Magnetocaloric effect and magnetic cooling near a field-induced quantum-critical point

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Abstract

The presence of a quantum critical point (QCP) can significantly affect the thermodynamic properties of a material at finite temperatures $T$. This is reflected, e.g., in the entropy landscape $S(T, r)$ in the vicinity of a QCP, yielding particularly strong variations for varying the tuning parameter $r$ such as pressure or magnetic field $B$. Here we report on the determination of the critical enhancement of $\partial S/\partial B$ near a $B$-induced QCP via absolute measurements of the magnetocaloric effect (MCE), $(\partial T/\partial B)_S$, and demonstrate that the accumulation of entropy around the QCP can be used for efficient low-temperature magnetic cooling. Our proof of principle is based on measurements and theoretical calculations of the MCE and the cooling performance for a Cu$^{2+}$-containing coordination polymer, which is a very good realization of a spin-$1/2$ antiferromagnetic Heisenberg chain – one of the simplest quantum-critical systems.
Introduction

The magnetocaloric effect (MCE), i.e. a temperature change in response to an adiabatic change of the magnetic field, has been widely used for refrigeration. Although, up until now applications have focussed on cryogenic temperatures (1-3), possible extensions to room temperature have been discussed (4). The MCE is an intrinsic property of all magnetic materials in which the entropy $S$ changes with magnetic field $B$. Paramagnetic salts have been the materials of choice for low-temperature refrigeration (1), including space applications (5-7), with an area of operation ranging from about one or two degrees Kelvin down to some hundredths or even thousandths degree Kelvin. Owing to their large $\Delta S/\Delta B$ values, the ease of operation, and the applicability under microgravity conditions, paramagnets have matured to a valuable alternative to $^3$He-$^4$He dilution refrigerators, the standard cooling technology for reaching sub-Kelvin temperatures.

A large MCE also characterizes a distinctly different class of materials, where the low-temperature properties are governed by pronounced quantum many-body effects. These materials exhibit a $B$-induced quantum-critical point (QCP) – a zero-temperature phase transition –, and the MCE has been used to study their quantum criticality (8, 9, 10, 11, 12, 13, 14) or to determine their $B$-$T$ phase diagrams (15, 16, 17, 18, 19). The aim of the present work is to provide an accurate determination of the enhanced MCE upon approaching a $B$-induced QCP both as a function of $B$ and $T$ and to explore the potential of this effect for magnetic cooling.
Materials in the vicinity of a QCP have been of particular current interest, as their properties reflect critical behavior arising from quantum fluctuations instead of thermal fluctuations which govern classical critical points (20). Prominent examples of findings made here include the intriguing low-temperature behaviors encountered in some heavy-fermion metals, itinerant transition metal magnets (21 and references cited therein, 22) or magnetic insulators (23, 24) and the occurrence of new quantum phases near a QCP (25, 26, 10). Generally, a QCP is reached upon tuning an external parameter \( r \) such as pressure or magnetic field to a critical value. Although the critical point is inaccessible by experiment, its presence can affect the material’s properties at finite temperature significantly. The thermodynamic properties are expected to show anomalous power-laws as a function of temperature and, even more spectacular, to exhibit an extraordinarily high sensitiveness on these tuning parameters (27, 28). Upon approaching a pressure \((p)\)-induced QCP, for example, the thermal expansion coefficient \( \alpha \propto \partial S/\partial p \) is more singular than the specific heat \( C = T \cdot \partial S/\partial T \), giving rise to a diverging Grüneisen ratio \( \Gamma_p \propto \alpha/C \) (27, 29). Likewise, for a \( B \)-induced QCP, a diverging \( \Gamma_B = -C^{-1} \cdot (\partial S/\partial B)_T \) is expected (27, 28, 30) and has recently been reported (14). Unlike \( \Gamma_p \), however, where experimental access, apart from measurements of the critical contributions of the two quantities \( \alpha \) and \( C \), is very difficult, \( \Gamma_B \) can be determined directly via the magnetocaloric effect (MCE) by probing temperature changes of the material in response to changes of the magnetic field \( B \) under adiabatic conditions,

\[
\Gamma_B = \frac{1}{T} \left( \frac{\partial T}{\partial B} \right)_S . \tag{1}
\]

In this work, we directly measure the critical enhancement of \( \Gamma_B \) near a \( B \)-induced QCP and demonstrate that this effect can be used for magnetic refrigeration over an extended range.
of temperatures. Our results indicate that quantum critical materials open up new possibilities for realizing very efficient and flexible low-temperature coolants.

**The model substance**

For the proof of principle, it is helpful to focus on simple model substances, characterized by a small number of material parameters which are well under control. One of the simplest quantum-critical systems, where an enhanced MCE (30, 28) and a potential use for magnetic cooling (31) have been predicted, is the uniform spin-$\frac{1}{2}$ antiferromagnetic Heisenberg chain (AFHC), described by

$$H = J \sum_i \vec{S}_i \cdot \vec{S}_{i+1}. \quad (2)$$

This model, and its highly non-trivial physics, contains a single parameter $J$ – the Heisenberg exchange interaction – by which nearest-neighbor spins interact along one crystallographic direction. In fact, by exploring the quasi-1D spin-$\frac{1}{2}$ Heisenberg antiferromagnet KCuF$_3$, it has been demonstrated that quantum-critical Luttinger liquid (LL) behavior governs the material’s properties over wide ranges in temperature and energy (24). The AFHC remains in the LL quantum-critical state also in magnetic fields (32) up to the saturation field $B_s$, given by $g\mu_B B_s = 2J$, with $g$ the spectroscopic $g$ factor and $\mu_B$ the Bohr magneton. Since the fully polarized state above $B_s$ is a new eigenstate of the system, different from that of the LL, $B_s$ marks the endpoint of a quantum-critical line in the $B$-$T$ plane.

