Signatures of a quantum Griffiths phase in a d-metal alloy close to its ferromagnetic quantum critical point

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Abstract

We report magnetization measurements close to the ferromagnetic quantum phase transition of the d-metal alloy Ni₁₋ₓVₓ at a vanadium concentration of x_c ≈ 11.4%. In the diluted regime (x > x_c), the temperature (T) and magnetic field (H) dependences of the magnetization are characterized by nonuniversal power laws and display H/T scaling in a wide temperature and field range. The exponents vary strongly with x and follow the predictions of a quantum Griffiths phase. We also discuss the deviations and limits of the quantum Griffiths phase as well as the phase boundaries due to bulk and cluster physics.

(Some figures in this article are in colour only in the electronic version)

1. Introduction

Magnetic quantum phase transitions (QPT) have been studied in transition metal alloys and in heavy-fermion compounds tuned, e.g., by pressure or chemical substitution. They still offer challenges to theory and experiment (see [1] for a recent review). Quantum critical behavior is signified by singularities in thermodynamic and transport properties. Usually, specific power laws with characteristic exponents have been predicted at the quantum critical point (QCP) for ‘clean’ homogeneous systems, while ‘disordered’ inhomogeneous systems, driven, e.g., by chemical substitution, may show different behavior [2]. In the case of metallic (itinerant) Heisenberg magnets, a strong-disorder renormalization group [3] predicts an exotic infinite-randomness QCP accompanied by quantum Griffiths singularities [4]. At such a QCP, thermodynamic observables are expected to be singular not just at criticality but in a finite region around the QCP called the quantum Griffiths phase (GP). This region features power laws (e.g., in the magnetic susceptibility, χ ∼ T^{λ−1}, and the magnetization, M ∼ H^λ) characterized by a nonuniversal Griffiths exponent λ which varies with distance to the QCP. Quantum Griffiths singularities have attracted a lot of attention. Many heavy-fermion compounds display anomalous power laws in specific heat C(T) and χ(T) [5]; and quantum Griffiths behavior was suggested as an explanation [6]. Recently, a more systematic variation of the exponents could be found at the ferromagnetic QPT of CePd₁₋ₓRhₓ [7].

To avoid additional complications due to the Kondo effect and to study a larger energy scale we recently investigated the simple fcc transition metal alloy Ni₁₋ₓVₓ [8] as an example of an itinerant ferromagnet (FM) in which the transition temperature (T_c = 630 K for pure Ni) can be tuned to zero by chemical substitution. As explained by Friedel [9] the ‘disorder’ is introduced because the charge contrast of the replacing vanadium atoms creates large defects yielding an inhomogeneous magnetization density. In contrast, diluting Ni with isoelectronic Pd does not lead to a strongly disordered scenario: Ni₁₋ₓPdₓ remains ferromagnetic up to x_c = 0.975 where it rather shows the signatures of a clean quantum critical point [10]. We showed in [11] that magnetization and susceptibility above the critical vanadium concentration x_c ≈ 11.4% where T_c is suppressed to 0 indeed follow simple power laws with nonuniversal exponents that confirm the quantum Griffiths scenario over a wide temperature and magnetic field regime. At very low temperatures, deviations from the quantum Griffiths scenario hint at a cluster glass phase. Here, we provide additional details not shown...
concentration line marks the onset of the CG. The inset shows the strong line is a tail fit describing the FM boundary, while the straight dotted line marks the onset of the CG. The inset shows the strong concentration x-variation of the exponent $\gamma$ from $\chi(T)$, $\alpha - 1$ from $M(H)$ and $\gamma$ from the $H/T$-scaling plot in the GP.

We demonstrate that $H/T$-scaling holds for a wide concentration regime and show the scaling plots. In addition, we reveal how the impact of disorder is manifest in the concentration regime and show the scaling plots. In addition, we reveal how the impact of disorder is manifest in the concentration regime and show the scaling plots. In addition, we reveal how the impact of disorder is manifest in the concentration regime and show the scaling plots.

