Small-angle neutron scattering study of mesoscale magnetic disordering and skyrmion phase suppression in the frustrated chiral magnet Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$

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1. Introduction

Small-angle neutron scattering (SANS) is an established technique for scrutinizing mesoscale incommensurately modulated magnetic structures in solids (Mühlbauer et al., 2019). Nowhere is this clearer than in the study of spiral, vortex, meron and skyrmion phases in various non-centrosymmetric magnets, ranging from the itinerant B20s MnSi (Mühlbauer et al., 2009) and FeGe (Moskvin et al., 2013), to the intermetallics Co–Zn–Mn (Tokunaga et al., 2015), Co–Zn–Mn chiral cubic magnets display versatile magnetic skyrmion phases, including equilibrium phases stable far above and far below room temperature, and the facile creation of robust far-from-equilibrium skyrmion states. In this system, compositional disorder and magnetic frustration are key ingredients that have profound effects on the chiral magnetism. Reported here are studies of the magnetism in Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ by magnetometry, small-angle neutron scattering (SANS), magnetic diffuse neutron scattering and Lorentz transmission electron microscopy (LTEM). While features in magnetometry and LTEM often give standard indications for skyrmion formation, they are not readily observed from the measurements on this system. Instead, skyrmion lattice correlations are only revealed by SANS, and they are found to form an orientationally disordered structure in a minority fraction of the sample. The majority fraction of the sample always displays orientationally disordered helical spin correlations, which undergo further disordering along the radial direction on cooling below the critical temperature ($T_c \approx 102$ K). The near-complete suppression of the skyrmion phase, and the process of disordering on cooling, are attributed to competing magnetic interactions that dominate over the ferromagnetic interaction expected to favour chiral magnetism in this system. These competing interactions start to develop above $T_c$ and become further enhanced towards low temperatures. The present observations of co-existing and disordered magnetic correlations over multiple length scales are not unique to Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ but are seemingly common to the family of Co–Zn–Mn compounds with finite Mn, and their accurate description presents a challenge for theoretical modelling. In addition, this study highlights a need for neutron instrumentation capable of the comprehensive measurement of magnetic correlations over expanded ranges of momentum transfer in such multiple-length-scale magnets.
CeAlGe (Puphal et al., 2020) and Y3CoSn4 (Takagi et al., 2018), to the insulators Cu2OSeO3 (Seki et al., 2012), GaV2Se8 (Bordács et al., 2017) and VOSe2O3 (Kurumaji et al., 2017), to name a few. In itinerant chiral cubic magnets, the antisymmetric Dzyaloshinskii–Moriya interaction (DMI) \( D \) arises from the spin–orbit interaction in the presence of broken space inversion, and competes microscopically with a dominant ferromagnetic exchange \( J \). This competition leads to the formation of a long-period helical spiral ground state that modulates with a real-space period \( \lambda \propto J/D \) along directions determined by magnetic anisotropy. Typical length scales for the helical modulation range from a few to a few hundred nanometres (Kanazawa et al., 2017; Tokura & Kanazawa, 2021), lying within the range accessible by SANS. Such a length scale is also inherited by topological skyrmions that form out of helical order under a small magnetic field \( H \) and near the critical temperature \( T_c \). Since fluctuations are crucial for skyrmion phase stability (Mühlbauer et al., 2009; Kruchkov et al., 2018), the equilibrium skyrmion phase is correspondingly narrow in temperature span, being just a few percent of \( T_c \) wide directly below \( T_c \). Otherwise, the non-topological helical or field-induced helical cone (‘conical’) phases dominate the rest of the phase diagram (Kanazawa et al., 2017; Tokura & Kanazawa, 2021).

Recently, Co–Zn–Mn compounds have emerged as a novel class of skyrmion-hosting chiral magnets (Tokunaga et al., 2015; Karube et al., 2016, 2017, 2018, 2020; Henderson et al., 2021). This system crystallizes in the \( \beta \)-Mn chiral cubic structure with either a \( P4_1/2m \) or \( P4_1220 \) space group, and with 20 atoms distributed across the 8\( c \) and 12\( d \) Wyckoff sites [

![Figure 1](https://example.com/figure1.png)

**Figure 1**

(a) The two \( \beta \)-Mn-type crystal enantiomers of Co0.75Zn0.25Mn1.5 viewed along the [111] direction. The 8\( c \) and 12\( d \) Wyckoff sites are coloured according to their average occupation of Co (blue), Zn (green) and Mn (red). (b) The low-temperature \( (T) \) versus Mn concentration \( x \) in \( \text{Co}_x\text{Zn}_{1-x}\text{Mn}_{1.5} \), for \( 0 \leq x \leq 8 \), determined from magnetization measurements. Circle and triangle symbols denote magnetic transitions identified previously from measurements on polycrystalline samples (Karube et al., 2018). Star symbols correspond to new data obtained from the Co0.75Zn0.25Mn1.5 crystals studied here. The shaded purple region denotes a \( T \) range for which the magnetization at \( H = 20 \text{ Oe} \) decreases, and concomitantly the helical wavevector \( q \) observed by SANS increases, for decreasing \( T \). The square (diamond) symbols indicate temperatures where \( q \) starts increasing (stopping increasing) on cooling, as denoted by \( T_q (T_d) \). (c) The temperature dependence of the field-cooled (FC) and zero-field-cooled (ZFC) d.c. magnetization in single-crystal \( \text{Co}_{0.75}\text{Zn}_{0.25}\text{Mn}_{1.5} \). (d) The temperature and frequency dependence of, respectively, the real \((\chi')\) and imaginary \((\chi'')\) parts of the a.c. susceptibility of \( \text{Co}_{0.75}\text{Zn}_{0.25}\text{Mn}_{1.5} \) in the low-\( T \) range.

Introduction of Mn leads to a rapid suppression of \( T_c \) to far below room temperature [Fig. 1(b)] and a decrease in the helical periodicity near \( T_c \) – and hence in skyrmion size – from \( \lambda \simeq 150 \text{ nm} \) below the transition temperature \( T_c \simeq 420 \text{ K} \) [beyond the upper temperature limit of Fig. 1(b)] and form an equilibrium skyrmion phase that is stable far beyond room temperature (Karube et al., 2020). In this low-\( x \) limit, the general paradigm of DMI chiral magnetism observed in structurally simpler chiral cubic magnets like the \( B20s \) broadly applies (Kanazawa et al., 2017).

