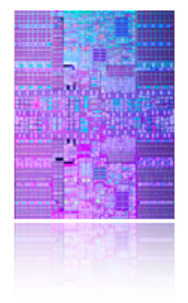
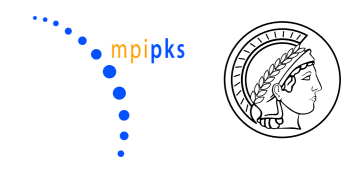
$\mathcal{H}|\psi\rangle = E|\psi\rangle$



Third Lecture: Introduction to Exact Diagonalization

Andreas Läuchli, "New states of quantum matter" MPI für Physik komplexer Systeme - Dresden

http://www.pks.mpg.de/~aml aml@pks.mpg.de



Lecture Notes at http://www.pks.mpg.de/~aml/LesHouches

"Modern theories of correlated electron systems" - Les Houches - 20/5/2009



Outline

 $\mathcal{H}|\psi\rangle = E|\psi\rangle$

- Main Idea
- Examples of typical applications
- Structure of an Exact Diagonalization package
 - Hilbertspace, Symmetries
 - Hamiltonian
 - Linear Algebra
 - Observables
 - Parallelization
- Applications



 $\mathcal{H}|\psi\rangle = E|\psi\rangle$



Solve the Schrödinger equation of a quantum many body system numerically

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Sparse matrix, but for quantum many body systems the vector space dimension grows exponentially!



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But if you want to get a maximum of physical information out of a finite system there is a lot more to do and the reward is a powerful:



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Quantum Mechanics Toolbox





 Quantum Magnets: nature of novel phases, critical points in 1D, dynamical correlation functions in 1D & 2D



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- Full Configuration Interaction in Quantum Chemistry





Spin S=1/2 models:

40 spins square lattice, 39 sites triangular, 42 sites star lattice at S^z=0 64 spins or more in elevated magnetization sectors up to 1.5 billion(=10⁹) basis states with symmetries, up to 4.5 billion without



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Hubbard models
 20 sites square lattice at half filling, 20 sites quantum dot structure
 22-25 sites in ultracold atoms setting w.o. spatial symmetries
 up to 160 billion basis states

Structure of an Exact Diagonalization code





Hilbert space



Hilbert space

Basis represention, Lookup techniques



Hilbert space

- Basis represention, Lookup techniques
- Symmetries



Hilbert space

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- Hamiltonian Matrix



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 - Static quantities (multipoint correlation functions, correlation density matrices,...)
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 - Real-time evolution

Hilbert Space



Basis representation

States of the Hilbert space need to be represented in the computer.

Choose a representation which makes it simple to act with the Hamiltonian or other operators on the states, and to localize a given state in the basis

Simple example: ensemble of S=1/2 sites in binary coding

 $|\uparrow\uparrow\downarrow\downarrow\uparrow\rangle \rightarrow [1\ 1\ 0\ 1]_2 = 13$

detection of up or down spin can be done with bit-test. transverse exchange $S^+S^- + S^-S^+$ can be performed by an XOR operation:

$$\begin{bmatrix} 1 \ 1 \ 0 \ 1 \end{bmatrix}_2 \text{ XOR } \begin{bmatrix} 0 \ 1 \ 1 \ 0 \end{bmatrix}_2 = \begin{bmatrix} 1 \ 0 \ 1 \ 1 \end{bmatrix}_2$$

initial config bit 1 at the two sites coupled final config

For S=1, one bit is obviously not sufficent. Use ternary representation or simply occupy two bits to label the 3 states.



Basis representation

- For t-J models at low doping it is useful to factorize hole positions and spin configurations on the occupied sites.
- For Hubbard models one can factorize the Hilbert space in up and down electron configurations.
- For constrained models such as dimer models the efficient generation of all basis states requires some thought.

- One of the key challenges for a fast ED code is to find the index of the new configuration in the list of all configurations (index f in H_{i,f}).
- Let us look at the example of S=1/2 spins at fixed S^z



Basis lookup procedures (Lin tables)

One of the key problems for a fast ED code is to find the index of the new configuration in the list of all configurations (index f in H_{i,f}).

$$[1 \ 0 \ 1 \ 1]_2 = 11_{10}$$

- But is 11 the index of this configuration in a list of all S^z=3/2 states ? no !
- Use Lin tables to map from binary number to index in list of allowed states: (generalization of this idea works for arbitrary number of additive quantum numbers)
- Two tables with 2^(N/2) [=sqrt(2^N)] entries, one for MSBs and one for LSBs

$[0 \ 0]$	=	X	
$[0 \ 1]$	=	0	
$[1 \ 0]$	=	1	
$[1 \ 1]$	=	2	
MSB			

$$\begin{bmatrix} 0 & 0 \end{bmatrix} = X \\
\begin{bmatrix} 0 & 1 \end{bmatrix} = 0 \\
\begin{bmatrix} 1 & 0 \end{bmatrix} = 1 \\
\begin{bmatrix} 1 & 1 \end{bmatrix} = 0$$

LSR

$$Ind([0 \ 1 \ 1 \ 1]) = 0 + 0 = 0$$

$$Ind([1 \ 0 \ 1 \ 1]) = 1 + 0 = 1$$

$$Ind([1 \ 1 \ 0 \ 1]) = 2 + 0 = 2$$

$$Ind([1 \ 1 \ 0]) = 2 + 1 = 3$$



Basis lookup procedures (Lin tables)

- Lookup can therefore be done with two direct memory reads. This is a time and memory efficient approach (at least in many interesting cases).
- An alternative procedure is to build a hash list [const access time] or to perform a binary search [log access time].
- This becomes somewhat more involved when using spatial symmetries...



 $H = \sum_{i,j} J_{i,j}^{xy} (S_i^x S_j^x + S_i^y S_j^y) + J_{i,j}^z S_i^z S_j^z$



$$H = \sum_{i,j} J_{i,j}^{xy} (S_i^x S_j^x + S_i^y S_j^y) + J_{i,j}^z S_i^z S_j^z$$

Consider a XXZ spin model on a lattice. What are the symmetries of the problem ?

$$H = \sum_{i,j} J_{i,j}^{xy} (S_i^x S_j^x + S_i^y S_j^y) + J_{i,j}^z S_i^z S_j^z$$

The Hamiltonian conserves total S^z, we can therefore work within a given S^z sector This easily implemented while constructing the basis, as we discussed before.

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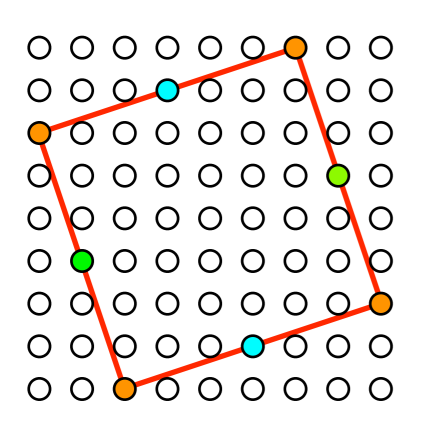
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- At S^z=0 one can use the spin-flip (particle-hole) symmetry which distinguishes even and odd spin sectors at the Heisenberg point. Simple to implement.



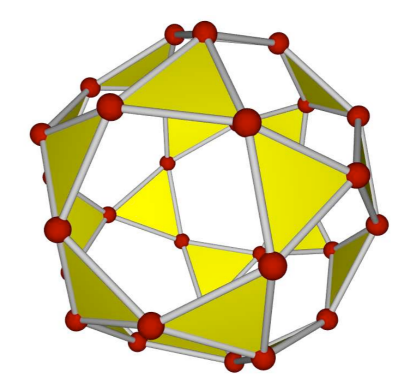
Spatial Symmetries

- Spatial symmetries are important for reduction of Hilbert space
- Symmetry resolved eigenstates teach a lot about the physics at work, dispersion of excitations, symmetry breaking tendencies, topological degeneracy, ...

40 sites square lattice $T \otimes PG = 40 \times 4$ elements



Icosidodecahedron (30 vertices) I_h:120 elements



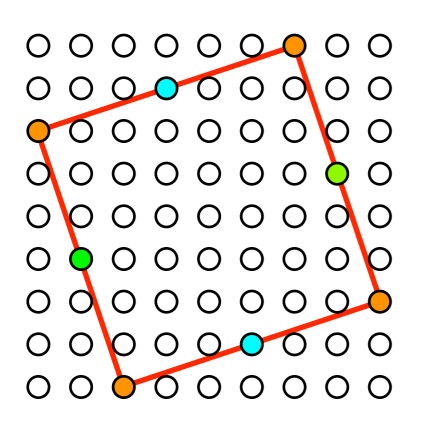


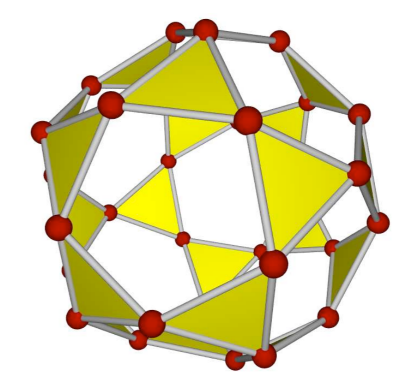
Spatial Symmetries

- Symmetries are sometimes not easily visible, use graph theoretical tools to determine symmetry group [nauty, grape].
- In an ED code a spatial symmetry operation is a site permutation operation. (could become more complicated with spin-orbit interactions and multiorbital sites)

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Icosidodecahedron (30 vertices) I_h:120 elements







- Build a list of all allowed states satisfying the "diagonal" constraints, like particle number, total S^z, ...
- for each state we apply all symmetry operations and keep the state as a representative if it has the smallest integer representation among all generated states in the orbit.
 Example: 4 site ring with cyclic translation *T*, S^z=3/2 sector

$$T^{0}([0\ 1\ 1\ 1]) \rightarrow [0\ 1\ 1\ 1]$$
$$T^{1}([0\ 1\ 1\ 1]) \rightarrow [1\ 0\ 1\ 1]$$
$$T^{2}([0\ 1\ 1\ 1]) \rightarrow [1\ 1\ 0\ 1]$$
$$T^{3}([0\ 1\ 1\ 1]) \rightarrow [1\ 1\ 1\ 0]$$

keep state

 $T^{0}([1 \ 0 \ 1 \ 1]) \rightarrow [1 \ 0 \ 1 \ 1]$ $T^{1}([1 \ 0 \ 1 \ 1]) \rightarrow [1 \ 1 \ 0 \ 1]$ $T^{2}([1 \ 0 \ 1 \ 1]) \rightarrow [1 \ 1 \ 1 \ 0]$ $T^{3}([1 \ 0 \ 1 \ 1]) \rightarrow [0 \ 1 \ 1 \ 1]$

discard state



$$\begin{split} |\tilde{r}\rangle &= \frac{1}{\mathcal{N}\sqrt{|G|}}\sum_{g\in G}\chi(g)|g(r)\rangle\\ \mathcal{N} &= \sqrt{\sum_{g\in G, g(r)=r}\chi(r)} \end{split}$$



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angle$

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- The norm (and therefore the state itself) can vanish if it has a nontrivial stabilizer combined with a nontrivial representation χ .
- Example: 4 site S=1/2 ring with cyclic translations:

$$\begin{array}{l} K = 0 \\ {\rm S}^{\rm Z}=2 & |1 \ 1 \ 1 \ 1 \rangle, {\cal N}=2 \\ {\rm S}^{\rm Z}=1 & |0 \ 1 \ 1 \ 1 \rangle, {\cal N}=1 \\ {\rm S}^{\rm Z}=0 & |0 \ 1 \ 0 \ 1 \rangle, {\cal N}=\sqrt{2} \\ |0 \ 0 \ 1 \ 1 \rangle, {\cal N}=1 \end{array}$$



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$$K = 0$$
 $K = \pm \pi/2$ $S^{z}=2$ $|1\ 1\ 1\ 1\rangle, \mathcal{N} = 2$ $|0\ 1\ 1\ 1\rangle, \mathcal{N} = 1$ $S^{z}=1$ $|0\ 1\ 1\ 1\rangle, \mathcal{N} = 1$ $|0\ 1\ 1\ 1\rangle, \mathcal{N} = 1$ $S^{z}=0$ $|0\ 1\ 0\ 1\rangle, \mathcal{N} = \sqrt{2}$ $|0\ 0\ 1\ 1\rangle, \mathcal{N} = 1$



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- Example: 4 site S=1/2 ring with cyclic translations:

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 $K = \pm \pi/2$
 $K = \pi$
 $S^{z}=2$
 $|1\ 1\ 1\ 1\rangle, \mathcal{N} = 2$
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The Hamiltonian Matrix



The Hamiltonian Matrix

- Now that we have a list of representatives and their norms, can we calculate the matrix elements of the Hamiltonian ? $\langle \tilde{s} | H | \tilde{r} \rangle =$?
- Let us look at an elementary, non-branching term in the Hamiltonian:

$$h^{\alpha}|r\rangle = h^{\alpha}(r)|s\rangle$$

• We can now calculate the matrix element $\langle \tilde{s} | h^{\alpha} | \tilde{r} \rangle$ without double expanding the Bloch states:

$$\langle \tilde{s} | h^{\alpha} | \tilde{r} \rangle = \frac{\mathcal{N}_s}{\mathcal{N}_r} \chi(g^*) h^{\alpha}(r)$$

• key algorithmic problem: given a possibly non-representative $|s\rangle$, how do we find the associated representative $|\tilde{s}\rangle$, as well as a symmetry element g^* relating $|s\rangle$ to $|\tilde{s}\rangle$?



