Specific heat study of single crystalline $\text{Pr}_{0.63}\text{Ca}_{0.37}\text{MnO}_3$ in presence of a magnetic field

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We present the results of a study of specific heat on a single crystal of $\text{Pr}_{0.63}\text{Ca}_{0.37}\text{MnO}_3$ performed over a temperature range $3\text{K}-300\text{K}$ in presence of 0 and 8T magnetic fields. An estimate of the entropy and latent heat in a magnetic field at the first order charge ordering (CO) transition is presented. The total entropy change at the CO transition which is $\approx 1.8\text{ J/mol K}$ at 0T, decreases to $\approx 1.5\text{ J/mol K}$ in presence of 8T magnetic field. Our measurements enable us to estimate the latent heat $L_{CO} \approx 235\text{ J/mol}$ involved in the CO transition. Since the entropy of the ferromagnetic metallic (FMM) state is comparable to that of the charge-ordered insulating (COI) state, a subtle change in entropy stabilises either of these two states. Our low temperature specific heat measurements reveal that the linear term is absent in 0T and surprisingly not seen even in the metallic FMM state.

75.30.Kz 65.40.+g 75.30.Vn

The discovery of a number of fascinating properties like colossal magneto- resistance, charge / orbital ordering and electronic phase separation in manganites with generalized formula $\text{Re}_{1-x}\text{Ae}_x\text{MnO}_3$ ($\text{Re}$ being a trivalent rare-earth ion, $\text{Ae}$ a divalent alkaline earth element) has resulted in a spurt of research activities. For certain values of $x$, close to 0.5, these manganites undergo a first order transition at certain temperature $T_{CO}$ to a Charge Ordered Insulating (COI) state where the Mn$^{3+}$ and Mn$^{4+}$ species arrange themselves in a commensurate order in the lattice. The charge ordering transition in these oxides is accompanied by a large change in volume and hysteresis in resistivity and is believed to be a first order transition. A fascinating aspect of the COI state (which is also accompanied by orbital ordering) is that it is unstable under an applied magnetic field and there is an insulator-metal transition (melting) of the COI state to a ferromagnetic metal state (FMM) below a temperature $T_{MH}$.

In this paper we have investigated the specific heat and related thermodynamic quantities in a single crystalline CO system over a wide temperature range ($3\text{K} < T < 300\text{K}$) and in a magnetic field upto 8T which can melt the CO order. A study of specific heat over an extensive temperature range is crucial in understanding the nature of the CO transition and in addition it can provide values of various fundamental parameters of manganites like the density of states at the Fermi level $N(E_F)$, the Debye temperature $\theta_D$, ferromagnetic / antiferromagnetic spin wave stiffness constant etc.

Measurements of specific heat in manganites particularly with CMR composition has been reported before, particularly in the low temperature region. In the CO systems there are reports of specific heat measurements both in single and polycrystalline samples mainly at low temperatures. Specific heat measurements in CO systems near $T_{CO}$ have been done in polycrystalline samples and no magnetic field data have been reported. There exists no data on thermodynamics associated with the transition at $T_{MH}$.

The charge ordering transition in these oxides is believed to be a first order transition, as mentioned before. Measurements across the CO transition should therefore show a discontinuous jump in entropy as the compound absorbs latent heat from the bath to transform to a new phase. However, till date there exists no proof from calorimetry that the transitions are indeed first order and whether a latent heat is released at the transition. There has been no calorimetric investigation of the melting of COI state to a FMM state, the transition which is also believed to be a first order transition.

In this paper we investigate this fundamental issues related to changes in entropy and latent heat across the charge-ordering transition near $T_{CO}$ and $T_{MH}$ in a single crystal of $\text{Pr}_{0.63}\text{Ca}_{0.37}\text{MnO}_3$. $\text{Pr}_{1-x}\text{Ca}_x\text{MnO}_3$ happens to be a prototypical and perhaps the most studied charge-ordered system. Due to its low tolerance factor it remains insulating for all values of $x$. For $x = 0.37$ composition, charge and orbital ordering occurs at $T_{CO}$ ($\approx 235\text{K}$), while a long range AFM order sets in only below $T_N$ ($\approx 170\text{K}$). A small ferromagnetic component appears with the canting of AFM spins at a lower temperature, $T_{CA}$ ($\approx 30\text{K}$).

Though there have been a number of studies on low
temperature specific heat of this particular system ($T < 20K$), a clear picture is yet to be emerge on the presence (or absence) of a linear term in specific heat. Recently there have been reports of large linear contribution to specific heat and appearance of excess specific heat associated with charge ordering in this system. Since such a linear term is often associated with a electronic contribution to the specific heat, appearance of this term in an insulating sample is intriguing. In this paper, we have also investigated the low temperature region and reached certain definite conclusions about the linear term.