For exploring the MCE around $B_s$, we use a copper-containing coordination polymer $[\text{Cu}(\mu-C_2\text{O}_4)(4\text{-aminopyridine})_2\text{(H}_2\text{O})]_n$ (abbreviated to CuP henceforth), a good realization of a spin-$\frac{1}{2}$ AFHC, and compare the results with model calculations for the idealized system. In
CuP, first synthesized by Castillo et al. (33), Cu²⁺ (spin-½) ions, bridged by oxalate molecules (C₂O₄), form chains along the crystallographic c-axis (33, 34). High-quality single crystals of CuP with a small concentration n = 0.4 – 1 % of uncoupled spin-½ impurities, made recently available by a slow-diffusion technique (34), were used for the present investigations (see SI Text I). The magnetic susceptibility of these single crystals for \( T \geq 0.055 \) K, are well described by the model of a spin-½ AFHC (35) with an intra-chain coupling \( J/k_B = (3.2 \pm 0.1) \) K (34).

In order to fully characterize the magneto-thermal properties of CuP, measurements of the specific heat \( C_B(T) \) and low-temperature magnetic susceptibility \( \chi_T(B) \) have been carried out (Fig. 1). The excellent agreement of these results with model calculations (solid lines in Fig. 1) (see SI Text II) confirms the assertion (33, 34) that CuP is a very good realization of a uniform spin-½ AFHC. Compared to other model substances of this kind, such as copper pyrazin dinitrate (36), however, CuP excels by its moderate size of \( J \) and the correspondingly small saturation field of 4.09 T (for \( B \parallel b \)), enabling the MCE to be studied in the relevant field range below about twice (§) the saturation field by using standard laboratory magnets.

**Absolute measurements of the magnetocaloric effect and comparison with theory**

For a quantitative determination of \( \Gamma_B \), a novel step-like measuring technique was employed (see SI Text III and IV) ensuring, to good approximation, adiabatic conditions. The \( \Gamma_B \) data obtained this way for two different temperatures exhibit negative values for \( B < B_s \) (Fig. 2A), implying that here cooling is achieved through magnetization! Upon increasing the

\[ \section{Footnote} \]

§ At high magnetic fields, i.e. \( B > 2B_s \), the spin-½ AFHC behaves essentially like a paramagnet, where \( T \) varies linearly with \( B \) at \( S = \) const. For fields lower than about twice \( B_s \) deviations from linearity become visible (30), indicating the influence of quantum critical fluctuations.
field, \( \Gamma_B \) passes through a weak minimum, changes sign and adopts a pronounced maximum at fields somewhat above 4 T. The large positive \( \Gamma_B \) values here indicate that in this \( B-T \) range, a pronounced cooling effect is obtained through demagnetization. The sign change in \( \Gamma_B \propto \partial S / \partial B \) within a narrow \( B \)-interval centered near \( B_s \) implies the presence of a distinct maximum in \( S_T(B) \) – a clear signature of a nearby \( B \)-induced QCP (28). It reflects the accumulation of entropy due to the competing ground states separated by the QCP. The data taken at 0.32 K show extrema in \( \Gamma_B \) which are distinctly sharper and more pronounced as compared to the signatures in the 0.97 K data, suggesting incipient divergencies of \( \Gamma_B \) on both sides of \( B_s \).

Theoretical calculations of \( \Gamma_B \) have been performed for the ideal spin-\( \frac{1}{2} \) AFHC with \( J/k_B = 3.2 \) K (solid lines in Fig. 2A). The model curves capture the essential features of the experimental results and even provide a good quantitative description for the data taken at 0.97 K. However, some systematic deviations become evident which increase with decreasing temperature: the extrema in \( |\Gamma_B| \) as a function of field at fixed temperature (Fig. 2A) are less strongly pronounced compared to theoretical results and the experimental data at \( T = 0.32 \) K exceed the theory curves at high fields (\( B \geq 5 \) T). To follow the evolution of the extrema as a function of temperature, measurements at \( B = \) const. have been performed (insert of Fig. 2B). The deviations from the model curves suggest that by the application of a magnetic field, the divergence in \( \Gamma_B(T \rightarrow 0) \) becomes truncated by the occurrence of a small energy scale, accompanied by a shift of entropy to higher fields. A plausible explanation for these observations would be the opening of a small field-induced gap. It may result, e.g., from a finite Dzyaloshinskii-Moriya (DM) interaction (38, 39) permitted by the lack of a center of inversion symmetry in CuP (33). In fact, indications for a finite DM interaction can be inferred
from measurements performed under different field orientations (**). According to ref. 38 the value of the DM-vector $\mathbf{D}$ can be estimated by $|\mathbf{D}| \sim (\Delta g/g)J$, where $g$ is the material’s g-factor and $\Delta g$ its deviation from the free electron value. For CuP the corresponding energy scale is estimated to $|\mathbf{D}| \sim 0.07 J \sim 0.22 \text{ K} \cdot k_B$, which exceeds the one resulting from dipole-dipole interactions by more than one order of magnitude (††). In addition, part of the deviations from the model curves at low temperatures can also be due to the presence of weak inter-chain interactions and the accompanied dimensional crossover, inevitable in any three-dimensional material. It may also in part reflect the influence of an enhanced thermal boundary resistance between sample and thermometer due to an enhanced spin-phonon interaction caused by a nearby phase transition (40).