The Hamiltonian Matrix

- key algorithmic problem: given a possibly non-representative $|s\rangle$, how do we find the associated representative $|\tilde{s}\rangle$, as well as a symmetry element g^* relating $|s\rangle$ to $|\tilde{s}\rangle$?
 - Brute force: loop over all symmetry operations applied on $|s\rangle$ and retain $|\tilde{s}\rangle$ and g^* . This is however often not efficient (many hundred symmetries).
 - Prepare a lookup list, relating each allowed configuration with the index of its representative, and also the associated group element linking the two. Gives fast implementation, but needs a list of the size of the non spatially-symmetrized Hilbert space.
 - For specific lattices and models (Hubbard models) clever tricks exist which factorize the symmetry group into a sublattice conserving subgroup times a sublattice exchange. They give $|\tilde{s}\rangle$ fast, then a hash or binary search is needed to locate $|\tilde{s}\rangle$ in the list of representatives in order to get its index.



Hamiltonian Matrix Storage

- Different possibilities exist:
 - Store hamiltonian matrix elements in memory in a sparse matrix format Fast matrix vector multiplies, but obviously limited by available memory.
 - Store hamiltonian matrix elements on disk in a sparse matrix format. In principle possible due to the vast disk space available, but I/O speed is much slower than main memory access times. Difficult to parallelize.
 - Recalculate the hamiltonian matrix elements in each iterations "on the fly". Needed for the cutting edge simulations, where the whole memory is used by the Lanczos vectors. Can be parallelized on most architectures.

The Linear Algebra Backend

Linear Algebra: The most popular: Lanczos Algorithm

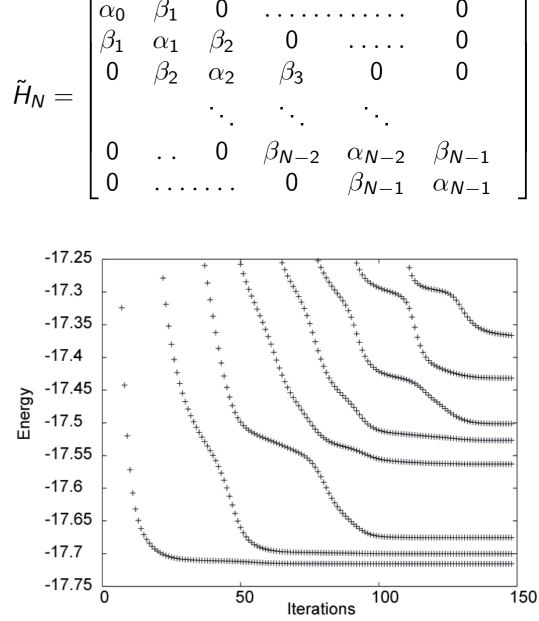


Lanczos Algorithm (C. Lanczos, 1950)

Three vector recursion

n
$$|\phi'\rangle = H|\phi_n\rangle - \beta_n |\phi_{n-1}\rangle$$
,
 $\alpha_n = \langle \phi_n | \phi' \rangle$,
 $|\phi''\rangle = |\phi'\rangle - \alpha_n |\phi_n\rangle$,
 $\beta_{n+1} = ||\phi''|| = \sqrt{\langle \phi'' | \phi'' \rangle}$,
 $|\phi_{n+1}\rangle = |\phi''\rangle / \beta_{n+1}$,

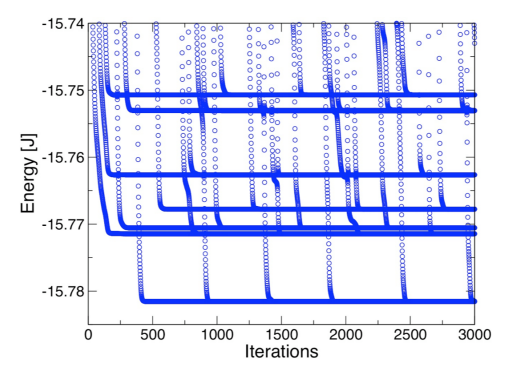
- Eigenvalues of H_N converge rapidly towards eigenvalues of H.
- Once desired eigenvalue is converged, restart recursion and assemble the eigenvector.



very quick convergence for extremal eigenvalues !

Linear Algebra: Lanczos Algorithm

- Once the ground state has converged, the vectors in the recursion tend to lose their orthogonality. As a consequence fake new eigenvalues show up in the approximate spectrum. These can be removed by heuristic techniques



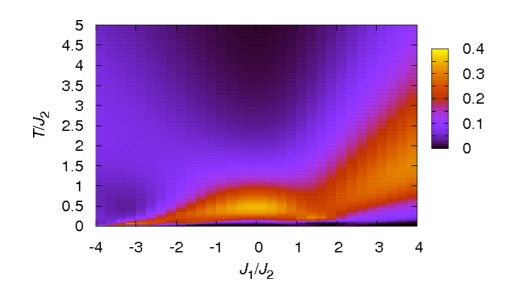
- Degeneracies of eigenvalues can not be resolved by construction. For this task one would need a band lanczos or the (Jacobi-)Davidson technique. However multiply degenerate eigenvalues are converged.
- Checkpointing is useful when performing large-scale simulations.



Full Diagonalization: Thermodynamics

- Lapack / Householder complete diagonalization of the spectrum.
- Calculate partition function and all the thermodynamic quantities you want, often the only pedestrian method available for frustrated systems.
- Symmetries are also very important, because the computational requirements scale as O(D³), where D is the dimension of the block Hilbert space. Typical D's for a workstation are a few 1'000, up to a few 100'000 on supercomputers.

S(N



F. Heidrich-Meisner, A. Honecker, T. Vekua, Phys. Rev. B 74, 020403(R) (2006).

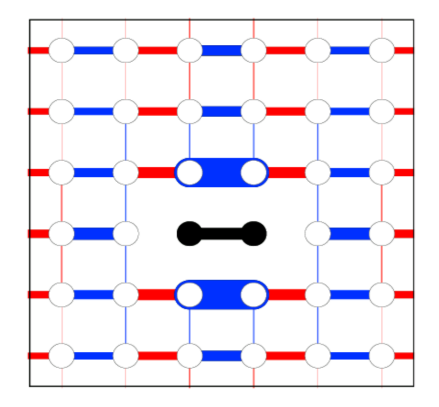
Observables



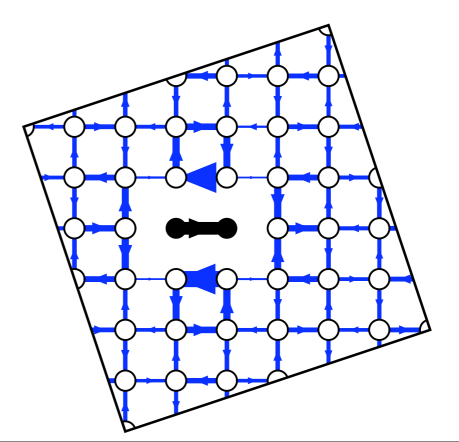
Observables

- In principle once can calculate any correlation function, since one has access to the full many body wave functions. When using spatial symmetries, the correlation functions need to be properly symmetrized too.
- Complicated correlation functions occur in frustrated systems:

Dimer-dimer correlations



Spin current correlations



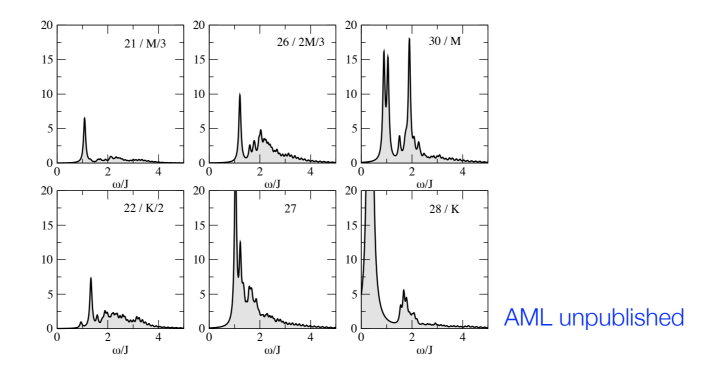


Frequency Dynamics

•
$$G_A(\omega + i\eta) = \langle \psi | A^{\dagger} \frac{1}{E_0 + \omega + i\eta - H} A | \psi \rangle$$
 $A = S^{\alpha}(\mathbf{q}), c_{\mathbf{k}}, \dots$

• Generate Krylov space of $A|\psi\rangle$ Use continued fraction used to invert $(E_0 + \omega + i\eta - H)$

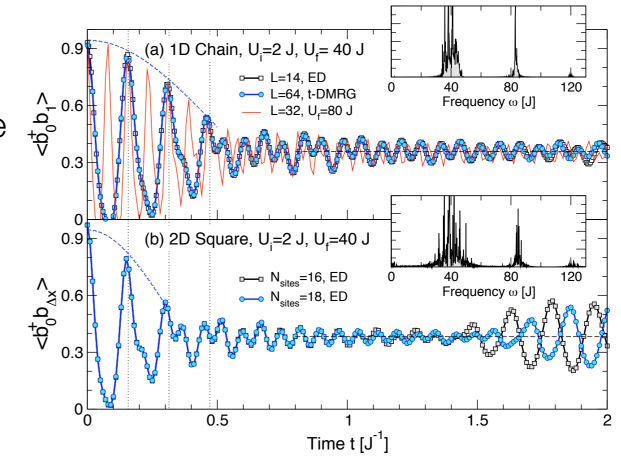
Triangular Lattice Spin Dynamics in zero field



Exact Diagonalization Real-Time Dynamics

ullet It is expensive to obtain the full propagator $\exp[-itH]$

- Krylov methods exist to approximate the propagator for a given state $|\psi(0)\rangle$ One can get the time propagated state $|\psi(t)\rangle$ with only $|v\rangle = H|u\rangle$ operations.
- Example: time evolution of a strongly correlated quantum systems after an abrupt change in the parameters in the Hamiltonian. Revivals and Relaxation.