In our experiments we have specifically asked the following questions:

1. Is it possible to identify the relevant transitions in zero and finite magnetic fields through calorimetric measurements?
2. What latent heat is released and the entropy change across the first order transitions, one at $T_{CO}$ in zero magnetic field and the other at $T_{MH}$ in 8T magnetic field?
3. When the FMM state is obtained from the COI state in presence of a magnetic field, do we obtain a linear term (arising from electronic contribution) in the specific heat?
4. Do we see a linear term in specific heat in COI state as reported by some investigators?

The remainder of the paper is divided into two principal sections. In the first section after the experimental section, we present and discuss the experimental data in the region $T \geq 50K$ which essentially encompasses the region where most of the transition occurs and this also happens to be the region where no experimental data have been reported in the past either in presence of magnetic field or in a single crystal. The second major section refers to the data at low temperatures where issues like linear term in the heat capacity, the Debye term, spin wave contributions etc. are looked into. In this region there are past studies, as mentioned earlier and we compare our results on these material with results from other CO systems.

**I. EXPERIMENTAL**

The crystal used in our experiment has been grown by float zone technique and has been used in a number of previous experiments by our group. The resistivity ($\rho$) vs. $T$ curve in zero field and in a field of H = 8T are shown in fig. 1. The COI state can be melted to a FMM state by application of a magnetic field of 8T at $T_{MH} \approx 90K$. In presence of magnetic field the region $T_{MH} < T < T_{CO}$ is termed as “mixed charge order” (MCO) region, where the COI phase coexist with the FMM phase. We have used a semi-adiabatic heat pulse technique to measure the specific heat in a wide temperature range of 2K - 300K and in 0T and 8T magnetic fields.

**II. SPECIFIC HEAT AT HIGH ($T \geq 50K$)**

In figure 2 we show the specific heat ($C_p$) data over an extended temperature range 3K - 300K for both 0 and 8T. The most prominent feature is the sharp peak at $T_{CO}$ both for H = 0 and 8T. The peak is much larger and sharper compared to that seen in polycrystalline materials (which is also plotted in the same graph). The peak at $T_{CO}$ retains its narrowness and sharpness at H = 8T while it shifts to lower T with $dT_{CO}/dT \approx -1K/T$. In figure 3 we show the region close to $T_N$. The graph shows the actual observed step like feature in $C_p$ (at H = 0T) at $T_N$. In the same graph we show the $C_{exc}$ after subtraction of the lattice contribution (the procedure for subtraction of the lattice contribution is given in the next sub-section). A peak in $C_{exc} \approx 6J/moleK$ at $T_N$ is visible. This feature is suppressed by application of a magnetic field of 8T. The entropy change, $\Delta S_{exc}(T_N)$, associated with this transition has been estimated to be $\approx 0.5 - 0.8J/mole.K$. Lee et. al. reported for $Pr_{0.6}Ca_{0.4}MnO_3$(polycrystal) $\Delta S_{exc}(T_N) \approx 0.6J/mole K$ at $T_N = 160K$. Ramirez et. al. for $La_{0.35}Ca_{0.65}MnO_3$(polycrystal) obtained $\Delta S_{exc}(T_N) \approx 1J/moleK$. It can be seen that all these are much less than that what one would expect from a complete spin ordering. The calorimetry data therefore points towards an incomplete spin ordering at $T_N$.

Another very interesting feature in our calorimetry data is a clear signature of the magnetic field induced transition at $T_{MH} \approx 78K$ (see fig. 2 and 9). This small yet distinct feature, discussed in detail later on, is the first signature of field-induced melting in a calorimetry experiment.

**A. Estimation of the Lattice contribution**

In this subsection we describe the procedure to estimate the lattice contribution to specific heat, the background on which the specific heat contribution by the other degrees of freedom add up. Proper estimation of the lattice contribution will thus allow us to get the contribution of CO and magnetic ordering to $C_p$. We define $C_{exc}$ as:

$$C_{exc} = C_p - C_{lattice}$$

According to Dulong and Petit’s law, the limiting heat capacity at high temperature for a compound with $r$ atoms per molecule is expected to be $3rR$, where R is the
gas constant \[13\]. In our case \((r = 5)\) this limiting value of the lattice (vibrational) heat capacity turns out to be \(\approx 125 \text{ J} / \text{ mole K}\). From our data we find that at \(T = 300K\), observed \(C_p\) is \(\approx 90\%\) of the Dulong Petit value. For most published data on manganites, \(C_p\) at room temperature reaches this value \(\sim 100 - 120 \text{ J} / \text{ mole K}\). This signifies that bulk of the contribution indeed comes from the vibrational heat capacity in this temperature range. This contribution must be subtracted from the observed data to obtain the excess specific heat.