**Magnetic cooling and performance characteristics**

Despite these deviations from the idealized system, the enhanced MCE (Fig. 2A) highlights distinct quantum-critical behavior in CuP over extended ranges of temperature and magnetic field. The strongly enhanced values of $|\Gamma_B|$, implying large temperature changes in

** Besides lacking inversion symmetry, CuP has a two-fold rotation axis parallel to the $b$-axis which constrains the DM-vector $\mathbf{D}$ to lie within the $ac$ plane. A finite DM interaction in CuP is consistent with the results of magnetic cooling experiments for two different orientations of the magnetic field with respect to $\mathbf{D}$: the accessible lowest temperature $T_f$ raises by about $(10 \pm 2)\%$ for $B \parallel c$ as compared to $B \parallel b$.

†† Using the distance between adjacent Cu ions, ranging from 6.357 Å to 8.908 Å, the dipole-dipole interaction energy is estimated to $J_{d-d}/k_B \leq 0.012 \text{ K}$.
response to small field variations, suggest that the system could be suitable for low-temperature magnetic refrigeration. To investigate its potential as a coolant, demagnetization experiments were carried out on CuP under near adiabatic conditions (see SI Text III) while simultaneously recording the sample temperature $T_s$ (Fig. 2B). While the cooling process approaches an in-$B$ linear behavior at higher temperature near $(T_i, B_i)$ – such is seen in simple paramagnets where $T_s \propto B$ –, it becomes superlinear upon decreasing the temperature. This enhanced cooling effect, which is in accordance with the model calculations for the ideal system (broken lines in Fig. 2B), is a direct manifestation of quantum criticality. Upon further cooling, $T_s(B)$ passes through a rounded minimum, assigned $T_0$, for $B$ close to $B_s$. The quantitative deviations from the theory curves, in particular an experimentally revealed $T_0$ staying clearly above the theoretically expected value, can be attributed only in part to non-ideal adiabatic conditions in the cooling experiments. Estimates of the effect of the parasitic heat flow onto the sample (see SI Text V) show that for an improved thermal isolation, $T_0$ can be reduced from 179 mK to about 132 mK (solid line in Fig. 2B). This implies that the main source for the deviations from the theoretical expectations lies in the presence of the above-mentioned perturbing interactions in CuP. The good agreement with the theory curves at higher temperatures, where these interactions are irrelevant, however, implies that for a better realization of the spin-$\frac{1}{2}$ AFHC, cooling to much lower temperatures should be possible.

In order to assess the principle limits of cooling near a QCP and its potential for applications, we compare some performance characteristics for the ideal spin-$\frac{1}{2}$ AFHC with those of two state-of-the-art adiabatic demagnetization refrigerator (ADR) materials (Fig. 3). These are CrK(SO$_4$)$_2$$\cdot$12H$_2$O (chrome potassium alum CPA) and FeNH$_4$(SO$_4$)$_2$$\cdot$12H$_2$O (ferric ammonium alum FAA), which cover the typical temperature range of applications for
paramagnetic salts and, due to their high efficiency, are also used in space applications (5-7). Representative parameters characterizing the performance of an ADR material include (i) the operating range, in particular its lower bound $T_{\text{min}}$. Standard ADR systems operate predominantly in a narrow temperature window near $T_{\text{min}} \approx 30-60$ mK, while extensions down to only a few millikelvin are possible. Such low temperatures are otherwise accessible only by using elaborated $^3$He-$^4$He dilution refrigerators, which, however, are much bigger compared to ADRs and cannot operate in a microgravity environment. Another important parameter is (ii) the “hold time” of the coolant, which is inversely proportional to its cooling power $\dot{Q}$. This quantity measures the ability of the refrigerant to absorb heat without warming up too rapidly. For some applications, (iii) the efficiency can be of importance. This includes the cooling capability per unit mass of refrigerant material and the ratio $\Delta Q_c/\Delta Q_m$. Here $\Delta Q_c$ is the heat the material can absorb after demagnetization to the final field $B_f$, and $\Delta Q_m$ the heat of magnetization released to a precooling stage held at a temperature $T_i$, the initial temperature of the magnetic cooling process. The efficiency can be an issue for modern multi-stage single-shot or continuous ADRs, such as the ones used in space (5), where the entire system has to be optimized with regard to precooling requirements and weight.

In principle, the cooling performance of a material is determined by the low-energy sector of its magnetic excitation spectrum, reflected in the low-temperature specific heat $C(T)$, and its variation with magnetic field. For paramagnetic salts, $C(T)$ at $B = 0$ is of Schottky type (see, e.g., ref. 40) due to an energy-level splitting arising from residual magnetic interactions. The resulting $C(T)$ maximum actually enables the material to be used as a coolant as it ensures a certain amount of cooling power during demagnetization at low temperatures. At the same time, the $1/T^2$-like decrement of the $C(T)$ anomaly at high temperatures and the accompanied
rapid reduction of the hold time (Fig. 3) constrain the operating range to a narrow temperature window around the position of the maximum. Thus, ADRs based on paramagnets are particularly well suited for applications within a narrow temperature range of operation.