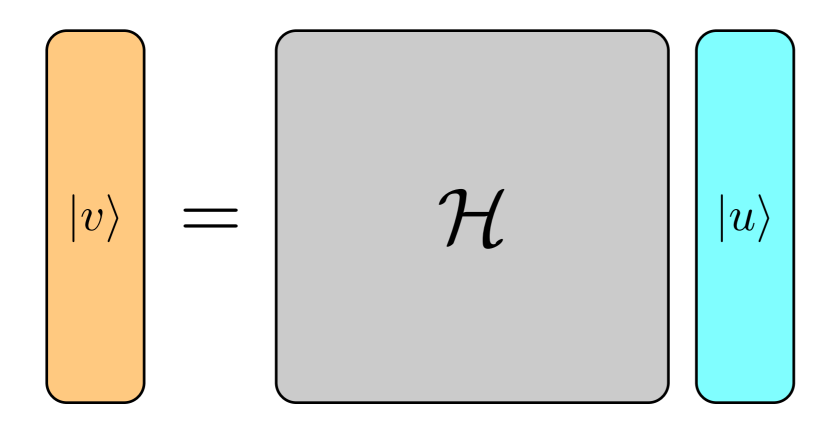
C. Kollath, AML, E. Altman, PRL 2007



Parallelization Strategies

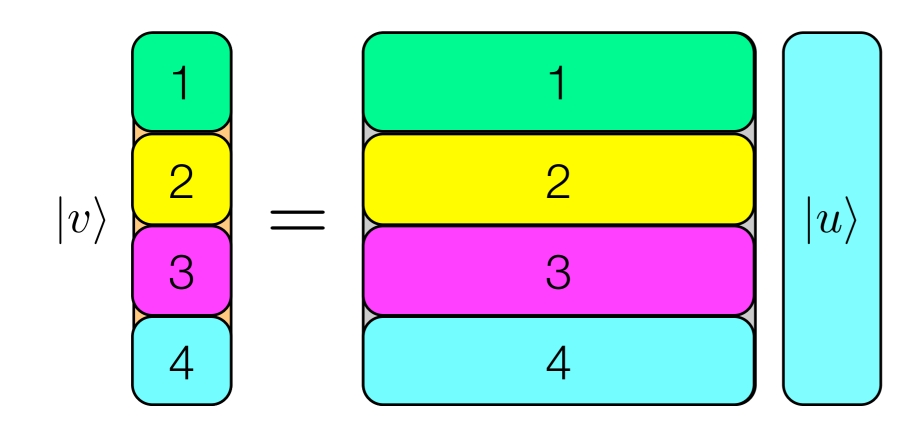
Parallelization: Shared memory nodes

- In the Lanczos algorithm the heaviest part is the elementary matrix-vector multiplication.
- In a matrix-free formulation this part can easily be parallelized using OpenMP pragmas in the code, even on your multi-core workstation. Choose the right strategy between pull and push !



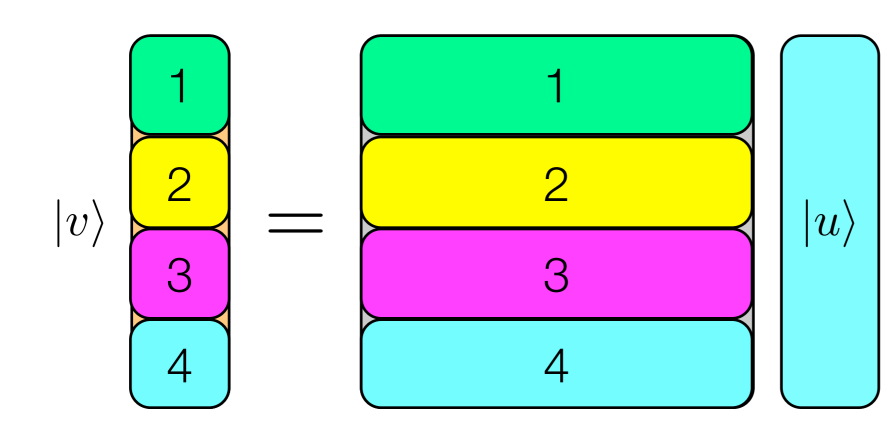
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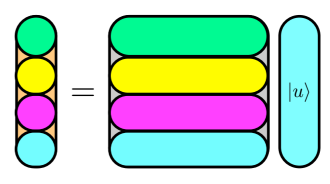
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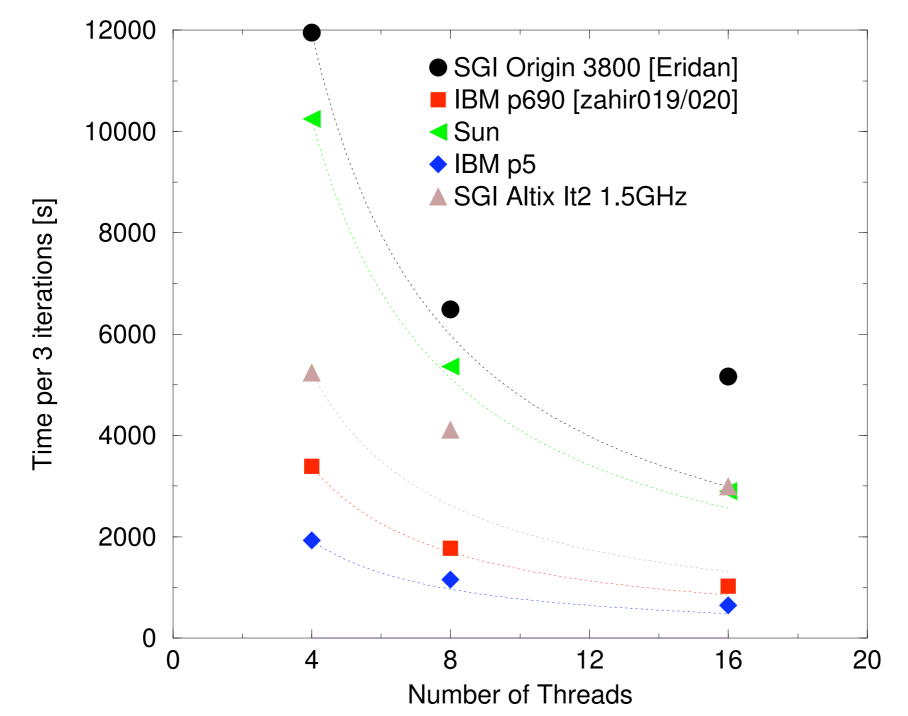


In this parallelization we have uncritical concurrent reads, but no concurrent updates of vector v.

Parallelization: Shared memory nodes



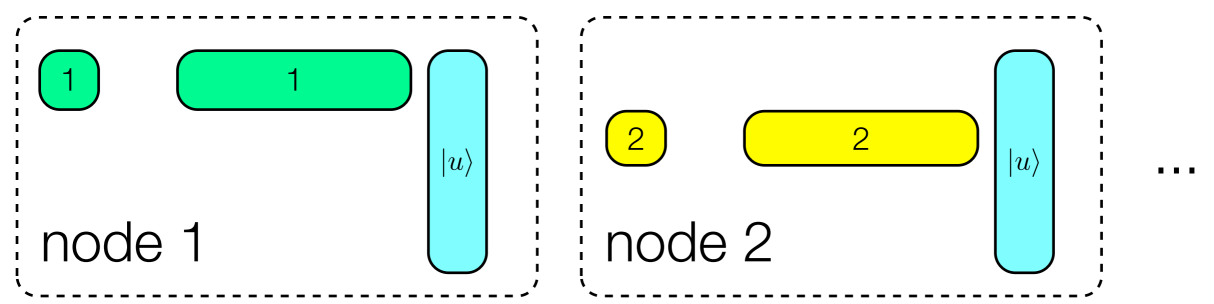




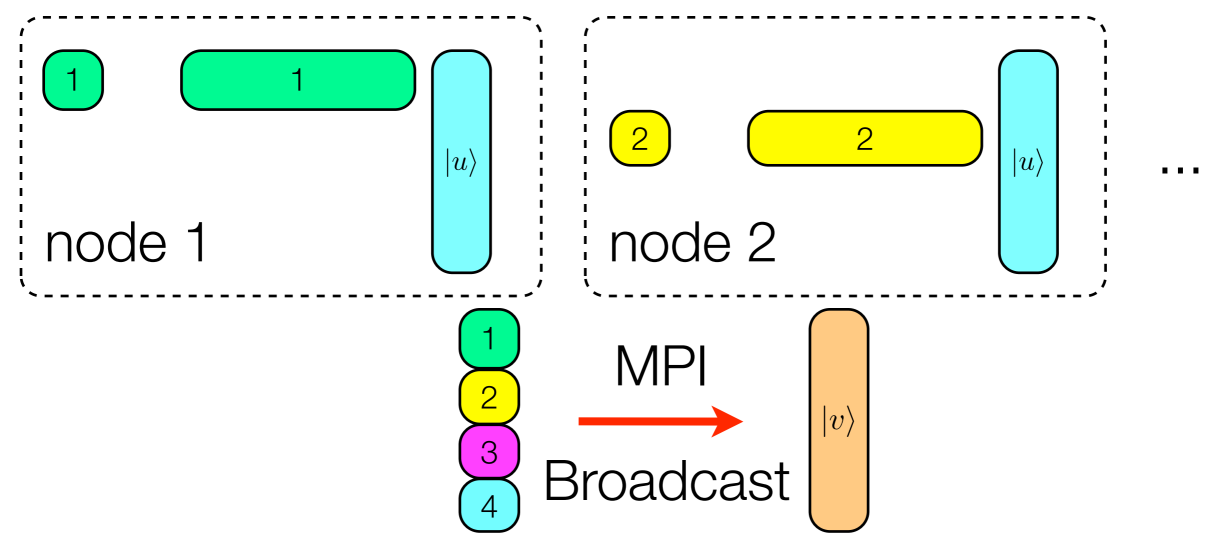
scales well up to a few ten threads on "memory uniform" SMP machines.

- For some classes of problems the Hilbert space size is not too big, but the vast number of matrix elements is a challenge.
 [ED in momentum space formulation & Quantum Hall problems]
- These problems can be OpenMP parallelized, but are also suitable for large scale Message passing parallelization.

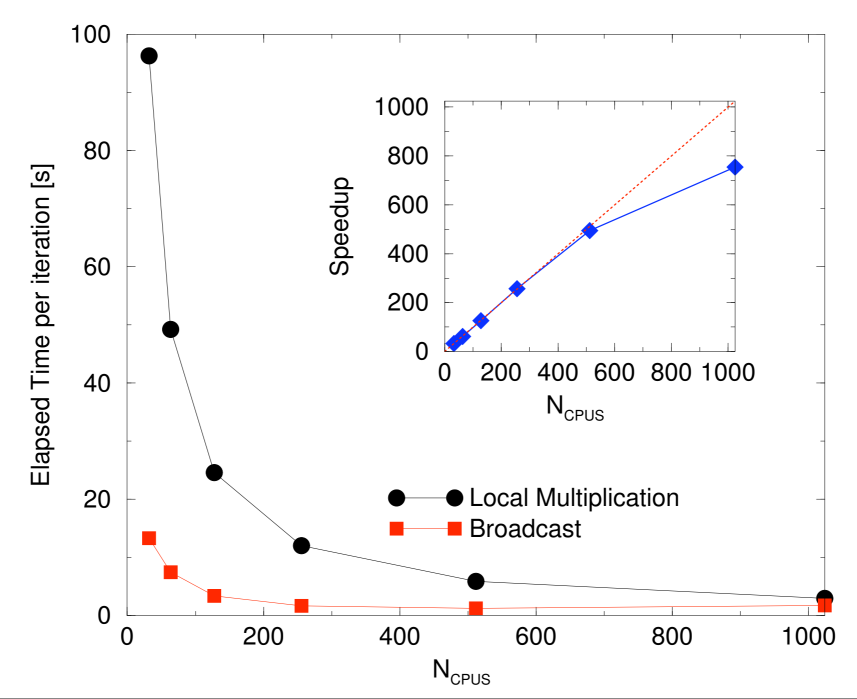
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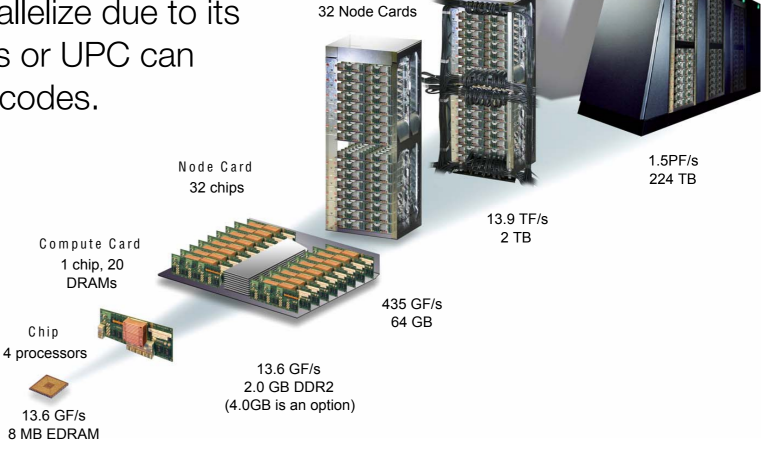


Strong scaling example RG-ED: matrix dimension 10 million performed on a 1024 node Cray XT-3 machine: speedup of ~ 800 on 1024 procs



Parallelization: How to harness the petaflop computers ?

- Cutting edge petaflop systems have a huge number of core, but only a moderate amount of node-local memory.
- Next generation ED codes need to be developed in order to attack e.g. the 80 billion Hilbert space of a 48 site kagome antiferromagnet.
- Problem remains difficult to parallelize due to its all-to-all structure. Global Arrays or UPC can help developing distributed ED codes.