We have obtained the lattice contribution \(C_{\text{lattice}}\) by fitting \(C_p\) in a region \(40K \leq T \leq 150K\) using three standard models as described below. The calculated \(C_{\text{lattice}}\) is then extrapolated to \(T > 150K\). Since, we are interested in \(C_p\) in the range \(80K - 300K\), the lower temperature for lattice contribution has been limited only to 40K. The upper limit of 150K was considered since it was lower than the AFM and CO transition temperatures. We have not estimated the background by including the data about \(T_{\text{CO}}\). We are of the opinion that there is structural transition/ modification associated with the CO transition. As a result it is not advisable to include the data above \(T_{\text{CO}}\) in the estimation of lattice heat capacity below \(T_{\text{CO}}\).

The following models were used for estimating the background lattice contribution:

**The Einstein model \[13\]:**

\[ C_{\text{Einstein}} = 3rR \sum a_i x_i^2 e^{x_i} / (e^{x_i} - 1)^2 \]

where \(x_i = h\nu E_i / K_B T_i\). In this model, all the \(3rN\) independent oscillators populate three optical modes in ratios \(a_1 : a_2 : a_3\) having Einstein frequencies \(h\nu E_i\). The best fit to the data is given the three Einstein modes with \(h\nu / K_B = 145\ K\), \(410\ K\) and \(625\ K\). It is interesting to note that a recent Raman measurement on \(x = 0.37\) composition observed optical modes at \(\approx 360\ K\), \(417\ K\), and \(648\ K\). The \(417K\) and \(648K\) modes nicely matches with those seen in our specific heat measurements. \[13\]

**The Debye model \[13\]:**

\[ C_{\text{Debye}} = 9rR / x_D^3 \int_0^\infty x_D^4 e^{x_D} / (e^{x_D} - 1)^2 dx_D \]

where \(x_D = h\nu D / K_B T\). The specific heat is due to collective low-frequency oscillations of phonons with a cut off frequency given by \(h\nu D / K_B\). For our sample, the best fit was obtained with \(r_D = 470K\). This is typically the value of \(r_D\) seen in most oxides.

**The Thirring model \[13\]:**

\[ C_{\text{Thirring}} = 3rR \sum \infty n = 0 B_n u^{-n} \]

where \(u = [(T/T_n)^2 + 1]\). The harmonic portion of the lattice specific heat can be expressed in a series with the above form where \(T_b \approx \theta_D / 2\pi\). \[13\] The above expansion permits the harmonic portion of the lattice specific heat to be fitted reasonably well down to temperatures \(\sim 50K\) even when the Debye temperature is \(\sim 500K\). In our case, \(T_b = 65K\) and we used \(n\) up to 50. The \(T_b\) gives an estimated \(\theta_D \approx 410K\), which is close to that obtained from fitting the data with Debye model.

In figure 4 we have shown the deviation of the observed data from the fit (i.e. \(\Delta C/C = (C_{\text{obs}} - C_{\text{calc}}) / C_{\text{obs}}\)) in order to ascertain the extent of uncertainties involved in the background subtraction. We find that for \(T < 100K\) the Debye model shows large systematic deviation. The Thirring model shows a systematic deviation above \(150K\) (not shown in graph). We find that over the whole range the least uncertainty is shown by the Einstein model (maximum deviation from fit \(\pm 5\%\)) and this is a random deviation.

We have used the same \(C_{\text{lattice}}\) for both \(H = 0\) and \(H = 8T\). We found that if we fit the \(C_p\) at \(H = 8T\) from \(50K\) to \(140K\) to the above models as we have done for the \(C_p\) at \(H = 0T\) case, we end up in getting essentially the same \(C_{\text{lattice}}\) using the Einstein model. There is a small \((< 10\%)\) systematic deviation (for the Einstein model) which is not numerically significant to affect our results. This is shown in the inset of figure 4. For compounds containing \(P_r^{3+}\) \(C_p\) may contain a contribution coming from the crystal field. We have estimated this crystal field contribution or \(C_{\text{cystal}}\) the crystal field data available for iso-structural compounds \(\text{PrNiO}_3, \text{PrGaO}_3\). We find that in the range of interest \((100K < T < 300K)\) this crystal field contribution \(< 7\%\) at \(100K\) and \(< 2\%\) at \(300K\) \[15\].

