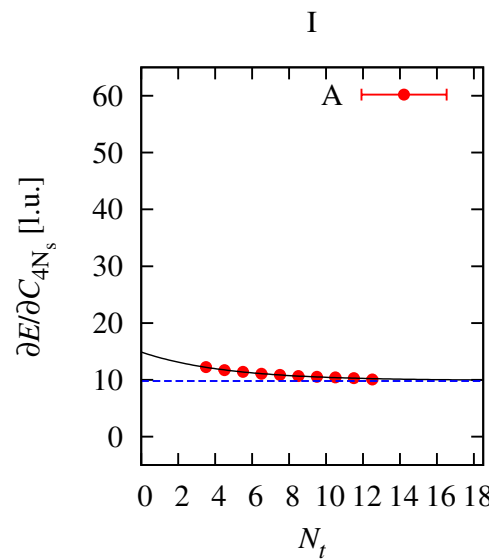
$\Rightarrow 0_2^+$ excited state: mostly square configs

2_1^+ excited state: rotational excitation of the 0_2^+

overlap w/ tetrahedral config.



overlap w/ square configs.



RESULTS for ^{16}O

- Spectrum:

	LO	NNLO(2N)	NNLO(3N)	4N_{eff}	Exp.
0_1^+	-147.3(5)	-121.4(5)	-138.8(5)	-131.3(5)	-127.62
0_2^+	-145(2)	-116(2)	-136(2)	-123(2)	-121.57
2_1^+	-145(2)	-116(2)	-136(2)	-123(2)	-120.70

- LO charge radius: $r(0_1^+) = 2.3(1) \text{ fm}$ Exp. $r(0_1^+) = 2.710(15) \text{ fm}$

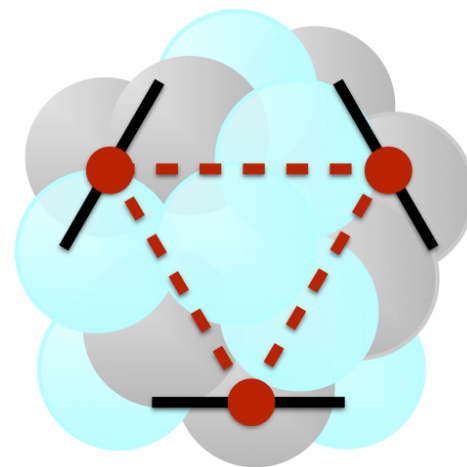
⇒ compensate for this by rescaling with appropriate units of r/r_{LO}

- LO EM properties:

	LO	LO(r-scaled)	Exp.
$Q(2_1^+) [\text{e fm}^2]$	10(2)	15(3)	—
$B(E2, 2_1^+ \rightarrow 0_2^+) [\text{e}^2 \text{ fm}^4]$	22(4)	46(8)	65(7)
$B(E2, 2_1^+ \rightarrow 0_1^+) [\text{e}^2 \text{ fm}^4]$	3.0(7)	6.2(1.6)	7.4(2)
$M(E0, 0_2^+ \rightarrow 0_2^+) [\text{e fm}^2]$	2.1(7)	3.0(1.4)	3.6(2)