This is different for the spin-$\frac{1}{2}$ AFHC near $B_c$: due to its peculiar excitation spectrum (35), i.e. the abundance of low-energy excitations above the QCP, (in principle) arbitrarily small values of $T_{\text{min}}$ can be reached, while keeping the cooling power large. At the same time, due to the extraordinarily large specific heat above the QCP, with a peak centered at a temperature around $J/2k_B$ (Fig. 1), the system affords extended hold times also for temperatures largely exceeding the maximum position (Fig. 3). Thus, materials near a $B$-induced QCP can be an excellent alternative to paramagnets for those applications where the temperature has to be varied over an extended range. This includes the possibility to reach very low temperatures, albeit with reduced hold times. In addition, the present quantum critical system, despite its low spin value, can absorb an amount of energy $\Delta Q_c$ which is almost comparable to that of the spin-3/2 (CPA) and spin-5/2 (FAA) paramagnets (Fig. 4). The quantum critical system excels, however, by its high efficiency $\Delta Q_c/\Delta Q_m$, which exceeds the corresponding numbers for the paramagnets by a factor 2-3 (Fig. 4). This is due to the system’s ability to absorb energy even at temperatures as high as $T_i$, in contrast to the paramagnets, where the absorption essentially occurs at low temperatures $T < T_i$ (Fig. 4). Besides these performance characteristics, the applicability of a material as a coolant may also depend on other material-specific features such as its thermal conductivity and the possibility to realize a good thermal contact to the body to be cooled. In terms of heat transport, the spin-$\frac{1}{2}$ AFHC near the $B$-induced QCP is particularly favorable as it shows a comparatively large spin thermal conductivity along the spin chains, which even dominates the material’s thermal transport at low temperatures, see,
e.g. ref. 42. In contrast, a small thermal conductivity and a weak thermal contact between the coolant and another body are principal concerns for paramagnetic ADR materials (40). Here, problems can arise due to the extreme hydration of these materials, required to reduce the mutual interactions between the magnetic centers, and the materials’ sensitivity to decompose if water is allowed to evaporate through imperfections in the materials’ housing.

**Conclusions and outlook**

We have demonstrated that the strong enhancement of the magnetocaloric effect, arising from quantum fluctuations near a $B$-induced quantum-critical point, can be used for realizing an efficient and flexible magnetic cooling with a good cooling performance over an extended range of temperatures. While the spin-$\frac{1}{2}$ AFHC, thanks to its simplicity, has enabled the provision of a proof-of-principle demonstration, extensions of this concept are obvious and may guide the search for materials with further improved cooling performance. On the one hand, this includes low-dimensional spin systems with geometric frustration, such as the saw-tooth chain discussed in ref. 30. In contrast to the uniform spin-$\frac{1}{2}$ AFHC discussed here, where the enhanced MCE results from large relative changes of the entropy across $B_s$, while $S$ vanishes for $T \to 0$, certain geometrically frustrated spin configurations in one dimension support a significant zero-temperature entropy (43). As a consequence, in these systems there is a large absolute variation of entropy across $B_s$ and hence a large $\partial T/\partial B$ down to, in principle, arbitrarily low temperatures, limited by weak residual interactions, such as dipole-dipole interactions. Consequently, a further enhancement of the MCE can be expected for those frustrated magnetic systems. The combination of quantum criticality with (i) low dimensionality, assuring a high density of low-energy excitations, and (ii) geometric
frustration, opens up a promising route to search for materials with further enhanced MCE. Of particular interest are highly symmetric (yet low-dimensional) spin systems with gapped ground states in zero field and $B$-induced XY antiferromagnetic states, since the high symmetry in spin space prevents the system from developing $B$-induced gaps. Likewise, as a large spin entropy can cause a large thermopower in conducting materials (44), conducting magnets driven close to a field-induced QCP would be a very interesting class of materials for the search for thermoelectric (Peltier) cooling systems with high efficiency. Finally, the concept of using quantum criticality for magnetic cooling might also be applicable to cold atom experiments and help there to realize low-temperature magnetic and superfluid phases.

Materials and methods

The specific heat was measured as a function of temperature by employing a compensated heat-pulse technique in combination with a $^3$He-$^4$He dilution refrigerator (see SI Text I). For the magnetic susceptibility measurements a state-of-the-art compensated-coil ac-susceptometer adapted to a top-loading $^3$He-$^4$He dilution refrigerator was used (see SI Text I). The MCE was determined by employing a specially-designed calorimeter (see SI Text III). For
the magnetic cooling experiments, the setup was modified to ensure quasi-adiabatic conditions (see SI Text IV).