**B. \(C_{\text{exc}}\)**

In figure 5 we have plotted the value of \(C_{\text{exc}}\) for both \(H = 0\) and \(H = 8T\) for \(T > 120K\). The data show the \(C_{\text{exc}}\) using all the three models. The \(C_{\text{exc}}\) data in the region close to \(T_{\text{CO}}\) is truncated in figure 5, because in this region the \(C_{\text{exc}}\) is very large. In this temperature range the differences in the \(C_{\text{exc}}\) obtained after the subtraction of the lattice contribution is well within 1 - 2 J/mole.K for the Debye and the Einstein model. The Thirring model shows systematic deviation for \(T > 150K\). Considering all the models we find that near \(T_N\) the uncertainty is the largest since \(C_{\text{exc}}\) is low and this can be as large as \(\pm 25\%\). However close to the CO transition when \(C_{\text{exc}}\) shots upto \(\approx 220J/mole.K\). The uncertainty in the background estimation falls below \(\pm 2\%\). The \(C_{\text{exc}}\) data also show clearly that barring the small region around \(T_N\) (where \(C_{\text{exc}} \approx 10\%\) of \(C_p\)) and near \(T_{\text{CO}}\) (where \(C_{\text{exc}} > 50\%\) of \(C_p\)) the contribution by other degrees of freedom compared to the lattice contribution is negligible in the temperature range \(T > 50K\).

**C. Entropy change near \(T_{\text{CO}}\) in \(H = 0\) and \(H = 8T\)**

A phase transition is signaled by a singularity in a thermodynamic potential such as free energy. If there is a finite discontinuity in one or more of the first derivatives of the free energy, the transition is first order. At
a first order transition one expects a discontinuous jump in the entropy. At the charge ordering transition, (which is conjectured as first order transition based on observed hysteresis in transport data on field and temperature cycling), our system is expected to absorb the latent heat of transformation to transform from charge-ordered phase to a charge-disordered phase as it is heated through \( T_{CO} \). The temperature should remain constant till this process is complete and the entropy change is given as:

\[
\Delta S_{21} = S_2 - S_1 = L_{21}/T
\]

where \( S_1, S_2 \) are the entropies of phases 1 and 2, \( L_{21} \) is the latent heat associated with the phase transformation. At a first order transition, since the entropy changes discontinuously, specific heat is actually undefined. Because at the transition point, the temperature does not change when heat is applied one would expect a \( \delta \) function like behaviour. In reality, however, one rarely sees such a \( \delta \) function like behaviour of \( C_p \) because the transition can get broadened either by the process of measurement or by the quality of the sample. This makes a part of the expected entropy change to become continuous and it shows up as a finite measurable specific heat. Interestingly, it is only very recently the expected features in specific heat in a first order transition has been seen experimentally in rare-earth compounds. In the following discussion, we have to keep in mind the above special features since in a first order transition has been seen experimentally in rare-earth compounds \[22\]. In the following discussion, we have to keep in mind the above special features since at a first order transition one expects a discontinuous jump in the entropy being the difference of two quantities with similar background, the error arising from the estimation of the background is not severe.

The charge-ordered phase is expected to have a lower entropy than the charge-disordered insulating phase at \( T > T_{CO} \). This implies that the sample absorbs latent heat to transform from the charge-ordered phase to the disordered phase. We would expect that at least a part of the entropy change \( \Delta S_T \) is released as latent heat is absorbed from the total heat \( \Delta Q \) supplied during the heat-pulse experiment and the sample temperature remains quite constant throughout the process. Since the change in sample temperature \( (\Delta T) \) is quite small, the specific heat \( C_p = (dQ/dT)_p \) shows a very sharp peak near \( T_{CO} \). As discussed before at a first-order transition \( \Delta T \to 0 \) and ideally \( C_p \) should be a \( \delta \) function at the transition temperature.

The peak however is broadened due to two reasons:

(a) Crystal quality : In most cases the expected \( \delta \) function gets broadened by sample quality. The real crystal contains some defects / inhomogeneities which would lead to broadening of the peak. With improved quality of the sample, the peak would be larger and sharper. For oxides containing multiple chemical constituents it is quite likely that such is the case. As shown in figure 1, where we have compared the \( C_p \) of a ceramic pellet and a single crystal, the single crystal has a much higher and narrower \( C_p \) at \( T_{CO} \) as compared to the ceramic sample.

(b) Measurement broadening : The second reason for broadening of the peak is connected to the measurement procedure itself. \( C_p \) measurement involves measuring a finite temperature jump \( \Delta T \) following the heat pulse \( \Delta Q \). Since \( C_p \) is well behaved in the region away from \( T_{CO} \), it does not depend much on the size of the temperature rise \( \Delta T \) (as long as \( C_p \) is not a very steep function of \( T \)). However, close to \( T_{CO} \), the height as well as the width \( \delta T \) of \( C_p \) sensitively depends on \( \Delta T \).