⇒ gives credit to the interpretation of the 2_1^+ as rotational excitation

Continuum EFT: new developments

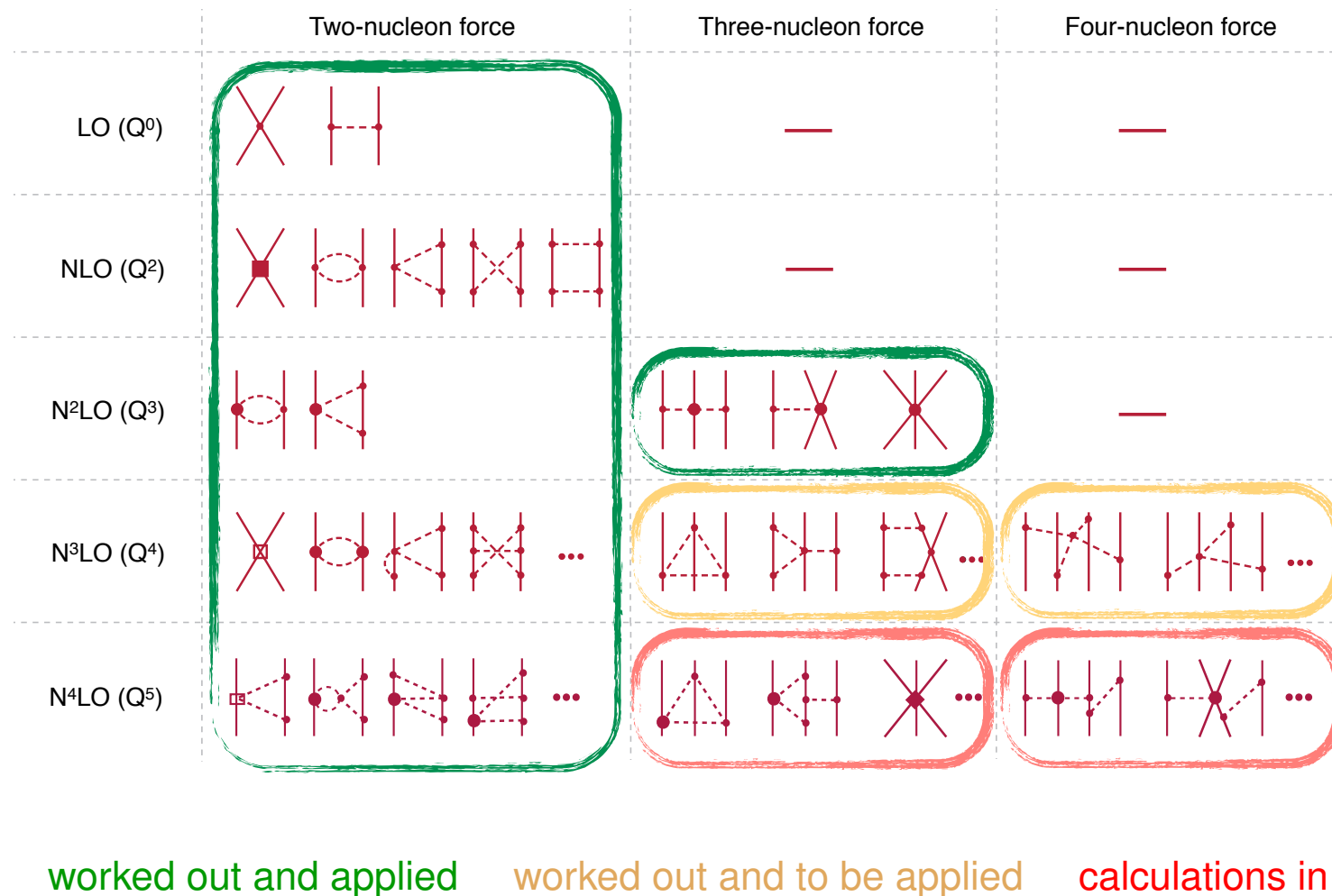


LENPIC

www.lenpic.org

NUCLEAR FORCES in CHIRAL NUCLEAR EFT

- expansion of the potential in powers of Q [small parameter]
- explains observed hierarchy of the nuclear forces



NN FORCES to FOURTH ORDER

Epelbaum, Krebs, UGM, Eur. Phys. J. A 51: 53 (2015)

- new regularization of long-range physics [coordinate space cut-off]:

$$V_{\text{long-range}}^{\text{reg}}(\vec{r}) = V_{\text{long-range}}(\vec{r}) f_{\text{reg}} \left(\frac{r}{R} \right), \quad f_{\text{reg}} = \left[1 - \exp \left(-\frac{r^2}{R^2} \right) \right]^6$$

- ⇒ No distortion of the long-range potential → better at higher energies
- ⇒ No additional spectral function regularization in the TPEP required
- ⇒ Study of the chiral expansion of multi-pion exchanges: $R = 0.8 \cdots 1.2 \text{ fm}$