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Figure Legends

Fig. 1. (A) Specific heat $C_B(T)$ measured in a constant magnetic field of $B = 0.15$ T (red symbols) and $7.14$ T (green symbols). While the data at $B = 0.15$ T $< B_s$ (saturation field) are characterized by an $T$-linear contribution $C/T \propto J^{-1}$ for $T < J/k_B$ (35), those at $B = 7.14$ T $> B_s$ show an activated behavior, reflecting the opening of an excitation gap in the field-induced ferromagnetic state. The red solid line is the result of Bethe Ansatz calculations (BAC) (35) for a spin-$\frac{1}{2}$ AFHC with $J/k_B = 3.2$ K, $g = 2.087$, $B/B_s = 0.0329$, representing the magnetic contribution $C_{mag}$, in addition to a small phonon contribution $C_{ph} = \beta T^3$ (black solid line). The coefficient $\beta$, the only adjustable parameter, was determined from a least-squares fit of $C = C_{ph} + C_{mag}$ to the experimental data taken at $B = 0.15$ T. The green solid line corresponds to $C_{ph} + C_{mag}$, with $C_{mag}$ obtained from BAC for $B/B_s = 1.564$, corresponding to $B = 7.14$ T. The excellent agreement of the $C_B(T)$ data with the model calculations at small fields and the moderate deviations at $7.14$ T indicate the presence of additional field-dependent terms in the material’s Hamiltonian which become relevant at high fields. (B) Blue open circles represent the magnetic susceptibility $\chi = \partial M/\partial B$, taken on single-crystalline material immersed in the $^3$He-$^4$He mixture kept at a constant temperature $T = 0.055$ K with $B || b$-axis. The field was swept at a low rate of 0.1 T/min, ensuring constant-temperature conditions. The growth of $\chi$ upon increasing the field, the sharp peak at $4.09$ T and the disappearance of $\chi$ at higher fields reflect the transition from the antiferromagnetic ($B < B_s$) to the polarized ferromagnetic state ($B > B_s$) at $B_s = 4.09$ T. The solid line represents a parameter-free description of the data based on the above-mentioned BAC for $J/k_B = 3.2$ K and $g = 2.33$ (34).

Fig. 2. (A) Variation of $\Gamma_B$, approximated by $\Gamma_B = T^{-1} \cdot (\Delta T/\Delta B)_S \approx \text{const}$ (see SI Text IV), with field at $T = 0.32$ K (blue squares) and $0.97$ K (black triangles) for $B || b$-axis. For $\Gamma_B$ data at an intermediate temperature $T = 0.47$ K, see ref. 13. Solid lines are the results of Quantum-Monte-Carlo (QMC) simulations and exact diagonalization of a finite-size lattice of the spin-$\frac{1}{2}$ AFHC for the corresponding temperatures (same color code) by using $J/k_B = 3.2$ K. A $g$ value of 2.28 has been used in the calculations to account for a small misalignment of the crystals. Error bars (below symbol size for the data taken at 0.97 K) of experimental data correspond to statistical error. The systematic error, mainly originating from the internal time constant of the resistance bridge in combination with the sweep time applied, is estimated to be of the order of 1 %. (B) Sample temperature (symbols) measured by demagnetizing a collection of CuP single crystals ($B || b$-axis) of total mass of 25.23 mg under near adiabatic conditions (see SI Text III). The initial parameters were set to $B_i = 7$ T and $T_i = 1.40$ K (green symbols) and 1.21 K (red symbols). The field was swept with a rate $\Delta B/\Delta t = -0.3$ T/min for $B \geq 6$ T and $0.5$ T/min for $B < 6$ T. The red solid line represents the experimental data (for $T_i = 1.21$ K) corrected for the effect of a parasitic heat flow (see SI Text V). Broken lines are those ideal isentropes, derived from the exact result for the entropy of the spin-$\frac{1}{2}$ AFHC (37), which correspond to the same initial conditions ($B_i$, $T_i$). The insert in (B) shows the temperature dependence of $\Gamma_B$, approximated by $\Gamma_B = T^{-1} \cdot (\Delta T/\Delta B)_S \approx \text{const}$ (see SI Text IV), at constant fields below ($B = 3.5$ and
4.0 T) and above (4.3 T) \( B_s \). The deviations of the experimental data from the model calculations for the ideal system (solid lines, same color code as used for the experimental data) grow with decreasing the temperature. The data reveal an anomaly around 0.22 K, the position of which shows no significant field dependence within the field range investigated. This feature is likely to be related to the opening of a field-induced gap in the magnetic excitation spectrum.

**Fig. 3.** Increase of the sample temperature \( T_s \) with time after demagnetization to \( B = B_f \) calculated for a heat load of 5 \( \mu \)W applied under adiabatic conditions to 100 gram substance of a spin-1/2 AFHC with \( J/k_B = 3.2 \) K (red solid line), a spin-3/2 (CrK(SO_4)_2⋅12H_2O in short CPA) (magenta solid line) and a spin-5/2 (Fe(NH_4)(SO_4)_2⋅12H_2O in short FAA) (blue solid line) paramagnetic salt (see SI Text VI). For the paramagnets at \( B_f = 0 \), base temperatures of \( T_{min} \approx 0.01-0.015 \) K (CPA) and \( 0.04-0.05 \) K (FAA) can be reached, which lie close to the transition temperature to long-range antiferromagnetic order at \( T_N \approx 0.01 \) K (CPA) and \( 0.04 \) K (FAA). In contrast, for the spin-1/2 AFHC, cooling to an (in principle) arbitrarily small \( T_{min} \) is possible. In practice, however, a finite \( T_{min} \) will result from the presence of small perturbing interactions such as weak inter-chain couplings, dipole-dipole interactions or single-ion anisotropies. The calculations for \( T_s(time) \) are based on computed (spin-1/2 AFHC) and measured (CPA and FAA) (40) specific heat data. The paramagnets exhibit long hold times only in a narrow temperature range around \( T_{min} \), defined by the position of their Schottky-type \( C(T, B=0) \) maximum. For temperatures somewhat above \( T_{min} \), \( T_s \) rapidly increases with exposure time \( t \), approximately as \( t^2 \) for FAA. This contrasts with the spin-1/2 AFHC, where \( T_s \) increases only moderately with time as \( t^{2/3} \). A further flattening of \( T_s(t) \), at not too low temperatures, can be achieved for \( B_f < B_s \); see, e.g. the warming curve for \( B_f = 0.75 B_s \) (broken orange line). Crossing \( B_s \) upon demagnetization from \( (B_i, T_i) = (7 \) T, 1.3 K) is, however, accompanied by a moderate warming to \( T_{min} = 0.112 \) K.