![Image](https://example.com/figure2.png)
SANS, and indicates a reduction in the ratio of $J/D$. This behaviour is interpreted as arising due to an onset of anti-ferromagnetic-like correlations between Mn moments on cooling that affect the ferromagnetic interactions and DMI between Co (Karube et al., 2018, 2020). At lower temperatures below $T_c$, a reentrant spin glass transition $T_g$ is observed for $3 < x < 6$, which is known to involve the 8c helical spin correlations (Karube et al., 2020). The origin of the spin-glass-like properties can be traced from the physics of the other end member, $\beta$-Mn ($x = 20$), a well known elemental spin liquid (Nakamura et al., 1997). In its pure form, $\beta$-Mn displays no spin glass transition, and the spin liquid property is due to the strong geometric frustration of antiferromagnetically coupled local Mn moments residing on the hyper-kagomé-coordinated 12d site. Spin-glass-like transitions emerge in lightly doped $\beta$-Mn alloys, due to combined geometric frustration and compositional disorder (Nakamura et al., 1997; Stewart et al., 2002, 2008, 2010; Stewart & Cywinski, 2009). In the presently studied composition line, no helical transition is seen for $6.5 < x < 19$, and instead only a spin-glass-like transition $T_g$ is observed (Karube et al., 2018).

Overall, the phase diagram of the present class of Co–Zn–Mn chiral magnets combines various key ingredients in a single system, namely mainly Co DMI chiral magnetism on the 8c site, magnetic frustration mainly due to antiferromagnetic correlations between Mn spins on the 12d site, magnetic anisotropy that varies with $T$ and $x$ (Preißinger et al., 2021), and compositional disorder. Tuning the composition controls the relative influence of each aspect and has thus far led to a series of novel findings. These include the observation of equilibrium skyrmion phases near $T_c$, both far above and far below room temperature (Tokunaga et al., 2015), and the facile disorder-assisted creation of supercooled metastable skyrmion states (Karube et al., 2016, 2017, 2018, 2020; Morikawa et al., 2017). These metastable skyrmion states are practically infinitely long lived, survive over the majority of the phase diagram, and display novel and reversible coordination transformations that preserve the topological charge carried by the skyrmions. Finally, in Co$_x$Zn$_y$Mn$_z$ the chiral magnetism and magnetic-frustration-induced fluctuations conspire to stabilize a second field-induced equilibrium skyrmion phase for temperatures just above $T_g$ (Karube et al., 2018; Ukleev et al., 2021).

In this study we explore the helical, skyrmionic and short-range magnetic correlations in a new composition Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ with $x = 6.5$ by SANS, magnetometry, magnetic diffuse neutron scattering and Lorentz transmission electron microscopy (LTEM) measurements. As seen in Fig. 1(b), the transitions at $T_c$ and $T_g$ are closer together for this composition compared with $3 < x < 6$, indicating that magnetic frustration and disorder compete more strongly with DMI chiral magnetism in this system. At the same time, in terms of a reduced temperature $\Delta T/T_c$, the thermal extent of the equilibrium skyrmion phase has been found to grow progressively with $x$ from 0.016 to 0.031 to 0.050 to 0.147 for $x = 0, 2, 4$ and 6, respectively (Karube et al., 2020), indicating an enhanced stability. Moreover, for $x = 6$, a second equilibrium skyrmion phase is observed far below $T_c$ (Karube et al., 2018; Ukleev et al., 2021), in accord with an enhanced stability of skyrmions. Therefore, it is important to understand how the helical magnetism and skyrmion phase stability evolve with $x$ in the presence of competing interactions.

Here, we show experimentally that, for $x = 6.5$, the effects of magnetic frustration and disorder are so strong that they almost completely suppress the formation of the skyrmion phase near $T_c$. Instead, SANS shows that the mesoscale spin correlations in the majority of the sample are dominated by strong magnetic disorder across the phase diagram. SANS nonetheless reveals that a $\sim10\%$ minority fraction of the sample does undergo a standard field-induced transition to a regime of skyrmion correlations near $T_c$ with the thermal window of stability $\Delta T/T_c \approx 0.16$. However, due to the generally strong magnetic disorder in the sample, the skyrmion correlations revealed by SANS are not clearly detectable by either magnetometry or LTEM. We discuss the results in the context of magnetic disorder induced on the 8c site through coupling with geometrically frustrated interactions on the 12d site. The important role of magnetic frustration is shown through direct measurement of short-range magnetic correlations in Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ by diffuse neutron scattering, from which characteristic couplings between Mn moments on 12d sites are estimated.

2. Experimental

2.1. Sample preparation

Bulk single crystals of Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ were prepared by the Bridgman method. Single crystals were identified using X-ray Laue diffraction and their structural quality characterized at room temperature using Cu $K\alpha$ radiation from a laboratory X-ray diffractometer (Rigaku, RINT-TTR III). Fig. 2 shows typical diffraction data obtained from the (110) structural peak of one crystal extracted from the growth. According to the $2\theta$ scan shown in Fig. 2(a) and the rocking

![Figure 2](image.png)

Figure 2. X-ray diffraction measurements of the (110) structural peak in a Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ single crystal. (a) A $2\theta$ scan of the (110) peak fitted by a single Gaussian line shape. (b) A rocking curve (i.e. $\theta$ scan) at fixed $2\theta$. The intensity distribution is fitted by two Gaussian line shapes.
2.3. Neutron scattering measurements

SANS measurements on a 57.2 mg single-crystalline sample of $\text{Co}_{6.75}\text{Zn}_{6.75}\text{Mn}_{6.5}$ were performed on the D33 beamline at the Institut Laue–Langevin (ILL), Grenoble, France (White et al., 2017, 2019). Neutrons with a wavelength of either 8 or 10 Å were collimated over a distance of 12.8 m before the sample. The scattered neutrons were detected by a two-dimensional position-sensitive multidetector positioned 12.8 m behind the sample. Data were collected over a range of momentum transfer $0.02 < q < 0.4 \text{ Å}^{-1}$ [$q = (4\pi/\lambda_n)\sin\theta$, where $\theta$ is half the scattering angle and $\lambda_n$ is the wavelength of the incident neutron beam].