We have measured the latent heat by measuring \( C_p \) with different \( \Delta T \) as suggested by Gschneider et. al. \[22\]. A sample of the data is shown in figure 8. We find that while away from \( T_{CO} \) there is no dependence of \( C_p \) on the rise of \( \Delta T \), as expected, the peak becomes narrower and higher when size of \( \Delta T \) decreases as we approach \( T_{CO} \). This continues till \( \Delta T \leq 0.5K \). For this value we reach a limiting width (at half maxima) \( \delta T \approx 2K \). We believe that for \( \Delta T \leq 0.5K \), the width of the peak is not determined by the measurement but by the crystal quality. This observation sets a limit to the height and width of \( C_p \). We find from our measurement that the latent heat \( \Delta L_{CO}/T_{CO} = \Delta S_{CO} \approx 1 \text{ J/mole K} \). Similar value of \( \Delta S_{CO} \) was obtained for \( H = 8T \). Our estimate of latent is a lower bound of the true latent heat and with a better crystal preferably with a transition width < 1K, \( \Delta S_{CO}/\Delta S_T \) will increase substantially and may
even → 1, as expected for a strong first order transition. We note that such a narrow peak might arise from a very narrow second order transition that has been broadened by defects or inhomogeneities. However we find that the applied field shifts the position of the peak but does not appreciably broaden the peak. This we take as a proof of a first order transition at $T_{CO}$.

D. Entropy change near $T_{MH}$ in 8T

In a field of 8T the low temperature phase is an FMM or a spin-aligned metal, allowing for spin canting. But the phase is metallic as can be seen from figure 1. On heating, the FMM state (8T) becomes unstable towards the COI state formation and makes transition to the mixed charge ordered (MCO) state at $T_{MH}$. This is accompanied by a jump in $\rho$. We would like to ask if there is a change in entropy associated with the melting of the COI state to FMM state at $T_{MH}$? This question is interesting because both the phases have ordering of a kind, in addition the FMM phase is expected to have extra entropy due to presence of free electrons. A close look at the region $T \approx 95K$, as shown in figure 9, shows a small dip in $C_p$ which as explained below can be considered as signature of this melting. It is important to note that, as expected in zero field, this feature is absent. As shown in fig. 9, $C_p(8T)$ starts to show change at $T = T_{MH} \approx 88K$ where $\rho$ shows a jump on heating and the specific heat transition is complete at 100K after a small dip in $C_p$, where the resistivity transition also stops.

The small dip in $C_p$ is associated with a small heat release of $\approx 10J$/mole at around 95K - 100K on heating. This suggests that on heating the FMM phase, stable in 8T for low T, starts to disorder. This probably destabilizes the FMM phase with regions of high resistivity COI regions appearing in it. Eventually at $T \approx T_{MH}$ the insulating regions increase in size and $\rho$ shows a jump in $\rho$. The resistivity transition has an element of percolation associated with it and thus occurs at a different volume fraction of the new phase in contrast to the specific heat transition which occurs mainly when the bulk of the sample is transformed. Nevertheless, the small but distinct change in $C_p$ close to $T_{MH}$ is clearly seen.

In the temperature range $T_{MH} < T < T_{CO}$ in a magnetic field (i.e., the MCO region) we find an interesting effect. The thermal relaxation time of the sample increases by more than one order of magnitude. This anomalously large relaxation is not observed at any temperature range in zero field or for $T > T_{CO}$ and $T < T_{MH}$ in $H = 8T$. The existence of this large thermal relaxation necessitated that the sample be properly equilibrated before the data are taken. We have done that and the data presented here are taken after the sample has been properly equilibrated thermally. The thermal relaxation, however, is interesting in its own right and has been discussed in a separate publication.

III. LOW TEMPERATURE $C_p$ IN 0 AND 8T MAGNETIC FIELD

As it has been pointed out before, that the issue of low temperature specific heat ($T < 10K$) is controversial in manganites showing charge ordering, particularly in Pr$_{1-x}$Ca$_x$MnO$_3$ system. A large number of observations on low temperature specific heat is available for different compositions of Pr$_{1-x}$Ca$_x$MnO$_3$, mostly in polycrystalline pellets and some in single crystals.

In figure 10 we show the specific heat data of our sample with that of different compositions as obtained by different groups. Such a comparison is meaningful because it brings out the essential similarity and differences in these materials. A large linear term ($\gamma_\approx 30.6$ mJ/mole-K$^2$ for $x = 0.3$ and $\gamma_\approx 15.7$ mJ/mole-K$^2$ for $x = 0.35$) appears in polycrystalline samples with $x = 0.3$ and $x = 0.35$ composition which decreases as $x \rightarrow 0.5$. We note that such a large linear $\gamma$ has not been seen any of the single crystal data. From a comparison of the data shown in figure 10 we can reach the following conclusions:

(i) The polycrystalline ceramic samples have higher specific heat than the single crystal samples,

(ii) There is a clear trend that the specific heat of the ceramic samples decreases as we approach $x = 0.5$. In particular, linear term in specific heat, is only observed for the ceramic samples and it decreases with increasing $x$.