**Fig. 4.** Calculated molar magnetic entropy \( S_{mag}(T, B=const.) \) as a function of temperature of the spin-1/2 AFHC with \( J/k_B = 3.2 \) K for an initial field of \( B_i = 7.14 \) T (red broken line) and a final field \( B_f = B_s = 4.09 \) T (red solid line). For comparison, \( S_{mag}(T, B=const.) \) of the spin-5/2 paramagnetic salt Fe(NH_4)(SO_4)_2⋅12H_2O is shown for \( B_i = 2 \) T (blue broken line) and \( B_f = 0 \) (blue solid line); the data were taken from refs. 40 and 41. In the cooling process, the materials are first isothermally magnetized (path AB for example), and then, after thermal isolation, adiabatically demagnetized (path BC) to the final temperature \( T_f \). For \( T_i = 1.3 \) K, for example, adiabatic demagnetization from 7.14 T to 4.09 T for the spin-1/2 AFHC results in \( T_f = 0.021 \) K, while demagnetization from 2 T to zero field for the spin-5/2 paramagnet leads to \( T_f = 0.026 \) K. The systems warm up along their entropy curves at the final demagnetization field \( S_{mag}(T, B=B_f) \). The heat of magnetization at \( T = T_i, \Delta Q_m = T_i \cdot [S_{mag}(B_f, T_i) - S_{mag}(B_i, T_i)] \), and the heat that the material is able to absorb after adiabatic
demagnetization, $\Delta Q_c = \int_{T_f}^{\infty} T \cdot (\partial S_{mag}/\partial T)_{B_f} \, dT$, can be read off the figure. For the spin-5/2 paramagnet (PM) Fe(NH₄)(SO₄)₂·12H₂O, for example, $\Delta Q_m = 14.3$ J/mole is given by the area of the rectangle ABDE, while $\Delta Q_c = 1.22$ J/mole corresponds to the hatched area in that rectangle. The efficiency factor $\Delta Q_c/\Delta Q_m$ in the temperature range indicated amounts to $26\%$ for the spin-1/2 AFHC as compared to only $9\%$ for the spin-5/2 paramagnet. For the spin-3/2 system CrK(SO₄)₂·12H₂O (see SI Text VII) one finds $\Delta Q_m = 10.01$ J/mole, $\Delta Q_c = 1.06$ J/mole and $\Delta Q_c/\Delta Q_m$ of $11\%$. 
Figure 3

Figure 4
Supporting Information for “Magnetocaloric effect and magnetic cooling near a field-induced quantum-critical point”

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I. Sample characterization. Single crystalline samples of CuP have been characterized by specific heat and magnetic susceptibility measurements. The specific heat was measured as a function of temperature for \(0.1 \, K \leq T \leq 4 \, K\) at varying magnetic fields by employing a compensated heat-pulse technique (1) in combination with a \(^3\)He-\(^4\)He dilution refrigerator. The specific-heat sample was composed of several single crystals with a total mass of 36.3 mg, the \(ac\)-planes of which were aligned parallel to the magnetic field. The magnetic susceptibility was measured on a single crystal of CuP down to \(T = 0.055 \, K\) and magnetic fields up to 8.5 T by using a state-of-the-art compensated-coil ac-susceptometer adapted to a top-loading \(^3\)He-\(^4\)He dilution refrigerator. The crystal was oriented with its \(b\)-axis running parallel to the applied magnetic field. Empty-coil measurements were performed to determine the magnetic background.

II. Model calculations. The spin-1/2 antiferromagnetic Heisenberg chain is exactly solvable, and, indeed, we used those exact results for calculating the specific heat (2), magnetic susceptibility (2), and entropy (3). The theory curves shown in Fig. 2(A) in the main text are based on exact diagonalization and Quantum-Monte-Carlo simulations which are essentially exact (see ref. (3) for details). Finally, in some cases we have used exact diagonalization for the spin-1/2 antiferromagnetic-Heisenberg-chain model with 20 sites, as outlined in ref. 4.