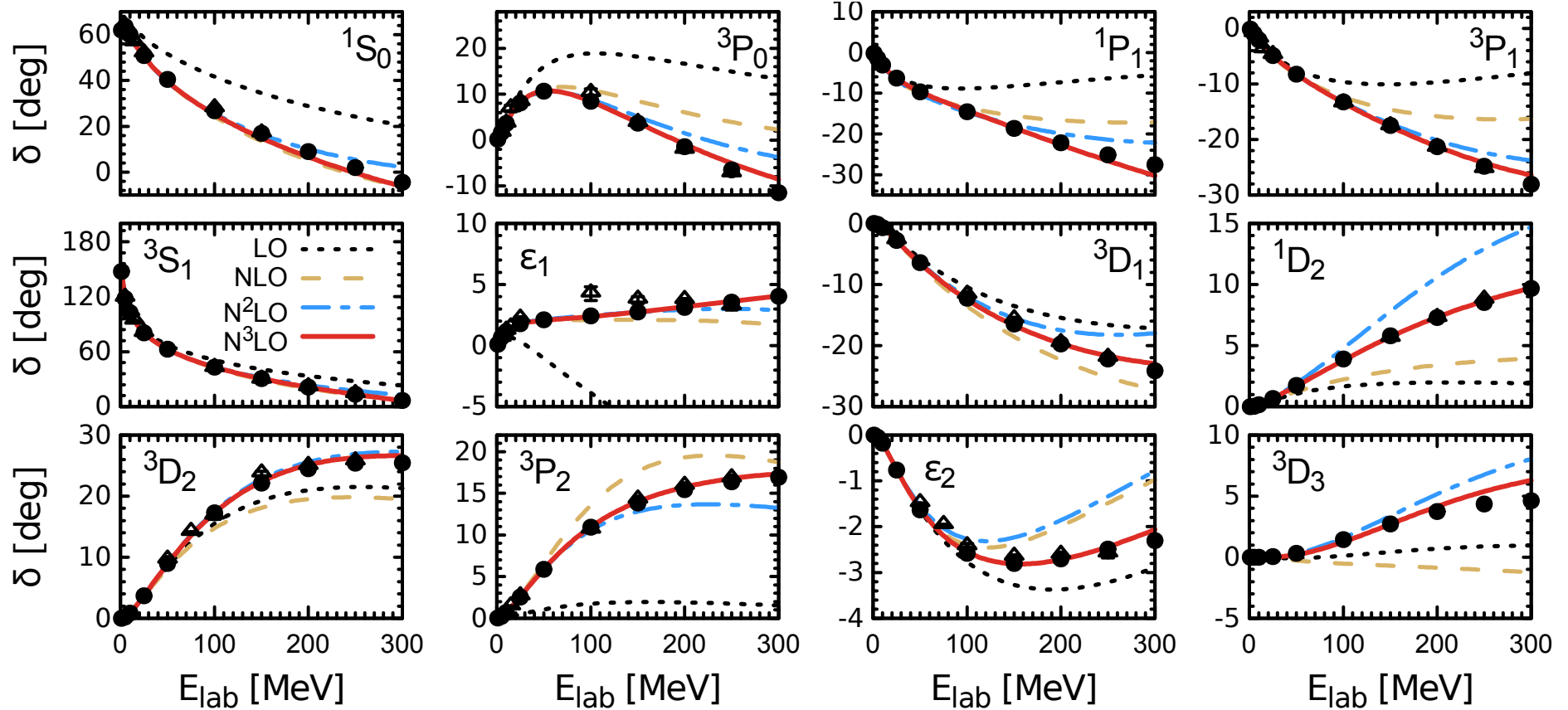
Baru et al., EPJ A48 (12) 69

- new way of estimation the theoretical uncertainty [before: only cut-off variations]
- ⇒ Expansion parameter depending on the region: $Q = \max \left(\frac{M_\pi}{\Lambda_b}, \frac{p}{\Lambda_b} \right)$
- ⇒ Breakdown scale $\Lambda_b = 600 \text{ MeV}$ for $R = 0.8 \cdots 1.0 \text{ fm}$

CONVERGENCE of the CHIRAL SERIES

39

- phase shifts show expected convergence [large N2LO corrections understood]