For SANS the single-crystal sample was installed in a horizontal-field cryomagnet at the sample position on the beamline. As shown in Fig. 3(b), two experimental geometries were studied, namely $\mathbf{H} \parallel \mathbf{k}_i$ and $\mathbf{H} \perp \mathbf{k}_i$, where $\mathbf{k}_i$ is the incoming neutron wavevector which was chosen to be parallel to [100] for both geometries. In either geometry, the SANS measurements were done by collecting data as the cryomagnet and sample were rotated together around the vertical axis over the neutron beam access range of ±10° provided by the cryomagnet windows. Detector measurements obtained at a range of rotation angles were summed together to produce a single image showing the distribution of magnetic scattering. Unless otherwise stated, SANS data were obtained at each temperature and magnetic field after an initial zero-field cooling through the critical temperature $\sim$102 K, and a subsequent $H$ ramp once at the target temperature. Further data obtained either in the paramagnetic regime at 120 K or in a field-polarized regime were used for background subtraction of the low-$T$/low-$H$ data to leave just the magnetic SANS signal of interest. SANS data reduction and analysis were performed using the GRASP software developed at the ILL (Dewhurst, 2003).

Magnetic diffuse neutron scattering (MDNS) experiments on a 17.9 g polycrystalline sample of $\text{Co}_{6.75}\text{Zn}_{6.75}\text{Mn}_{6.5}$ were carried out on the D7 beamline at the ILL (White et al., 2018). An $xyz$ polarization analysis allowed pure isolation of the magnetic scattering cross section from the total scattering (Stewart et al., 2009). The sample was loaded into a double-walled Al container and installed in a standard Orange He cryostat. For the experiment, an incoming neutron wavelength of 3.1 Å was used, and data were collected over a range of momentum transfer $0.6 < q < 3.8 \text{ Å}^{-1}$ (or equivalently $6 < q < 38 \text{ nm}^{-1}$).

2.4. LTEM measurements

LTEM measurements were performed with a transmission electron microscope (JEOL JEM-2800). A magnetic field was applied perpendicular to the plate, i.e. parallel to [001], and its

For the LTEM measurement, a thin plate sample with a 150 nm was prepared from a 17.9 g polycrystalline sample of $\text{Co}_{6.75}\text{Zn}_{6.75}\text{Mn}_{6.5}$ were carried out on the D7 beamline at the ILL (White et al., 2018). An $xyz$ polarization analysis allowed pure isolation of the magnetic scattering cross section from the total scattering (Stewart et al., 2009). The sample was loaded into a double-walled Al container and installed in a standard Orange He cryostat. For the experiment, an incoming neutron wavelength of 3.1 Å was used, and data were collected over a range of momentum transfer $0.6 < q < 3.8 \text{ Å}^{-1}$ (or equivalently $6 < q < 38 \text{ nm}^{-1}$).
magnitude was controlled by tuning the objective lens current of the microscope.

2.5. Demagnetization calibration
For single-crystal measurements, the relative demagnetization factors between measurement configurations with \( \mathbf{H} \) parallel to the sample plate (d.c. and a.c. susceptibility, and SANS in the \( \mathbf{H} \perp \mathbf{k} \), geometry) and \( \mathbf{H} \) perpendicular to the sample plate (SANS in the \( \mathbf{H} \parallel \mathbf{k} \), geometry) are different. To cater for the difference in demagnetization, data presented throughout this article with \( \mathbf{H} \) parallel to the sample plate have had the field scale calibrated higher so that \( H' = H \times N \), where \( N = 3.0 \).

3. Results and discussion
3.1. Magnetometry measurements
Fig. 1(c) shows the temperature \( (T) \)-dependent magnetization \( (M) \) in single-crystal Co\(_{0.75}\)Zn\(_{0.75}\)Mn\(_{6.5}\). The data are obtained in a small magnetic field of 20 Oe, either on field cooling (FC) or on field warming after an initial zero-field cooling (ZFC). In the high-\( T \) region for both curves, \( M(T) \) clearly increases (decreases) on the FC (ZFC) process. A transition temperature at \( T_c \approx 102 \text{ K} \) between helical and paramagnetic regimes is estimated from the inflection point of \( M_{ZFC}(T) \), though it is noted that the transition is smeared over a broad \( T \) range compared with lower-\( x \) compounds (Karube et al., 2020). Below 80 K, \( M_{FC}(T) \) falls with decreasing \( T \), and a clear separation between the \( M_{FC}(T) \) and \( M_{ZFC}(T) \) curves at \( \approx 65 \text{ K} \) can be interpreted as the onset of spin-glass-like behaviour.

To investigate the low-\( T \) behaviour further, Figs. 1(d) and 1(e) show the temperature and frequency dependence of, respectively, the real \( (\chi') \) and imaginary \( (\chi'') \) parts of the a.c. susceptibility. A frequency dependence of \( \chi' \) and \( \chi'' \) is a common spin glass characteristic, though in the present sample it is modest and exists over a broad thermal range. At the same time, no thermally sharp spin glass transition is seen. Therefore, while the material displays some tendency towards glassy behaviour at low temperature, further investigations are needed to determine if the low-temperature regime can be properly allocated as a spin glass. Here, for consistency with previous work on lower-\( x \) compounds (Karube et al., 2020), in Figs. 1(b) and 1(c) we use the label \( T_g \) to denote the temperature where \( M_{ZFC}(T) \) shows an inflection point, and thus the temperature where spin-glass-like behaviour is most clearly inferred in the data. For the present compound \( T_g \approx 53 \text{ K} \) can be compared with \( \approx 65 \text{ K} \), which is where the magnetization hysteresis vanishes.

Overall, the bulk magnetic measurements are hallmarked by transition features – labelled \( T_g \) and \( T_c \) – that are thermally smeared. This contrasts strongly with sharper transitions observed at lower \( x \), including the nearby composition Co\(_{0.75}\)Zn\(_{0.75}\)Mn\(_{6} \) (\( x = 6 \)), indicating comparatively stronger disorder of the magnetically correlated phases in the present \( x = 6.5 \) composition which is likely to be manifested in the low-\( T \) magnetic spin configurations.