Rack

Cabled

System 112 Racks





 Quantum Magnets: nature of novel phases, critical points in 1D, dynamical correlation functions in 1D & 2D



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Frequency Dynamics (in 2D)

Exact Diagonalization:

numerically determine the low-lying eigenstates of the full many-body Schrödinger equation using Krylov-space techniques.

- Ground state at different total S^z obtained by the Lanczos method
- Dynamical correlations by the continued fraction method $S(Q,\omega)_{\eta} = -\frac{1}{\pi} \text{Im} \langle GS | S(Q)^{\dagger} \frac{1}{\omega - H + E_{GS} + i\eta} S(Q) | GS \rangle$ $S(Q,\omega) = \sum_{n} |\langle \psi_{n} | S(Q) | GS \rangle|^{2} \delta(\omega - E_{n})$
- Typical dimensions dim=10⁸ states, i.e. 64 sites and 200-500 iterations give a good spectrum

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$$S(Q, \omega) = \sum_{\substack{n \\ n \\ n \\ (a)}} \langle \psi_{\pi}|S(Q)|GS \rangle \langle \psi_{\pi}|S(Q)|GS \rangle \langle \psi_{\pi}|S(Q)|GS \rangle$$

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Numerical Approaches (in 2D)

Quantum Monte Carlo:

Highly efficient sampling of the partition function for unfrustrated quantum magnets using e.g. Stochastic Series Expansion (SSE) Sandvik '91,'99

Stochastic Analytical Continuation MC Measurements Measures correlation functions in imaginary time ⁵⁰ Correlation C(Q,ω) Correlation $C(Q, \tau)$ 0. Analytical continuation to real frequency needed (inverse Laplace transform): 0.01 Maximum Entropy, Stochastic Analytical Continuation Jarrell & Gubernatis '96, Sandvik '98, Beach '04 0.001<u></u>∟ 6⁰ 0 2 Imaginary time τ

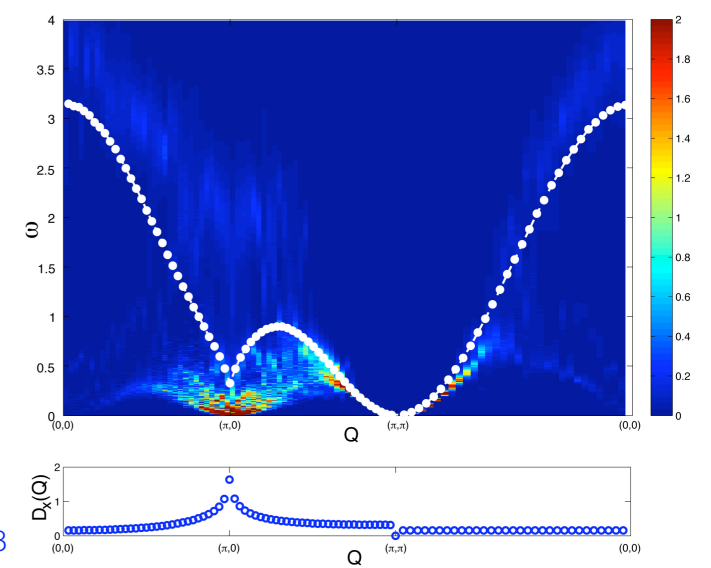
Frequency ω

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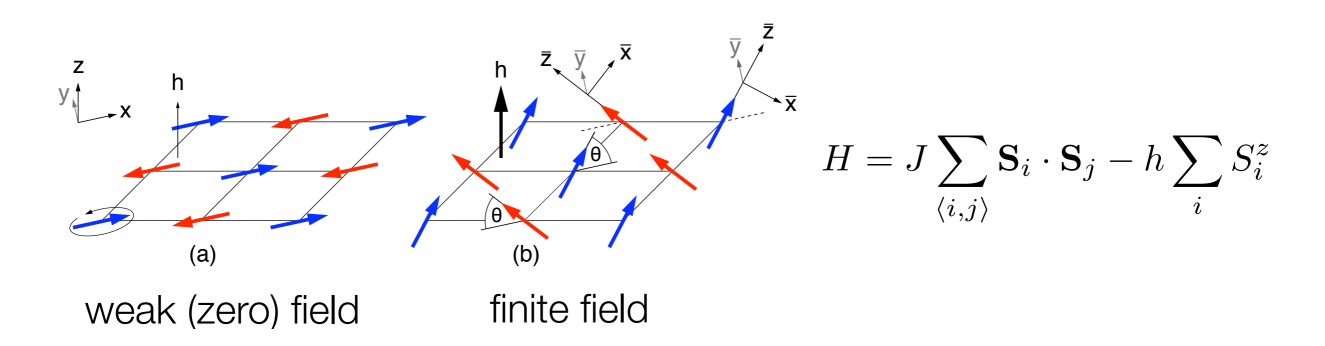
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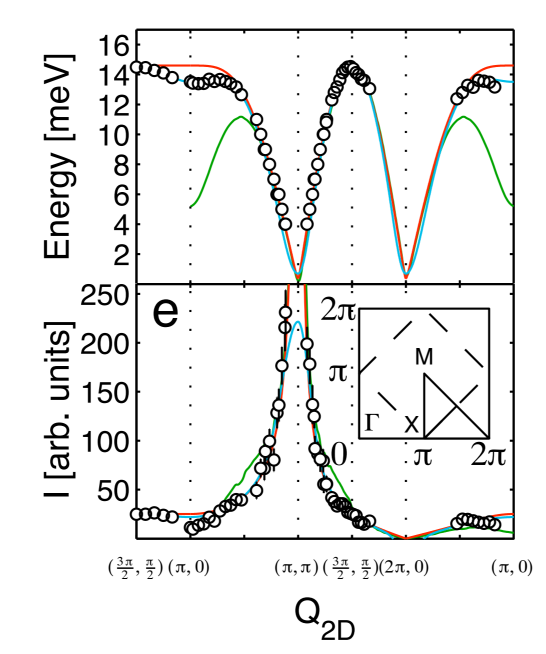
AML, Capponi, Assaad, JSTAT 08

Square Lattice Heisenberg Antiferromagnet



A. Lüscher, AML, PRB '09

 The zero field case is well understood both theoretically and experimentally (Series Expansions; QMC + MaxEnt; Experiments on CTFD, Ronnow '02/Christensen '07)



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- In a magnetic field the situation is much less clear:
 - Theory: two conflicting predictions:
 M.E. Zhitomirsky & A.L. Chernyshev (PRL 99)
 interacting spin wave theory → magnons decay above a threshold field of approximately 3/4 of the saturation field.
 - O. Syljuåsen & P.A. Lee (PRL 02)
 - π flux state mean-field calculations \rightarrow no evidence for magnon decay, however low energy spectral weight in a region where spin wave theory predicts none.

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 - π flux state mean-field calculations \rightarrow no evidence for magnon decay, however low energy spectral weight in a region where spin wave theory predicts none.
 - Experiments: not yet performed (or on the way ?) ...
 - Numerical simulation can help to settle this issue

Dynamical Spin Correlations in a Field

In a magnetic field the SU(2) symmetry is reduced to U(1)

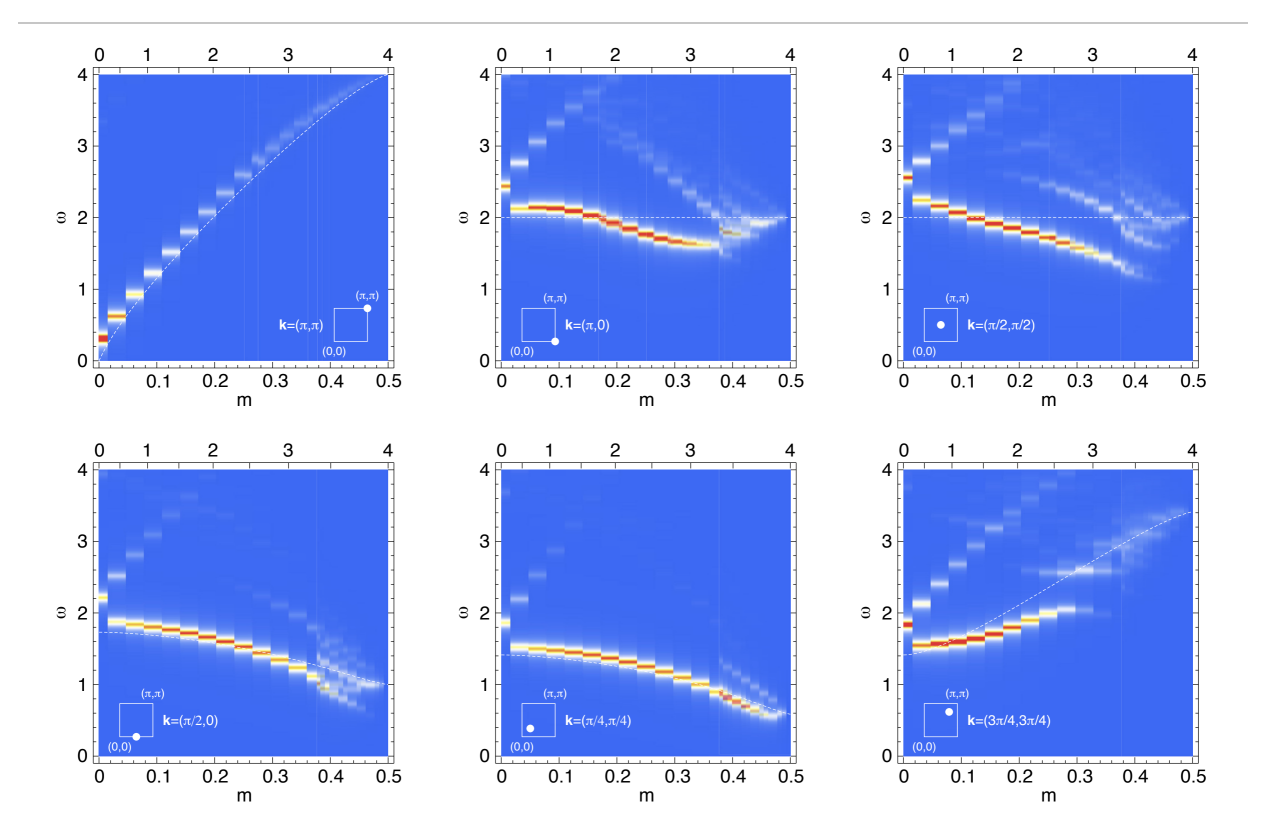
The relevant spin correlators are

- The longitudinal response: $S^{zz}(Q,\omega)$
- The transverse response:

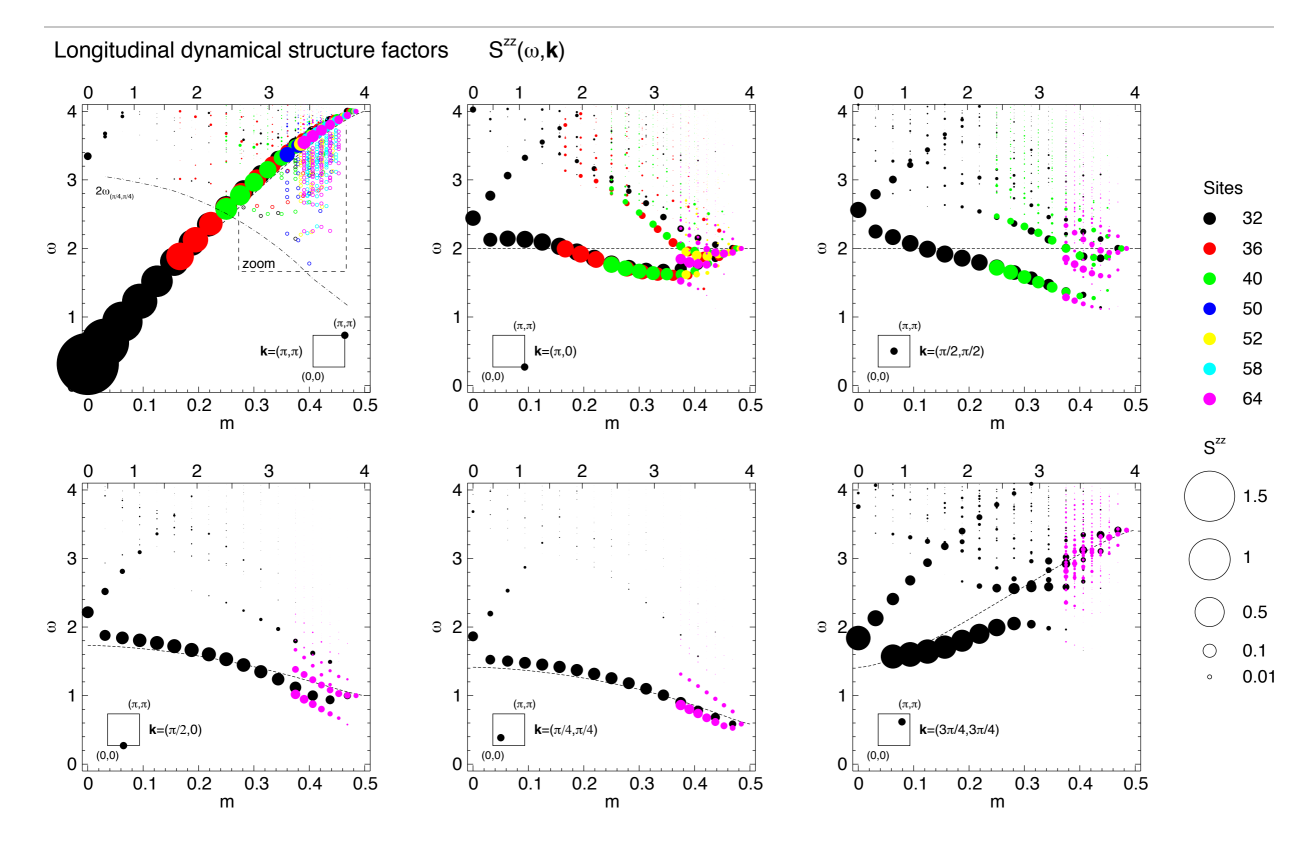
$$S^{xx}(Q,\omega) = S^{yy}(Q,\omega) = \frac{1}{4} \left[S^{+-}(Q,\omega) + S^{-+}(Q,\omega) \right]$$