(iii) The specific heat of the single crystal samples with compositions $x = 0.37$ and 0.5 are very similar for $T < 10K$.

In figure 11 we have compared the specific heat of two close compositions: a polycrystalline $x = 0.35$ sample and a single crystal sample with $x = 0.37$ composition. The comparison is to elucidate how much of the excess specific heat is due to polycrystallinity and can we infer a likely origin of the excess specific heat. There are two mechanisms that can contribute to the excess specific heat: (1) Due to grains of small dimensions and (2) Due to two-level-systems(TLS) arising due to disorder as in amorphous solids and several disordered crystals. If the excess specific heat, is due to grains then $\delta C$ can be expressed as $\delta C = C_1T + C_2T^2$, where the coefficients $C_1$ and $C_2$ are related to the average grain diameter $< R >$ as $< R > \approx \frac{k_B}{6\nu_{sound}}C_1^{-0.5}$, where $\nu_{sound}$ is the sound velocity in the crystal. We can fit the experimentally observed $C_{poly}$ to the equation $C_{poly} = C_{sc} + \delta C$. Using $C_{sc}$ from our observed data and the above expression of $\delta C$ we obtain an estimate of the average grain size $< R >$ and we obtain $< R > \approx 10^{-4} \mu m$ which is far too small compared to the
typical \( < R >> 1 \mu m \) seen in most polycrystalline samples. We can thus rule out finite grain size as the source of excess specific heat.

If the excess specific heat \( \delta C \) arises from TLS then it can be fit to a form \( \delta C = C_1 T + C_2 T^3 \). This would arise from TLS with density of state \( P(E) = \tilde{a} + b E^2 \), where \( \tilde{a} \) and \( b \) are constants [21]. Using this expression for \( \delta C \) we then fitted the observed \( C_{\text{poly}} \) as had been done before. The fit is shown in figure 11. From the fitted parameters \( C_1 \) and \( C_2 \), we arrive at values of \( \tilde{a} \approx 2.3 \times 10^{35} \text{ erg}^{-1}/\text{cc} \) and \( b \approx 1.5 \times 10^{35} \text{ erg}^{-3}/\text{cc} \). These values are quite comparable to but some what larger than those obtained in glasses. We conclude that extra specific heat of the polycrystalline samples arise from excitations which behave as TLS. We have no clear understanding of the origin of these TLS but we can speculate that they will arise from incomplete orbital /charge order that can happen in polycrystalline samples due to random strains. What is needed for TLS to occur is to have multitude of ground states with almost degenerate energy. This can actually be due to the predominant incommensurate charge ordering as seen in both \( x = 0.35 \) and \( x = 0.37 \) compositions. As the charge-ordering becomes commensurate with the lattice with \( x \rightarrow 0.5 \), the difference in the specific heat between the polycrystalline and the single crystal samples decrease. We thus suspect that the disorder in polycrystalline samples give rise to TLS low energy excitations typically with energy \( \leq 10 K \) and as \( x \rightarrow 0.5 \), the \( P(E) \rightarrow 0 \).

Next we attend to the low temperature data taken on our single crystal sample. Figure 12 shows the specific heat data for the \( x = 0.37 \) sample plotted as \( C_p/T \) vs. \( T^2 \), a customary way to plot the data in anticipation of a linear term. It is clear that the linear term is absent in the COI state. This is not surprising for an insulating sample, the linear term was also absent in the specific heat data observed in the ceramic samples of \( x = 0.4 \) by Lees et. al [3]. Neglecting the linear term, we have fitted our observations to the following relation:

\[
C_p = aT^{-2} + \beta_3 T^3 + \beta_5 T^5
\]

where, \( aT^{-2} \) hyperfine contribution caused by the local magnetic field at the Mn nucleus due to electrons in unfilled shells, \( \beta_3 T^3 \) and \( \beta_5 T^5 \) are the lattice contribution to the specific heat, arising from phonons. A part of the \( T^3 \) contribution to \( C_p \) can also arise from AFM spin waves. The \( T^3 \) contribution is likely to be enhanced over and above the actual Debye contribution because of the presence of AFM spin waves, since our COI sample has canted AF order in this temperature range. The results of fitting the data to the above equation are shown in table 1. The parameters obtained by us are very similar to that seen by Lees et. al [3]. In table 1 we have collected the parameters from different published data for comparison.