3.2. SANS measurements of helical and skyrmion correlations
3.2.1. Helical disordering during zero-field cooling. To shed more light on the magnetometry data, we next turn to SANS investigations of the mesoscale magnetic spin correlations in a Co\(_{0.75}\)Zn\(_{0.75}\)Mn\(_{6.5}\) single crystal. Figs. 3(c)–3(g) show SANS patterns collected at selected temperatures during ZFC. Just below \( T_c \) at 100 K, the pattern shown in Fig. 3(c) is composed of two peaks with propagation vectors aligned with the [010] direction co-existing with an isotropic ring of scattering intensity. The intensity ring is indicative of incommensurate helical spin correlations with full orientational disorder, while the peaks are more typical for an incommensurate helical order with finite orientational order. The alignment of a helical propagation vector with the cubic axis is consistent with that observed in other Co–Zn–Mn compounds with finite Mn content (Karube et al., 2016, 2017, 2018, 2020), perhaps indicating that the sign of magnetic anisotropy in \( x = 6.5 \) is the same as that observed for \( 2 \leq x \leq 6 \) (Karube et al., 2020; Preißinger et al., 2021). According to the cubic symmetry, additional peaks would be expected in this scattering plane due to a helical domain with propagation vectors aligned with the [001] direction. The observation of only two peaks indicates a preferential helical domain selection effect, which may be due to a residual sample strain.

On cooling to lower \( T \), the two peaks seen at 100 K quickly broaden azimuthally, so that by 80 K the full decay of the orientational order of all helical spin correlations in the sample is complete and the entire SANS intensity appears as a ring around the origin of reciprocal space. On cooling further to the base temperature of 1.5 K, the intensity of the ring becomes weaker compared with higher \( T \) and is distributed over an extended range in \( q \), thus reflecting a reduction in the radial correlation length. At all temperatures, the correlation length normal to the detector plane determined by sample rotation-angle-dependent (i.e. rocking curve) measurements remains nearly angle independent (data not shown). Overall, the ZFC data show that the sample always displays mainly three-dimensionally disordered helical spin correlations, with a clear reduction in the radial correlation length taking place on cooling.

With Fig. 4, we analyse more quantitatively the magnetic SANS intensity observed during ZFC. Fig. 4(a) shows the radial \( |q| \) dependence of the SANS intensity integrated over all azimuthal angles. At all temperatures, the profiles are fitted well by a Lorentzian function, indicating the exponential decay of the radial correlation function with distance. The fitted function at each \( T \) is \( I(q) = I_0 + (2A/\pi r)[w(4(q - q_0)^2 + w^2)] \), where \( I_0 \) is a fitted constant background, \( A \) is the fitted integrated intensity of the peak, \( w \) is the fitted peak FWHM and \( q_0 \) is the fitted peak centre. The temperature dependence of parameters \( A \), \( w \) and \( q_0 \) is shown in Figs. 4(b)–4(d).

From Fig. 4(b) we observe the integrated intensity \( A \) to rise initially on cooling below \( T_c \), before decreasing over the range from \( \approx 80 \) to \( \approx 40 \text{ K} \) and then remaining approximately constant down to 1.5 K. The reduction in \( A \) over the intermediate-temperature range largely coincides with significant enhancements of both \( q_0 \) [Fig. 4(c)] and \( w \) [Fig. 4(d)]. Finally,
A dependence of are needed for the accurate determination of the temperature investigations designed to measure the full scattering intensity extend far beyond the experimentally accessible range the intensity distribution normal to the detector plane to short longitudinal magnetic correlation lengths, which cause it was not possible to measure the latter due the inherently measurement of the scattering intensity distribution both the temperature dependence of implied by our analysis, but we stress that a full treatment of development of antiferromagnetic-like correlations on cooling.

Turning to the fitted values of \( w \) shown in Fig. 4(d), they are seen to be always significantly larger than the instrumental resolution, allowing a simple estimation of the radial correlation length using \( \xi \approx 1/\omega \). We find that the radial correlation length near to \( T_c \) is \( \sim 25 \text{ nm} \), i.e. of the order of 40 cubic unit-cell spacings, before it falls on cooling to \( \sim 7 \text{ nm} \) below \( T_g \), which is just above 10 cubic unit-cell spacings. It is notable that \( \xi \) is always shorter than \( \lambda \), which indicates that the helical spin correlations are typically only short-range ordered. Such a tendency towards helical short-range order can reflect competing magnetic interactions on the mesoscale in this system, while we also note that, according to theory, defect-induced disorder can contribute further to the broadening of the helical Bragg peak (Utesov et al., 2015).

The general behaviour observed in the ZFC temperature dependence of the \( A, w \) and \( q_0 \) parameters in \( \text{Co}_{6.75}\text{Zn}_{6.75}\text{Mn}_{6.5} \) \((x = 6.5)\) bears qualitative similarity to that observed in the \( 2 < x \leq 6 \) compounds (Karube et al., 2016, 2017, 2018, 2020). In comparison with the lower-\( x \) compositions however, the magnetic disorder and disordering process on ZFC in the present \( x = 6.5 \) sample is clearly more pronounced. Firstly, by 80 K, the minority sample fraction of helical-like order showing up as SANS peaks at 100 K undergoes a full loss of orientational order by 80 K. The full loss of helical orientational order is not observed in any sample from 0 \( \leq x \leq 6 \) on ZFC down to base temperature. Secondly, this is followed by the strongest reduction in the radial correlation length amongst the studied compositions (Karube et al., 2016, 2017, 2018, 2020). The present results thus extend to \( x = 6.5 \) the empirically observed inverse correlation between the increase in Mn concentration \( x \) and the stability of helical order at low \( T \) for \( 0 < x < 6 \). In addition, the present data imply that the \( x = 6.5 \) sample provides a future opportunity for detailed studies of an unusual melting of the helical correlations on cooling from \( T_c \) to \( T_g \), and potentially spiral spin liquid characteristics (Gao et al., 2017).