In the present case the transverse response is to a very good approximation equal to the longitudinal response shifted by (π, π) .

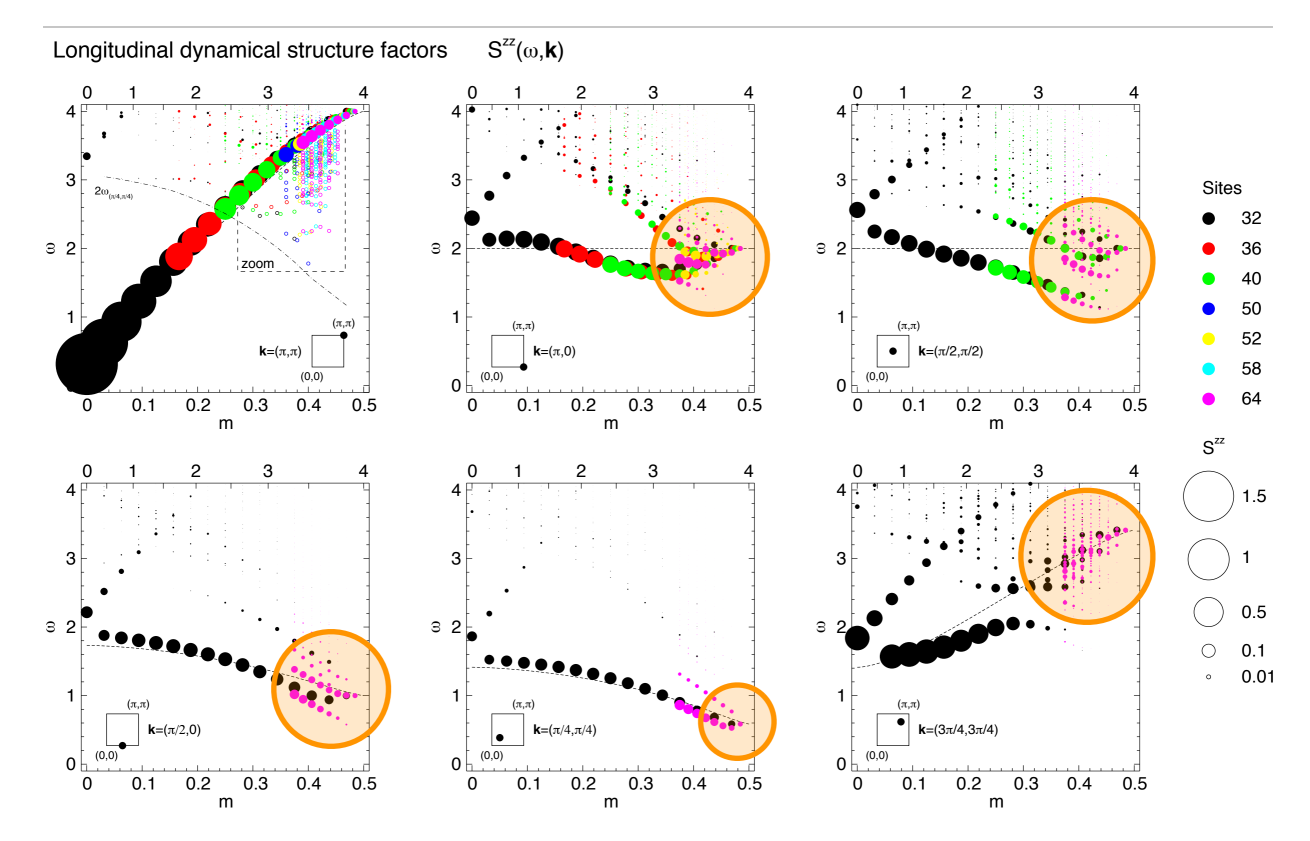
Predicted INS Spectra as a function of field



Field dependence: Finite size pole structure



Field dependence: Finite size pole structure



Field dependence: Finite size pole structure

 $S^{zz}(\omega, \mathbf{k})$ Longitudinal dynamical structure factors 2 2 3 2 3 0 3 4 Λ 4 Ω **4** P 3 З 2ω_(π/4,π/4) з 2 З З zoom 1 1 (π,π) (π,π) (π,π) **k**=(π/2,π/2) **k**=(π,0) **k**=(π,π) • (0,0)(0,0) (0,0) 0 0 0.1 0.2 0.3 0.4 0.5 0.2 0.3 0.4 0.5 0.1 0.2 0.3 0.4 0.5 0 0 0.1 0 m m m 0 2 3 0 2 3 3 n 4 4 4 3 3 3 з 2 з 2 З 1 1 (π,π) (π,π) (π,π) $\mathbf{k} = (3\pi/4, 3\pi/4)$ $k = (\pi/2.0)$ $k = (\pi/4, \pi/4)$ (0,0) (0,0)(0,0) 0 0 h 0 0 0.1 0.2 0.3 0.4 0.5 0 0.1 0.2 0.3 0.4 0.5 0 0.1 0.2 0.4 0.5 0.3 m m m