However, the interesting question is whether in a magnetic field of 8T where the COI is melted into a FMM phase, we see appearance of a linear specific heat term. In a past investigation on polycrystalline samples of \( \text{La}_{0.5} \text{Ca}_{0.5}\text{MnO}_3 \) in a magnetic field of 8.5 T [10] (which is high enough to melt the COI state) no linear term had been observed. The issue of absence of a linear term in the FMM phase obtained after melting the COI state is thus real. Particularly, when one compares it with the low temperature specific heat of the FMM phase as seen in the CMR region \( \gamma \approx 7.8 \text{ mJ/mole K}^2 \) in \( \text{La}_{0.8} \text{Ca}_{0.2}\text{MnO}_3 \) [21], \( \approx 5.2 \text{ mJ/mole K}^2 \) in \( \text{La}_{0.7} \text{Ca}_{0.3}\text{MnO}_3 \) [20].

The \( C_p/T \) vs. \( T^2 \) for our sample in presence of an 8T magnetic field are shown in figure 12. The specific heat data have been fitted using the eqn. 1, with the parameters as shown in table 1. Interestingly, the specific heat in presence of magnetic field is distinctly lower than the specific in 0T for all \( T < 40K \). Lowering of specific heat in presence of magnetic field has been reported Smolyaninova et. al [3]. Since resistivity measurements indicate the sample to be in a metallic state, one would expect an extra linear term appearing in the eqn.1 in presence of magnetic field. As seen in fig. 12, we could not detect the presence of any linear term even in presence of 8T. From the uncertainty in the data we find an upper limit of \( \gamma \leq 0.1 \text{ mJ/mole K}^2 \), which is too small.

It is apparent from table 1, that contribution of the \( \beta_3 \) term is halved in presence of magnetic field. This appears to be the main reason for lower \( C_p \) in \( H = 8T \) compared to that in absence of magnetic field. This can be explained as follows. The \( T^3 \) term contains an additional antiferromagnetic spin wave contribution for the COI sample. For an antiferromagnetic spin wave spectrum where \( E = D_q \), the magnetic contribution to specific heat is \( \propto T^3 \). The melting of COI state to a FMM state leads to collapse of this AFM order and thus there will be no AFM spin wave contribution to the specific heat. This would result in a decrease of the \( T^3 \) term. However, presence of FM order should give a ferromagnetic spin wave contribution \( \propto T^{3/2} \). We have attempted fitting a \( T^{3/2} \) term to the 8T specific heat data. However, we could not detect any \( T^{3/2} \) term. Thus, it appears that the specific heat as observed in presence of magnetic field does not have a ferromagnetic spin wave contribution.

Briefly, we find that the specific heat at low temperature needs more investigation in single crystalline CO systems and in high magnetic field. The existing data (which includes too few single crystal data) both in zero field and magnetic field (high enough to melt the CO state) does not allow us to reach much definite conclusion other than the fact that in the FMM phase obtained by melting COI state in a high magnetic field does not have a large enough \( \gamma \) that is comparable to FMM phases seen in manganites showing CMR behavior.
IV. CONCLUSION

Our study of specific heat in a CO system done over an extended temperature range of 3K - 300K in presence of 0 and 8T magnetic field gives us a number of useful and new information regarding the thermodynamics of CO transition. This paper gives the first clear measurement of the entropy in a single crystal and also the latent heat (at T_{CO}) in zero field and in a magnetic field. The latent heat released proves the strong first order nature of the transition, which had never been shown to be so in actual measurement of specific heat. An estimate of the numbers like the latent heat in 0 and 8T field sets clearly the bound and scale within which any theoretical models must work. It is a very important number for any phase transition which gives the CO transition (both in 0T and 8T) a thermodynamic basis. Important is also the observation of a finite entropy change at the CO melting in a magnetic field. The result shows that a small but finite and qualitative difference in entropy exists between the FMM phase and the CI phase at the melting transition in the magnetic field where the former (FMM phase) has a lower entropy than the other phase (which is the high T phase). Interesting also is the smallness of the entropy. The low entropy than the other phase (which is the high T phase). (FMM) phase (which is the low T phase) has a lower entropy at the small but finite and qualitative difference in entropy at the temperature.

An attempt had been made to understand the observed discrepancy between the specific heat of a single crystal and a polycrystalline sample in the low temperature regime.

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Figure captions:

**Figure 1:** The temperature dependence of resistivity of Pr$_{0.63}$Ca$_{0.37}$MnO$_3$ in presence of 0 and 8T magnetic fields.

**Figure 2:** $C_p$ of single crystalline Pr$_{1-x}$Ca$_x$MnO$_3$ (x = 0.37) over 3K-300K in presence of 0 and 8T magnetic fields. Also plotted the $C_p$ of the polycrystalline x = 0.4 composition.