3.2.2. Skyrmion correlations in a finite magnetic field. Next, we turn to measurements in a finite magnetic field. Figs. 5(a) and 5(b) show SANS patterns obtained in the \( \mathbf{H} \parallel \mathbf{k}_i \) geometry at 90 K, and in zero field and at 18 mT, respectively. The pattern in zero field is consistent with that shown in Fig. 3(d), where the SANS intensity forms a ring-like structure with more intensity on the right- and left-hand sides due to azimuthally broadened helical spots. Application of the magnetic field leads to a homogenization of the intensity distribution such that it forms a near isotropic intensity ring by 18 mT [Fig. 5(b)]. As will be supported from measurements in the \( \mathbf{H} \perp \mathbf{k}_i \) geometry, this \( H \)-driven rearrangement of the intensity is consistent with a field-induced transition of helical spin correlations into orientationally disordered skyrmion lattice correlations, as opposed to an \( H \)-driven disordering process.

Fig. 5(c) shows the magnetic field dependence of the summed SANS intensities within the two top/bottom (blue) 90° sectors and the two left/right (red) 90° sectors for the \( \mathbf{H} \parallel \mathbf{k}_i \) geometry. Closed (open) symbols denote \( H \)-increasing (\( H \)-decreasing) measurements with little intensity hysteresis.
observed. In zero field, the intensity in the red sector box pair is only 12 (2)% larger than that in the blue sector box pair, showing that the majority of the observed SANS intensity is due to the ring-like intensity component. The difference between the red and blue curves falls monotonically as the field is increased, eventually forming a fully azimuthally isotropic intensity distribution that survives until saturation.

Figs. 5(d)–5(e) show the SANS data obtained at 90 K in the \( \mathbf{H} \perp \mathbf{k} \), geometry, in zero field and at 18 mT. In this geometry, the pattern obtained at 18 mT shows the magnetic field to drive the partial rearrangement of the ring-like intensity in zero field so that there are two clear intensity maxima aligned with the vertical [001] direction. Since, in chiral cubic magnets, the skyrmion lattice forms with propagation vectors distributed in the plane perpendicular to the magnetic field, the emergence of additional top/bottom SANS intensity seen clearly in Fig. 5(e) corresponds to a hallmark signature for two-dimensional skyrmion lattice correlations (Mühlbauer et al., 2009; Karube et al., 2016). In Fig. 5(f) the formation of the skyrmion tubes aligned with \( \mathbf{H} \) is manifested as a magnetic field range for which the combined intensity in the two top/bottom (blue) 90° sectors is maximally \(~20\)% larger than those of the two left/right (green) 90° sectors. On further increase in the field, the relative intensity of the green sector box pair becomes larger than that of the blue sector box pair before saturation. This can be understood in terms of the generally enhanced susceptibility of helical (or conical) spin correlations when the field is applied in the direction of propagation. Therefore, the SANS intensity due to helical spin correlations propagating relatively close to the applied field direction survives to higher fields compared with helical and skyrmion correlations propagating along directions far away from the direction of \( \mathbf{H} \).

By 70 K, no magnetic-field-driven rearrangement of the SANS intensity is observed that is consistent with a conventional skyrmion formation signature in chiral magnets. Fig. 6 shows the relevant SANS data and data analysis at this temperature, which are analogous to those shown in Fig. 5 obtained at 90 K. In the \( \mathbf{H} \parallel \mathbf{k} \) geometry, Figs. 6(a) and 6(b), respectively, show the SANS patterns obtained in zero field and at 28 mT. In each case, the azimuthal distribution of the SANS intensity appears isotropic around the origin, with this confirmed at all fields in the \( H \) scans by the quantitative analysis presented in Fig. 6(c). For the \( \mathbf{H} \perp \mathbf{k} \), geometry, Figs. 6(d) and 6(e) show that the intensity ring in zero field evolves with increasing \( H \) so that there is slightly more intensity on the left- and right-hand sides at 30 mT. This is borne out by the quantitative analysis shown in Fig. 6(f), which shows that as \( H \) increases the SANS intensities extracted from the green left/right sector boxes [inset Fig. 6(f)] remain higher until saturation. Similar to the behaviour at 90 K, this can be understood in terms of the higher susceptibility for helical correlations propagating closer to the direction of \( \mathbf{H} \). Importantly, the analysis shows that, by 70 K, no standard signature of equilibrium skyrmion formation is detected, and the magnetic state is always predominantly

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**Figure 5**
SANS patterns obtained in the \( \mathbf{H} \parallel \mathbf{k} \), geometry at \(-90 \text{ K}\). (a) in zero field and (b) at 18 mT. (c) The \( H \) dependence of the SANS intensities integrated over the 90° wide sector boxes indicated by the inset. Blue (red) symbols correspond to the sum of intensities in the two blue (red) 90° sectors. (d), (e) SANS patterns obtained at \( H = 0 \) and 18 mT, respectively, for the \( \mathbf{H} \perp \mathbf{k} \), geometry at 90 K. (f) The \( H \) dependence of the SANS intensities integrated over the 90° wide sector boxes indicated by the inset. Blue (green) symbols correspond to the sum of intensities in the two blue (green) 90° sectors. In panels (c) and (f), closed (open) symbols denote data obtained in an \( H \)-increasing (\( H \)-decreasing) scan. For the \( \mathbf{H} \perp \mathbf{k} \), geometry, the magnetic field scale is corrected for demagnetization effects and calibrated according to \( H^\prime = H \times 3.0 \).

**Figure 6**
SANS patterns obtained in the \( \mathbf{H} \parallel \mathbf{k} \), geometry at \(~70 \text{ K}\). (a) in zero field and (b) at 28 mT. (c) The \( H \) dependence of the SANS intensities integrated over the 90° wide sector boxes indicated by the inset. Blue (red) symbols correspond to the sum of intensities in the two blue (red) 90° sectors. (d), (e) SANS patterns obtained at \( H = 0 \) and 30 mT, respectively, for the \( \mathbf{H} \perp \mathbf{k} \), geometry at 90 K. (f) The \( H \) dependence of the SANS intensities integrated over the 90° wide sector boxes indicated by the inset. Blue (green) symbols correspond to the sum of intensities in the two blue (green) 90° sectors. In panels (c) and (f), closed (open) symbols denote data obtained in an \( H \)-increasing (\( H \)-decreasing) scan. For the \( \mathbf{H} \perp \mathbf{k} \), geometry, the magnetic field scale is corrected for demagnetization effects and calibrated according to \( H^\prime = H \times 3.0 \).
three-dimensionally disordered. Further measurements confirm that this picture prevails down to the base temperature.