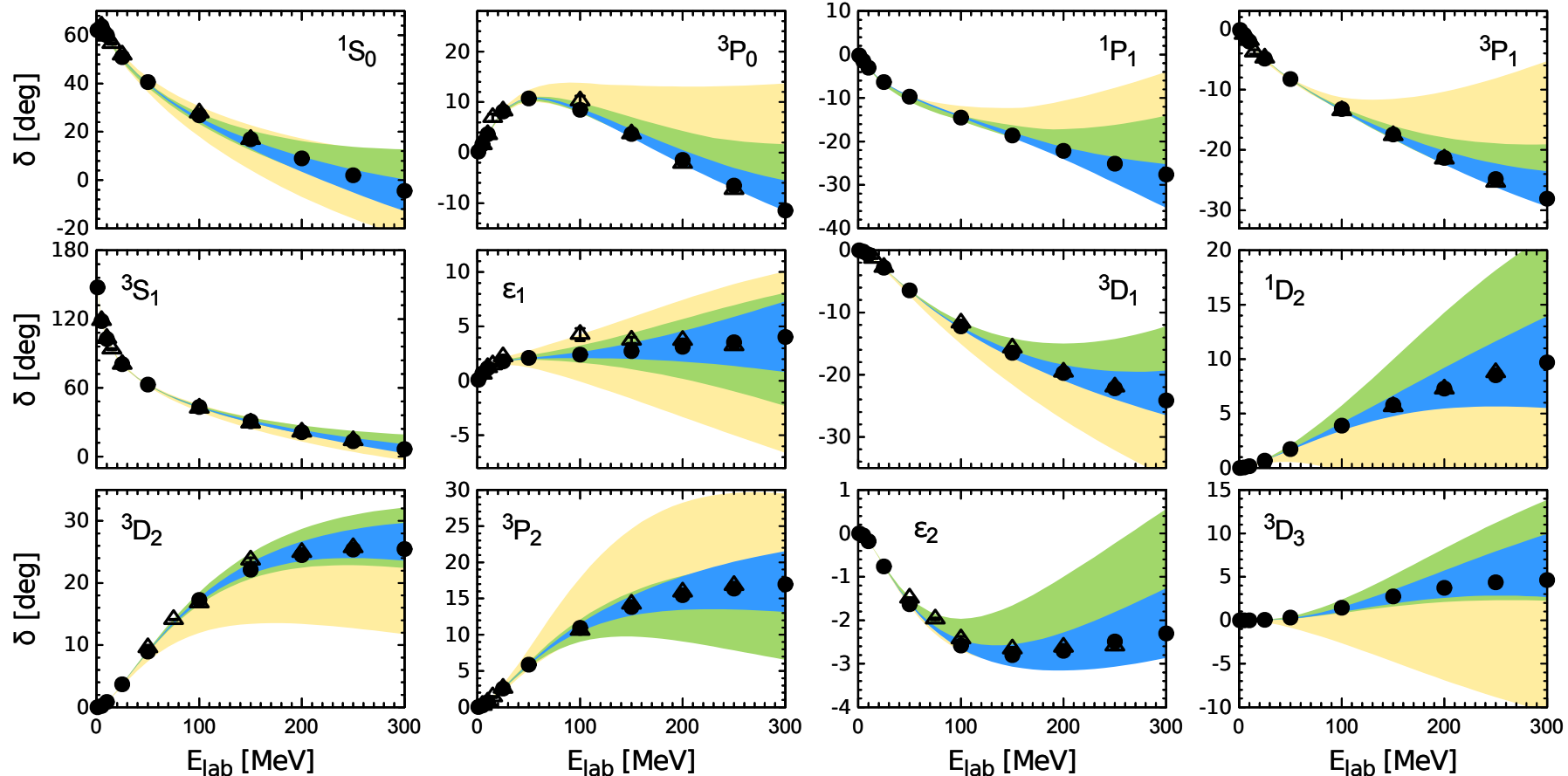


⇒ clear improvement comp. to earlier N3LO potentials [momentum space reg.]

Entem, Machleidt; Epelbaum, Glöckle, UGM

UNCERTAINTIES

- uncertainties show expected pattern



NLO

N2LO

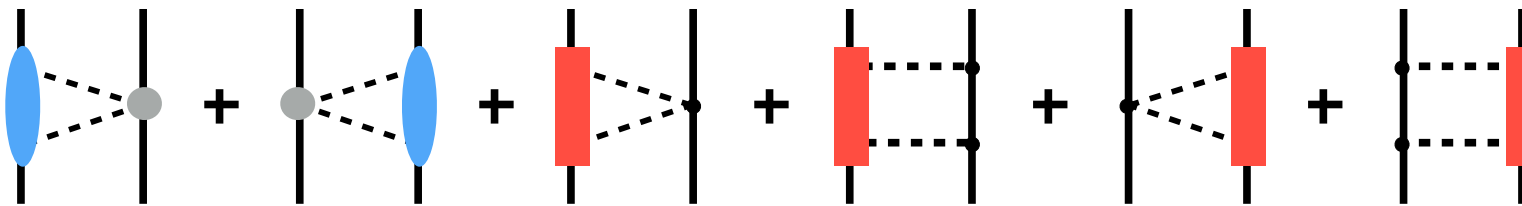
N3LO

NN FORCES to FIFTH ORDER

Epelbaum, Krebs, UGM, arXiv:1412.4623

- No contact interactions at this order - odd in Q
- New contributions fixed from πN scattering, LECs c_i, d_i, e_i :