Sites

32

36

40

50

52

58

64

1.5

1

0.5

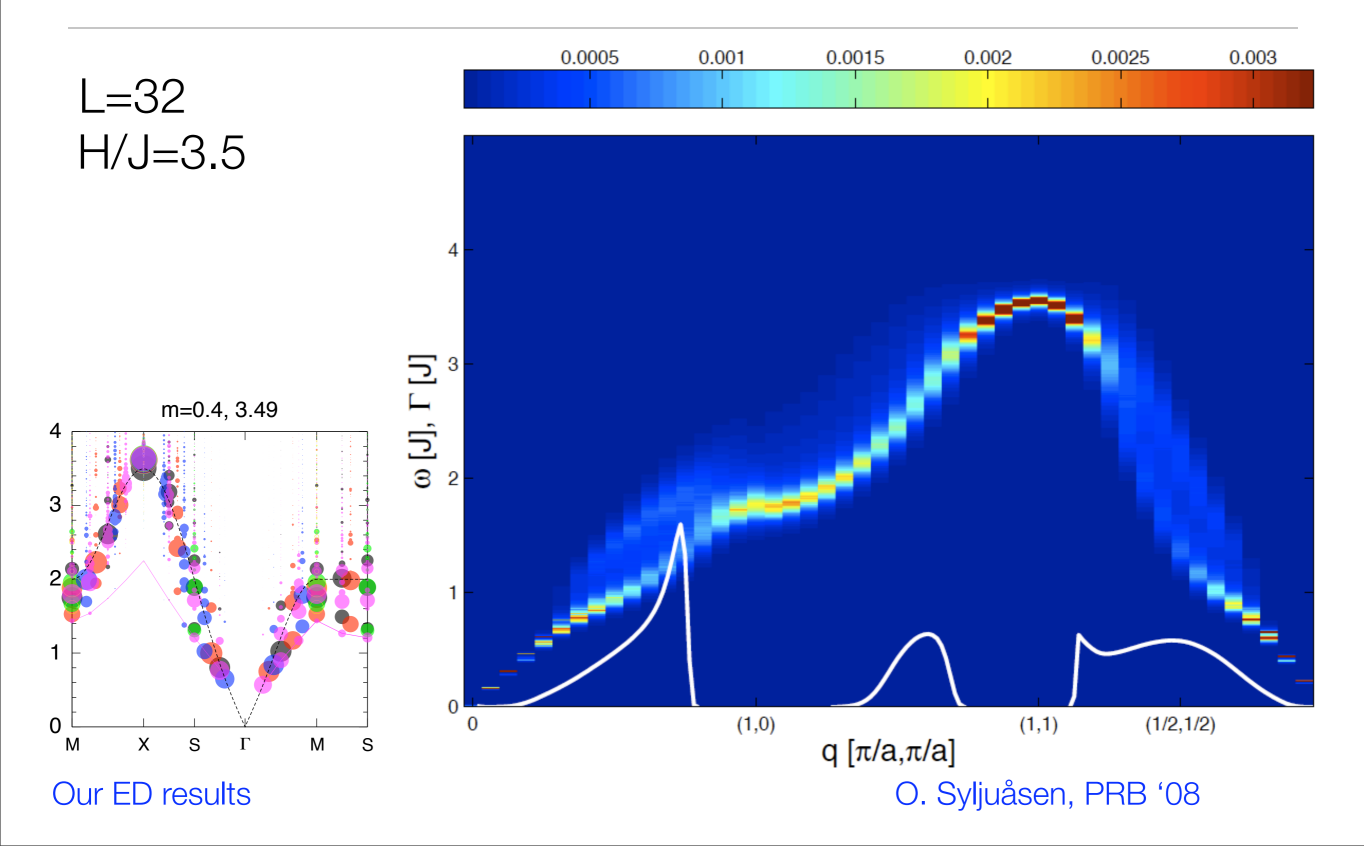
0.1

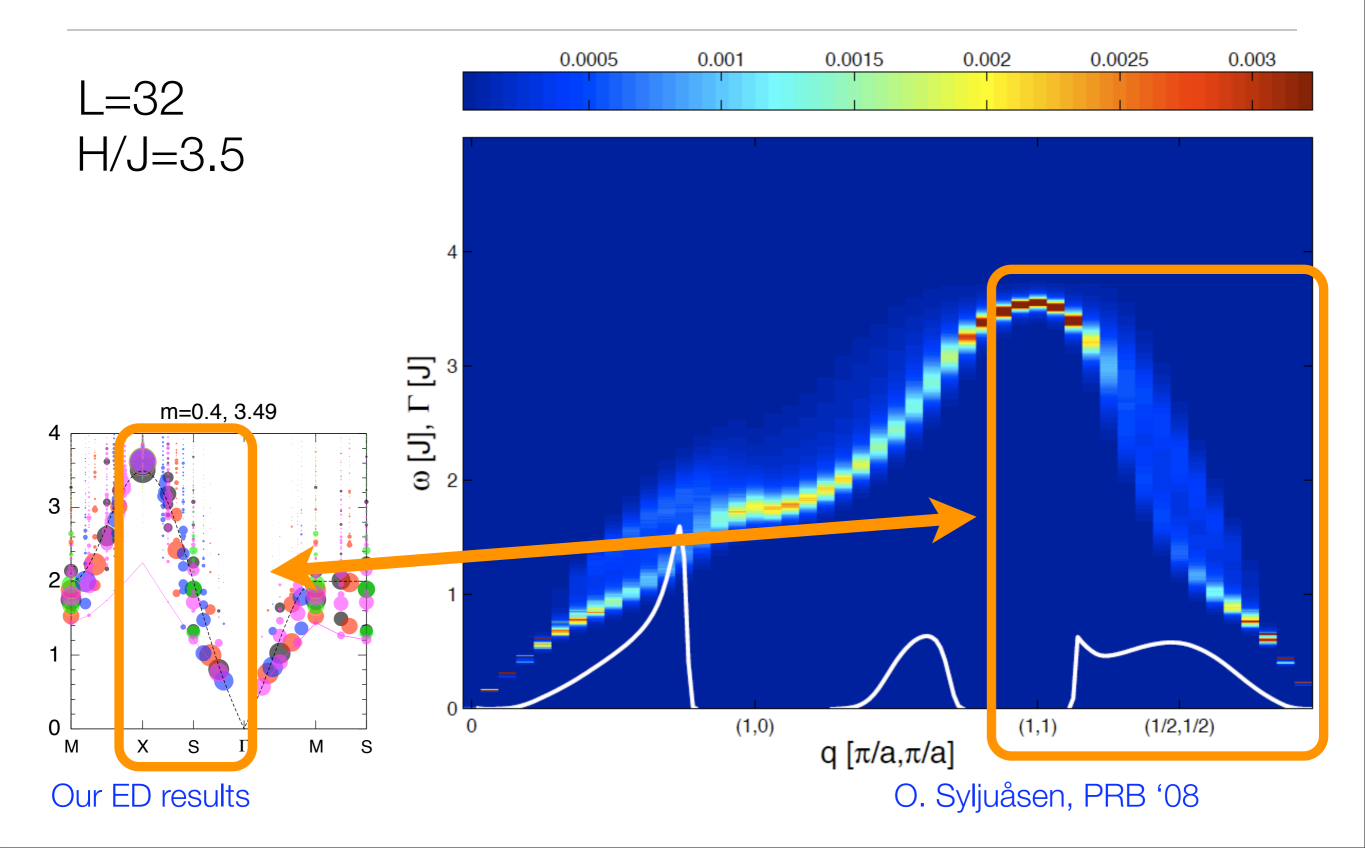
0.01

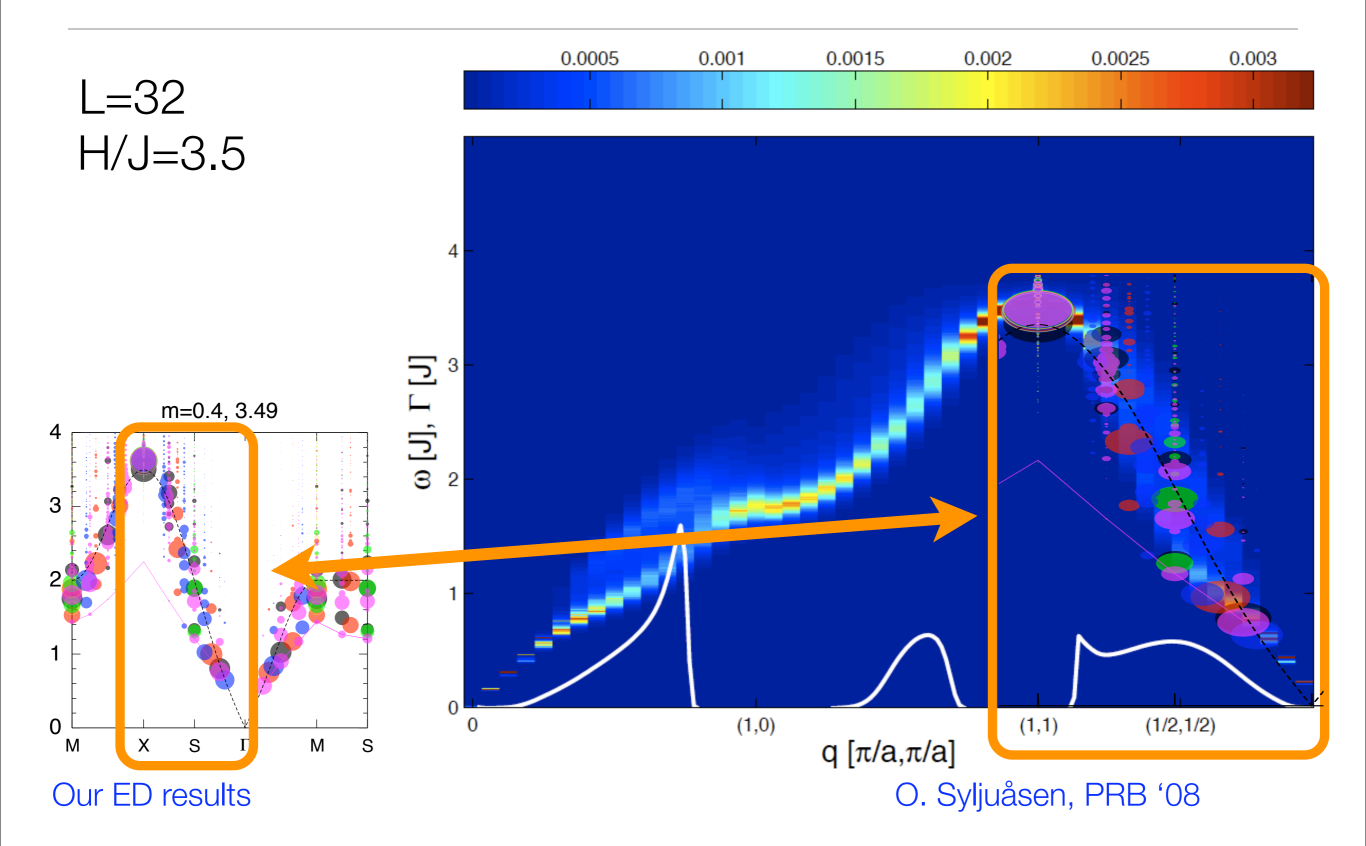
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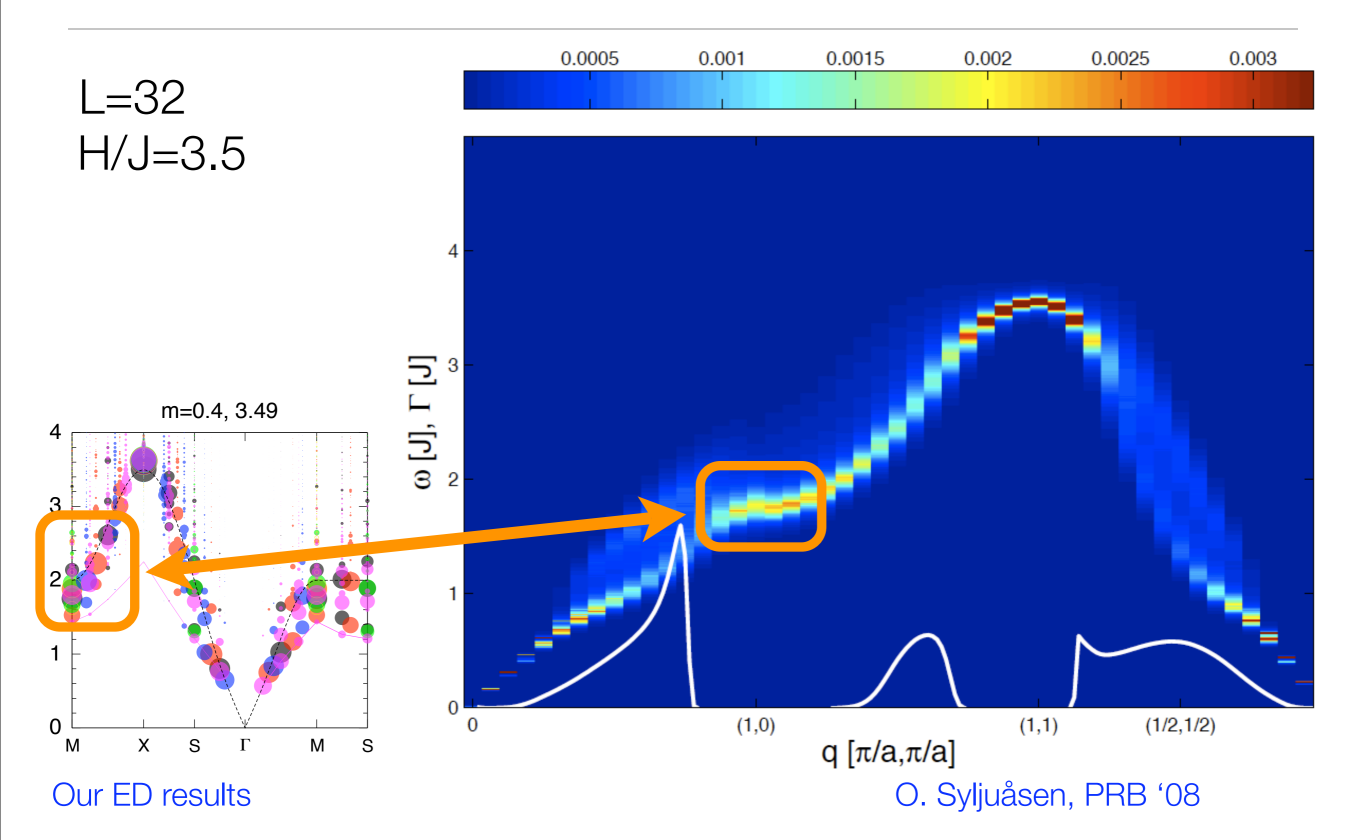
0