To summarize the $H$-dependent measurements on single-crystal $\text{Co}_{6.75}\text{Zn}_{6.75}\text{Mn}_{6.5}$, in Fig. 7 we present phase diagrams constructed from the real part of the a.c. magnetic susceptibility $\chi$ [Figs. 7(a) and 7(b)] and SANS [Figs. 7(c) and 7(d)] data. Figs. 7(a) and 7(b) show phase diagrams constructed from, respectively, $H$-increasing and $H$-decreasing scans done at constant temperature. These two phase diagrams are very similar, and bear little resemblance to those of either low-$x$ ($\text{Co}_{0.2}\text{Zn}_{0.8}$)$_{20-x}\text{Mn}_x$ compounds (Karube et al., 2020) or archetypal chiral magnets such as MnSi (Mühlbauer et al., 2009) and Cu$_2$OSeO$_3$ (Seki et al., 2012). In the latter-mentioned systems, $\chi$ measurements are particularly useful for identifying the extent in $T$ and $H$ of different phases (Bauer & Pfleiderer, 2012; Birch et al., 2020). In the present case, the most striking features are the $H$-smeared low-field transitions at $T_c$ and $T_g$, the vanishing of $\chi'$ for $T < T_g \simeq 50$ K, and the monotonic $H$ variation of $\chi'$ in the region of $T_g < T < T_c(H)$. In contrast to clear signatures in $\chi'$ revealing magnetic phase transition boundaries in chiral magnets like the B20s, no fine structure in $\chi'$ is identifiable from the present sample, and none indicating skyrmion formation. Instead, $\chi'$ is presumably dominated by the temperature evolution of the majority fraction of disordered helical spin correlations, and the spin-glass-like properties at low $T$.

In contrast, the signature for skyrmion formation shows up more clearly in phase diagrams constructed from SANS data obtained in the $\textbf{H} \perp \textbf{k}$ geometry, for both $H$-increasing [Fig. 7(c)] and $H$-decreasing [Fig. 7(d)] scans at constant temperature. The colour scale describes the ratio of summed SANS intensities measured in the blue top/bottom sector boxes [denoted $I(\varphi = 0^\circ)$] to that determined from the green left/right sector boxes [denoted $I(\varphi = 90^\circ)$]. In both phase diagrams, and in the field-reversible region $T_g < T < T_c(H)$, the red portion of the colour plots indicates the excess of intensity in the top/bottom sectors, providing the clearest indication for the thermodynamic conditions under which skyrmions may form in the system. The location of this red portion of the phase diagram is similar to that for the conventional two-dimensional skyrmion $A$ phase in chiral cubic magnets, namely occupying a narrow temperature region directly below $T_c$ and at finite $H$. In quantitative terms, the thermal extent of the $A$ phase in reduced temperature $T/T_c$ is $\sim 0.16$, which is comparable to that in $\text{Co}_2\text{Zn}_2\text{Mn}_6$ (Karube et al., 2018), but the intensity ratio between the blue and green pairs of integration sectors in the ‘skyrmion phase’ is only $\sim 1.2$ and the majority of the SANS intensity nonetheless remains attributable to the disordered helical correlations. From this perspective, the skyrmion phase is largely suppressed in the present $\text{Co}_{6.75}\text{Zn}_{6.75}\text{Mn}_{6.5}$ sample compared with lower $x$ compositions. Apart from the minority skyrmion phase, disordered helical configurations dominate the phase diagram above $T_g$ and for fields below saturation, with the green colours indicating ratios between blue and green sector box intensities that are closer to 1. Finally, in the region below $T_g$, there is a pronounced irreversibility in the intensity ratio defined by the colour scale between the $H$-increasing and $H$-decreasing phase diagrams, consistent with that expected from a spin-glass-like regime.

3.3. Lorentz transmission electron microscopy

As it is one of the key experimental tools for studying helical and skyrmion spin textures in chiral magnets (Kanazawa et al., 2017; Huang et al., 2018; Tokura & Kanazawa, 2021), we employed LTEM with the aim of direct visualization of the magnetic textures below $T_c$ in a thin plate sample of $\text{Co}_{6.75}\text{Zn}_{6.75}\text{Mn}_{6.5}$. Fig. 8(a) shows the phase diagram of Fig. 7(c), with labels indicating the $T$ and $H$ conditions under which the LTEM images shown in Figs. 8(b)–8(d) were obtained. In all images, neither periodic magnetic textures nor (isolated) topological skyrmions can be identified. This indicates the magnetic contrast to be extremely weak, which is presumably due to the strong magnetic disorder, not only within the plane but also along the sample thickness. This contrasts with LTEM studies on lower-$x$ compounds, including $x = 6$, where unambiguous imaging of helical order and skyrmions has been performed despite the existence of moderate magnetic disorder (Tokunaga et al., 2015; Morikawa et al., 2017; Karube et al., 2017, 2018).
3.4. Magnetic diffuse neutron scattering

Finally, we turn to magnetic diffuse neutron scattering measurements from a Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ powder sample taken using the D7 spectrometer at the ILL. With these measurements we explore magnetic neutron scattering at higher momentum transfers than afforded by SANS, which extend down to atomic length scales. In particular, we seek experimental evidence for persistent short-range magnetic correlations. Earlier work on $\beta$-Mn-type samples shows structured magnetic diffuse scattering to be a common experimental signature for short-range correlations of Mn moments on the 12$d$ site (Stewart et al., 2008, 2009, 2010; Stewart & Cywinski, 2009; Paddison et al., 2013). The origin of the scattering is ascribed to the strong geometric frustration of interacting Mn moments on the 12$d$ site with its frustration-inducing hyper-kagomé geometry. It is notable that qualitatively similar magnetic diffuse scattering was reported from measurements on the Co$_2$Zn$_3$Mn$_6$ ($x = 6$) compound (Ukleeve et al., 2021). Since the average 12$d$ occupation for $x = 6$ is Co:Zn:Mn = 0.2:7.0:4.8 (Nakajima et al., 2019), it becomes clear that only partial average occupation of the 12$d$ site by magnetic Mn ions is needed for magnetic diffuse scattering to show up. In this context it can be expected that a similar experimental signature is observed for the present $x = 6.5$ composition, since the average Mn content of the 12$d$ site is only slightly higher, namely Co:Zn:Mn = 0.25:6.75:5.0 (Nakajima et al., 2019).