2. Results

Magnetization and ac-susceptibility measurements were performed on polycrystalline Ni$_1-x$V$_x$ samples with $x = 9–15\%$ as described in [11]. An orbital contribution of $\chi_{orb} = 6 \times 10^{-5}$ emu mol$^{-1}$ has been subtracted from all data shown ($M_{in} = M - \chi_{orb}H$).

Figure 1 shows the temperature-phase concentration diagram. For $x \leq 11\%$, the critical temperature $T_c$ was determined by the standard Arrott analysis. Plots of $M^2$ versus $H/M$ as in figure 2 show straight parallel isotherms which implies $M^2 = M_0^2(T) + cH/M$ as is common for itinerant magnets (in figure 2, only low $T$ data are shown). $T_c$ is then extracted via the mean-field $T$ dependences of $M_0(T)$ and susceptibility ($-\chi(M_0^2(T))$) [12]. The resulting $T_c(x)$ can be simply extrapolated linearly (dashed line) from the high $T_c = 630$ K of nickel down to 0 at $x \approx 11\%$ [8].

For $x \geq 11\%$, the straightforward analysis does no longer work because the data in figure 2 are not described by straight lines. Introducing ‘exponents’ as in a classical critical regime leads to a ‘modified’ Arrott plot [13] implying the behavior $M^{1/\gamma} = M_0^{1/\gamma}(T) + c(H/M)^{1/\gamma}$. A good description for $x > 11\%$ of the $M(H > 0.5 T, T)$ data in a wide regime (outside any critical regime) can be achieved with $\beta = 0.5$ and $\gamma(x) < 1$ [12], as indicated by the dotted fit line in figure 2. The resulting transition temperatures $T_c^x$ of these modified Arrott plots remain finite up to $x = 15\%$, while other extrapolations in figure 2 using smaller $H/M$ values lead to smaller $T_c^x$.

In addition to Arrott plots, we analyze field-dependent maxima at $T_{max}(H)$ in the differential susceptibility $\chi(T) = dM(T)/dH$ indicating spin ordering or freezing as shown in figure 3(a). Figure 3(b) shows the linear extrapolation of $T_{max}(H)$ taken at 0.55–0.1 T to determine $T_{max} = T_{max}(H \to 0)$. In particular for $x = 12.25\%$ a frequency dependent maximum at $T_{max} = 0.19$ K was determined by $\chi_{ac}(T, \nu = 380$ Hz) in zero field with $H_{ac} = 0.1$ G which hints at the onset of a cluster glass [11]. $T_{max}$ increases by 0.018 K/decade in frequency [11] as shown in figure 3(c). Although a detailed study of the evolution with dilution $x$ of the cluster growth and dynamics is still outstanding, we can already note the qualitative effects of disorder on the ferromagnetic ordered state for $x > 11\%$. As is obvious in figure 1, the high and low-field extrapolation lead to different transition temperatures ($T_c, T_{c}^x > T_{max}$) hinting at cluster freezing for $x > 11\%$. The $x$-dependence of $T_{max}$ in the accessible temperature region is better described by an exponential (dotted line) rather than a power law. Also, a ‘tail’ fit to $(\ln(T/T_0) \sim (x_c - x)^{-\psi}$, see [14]) rather than a power law serves as a good description of the onset of FM order for data between about 9% and 11% leading to $x_c \approx 11.6\%$ (solid line). The discrepancies between the various methods and the spin-glass like features at the lowest temperatures suggest that the real QCP is masked at very low $T$ by ordering of clusters.

 Nonetheless, at sufficiently high temperatures (in the region $T_{max} < T < T_c(0%)$) cluster ordering does not seem to play a role, and various quantities display power laws. Figures 4(a) and (b) present the $H$ and $T$ dependences of the magnetization as $M/H$ for various $x$. Figure 4(b) shows essentially the susceptibility $\chi$, since $x = M/H = dM/dH$ for low fields ($H < 0.5 T$) and high $T (T > 20 K, T > T_c)$.
While for $x \leq 11\%$ the negative slope in the log–log plot $\gamma = -d \ln(\chi_{\text{m}})/d \ln(T)$ increases with falling $T$ towards $T_c$, for $x > 11\%$, $\chi(T)$ follows a simple power law for $20 \text{ K} < T < 300 \text{ K}$. The exponent decreases from $\gamma(x = 11\%) = 1$ to $\gamma(x = 15\%) = 0.04$. $M/H(H)$ follows a power law $M/H \sim H^{y-1}$ for high $H$. For $x < 11\%$, where $M(H)$ nearly saturates, the exponent $1-\alpha$ is close to 1, and therefore very different than $\gamma$. However, for $x > 12\%$, the high-field exponent $1-\alpha$ matches the susceptibility exponent $\gamma$. The deviations from a power law at low fields in figure 4(a) are due to the finite $T$ limitations.

