# Exact and quasi exact numerical methods for giant magnetic molecules

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Problem

# The problem

# You have got a molecule!



**Congratulations!** 

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#### You have got an idea about the modeling!

$$\begin{array}{lll} H &=& -2\sum_{i < j} \,\, J_{ij} \,\, \vec{\underline{s}}(i) \cdot \vec{\underline{s}}(j) & + \\ & & \\ &$$

$$g\,\mu_B\,B\,\sum_i^N \,\underline{s}_z(i)$$

Zeeman

#### In the end it's always a big matrix!



Fe<sup>III</sup><sub>10</sub>: N = 10, s = 5/2

Dimension=60,466,176. Maybe too big?

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#### Thank God, we have computers



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#### **★ ↔ → □** ? **×**

### Magnetic Molecules



### possess symmetries! Use them!

**★ ↔ → → □** ? **×** 

## Key question:

# Can we employ SU(2) symmetry together with arbitrary

point group symmetries?

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# SU(2) symmetry

#### Quantum chemists need to be much smarter since they have smaller computers!

(1) D. Gatteschi and L. Pardi, Gazz. Chim. Ital. 123, 231 (1993).

(2) J. J. Borras-Almenar, J. M. Clemente-Juan, E. Coronado, and B. S. Tsukerblat, Inorg. Chem. 38, 6081 (1999).

(3) B. S. Tsukerblat, *Group theory in chemistry and spectroscopy: a simple guide to advanced usage*, 2nd ed. (Dover Publications, Mineola, New York, 2006).

#### Irreducible Tensor Operator approach



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#### Spin rotational symmetry:

- $H_{\sim} = -2 \sum_{i < j} J_{ij} \, \vec{\underline{s}}_i \cdot \vec{\underline{s}}_j + g \mu_B \vec{\underline{S}} \cdot \vec{B}$ ;
- $\left[ \underbrace{H}_{\sim}, \underbrace{\vec{S}^2}_{\sim} \right] = 0, \left[ \underbrace{H}_{\sim}, \underbrace{S}_{z}_{\sim} \right] = 0;$
- Irreducible Tensor Operator (ITO) approach;
- Free program MAGPACK (2) available.

(1) D. Gatteschi and L. Pardi, Gazz. Chim. Ital. 123, 231 (1993).

(2) J. J. Borras-Almenar, J. M. Clemente-Juan, E. Coronado, and B. S. Tsukerblat, Inorg. Chem. 38, 6081 (1999).

#### Idea of ITO

$$\begin{array}{lll} \underset{\sim}{H} \text{Heisenberg} &=& -2\sum_{i < j} J_{ij} \; \vec{s}_i \cdot \vec{s}_j \\ &=& 2\sqrt{3} \sum_{i < j} J_{ij} \; \vec{T}^{(0)}(\{k_i\}, \{\overline{k}_i\} | k_i = k_j = 1) \end{array}$$

#### **Irreducible Tensor Operator approach**

- Express spin operators and functions thereof as ITOs;
- Use vector coupling basis  $| \alpha S M \rangle$  and recursive recoupling.
- (1) Gatteschi, Tsukerblat, Coronado, Waldmann, ...
- (2) R. Schnalle, Ph.D. thesis, Osnabrück University (2009)

Point group symmetry

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# Point group symmetry (Ph.D. of Roman Schnalle)

#### **Point Group Symmetry I**

$$|\alpha' S M \Gamma\rangle = \mathcal{P}^{(\Gamma)} |\alpha S M\rangle = \left(\frac{l_{\Gamma}}{h} \sum_{R} \left(\chi^{(\Gamma)}(R)\right)^* \tilde{G}(R)\right) |\alpha S M\rangle$$

#### Method:

- Basis function generating machine (1);
- Projection on irreducible representations  $\Gamma$  (Wigner);
- Orthonormalization necessary.
- (1) M. Tinkham, Group Theory and Quantum Mechanics, Dover.
- (2) D. Gatteschi and L. Pardi, Gazz. Chim. Ital. **123**, 231 (1993).
- (3) O. Waldmann, Phys. Rev. B 61, 6138 (2000).
- (4) R. Schnalle and J. Schnack, Int. Rev. Phys. Chem. 29, 403-452 (2010).