**Figure 3:** $C_p$ and $C_{exc}$ close to $T_N$ in H = 0T.

**Figure 4:** Deviation of $C_p$ from the fit for different models. Inset shows the same near $T_{MH}$ in H = 8T.

**Figure 5:** $C_{exc}$ obtained after subtracting the lattice background (as determined by the three models) for both H = 0 and 8T.

**Figure 6:** $C_{exc}/T$ around $T_{CO}$ in H = 0 and 8T.

**Figure 7:** Excess entropy $S_{exc}$ as calculated from numerically integrating $C_{exc}/T$ around $T_{CO}$ in H = 0 and 8T.

**Figure 8:** Varying sharpness and width of $C_p$ near $T_{CO}$ for two different temperature rise $\Delta T$ = 0.5 and 1K.

**Figure 9:** $C_p$ and resistivity near the $T_{MH}$. The arrow marks the melting of the COI state.

**Figure 10:** A comparison of $C_p$ of Pr$_{1-x}$Ca$_x$MnO$_3$ in the low temperature regime as observed by different groups, x = 0.5(sc),0.45(ce),0.35(ce) a, x = 0.4(ce) b and x = 0.37(sc) is our sample.

**Figure 11:** The excess specific heat, $\delta C$, as seen in the polycrystalline sample arising due to presence of two level states.

**Figure 12:** The low temperature $C_p$ of single crystalline Pr$_{0.63}$Ca$_{0.37}$MnO$_3$ in H = 0 and 8T. The solid line is the fitted equation.

**TABLE I.** The fitting results for the $C_p$ of Pr$_{1-x}$Ca$_x$ MnO$_3$ as obtained by various groups. The units of different quantities are : $\alpha$ (mJ K/mole), $\gamma$ (mJ/mole K$^2$), $\beta_3$ (ml/mole K$^3$) and $\beta_5$ ($\times 10^{-4}$ mJ/mole K$^6$)

| x (Pr$_{1-x}$Ca$_x$MnO$_3$) | $\alpha$ | $\gamma$ | $\beta_3$ | $\beta_5$ |
|---------------------------|---------|---------|----------|----------|
| 0.3$^a$                  | 63.0    | 30.6    | 0.30     |          |
| 0.35$^a$                 | 56.0    | 15.7    | 0.39     |          |
| 0.45$^a$                 | 28.0    | 3.1     | 0.31     |          |
| 0.5$^a$                  | 22.0    | 2.4     | 0.26     |          |
| 0.4$^b$                  | 28.0    |         | 0.54     | 4.5      |
| 0.37 (H = 0T)$^c$        | 85.0    | 0.54    | 4.5      |          |
| 0.37 (H = 8T)            | 400.0   | 0.16    | 18.0     |          |

$^a$from Ref. [1]

$^b$from Ref. [2]

$^c$our sample
$\rho$ (ohm cm) vs. $T$ (K)

- $T_{\text{MH}}$
- $T_N$
- $T_{\text{CO}}$

$\Pr_{0.63}\text{Ca}_{0.37}\text{MnO}_3$

Figure 1
$C_p (J \text{ / mole K})$

$T(K)$

$x = 0.4 \text{ (ce)}$

figure 2
\[ \frac{\Delta C}{C_p} = \frac{C_{\text{obs}} - C_{\text{cal}}}{C_{\text{obs}}} \]

Figure 4
figure 5

H = 0T

$C_{\text{thir}}$, $C_{\text{ein}}$, $C_{\text{deb}}$

$C_{\text{exc}}$ (J/mole K)

H = 8T

$C_{\text{thir}}$, $C_{\text{ein}}$, $C_{\text{deb}}$

$C_{\text{exc}}$ (J/mole K)

T(K)
Figure 6

The graph shows a plot of $C_{exc}/T$ (J/mole K$^2$) against $T$ (K) for two different magnetic fields: 0T and 8T. The graph highlights the peak behavior at different temperatures, indicating a phase transition or anomaly specific to the system under study.
Figure 7

The graph shows the variation of entropy ($S$) with temperature ($T$) for different magnetic fields. The curves marked $0T$ and $8T$ represent the entropy at zero and eight times the magnetic field, respectively. The changes in entropy, $\Delta S_T$ and $\Delta S_{CO}$, are indicated by arrows for the respective magnetic field conditions.
$H = 8T$

Figure 9
$\text{Pr}_{1-x} \text{Ca}_x \text{MnO}_3$

$x = 0.3 \ (\text{ce})$

0.35 (ce)

0.37 (sc)

0.4 (ce)

0.45 (ce)

0.5 (sc)

sc : single crystal

ce : poly crystalline
$C_p$ calculated from a 2 level state model

$\delta C$ calculated from a 2 level state model

$x = 0.35$ (ce)

$x = 0.37$ (sc)

Figure 11