Büttiker, Fettes, UGM, Steininger (1998-2000); Krebs, Gasparian, Epelbaum (2012)



$$\mathcal{L}_{\pi N} = \mathcal{L}_{\pi N}^{(1)} + \mathcal{L}_{\pi N}^{(2)}(c_i) + \mathcal{L}_{\pi N}^{(3)}(d_i) + \mathcal{L}_{\pi N}^{(4)}(e_i)$$

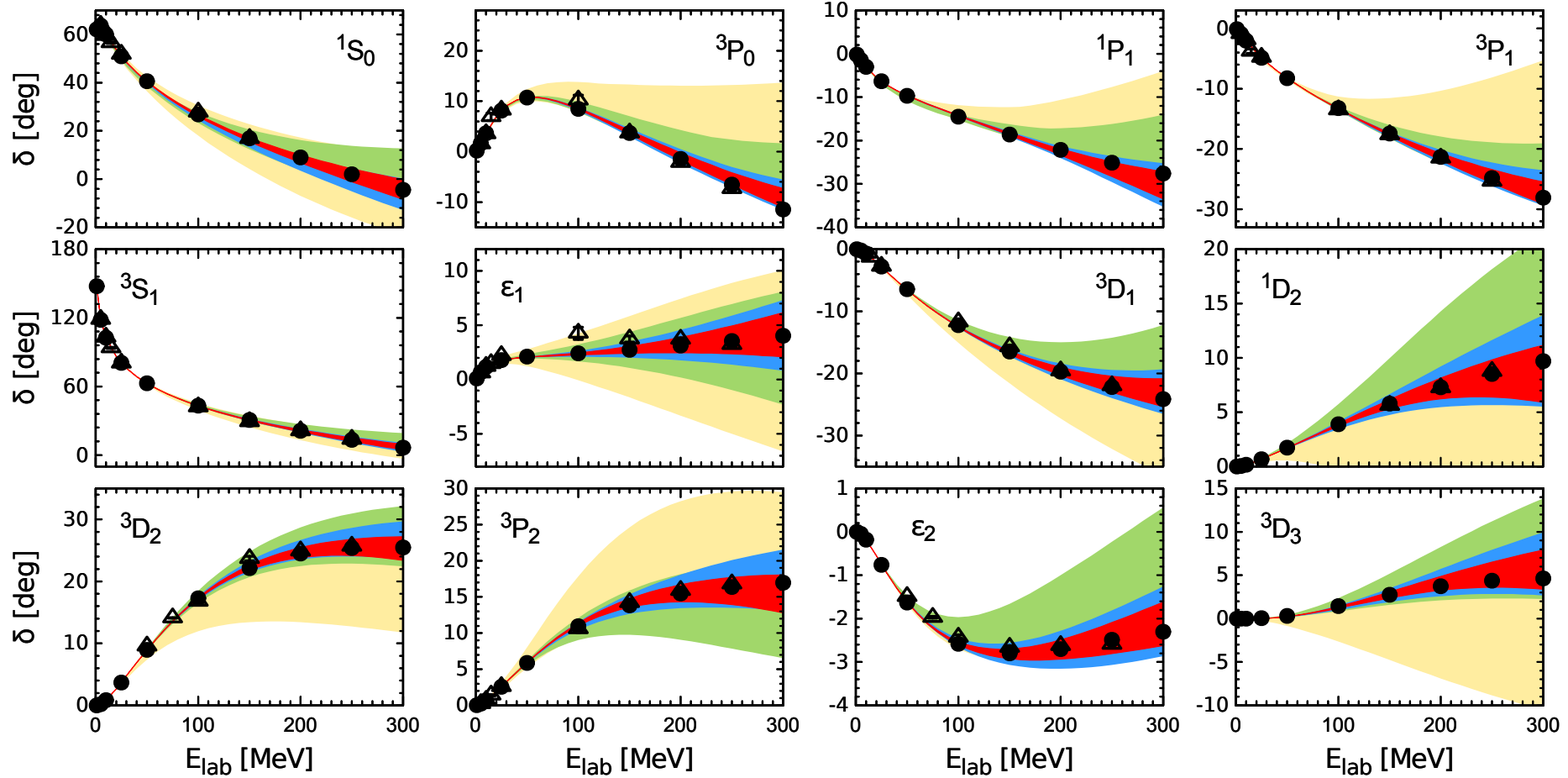
- Three-pion exchange can be neglected
 - explicit calculation of the dominant NLO contribution
 - no influence on phase shifts or deuteron properties

