Bose-Einstein Condensation of Triplons in Ba$_3$Cr$_2$O$_8$

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By performing heat capacity, magnetocaloric effect, torque magnetometry and force magnetometry measurements up to 33 T, we have mapped out the T-H phase diagram of the S = 1/2 spin dimer compound Ba$_3$Cr$_2$O$_8$. We found evidence for field-induced magnetic order between $H_{c1} = 12.52(2)$ T and $H_{c2} = 23.65(5)$ T, with the maximum transition temperature $T_c \sim 2.7$ K at $H \sim 18$ T. The lower transition can likely be described by Bose-Einstein condensation of triplons theory, and this is consistent with the absence of any magnetization plateaus in our magnetic torque and force measurements. In contrast, the nature of the upper phase transition appears to be quite different as our measurements suggest that this transition is actually first order.

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Quantum phase transitions (QPT) can be achieved by varying a non-thermal control parameter, such as pressure or applied magnetic field, while at a temperature of absolute zero. These transitions are driven by quantum fluctuations resulting from the uncertainty principle, as opposed to the thermal fluctuations that drive classical phase transitions. A particular type of QPT is realized in a Heisenberg spin dimer system, which possesses a non-magnetic spin-singlet ground state with a gap to the first triplet excited state. The excited triplets (triplons) can be considered as bosons with a hard core on-site repulsion. The repulsion condition is necessary in order to prevent more than one triplon from lying on a single dimer.

If one applies a magnetic field $H$ to close the spin gap, a critical field $H_{c1}$ is eventually reached which results in the generation of a macroscopic number of triplons. Above $H_{c1}$, the magnetic field can be varied to control the triplon density, and so it acts as a chemical potential. The system now consists of a series of interacting triplons with a ground state that critically depends on the balance between the kinetic energy and the repulsive interactions. Note that the kinetic energy of the interacting triplons arises from the $xy$-component of the Heisenberg interdimer interaction, while the nearest neighbour repulsive interaction (different from the on-site repulsion) arises from the Ising or $z$-component. The delicate balance between these two energies has led to interesting and diverse properties of QPTs in spin dimer systems.

If the repulsive interactions dominate, it is most crucial to minimize this contribution to the microscopic Hamiltonian. The easiest way to do this is to ensure that the triplon density per dimer is a simple rational fraction, as this allows the triplons to form a superlattice. These preferred fractional triplon densities result in plateaus in the magnetization as a function of field, and such behaviour has been observed in SrCu$_2$(BO$_3$)$_2$ (Ref. [5,6]).

When the kinetic energy terms dominate instead, this contribution will be minimized by allowing the triplons to have freedom to hop from dimer to dimer. The ground state then consists of a coherent superposition of singlets and triplets. No magnetization plateaus are observed in this case, but rather there is a continuous rise in the magnetization from $H_{c1}$ until saturation at $H_{c2}$. In many cases, the phase boundary at $H_{c1}$ satisfies a power law of the form: $T_c \propto (H - H_{c1})^{2/\delta}$ (d: dimensionality), which corresponds to a Bose-Einstein condensation (BEC) of triplons universality class. This type of phase transition generates a staggered magnetization transverse to the external field, creating a canted antiferromagnetic state in the intermediate regime between $H_{c1}$ and $H_{c2}$. This behaviour has been observed in BaCuSi$_2$O$_6$ (Ref. [7]), TiCuCl$_3$(Ref. [8]), and NiCl$_2$-4SC(NH$_2$)$_2$(Ref. [9]). An important property of these systems is that they must possess U(1) rotational symmetry as required by BEC theory, or at the very least the anisotropic spin terms must be small enough that they do not alter the universality class of the phase transition.

Recently, a new class of spin dimer compounds have been discovered with the general formula $A_3M_2O_8$ (Refs. [11-13]) where $A =$ Ba or Sr and $M =$ Cr or Mn. At room temperature, these compounds crystallize in the R-3m space group, and the crystal structure consists of MO$_4$ tetrahedra and isolated A$^{2+}$ ions. The magnetic M$^{3+}$ ions may carry spins of either $S = 1/2$ or 1, and these are arranged in double-stacked triangular lattices with three-fold periodicity and so form dimers along the c-axis. These systems are all described well by interacting dimer models, and so they provide a new opportunity to study field-induced quantum phase transitions.

In this work, we focus on the particular $S = 1/2$ system Ba$_3$Cr$_2$O$_8$. We have completed magnetic torque, magnetocaloric effect (MCE), and specific heat measurements at the National High Magnetic Field Laboratory (NHMFL)
on single crystals 1-5 mg in size. We have also performed magnetic force measurements at NHMFL, using a very small $\sim 1.3 \, \mu\text{g}$ sample. These measurements allowed us to map out the phase diagram for $\text{Ba}_3\text{Cr}_2\text{O}_8$ and to investigate the associated quantum phase transitions. We observed only two phase transitions as a function of field, and so we found no evidence for magnetization plateaus between $H_{c1}$ and $H_{c2}$ in our torque and force measurements. We also found that the lower phase transition can likely be described by BEC triplon theory, while the upper phase transition appears to be first order.

Single crystals of $\text{Ba}_3\text{Cr}_2\text{O}_8$ were grown by the traveling solvent floating zone method, and a detailed room temperature structure determination was completed as described in Ref. [15]. Fig. 1 depicts DC susceptibility measurements of our crystals for two different orientations. The large drop with decreasing temperature is characteristic of systems with non-magnetic spin-singlet excitations. The presence of only two critical fields and the lack of magnetization plateaus in our data suggests that the triplons are highly delocalized in $\text{Ba}_3\text{Cr}_2\text{O}_8$ and the kinetic energy terms dominate in the relevant microscopic Hamiltonian. These observations are consistent with the possibility that the lower transition can be described by BEC triplon theory.