III. The calorimeter – design features. The MCE was measured by attaching several single crystals with a total mass of 7.23 mg to a specially-designed calorimeter (see below) by applying a small amount of Apiezon N-grease. For the magnetic cooling experiments, where field sweeps from \(B_i = 7 \, T\) to \(B_f = 3 \, T\) were carried out with a rate of 0.3 – 0.5 T/min, the setup was modified in the following way to ensure quasi-adiabatic conditions: single crystals of total mass of about 25 mg were used and the thermal coupling between the calorimeter and the thermal bath was reduced by a factor 3-9, depending on the temperature. The calorimeter for measuring the MCE consists of a sapphire plate (5×5×0.3 mm\(^3\)) and a grinded RuO\(_2\) resistor, which serves as a thermometer. The heat capacity of the calorimeter,
amounting to less than $10^{-4}$ of that of the composite CuP sample over the whole field ($B < 7.2$ T) and temperature ($T \leq 2$ K) range investigated, can be neglected in the measurements of the MCE. Four thin Pt-W wires (25 μm in diameter and ~ 1.5 cm in length) were used to make electrical contacts to the thermometer. The calorimeter was suspended by two thin nylon strings, each 0.1 mm in diameter. The crystals were oriented such that the external magnetic field was parallel to the $b$-axis. The thermal conductance $\kappa_{\text{cal}}$ between the calorimeter and the surrounding thermal bath was determined from the MCE experiment through $\kappa_{\text{cal}} = C/\tau$. Here $C$ is the specific heat of the sample, determined independently, and $\tau$ the relaxation time by which the sample temperature relaxes back to the bath temperature $T_b$ after a field change $\Delta B$. These measurements yield $\kappa_{\text{cal}} \approx 2.5 \times 10^{-8}$ W/K at 0.32 K and $1.1 \times 10^{-7}$ W/K at 0.9 K. The relaxation time $\tau$ varies from about 1500 s (at low temperatures and small fields) to about 90 s (for higher temperatures and large fields).

IV. Quasi-adiabatic (step-like) measurements of the magnetocaloric effect. To reduce systematic errors in the data analysis, which may result from a too fast relaxation, see below, the measurements of the MCE were restricted to fields $B \leq 6$ T and $T \leq 1$ K, where $\tau \geq 300$ s. In this range of parameters, the MCE was determined by employing a step-like technique which ensures quasi-adiabatic conditions (for a setup operating in the opposite limit, i.e. under highly non-adiabatic conditions, see ref. 5): at a fixed bath temperature $T_b$, and the calorimeter initially in thermal equilibrium with the bath, the magnetic field was swept in a small step $\Delta B$ while simultaneously measuring the changes in the sample temperature $T_s$. The latter was detected with an ac-resistance bridge with a resolution of $\Delta R/R \sim 10^{-5}$ and an internal time constant of $\sim 0.3 \sim 0.4$ s, depending on the temperature range. $T_b$ during the field sweeps was stabilized within $\pm 0.025$ mK. The temperature of the calorimeter, prior to a field sweep, varies typically by $\pm 0.04$ mK. A field change of $\Delta B = 20$ mT was applied to ensure measurable temperature changes $\Delta T_s$. The time for sweeping the field was kept at $t_{\text{swp}} \sim 65$ s, which is short enough to guarantee quasi-adiabatic conditions for relaxation times $300$ s $\leq \tau \leq 1500$ s, while keeping effects due to finite internal time constants and eddy-current heating in the metallic parts of the calorimeter small. The latter effect was eliminated by using two
consecutive sweeps, an up sweep by $\Delta B$, followed by a down sweep by $-\Delta B$, and estimating 
$\Delta T_s = \Delta T^{+}_{AB} - \Delta T^{-}_{AB}$ (Fig. S1A). After each field change, $T_s$ relaxes back to $T_b$. This process 
can be well described by a single exponential $T_s \propto \exp(-t/\tau)$, except for $t \ll \tau$, indicating that all 
internal time constants $\tau_i$, corresponding to the thermal couplings of the sample to the 
calorimeter and the calorimeter to the thermometer, are much smaller than $\tau$. Occasional weak 
thermal drifts of the bath or the calorimeter, with $\Delta T_{drift} \ll \Delta T_s$, were accounted for by 
subtracting a small in-$t$ linear background from $T_s(t)$. To extract the temperature changes 
$\Delta T^{+}_{AB}$ and $\Delta T^{-}_{AB}$ from the data, an “equal-areas” construction was employed, whereby the 
gradual changes in $T_s$ throughout the field changes were replaced by idealized sharp ones (Fig. 
S1B). This procedure also compensates, to a good approximation, for uncertainties resulting 
from the internal time constants $\tau_i$.

V. Corrections for non-adiabaticity in the demagnetization experiments. The magnetic 
cooling experiments on CuP were performed under quasi-adiabatic conditions using the same 
calorimeter as employed for measuring the MCE but with a reduced thermal coupling between 
calorimeter and thermal bath. Even under these improved conditions, the finite heat flow 
$dQ/dt$ on the calorimeter (due to the wiring, measuring current and a finite pressure of 
residual gas) gives rise to a parasitic temperature drift $\dot{T}_{para}$. Thus, at a given time $t_0$ (which 
corresponds to a magnetic field $B$), the experimentally determined sample temperature $T_{exp}(B, 
t_0)$ has to be corrected by

$$T_{corr}(B, t_0) = T_{exp}(B, t_0) - \int_0^{t_0} \dot{T}_{para} dt, \quad (S1)$$