 S^{zz}

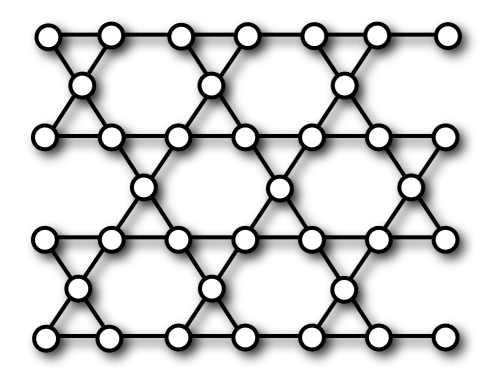








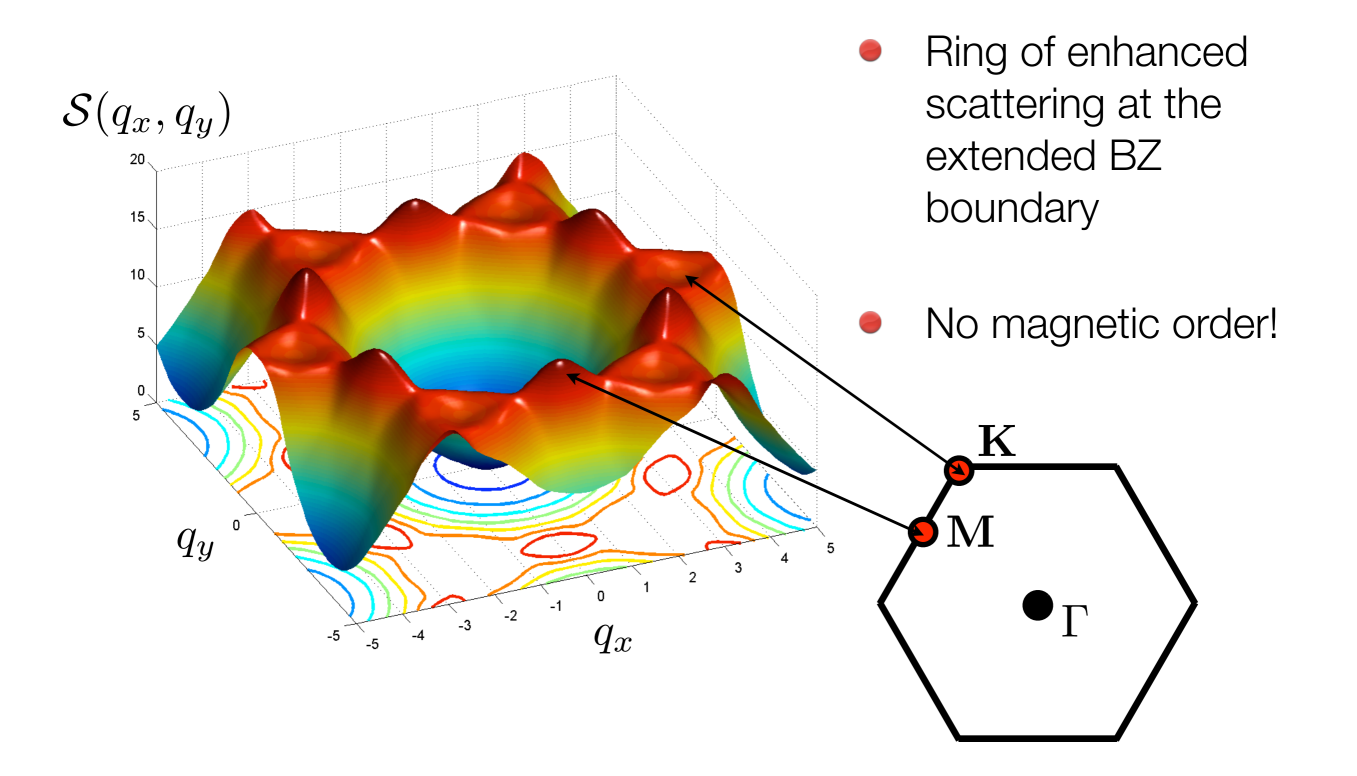
Kagome Antiferromagnet



$$H = J \sum_{\langle i,j \rangle} \mathbf{S}_i \cdot \mathbf{S}_j$$

AML & C. Lhuillier, arXiv:0901.1065

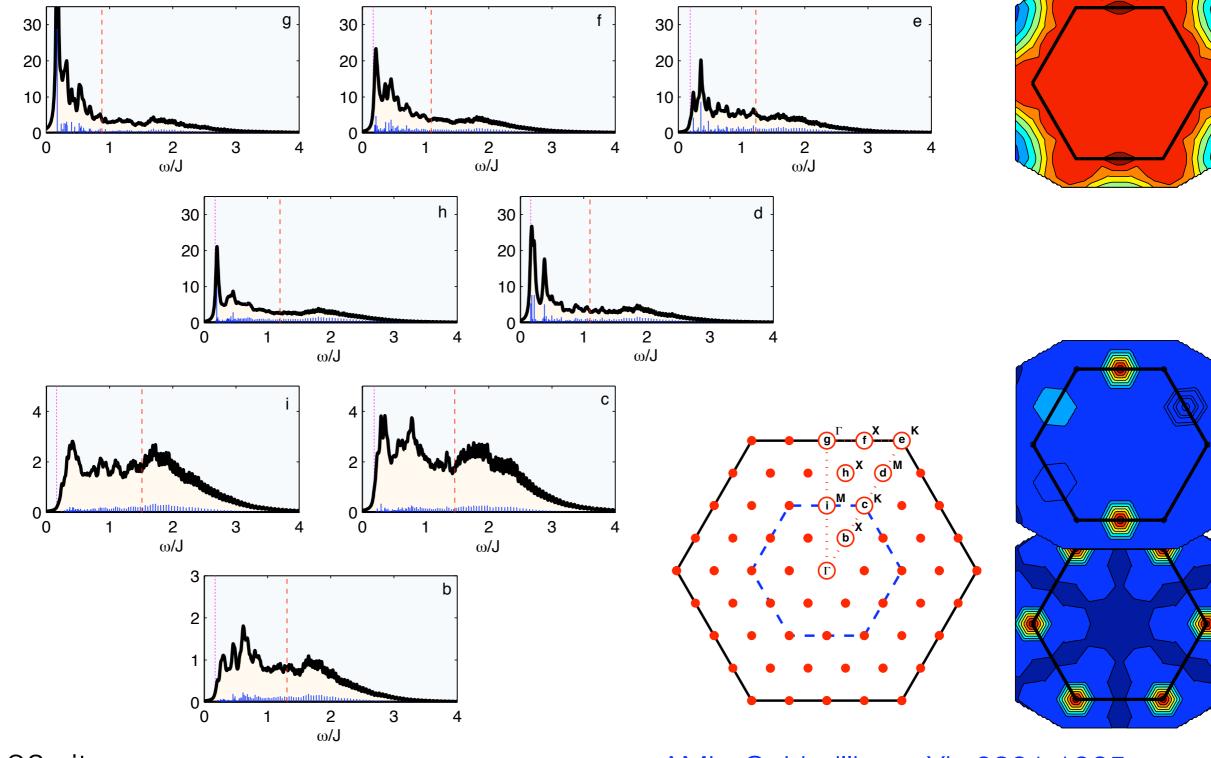
Kagome AFM Static Structure Factor







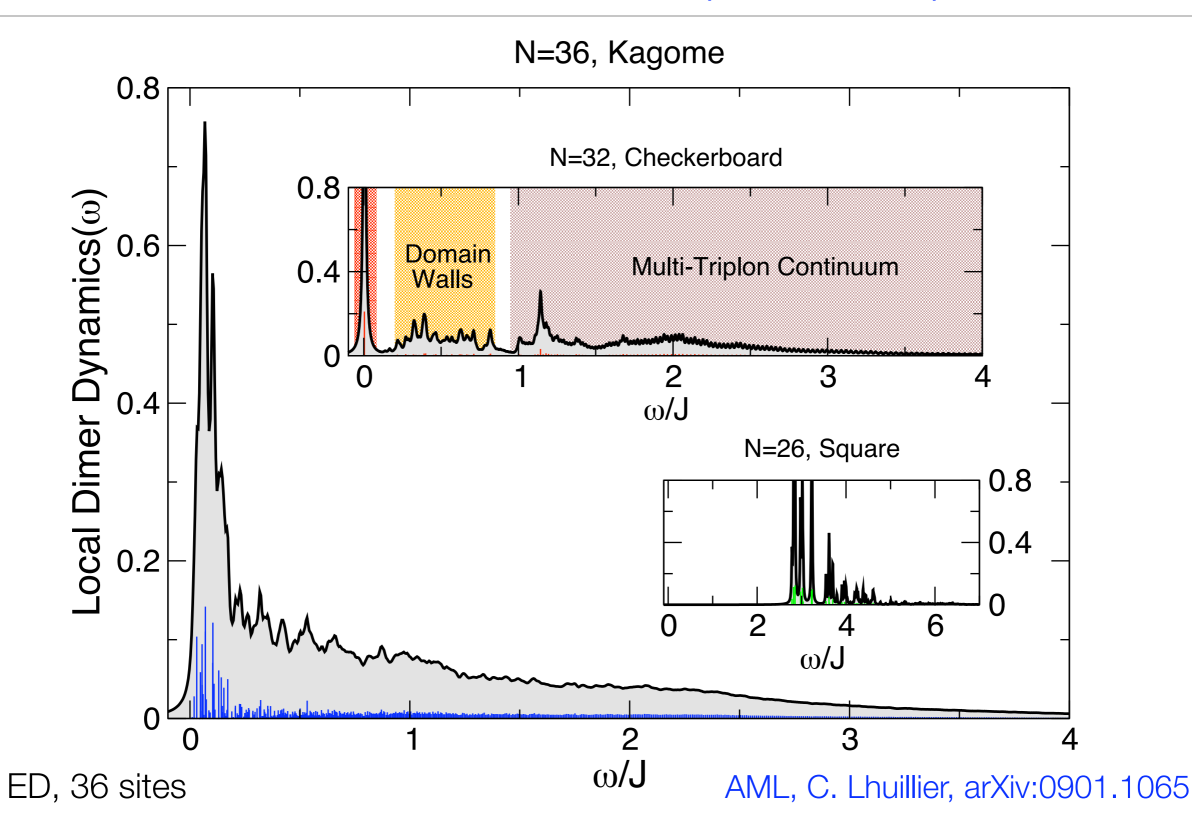
Kagome AFM Dynamical Spin Structure Factor (~ INS)



ED, 36 sites

AML, C. Lhuillier, arXiv:0901.1065

Kagome AFM Local Dimer Autocorrelations (~ Raman)





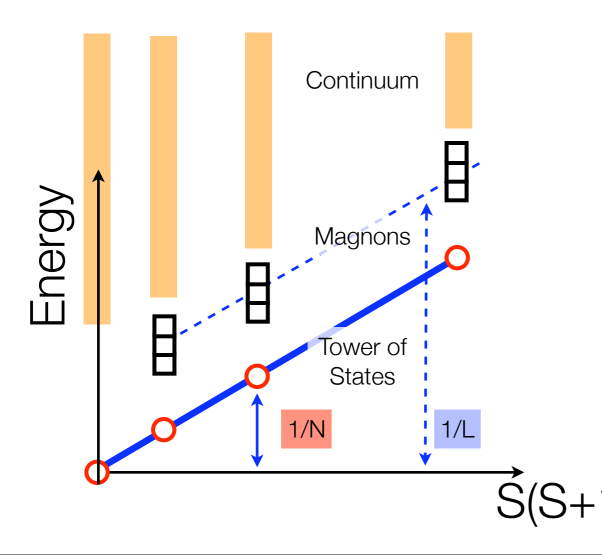
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"Tower of States" spectroscopy

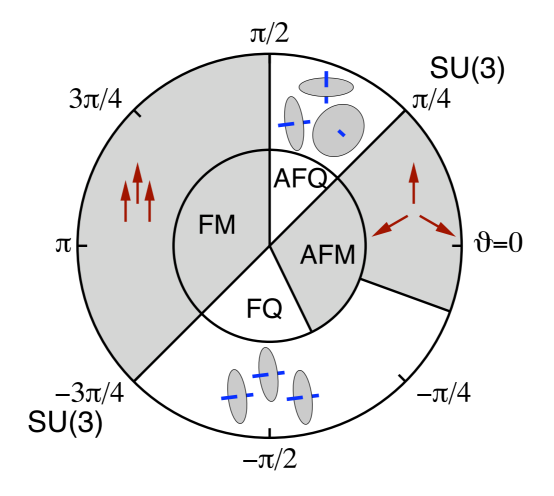
- What are the finite size manifestations of a continuous symmetry breaking ?
- Low-energy dynamics of the order parameter
 Theory: P.W. Anderson 1952, Numerical tool: Bernu, Lhuillier and others, 1992 -



- Dynamics of the free order parameter is visible in the finite size spectrum. Depends on the continuous symmetry group.
- U(1): $(S^z)^2$ SU(2): S(S+1)
- Symmetry properties of levels in the Tower states are crucial and constrain the nature of the broken symmetries.

Tower of States S=1 on triangular lattice

• Bilinear-biquadratic S=1 model on the triangular lattice (model for NiGaS₄).

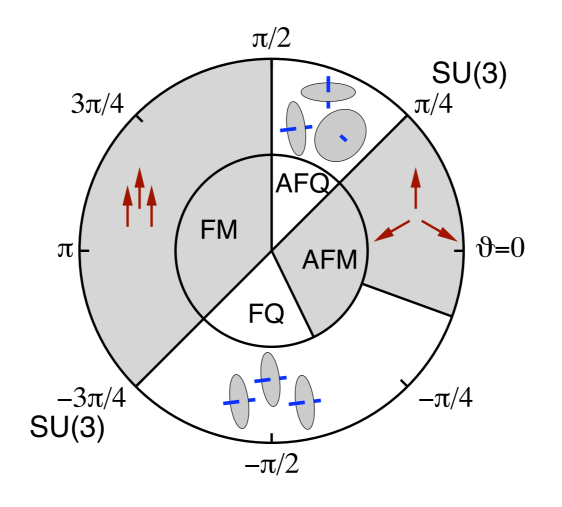


AML, F. Mila, K. Penc, PRL '06

Tower of States S=1 on triangular lattice

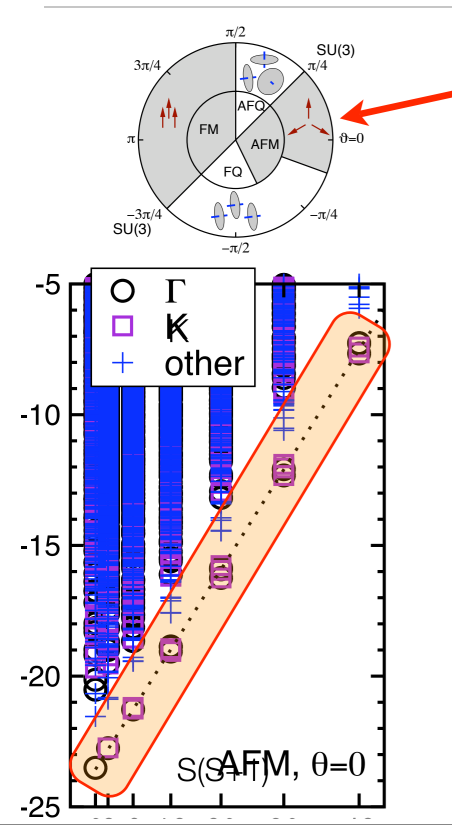
Bilinear-biquadratic S=1 model on the triangular lattice (model for NiGaS₄).