#### **Point Group Symmetry II**

$$\mathcal{G}(R) \mid \alpha \, S \, M \,\rangle_a = \mid \alpha \, S \, M \,\rangle_b = \sum_{\alpha'} \mid \alpha' \, S \, M \,\rangle_a \, _a \langle \, \alpha' \, S \, M \mid \alpha \, S \, M \,\rangle_b$$

- Serious problem: application of *G*(*R*), i.e. permutation of spins, leads to different coupling schemes: *a* ⇒ *b*;
- Solution: implementation of graph-theoretical results to evaluate recoupling coefficients  $_a\langle \alpha' S M | \alpha S M \rangle_b$ .

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Results

# Results

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Cuboctahedron, s = 3/2, Hilbert space dimension 16,777,216; symmetry  $O_h$  (1). Evaluation of recoupling coefficients very time consuming (1,2).

(1) J. Schnack and R. Schnalle, Polyhedron 28, 1620 (2009).
(2) R. Schnalle and J. Schnack, Phys. Rev. B 79, 104419 (2009).



Icosahedron, s = 3/2, Hilbert space dimension 16,777,216; symmetry  $I_h$ ; Evaluation of recoupling coefficients for s = 3/2 in  $I_h$  practically impossible (1).

(1) R. Schnalle and J. Schnack, Int. Rev. Phys. Chem. 29, 403-452 (2010).

Results III



Spin ring, N = 10, s = 5/2, Hilbert space dimension 60,466,176; symmetry  $D_2$ ; Symmetry  $C_{10}$  would lead to more complicated recoupling coefficients & complex representation (1).

(1) R. Schnalle and J. Schnack, Int. Rev. Phys. Chem. **29**, 403-452 (2010).

(2) C. Delfs et al., Inorg. Chem. **32**, 3099 (1993).

# This could be the end . . .

# ... but you have got another molecule!

#### N=30, s=1/2 DIMENSION = 1,073,741,824



**Congratulations!** 

#### The idea of finite-temperature Lanczos

$$Z(T,B) = \sum_{\nu} \langle \nu | \exp\left\{-\beta H\right\} | \nu \rangle$$
$$\langle \nu | \exp\left\{-\beta H\right\} | \nu \rangle \approx \sum_{n} \langle \nu | n(\nu) \rangle \exp\left\{-\beta \epsilon_{n}\right\} \langle n(\nu) | \nu \rangle$$
$$Z(T,B) \approx \frac{\dim(\mathcal{H})}{R} \sum_{\nu=1}^{R} \sum_{n=1}^{N_{L}} \exp\left\{-\beta \epsilon_{n}\right\} |\langle n(\nu) | \nu \rangle|^{2}$$

- $|n(\nu)\rangle$  n-th Lanczos eigenvector starting from  $|\nu\rangle$ ;
- Partition function replaced by a small sum:  $R = 1 \dots 10, N_L \approx 100$ .
- J. Jaklic and P. Prelovsek, Phys. Rev. B 49, 5065 (1994).

#### How good is finite-temperature Lanczos?



• Works very well, see e.g. cuboctahedron and icosahedron.

• N = 12, s = 3/2: Considered < 100,000 states instead of 16,777,216.

Exact results: R. Schnalle and J. Schnack, Int. Rev. Phys. Chem. **29**, 403-452 (2010). FTLM: J. Schnack and O. Wendland, Eur. Phys. J. B **78**, 535-541 (2010).



#### Exp. data: A. M. Todea, A. Merca, H. Bögge, T. Glaser, L. Engelhardt, R. Prozorov, M. Luban, A. Müller, Chem. Commun., 3351 (2009).



#### Summary

- One can indeed exploit SU(2) and point group symmetries together. Good for molecules, since they are of finite size.
- Problem: Recoupling coefficients.
- Finite-temperature Lanczos is a good approximate method for Hilbert space dimensions smaller than  $10^{10}$ .
- I believe that this is the future.

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# Thank you very much for your attention.

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