This correction procedure is applicable only for a sufficiently weak parasitic heat flow, which 
does not strongly affect the whole demagnetization process, see, e.g. ref. 6. The parasitic 
temperature drift is given by

$$\dot{T}_{para} = \frac{\dot{Q}_{para}(B, t)}{C(B, t)}, \quad (S2)$$
where $\dot{Q}(B(t), T(t))$ is the parasitic heat flow and $C(B(t), T(t))$ the specific heat at a magnetic field and a temperature corresponding to the time $t$. The heat flow $\dot{Q}$ is given by $\dot{Q} = \kappa \Delta T$, where $\Delta T$ is the temperature gradient between the bath and the calorimeter and $\kappa$ denotes the thermal conductance between calorimeter and bath. The quantity $\kappa(T)$ was determined from relaxation experiments at various temperatures and fields. The error bar of the so-derived $\kappa(T)$-values amounts to approximately $\pm 15\%$ for temperatures $0.250 \, \text{K} \leq T \leq 2.01 \, \text{K}$. In this temperature range, $\kappa(T)$ varies, to a good approximation, linear in temperature. For the determination of $C(B, t)$ in eq. S2, specific heat data on CuP, measured as a function of temperature at various fixed fields up to $7.14 \, \text{T}$, were used to determine the specific heat landscape in the relevant field and temperature range. The quantity $C(B, t)$ was then obtained by parameterization, using a 4th-order polynomial, of the data along the $(B(t), T(t))$ trajectory the sample took during the demagnetization experiment. After having transformed the variables $T$ and $B$ into time by using appropriate inverse functions, the numerical integration in eq. (S1) was carried out. For the demagnetization experiment starting at $T_i = 1.25 \, \text{K} \, (\, = T_b)$ and $B_i = 7 \, \text{T}$, for example, a minimal temperature of $T_0 = 179 \, \text{mK}$ was obtained at a field of $B_f = 3.97 \, \text{T}$. According to the above procedure, this value has to be corrected to $T_0 = (132 \, \pm 9) \, \text{mK}$ for idealized adiabatic conditions.

VI. Calculations of the warming rate under an external heat load for CrK(SO$_4$)$_2$·12H$_2$O and Fe(NH$_4$)(SO$_4$)$_2$·12H$_2$O – two standard magnetic coolants. The time evolution of a material’s temperature under adiabatic conditions and exposed to a given heat load $P$ is obtained from the internal energy $U(T)$, derived by integrating $C(T)$ and the equation $\frac{dU(T(t))}{dt} = P$. Integration then leads to $U(T(t_i)) – U(T(t)) = (t_i – t) \cdot P$. The paramagnetic salts CrK(SO$_4$)$_2$·12H$_2$O (CPA) and Fe(NH$_4$)(SO$_4$)$_2$·12H$_2$O (FAA), containing Cr$^{3+}$ ions with spin $3/2$ and Fe$^{3+}$ ions with spin $5/2$, respectively, represent state-of-the-art coolants for low-temperature magnetic refrigeration, including space applications (see, e.g., refs. 7, 8, 9). The salts exhibit long-range antiferromagnetic order below $10 \, \text{mK}$ (CPA) and $40 \, \text{mK}$ (FAA),
which sets a lower bound to the accessible base temperature $T_{\text{min}}$. For calculating the materials’ warming rate at $B_f = 0$, the specific heat data $C(T)$ reported in ref. 10 were used.

VII. Adiabatic demagnetization of the paramagnetic salt CrK(SO$_4$)$_2$-12H$_2$O. In the cooling process, the materials are first isothermally magnetized (path AB), and then, after thermal isolation, adiabatically demagnetized (path BC) to the final temperature $T_f$. For $T_i = 1.3$ K and $B_i = 2$ T, for example, for the spin-3/2 paramagnet reaches $T_f = 0.010$ K. The system warms up along its entropy curve at the final demagnetization field $S_{\text{mag}}(T, B=B_f)$. The heat of magnetization at $T = T_i$, $\Delta Q_m = T_i \cdot [S_{\text{mag}}(B_f, T_i) - S_{\text{mag}}(B_i, T_i)]$, and the heat that the material is able to absorb after adiabatic demagnetization, $\Delta Q_c = \int_{T_i}^{T_f} T \cdot (\partial S_{\text{mag}} / \partial T)_{B_f} dT$, can be read off figure S2. For the spin-3/2 paramagnet Cr(NH$_4$)(SO$_4$)$_2$-12H$_2$O, we obtain $\Delta Q_m = 10.1$ J/mole, corresponding to the area of the rectangle ABDE, and $\Delta Q_c = 1.06$ J/mole, corresponding to the hatched area in that rectangle. This results in an efficiency factor $\Delta Q_c/\Delta Q_m$ in the temperature range indicated of about 11%.

References


Figure Legends

**Fig. S1.** (A) Exemplary illustration of the measuring cycle for the determination of the MCE at $T \approx 0.324$ K and $B = 4.3$ T: Time dependence of the sample temperature $T_s$ (black spheres), initially in equilibrium with the bath temperature $T_b \approx 0.324$ K and at an external field of $B = 4.29$ T, in response to an up and subsequent down sweep of the magnetic field (red solid line) by $\Delta B = 20$ mT using a sweep rate of 0.31 mT/s. (B) Change of sample temperature $\Delta T_s = T_s - T_b$ around the time of the field sweep with duration $t_{sweep}$. Red solid line corresponds to an exponential fit $T_s \propto \exp(-t/\tau)$ with $\tau = 603$ s, performed over a time period of typically 3 – 4 times $\tau$. The blue vertical arrow, corresponding to $\Delta T + \Delta B$, is derived from an “equal-areas” construction, as indicated in the figure.

**Fig. S2.** Calculated molar magnetic entropy $S_{mag}(T, B=const.)$ of the spin-3/2 paramagnetic salt Cr(NH$_4$)(SO$_4$)$_2$·12H$_2$O shown for $B_i = 2$ T (orange broken line) and $B_f = 0$ (orange solid line); the calculations were based on data taken from refs. 10 and 11.