An additional feature of the data that is particularly interesting is the magnetic hysteresis observed in association with the upper transition at $H_{c2}$. This suggests that while the lower transition is likely of a second-order nature, the upper transition is more first-order-like and lattice coupling may play a crucial role there.

Magnetic force measurements were also performed in a resistive magnet at NHMFL using a Faraday balance micromechanical magnetometer. One advantage of this method over magnetic torque is that one can work with very small samples ($\sim 1 \, \mu\text{g}$). A plot of the resulting magnetization vs. field at 600 mK is depicted in Fig. 2(b), and it is apparent that the main qualitative features are in agreement with the torque measurement, including the magnetic hysteresis observed near the upper transition. Note that the critical fields were determined in an analogous way to the torque measurements - by locating extrema in the second derivative of the magnetization.

Further details of the phase transitions were uncovered by performing MCE and heat capacity measurements in a resistive magnet at NHMFL using a home-built calorimeter. All measurements were performed with $H \parallel \hat{c}$. Some representative MCE scans are shown in Fig. 3(a) and (b), for cases of sweeping the field both up (dotted lines) and down (solid lines) at $2 \, \text{T/min}$. Since the MCE is a quasi-adiabatic process, an abrupt change in the sample temperature is observed upon crossing an order-
disorder transition to ensure entropy conservation\textsuperscript{21}. If the phase transition is second order, this process should be reversible - the temperature will increase (decrease) by the same amount upon entering (leaving) the ordered state. However, if the phase transition is first order this will introduce a dissipative component to the temperature change that is always positive, and so the MCE will become irreversible. In the present case, we find the MCE traces are essentially reversible at the lower transition, but are highly irreversible at the upper transition, especially for lower temperatures. This provides further evidence that the lower transition is of a second-order nature, while the upper transition is first order. Note that the transition points were found by locating extrema in the first derivative of $T(H)$ in a similar way to what has been done previously for other systems\textsuperscript{8,18}.

Heat capacity measurements are shown for selected applied fields in Fig. 3(c) and (d). Both the standard thermal relaxation method (for 13, 22, and 23 T) and the dual slope method\textsuperscript{22} (for 15, 18, and 20 T) were used to estimate the heat capacity. In all cases, the zero field background contribution was subtracted. A large lambda anomaly is observed in the intermediate field cases, but as the field is decreased closer to $H_{c1}$ or increased closer to $H_{c2}$ this becomes much less prominent and the magnitude of the heat capacity drops off sharply. Finally, although the anomaly remains distinctly lambda-like even down to 13 T, at 23 T the anomaly starts to look more symmetric. This suggests that the phase transition becomes more first-order-like in this region, and is consistent with our other measurements.

Fig. 4 combines our results in a phase diagram. We attribute the small discrepancies in the phase boundaries amongst the various techniques to slight differences in sample orientation. The maximum transition temperature to the magnetically-ordered state was found to correspond to $\sim 2.7$ K at H$\sim 18$ T. The phase diagram is also very nearly symmetric, as is expected for a system with a much larger intradimer than interdimer interaction (i.e. the present case), and is due to particle-hole symmetry that comes about from the effective Hamiltonian describing these systems\textsuperscript{23}. The small asymmetry in the phase diagram is likely due to lattice coupling associated with the upper phase transition and possible contributions from the S$_2 = 0$ and -1 triplet states. The latter are neglected in the aforementioned Hamiltonian.

Dzyaloshinskii-Moriya (DM) interactions are often the source of spin anisotropy that leads to U(1) symmetry breaking and hence non-BEC behaviour in spin dimer systems. While recent elastic neutron scattering measurements\textsuperscript{18} have found evidence for a structural phase transition in Ba$_3$Cr$_2$O$_8$ at $\sim 70$ K from the highly symmetric space group R-3m to the monoclinic space group C2/c, it turns out that the dimer centers are inversion centers in both the low and high temperature structures of this material. This suggests that intradimer DM interactions should be negligible in Ba$_3$Cr$_2$O$_8$. However, ESR measurements\textsuperscript{17} were performed very recently to study the issue of spin anisotropy in Ba$_3$Cr$_2$O$_8$ further, and evidence was found for a weak DM interaction of less than 0.1 meV. The detection of this weak DM interaction suggests either that it is an interdimer effect or that the low temperature crystal structure of Ba$_3$Cr$_2$O$_8$ has an even lower symmetry than that of the space group C2/c. In any event, as a result of this finding the most important issue to address is whether the resulting spin anisotropy is negligible in the sense that the lower transition can still be described by BEC theory.

The best way to verify that the spin anisotropy is negligible is to ensure that the lower phase boundary obeys a
power law of the form: $T_c \propto (H - H_c)\nu$ with $\nu = 2/d$. With this in mind, we calculated the critical exponent pertaining to our lower transition to determine whether or not our data satisfied this criterion. More specifically, we used a windowing analysis technique first introduced in Ref. [18]. Our narrowest fitting window contained data points in the range 333 mK $\leq T \leq 891$ mK and yielded a critical exponent of 0.49(2). This is substantially different from any exponent pertaining to a BEC universality class, and actually agrees with the expected exponent of 0.5 corresponding to the Ising universality class for easy-axis magnetic systems. However, determining accurate critical exponents reliably is often quite tricky, as one needs to ensure that the experimental data lies in the universal regime. To this end, recent elastic neutron scattering and heat capacity measurements have determined the lower phase boundary down to 30 mK and a

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