Kaiser (2001)

PHASE SHIFTS at N4LO

42

⇒ Precision phase shifts with small uncertainties up to $E_{\text{lab}} = 300 \text{ MeV}$

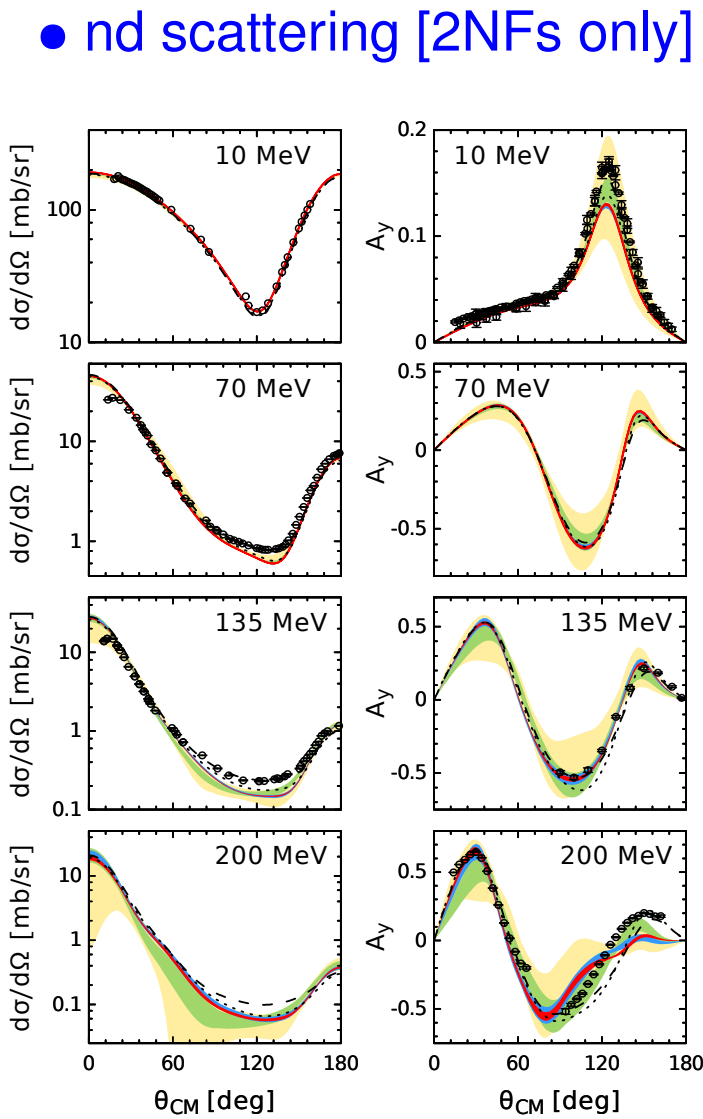
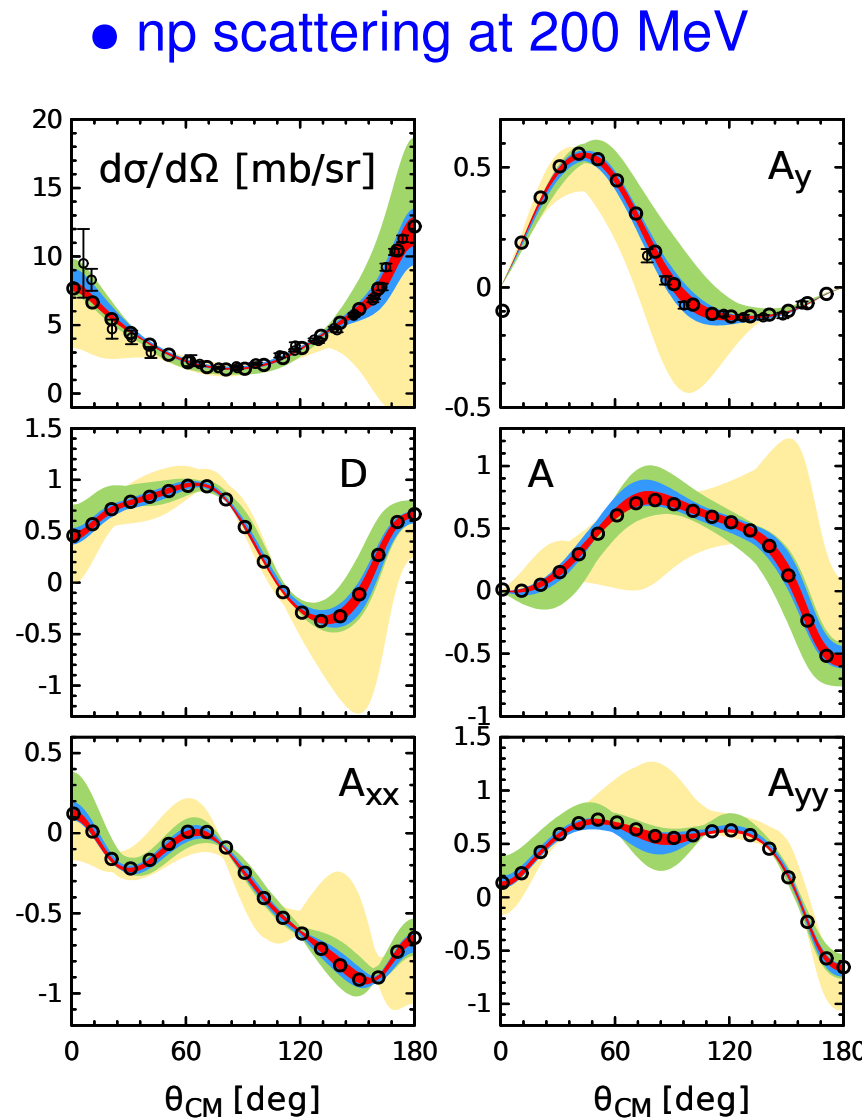