Since both $M(T)$ and $M(H)$ show power laws with the same exponent, simple $H/T$ scaling is expected for $x > 12\%$. Figure 5 shows the scaling plot using the form $M/H = H^\gamma Y(\mu_{\text{scal}} H/k_B T)$ where $Y$ is the scaling function and $\mu_{\text{scal}}$ is a scaling moment for several $x$. All $M(H,T)$ data for $T \geq 14 \text{ K}$ collapse, confirming $H/T$ scaling. The scaling function $Y$ is well approximated by the form $Y(\zeta) = A'(1 + \zeta^{-2})^{y/2}$ where $A' = A/\mu^y$ is a constant. This phenomenological form arises from simply combining the two limiting power laws with the same exponent $\gamma$ in the $H$–$T$ plane, $(M/H)^{-1} = H^\gamma Y^{-1} \sim [\mu_{\text{scal}} H]^2 + (k_B T)^y/2$. Close to $x = 11.6\%$, the quality of the collapse is less satisfactory. The resulting exponent $\gamma$ (which matches that obtained by a fit of $\chi(T)$ for all $x$ between 11.4\% and 15\%) is shown in the inset of figure 1. The scaling moment $\mu_{\text{scal}}$ and amplitude $A$ are shown in the inset of figure 5, demonstrating the growth of the typical cluster size and number with $x \rightarrow x_c$.

The consistent power laws, and in particular, the $H/T$ scaling of $M(H,T)$ are in excellent agreement with the predictions for a quantum Griffiths phase with Griffiths exponent $\lambda = \alpha = 1 - \gamma$. A critical concentration of $x_c = 11.4\%$ can be identified from the condition $\gamma(x_c) = 1$ (neglecting logarithmic terms). Fitting to the power law $1 - \gamma(x) \sim (x - x_c)^{\nu\psi}$ as predicted by theory [3] yields $x_c = 11.6\%$ with $\nu\psi = 0.42$ as shown in the upper inset in figure 1. This value is in close agreement with the ‘tail’ fit of $T_{\text{max}}(x)$.

3. Conclusions

On the one hand, our results confirm that Ni$_{1-x}$V$_x$ follows the scenario of an infinite-randomness QCP with a quantum Griffiths phase, as expected in an itinerant Heisenberg magnet [3, 4]. The QCP at $x_c \approx 11.6\%$ has been estimated by extrapolations from outside the critical region, where the
cluster ordering is less disturbing (through $\gamma(x_c) \to 1$ and $T_{\text{max}}(x_c) \to 0$). On the other hand, we see clear signs of cluster ordering towards $x_c$, in particular deviations from scaling at lower temperatures (such as the upturns in figure 4(a) as well as model dependent transition temperatures for $x > 11\%$). As discussed in [11], the magnetization $M(H, T > T_{\text{max}})$ for $x > 12\%$ can be well described by an additional 'Curie term' due to frozen clusters which exceeds the term due to the fluctuating (Griffiths) clusters below $T_{\text{cross}}$ (see figure 1). Such a change in low temperature behavior was predicted to occur in itinerant Heisenberg systems due to the RKKY interactions [15]. A Griffiths phase with nonuniversal power laws at higher $T$ (but below $T_c(0\%)$) combined with a cluster glass (CG) (indicated by maxima in $\chi(T)$) at very low $T$ has also been observed in other diluted compounds (CePd$_{1-x}$Rh$_x$ [7], URu$_{2-x}$Re$_x$Si$_2$ [16]) close to a ferromagnetic transition with much lower $T_c$ and can be understood as a generic feature of this disordered itinerant QPT [14].

Acknowledgments

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