$$H = \sum_{\langle i,j \rangle} \cos(\theta) \, \mathbf{S}_i \cdot \mathbf{S}_j + \sin(\theta) \, \left(\mathbf{S}_i \cdot \mathbf{S}_j\right)^2$$



AML, F. Mila, K. Penc, PRL '06

Tower of States S=1 on triangular lattice: Antiferromagnetic phase

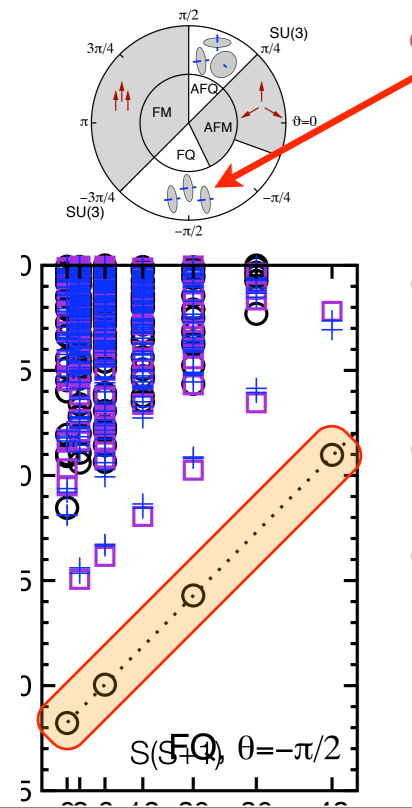


- 9=0 : coplanar magnetic order,
 - 120 degree structure
- Breaks translation symmetry. Tree site unit cell
 ⇒ nontrivial momenta must appear in TOS
- non-collinear magnetic structure
 \Rightarrow SU(2) is completely broken,

number of levels in TOS increases with S

Quantum number are identical to the S=1/2 case

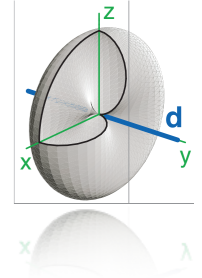
Tower of States S=1 on triangular lattice: Ferroquadrupolar phase



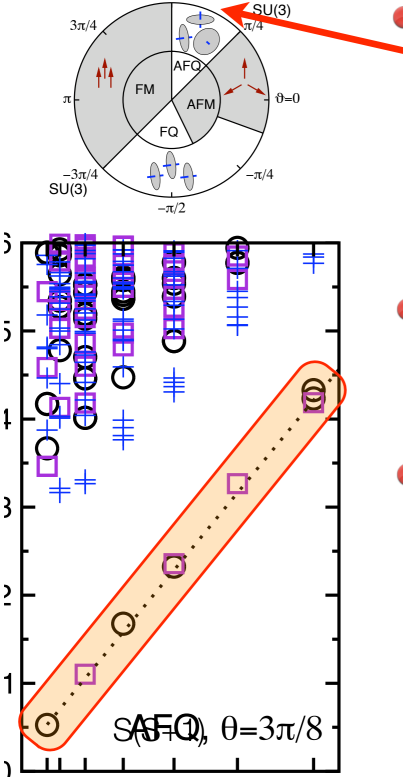
 $9=-\pi/2$: ferroquadrupolar phase, finite quadrupolar moment, no spin order

- No translation symmetry breaking.
 \Rightarrow only trivial momentum appears in TOS
- Ferroquadrupolar order parameter, only even S
- all directors are collinear
 - \Rightarrow SU(2) is broken down to U(1),

number of states in TOS is independent of S.



Tower of States S=1 on triangular lattice: Antiferroquadrupolar phase



 $\pi/2$

 $9=3\pi/8$: antiferroquadrupolar phase, finite quadrupolar moment, no spin order, three sublattice structure.

- Breaks translation symmetry. Tree site unit cell ⇒ nontrivial momenta must appear in TOS
- Antiferroquadrupolar order parameter, complicated S dependence. Can be calculated using group theoretical methods.

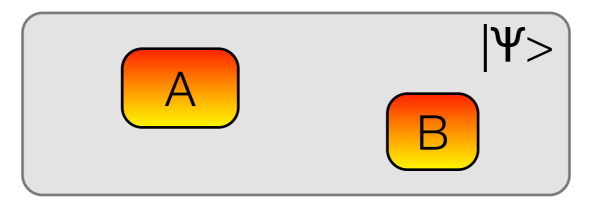


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The correlation density matrix (CDM)



- Is there a systematic way to detect important correlations between parts A and B of a larger system ?
- The correlation density matrix:

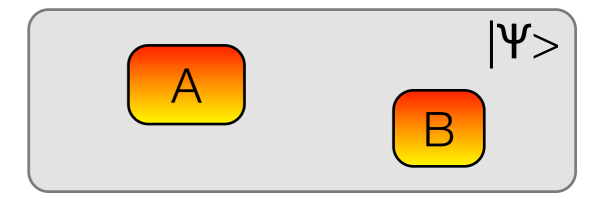
$$\rho_{AB}^{c} = \rho_{AB} - \rho_{A} \otimes \rho_{B}$$

contains all the required information



The correlation density matrix (CDM)

$$\rho_{AB}^{c} = \rho_{AB} - \rho_{A} \otimes \rho_{B}$$



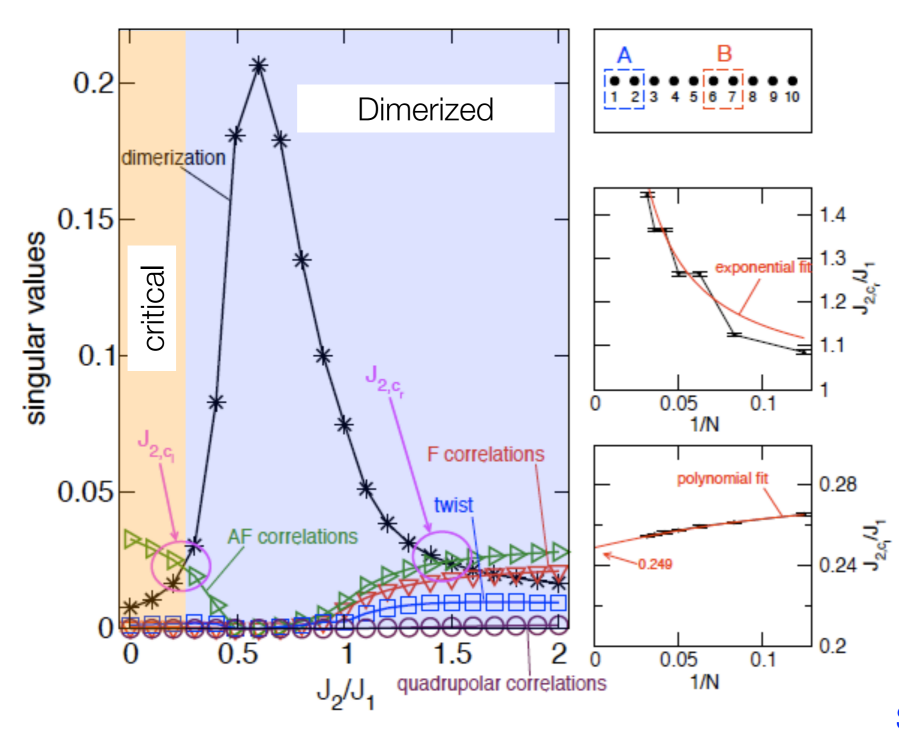
- Contains all information on any connect correlation function between A and B: $Tr(\rho_{AB}^{c}\widehat{O}_{A}\widehat{O}_{B}) = \langle \widehat{O}_{A}\widehat{O}_{B} \rangle - \langle \widehat{O}_{A} \rangle \langle \widehat{O}_{B} \rangle$
- The key step is to perform a singular value decomposition

$$\rho_{AB}^{c} = \sum_{i=1}^{c} \sigma_i X_i' Y_i'^{\dagger}$$

where the σ_i give the strength of the correlation i and the X_i and Y_i are the operators of the correlator acting in A and B.

S.-A. Cheong, C. Henley, arXiv:0809.0075

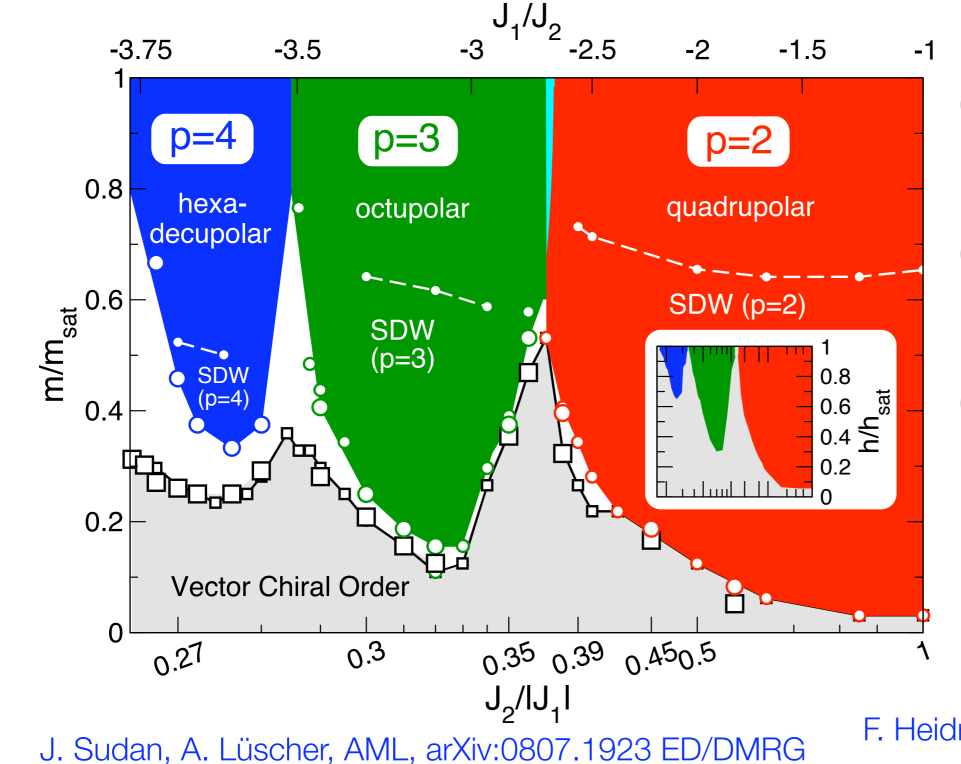
CDM J₁-J₂ frustrated Heisenberg Chain (all AF)



- Benchmark on existing phase diagrams.
- singular values
 respect SU(2)
 symmetry in S=0 GS
 (multiplicities).
- works very well for the well understood Majumdar-Ghosh chain.

Sudan & AML, unpublished

CDM J₁-J₂ frustrated Heisenberg Chain (F-AF)



- vector chiral phase at low m
- spin multipolar liquids at high m
- CDM helped us understand that spin multipolar phases are generically imprinted in close-by magnetically ordered states

F. Heidrich-Meisner et al. PRB '06 T. Hikihara et al., PRB '08



Conclusions

- Exact Diagonalization has an obvious disadvantage (finite size limitation), but when combined with physical concepts and ideas the method becomes a powerful Quantum Mechanics Toolbox, and can access systems which are difficult or impossible to solve otherwise.
- Dynamical correlation functions gave evidence for decay of spin waves in the square lattice antiferromagnet in a field, while the dynamical spin response of the kagome lattice is very incoherent, with possibly some VBC-triplon remnants at low energy.
- Tower of states spectroscopy is powerful tool to study continuous symmetry breaking.
- Correlation Density Matrices are a novel tool to study correlations (or the absence thereof) in unified framework. First applications to frustrated spin chains revealed new mechanisms for the appearance of spin nematic phases.



Exact Diagonalization Literature

N. Laflorencie & D. Poilblanc,

"Simulations of pure and doped low-dimensional spin-1/2 gapped systems" Lecture Notes in Physics 645, 227 (2004).

R.M. Noack & S. Manmana,

"Diagonalization- and Numerical Renormalization-Group-Based Methods for Interacting Quantum Systems", AIP Conf. Proc. 789, 93 (2005).

A. Weisse, H. Fehske

"Exact Diagonalization Techniques" Lecture Notes in Physics 739, 529 (2008).

A. Läuchli

"Numerical Simulations of Frustrated Systems"

to appear in "Highly Frustratred Magnetism", Eds. Lacroix Mendels, Mila, (2009).

Thank you !