Using the $xyz$ polarization analysis method on D7, the nuclear coherent, magnetic and nuclear spin-incoherent scattering cross sections are straightforwardly and cleanly isolated from the total scattering observed (Stewart et al., 2009). Fig. 9(a) shows these various scattering cross sections measured from Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ powder as a function of modulus wavevector $|q|$ at 2 K and zero magnetic field. The nuclear spin-incoherent scattering intensity displays no $|q|$ dependence as expected, while in the nuclear coherent scattering both structural diffraction peaks and diffuse scattering due to structural disorder are observed. Since the studied $q$ range is too limited to extract detailed information on the origin of the structural diffuse scattering, in what follows we focus on the magnetic diffuse scattering.

Fig. 9(b) shows a closer view of the observed magnetic diffuse scattering cross section versus $|q|$. The scattering is composed largely of a peak that is broad in momentum...
transfer around $|q| \approx 1.7 \text{ Å}^{-1}$, with additional weak peak-like features that are consistent with being due to the disordered helical correlations observed by SANS. For further data treatment and analysis of the magnetic diffuse scattering, we follow the same approach reported for $x = 6$ (Ukleev et al., 2021). Namely, the weak peaks are removed and the remaining diffuse scattering is fitted to a mean-field model that accounts for the paramagnetic scattering due to interactions between Mn–Mn moments occupying the 12$d$ site. Further technical details for the calculation method are reported by Ukleev et al. (2021). Here we mention that, for simplicity, similarly to the details for the calculation method are reported by Ukleev (2021), an average crystal structure model is assumed, as well as a 12$d$ site that is fully occupied by magnetic moments. The blue line in Fig. 9(b) shows the fit according to this minimal model that considers just a single nearest-neighbour antiferromagnetic exchange constant with fitted $J_1 = -0.53$ meV, and a non-interacting temperature-dependent spin susceptibility. The reasonable fit of the model makes it plausible that the diffuse scattering is due to frustrated nearest-neighbour antiferromagnetic Mn–Mn interactions on the 12$d$ site. The fitted value of $J_1$ is essentially identical to that found for $x = 6$ ($J_1 = -0.54$ meV), consistent with the observation that the diffuse scattering profiles from the two compounds are quantitatively similar.

Inclusion in the model of a second nearest-neighbour ferromagnetic interaction $J_2$ leads to an improvement in the fit, as shown by the black dashed line in Fig. 9(b). The fit yields $J_1 = -0.67$ meV and $J_2 = 0.67$ meV, which have the same magnitude but opposite sign, thus implying that the proper description for the distribution of magnetic diffuse scattering involves a combination of competing antiferromagnetic ($J_1$) and ferromagnetic ($J_2$) interactions. While future studies on single-crystal samples are needed to quantify the interactions more rigorously, at the level of the data and analysis shown here, it is reasonable to conclude that the origin of magnetic diffuse scattering involves a major contribution from nearest-neighbour antiferromagnetic interactions between Mn moments on the 12$d$ site.

4. Discussion

Our neutron scattering and magnetometry measurements reveal that the interactions and compositional disorder in the chiral magnet Cu$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ conspire to generate magnetically disordered phases at low temperature, over both atomic and mesoscopic length scales. As deduced in previous studies of Co–Zn–Mn compounds, mesoscale helices and skyrmions originate from the chiral magnetism of spins occupying the 8$c$ site, involving mainly ferromagnetic Co ions (Karube et al., 2016, 2017, 2018, 2020; Bocarsly et al., 2019; Ukleev et al., 2019, 2021). For $0 < x < 3$, antiferromagnetic Mn fills in exclusively onto the geometrically frustrated 12$d$ site, while for $x > 3$ Mn starts to fill in and replace Co on the 8$c$ site too (Nakajima et al., 2019). As borne out by the sharp suppression of $T_c$, the emergence of spin glass behaviour and the tendencies towards magnetic disorder, adding Mn is generally unfavourable for mesoscale chiral magnetism. At the same time, however, the reduced temperature ($T/T_c$) extent of the high-temperature skyrmion phase progressively increases with $x$ for the $3 < x < 6$ compounds, implying that Mn substitution effectively enhances skyrmion phase stability within the phase diagram (Karube et al., 2020). In the following, we discuss the origin of the disordered mesoscale magnetism and the suppression of the equilibrium high-temperature skyrmion phase in the $x = 6.5$ sample.

As seen in Fig. 1(b), compared with lower-$x$ compounds, an important aspect of the present $x = 6.5$ system is that the characteristic temperatures $T_c$ and $T_g$ are closer together, which already indicates a closer competition between chiral magnetism on the one hand, and atomic-scale frustration and disorder on the other. In addition, the location of $T_c$ itself is also important in relation to the characteristic temperatures $T_{H1}$ and $T_{L1}$, which bound the temperature range over which the helical $q$ and degree of mesoscale disorder both increase on cooling for $3 < x < 6$. Both of these effects are understood to reflect the developing role of correlations involving Mn moments on both the 8$c$ and 12$d$ sites (Karube et al., 2020). For $x = 6.5$, $T_c$ already lies between the values of $T_{H1}$ and $T_{L1}$ that would be expected for this composition according to an extrapolation of their lower $x$ dependences. In turn, this implies that, when helical correlations begin to form, Mn spin correlations detrimental to long-range order are already present, and thus mesoscale magnetism that is largely only short-range ordered is observed, together with the near-complete suppression of the high-temperature skyrmion phase.