NLO N2LO N3LO N4LO

EVIDENCE for THREE-NUCLEON FORCES

43

- Two-nucleon system under control, three-nucleon system requires 3NFs!
→ being implemented [LENPIC collaboration]

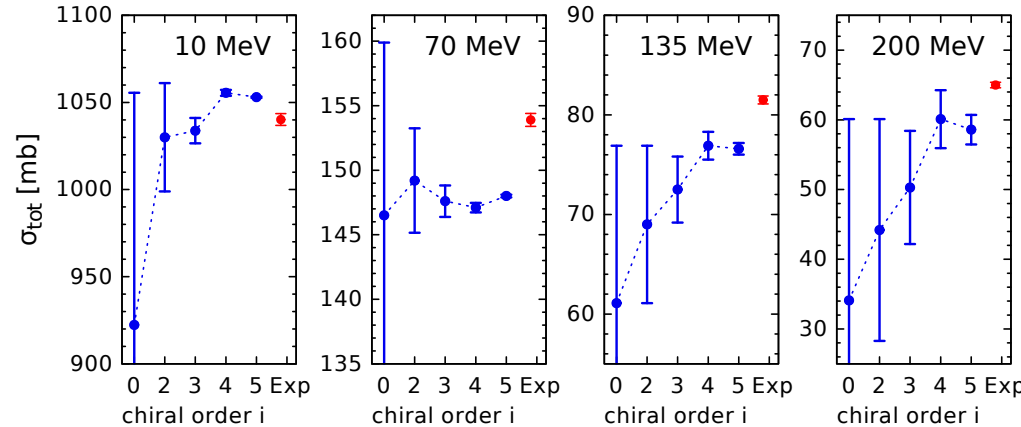


NLO
N2LO
N3LO
N4LO

MORE EVIDENCE for THREE-NUCLEON FORCES

Binder et al. [LENPIC collaboration], arXiv:1505.07218

- Total cross section for Nd scattering [2NFs only]



- Binding energy and rms radius of ^4He , lowest levels in ^6Li [2NFs only]