Overall, the data across all low-$x$ compositions indicate that when the Mn spin correlation onset temperature $T_{H1} < T_c$, i.e. for $3 < x < 6$ where helical long-range order occurs on cooling below $T_c$, progressive introduction of Mn enhances high-temperature skyrmion phase stability in terms of $T/T_c$, despite a concomitant increase in mesoscale disorder. This tendency ends when $T_c < T_{H1}$, and pre-existing inter-site Mn spin correlations dominate the helical ordering process such that skyrmions can no longer form. We also stress that the suppression of the skyrmion phase for $x = 6.5$ could not be expected by merely examining the $x$ dependence of $T_c$ and $T_g$ alone, with SANS measurements proving crucial to obtain the necessary microscopic picture.

Next we discuss the mesoscale magnetic disorder in Cu$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ ($x = 6.5$). As mentioned above, our SANS data show the mesoscale magnetism to be dominated always by a majority fraction of fully orientationally disordered helical spin correlations. On ZFC below $T_c$, further magnetic disordering via the reduction of the radial correlation length takes place, which is quantitatively stronger than that observed in lower-$x$ compounds, including the nearly composition Co$_{7}$Zn$_{7}$Mn$_{6}$ ($x = 6$) (Karube et al., 2018). While a detailed description of the ZFC helical disordering process presents an interesting challenge for theory, we point out that a key ingredient must lie with the degree of Co–Mn mixing on the 8$c$ site. According to neutron diffraction, the average 8$c$ occupation for Cu$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ is Co:Zn:Mn = 6.5:0:1.5, while for Co$_{7}$Zn$_{7}$Mn$_{6}$ it is Co:Zn:Mn = 6.8:0.1:2 (Nakajima et al.
In general, for the 8c site containing a majority Co/ minority Mn mixture, the ferromagnetic Co exchange field can be expected to drive a ferromagnetic alignment of small ordered Mn moments just below $T_c$, so that these Mn moments do indeed contribute to helically modulating spin correlations. This picture has been confirmed in both Co$_6$Zn$_6$Mn$_4$ (Ukleev et al., 2019) and Co$_7$Zn$_6$Mn$_6$ (Ukleev et al., 2021) by element-selective resonant X-ray scattering measurements. On cooling below $T_{11}$, the development of couplings between growing 8c Mn moments and erstwhile magnetically disordered 12$d$ Mn moments competes with the Co exchange field. This will result in disordering due to the random deviation of 8c Mn moments on cooling, as they tend towards a disordered ground state native to very Mn-rich compounds. In this qualitative picture, the tendency towards increasingly disordered helical textures will clearly be stronger in Mn-richer Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ compared with lower-x compounds, consistent with SANS observations (Karube et al., 2016, 2018). Further contributions to the mesoscale disorder are also expected to arise due to the compositional disorder which will modify locally the atomic scale interactions such as the DMI. Theory shows that such a bond disorder can distort helical and skyrmion correlations in the chiral magnet. In general, muon spin relaxation experiments detect pronounced megahertz magnetic dynamics in and around the skyrmion phases of a number of chiral magnets (Hicken, Holt et al., 2021; Hicken, Wilson et al., 2021; Ukleev et al., 2021). This makes the development of theoretical models for their origin at low temperature important, in order to test suggestions that they provide a route towards skyrmion phase stability.

Finally, we comment on the future perspectives for neutron scattering from quantum materials such as Co$_{6.72}$Zn$_{6.72}$Mn$_{6.5}$. In the present system, we observe both structural and magnetic correlations with short- and long-range ordered character, over a broad range of momentum transfer that includes the conventional SANS range. It is clear that in complex quantum materials such as the Co–Zn–Mn intermetallics the eventual development of accurate theoretical models requires comprehensive measurement of structural and magnetic correlations over multiple length scales. While in principle the relevant scattering data can be pieced together from measurements done on different instruments covering the different dynamic ranges, for reasons of absolute scaling of scattering intensities and the avoidance of systematic uncertainties it is desirable to develop instrumentation capable of providing insight on the equal time correlations over a continuous extended range of momentum transfer. The data obtained from such instruments will facilitate the construction of atomistic theoretical models for multiple-length-scale magnets (such as Co–Zn–Mn compounds), which are needed for describing simultaneously the existence of magnetic long- and short-range order on both mesoscopic and atomic scales.

5. Conclusions

Using small-angle neutron scattering, magnetometry, Lorentz transmission electron microscopy and magnetic diffuse neutron scattering, we have investigated the microscopic helical and skyrmion correlations in the chiral magnet Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ ($T_c \approx 102$ K). SANS experiments reveal that, below $T_c$, the mesoscopic phase diagram is dominated by orientationally disordered helical correlations that are largely unresponsive to applied magnetic fields below the saturation field. Close to $T_c$, a minority fraction of helical correlations in the sample undergo a magnetic-field-driven transformation into conventional two-dimensional skyrmion correlations. The signature for skyrmion formation is not obviously observed in magnetometry or LTEM measurements, highlighting the power of SANS to reveal the existence of skyrmions in this
particular case, due its ability to probe the entire sample volume. The characteristically strong magnetic disorder observed on both mesoscopic and atomic length scales is argued to arise from the interplay between compositional disorder, in particular the random mixing of magnetically distinct Co and Mn moments on the 8c Wyckoff site, and the magnetic frustration of Mn moments occupying the 12d Wyckoff site. While short-range magnetic correlations due to 12d frustrated Mn–Mn interactions lead to the observed magnetic diffuse scattering on atomic length scales, the mesoscale order observed by SANS is due to modulating moments on the 8c site. To explain the unusual temperature-dependent disordering process of helically modulating 8c moments, we invoke a temperature-dependent coupling between Mn moments on the 12d and 8c sites that grows on cooling and promotes increasing mesoscale magnetic disorder due to random reorientations of erstwhile helically modulating 8c Mn moments. It is also likely that this strong tendency towards magnetic disorder on cooling prevents the stability of a second, low-temperature, skyrmion phase in Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$, which is different from what is observed in the nearby composition Co$_{6.75}$Zn$_{6.75}$Mn$_{6}$.

Our study of Co$_{6.75}$Zn$_{6.75}$Mn$_{6.5}$ also highlights a general need for neutron instrumentation that can provide continuous access to an extended range of momentum transfer that can be measured comprehensively, thus facilitating the development of appropriate theoretical models.

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