Molecular beam epitaxy growth of [CrGe/MnGe/FeGe] superlattices: Toward artificial B20 skyrmion materials with tunable interactions

Adam S. Ahmed a, Bryan D. Esser b, James Rowland a, David W. McComb b, Roland K. Kawakami a,*

a Department of Physics, The Ohio State University, Columbus, OH 43210, United States
b Center for Electron Microscopy and Analysis, The Ohio State University, Columbus, OH 43212, United States

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ABSTRACT

Skyrmions are localized magnetic spin textures whose stability has been shown theoretically to depend on material parameters including bulk Dresselhaus spin orbit coupling (SOC), interfacial Rashba SOC, and magnetic anisotropy. Here, we establish the growth of a new class of artificial skyrmion materials, namely B20 superlattices, where these parameters could be systematically tuned. Specifically, we report the successful growth of B20 superlattices comprised of single crystal thin films of FeGe, MnGe, and CrGe on Si(111) substrates. Thin films and superlattices are grown by molecular beam epitaxy and are characterized through a combination of reflection high energy electron diffraction, X-ray diffraction, and cross-sectional scanning transmission electron microscopy (STEM). X-ray energy dispersive spectroscopy (XEDS) distinguishes layers by elemental mapping and indicates good interface quality with relatively low levels of intermixing in the [CrGe/MnGe/FeGe] superlattice. This demonstration of epitaxial, single-crystalline B20 superlattices is a significant advance toward tunable skyrmion systems for fundamental scientific studies and applications in magnetic storage and logic.

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1. Introduction

Topological spin textures in chiral magnetic materials have become an exciting research direction for condensed matter physics and magnetoelectronic applications [1–9]. In particular, skyrmions are localized spin textures ranging in size from 1 nm to 3 μm, with vortex-like structure as shown in Fig. 1. The topological character of this spin texture leads to new magnetic and electronic phenomena such as the topological Hall effect [7,10–16] and skyrmion Hall effect [17,18], as well as the so-called “topological protection” that permits these systems to be robust to perturbations and defects compared to conventional ferromagnetic domain walls [19]. Furthermore, the much lower critical current density necessary for depinning skyrmions (compared to domain walls) makes them promising for potential applications in ultra-high density magnetic memory including racetrack memory [9,20–25]. Typically, skyrmions are stabilized by the competition between two interactions. The first is the exchange interaction (J), which prefers neighboring spins to be parallel in a ferromagnet. The second competing interaction is the Dzyaloshinskii-Moriya interaction (DMI), which prefers neighboring spins to be 90° to one another. The DMI can only appear in systems that break bulk inversion symmetry or mirror symmetry. The first experimental evidence for the existence of skyrmions in condensed matter systems was shown in bulk MnSi crystals [26], where the bulk inversion symmetry breaking of its B20 crystal structure generates a Dresselhaus spin orbit coupling and produces a non-zero DMI (D0). Subsequent efforts continued on a variety of bulk B20 skyrmion crystals including FeGe [23,27–32]. With the development of B20 thin film systems, skyrmion stability was dramatically increased both in magnetic field and temperature range compared to their bulk counterparts [31]. For example, FeGe bulk crystals stabilize the skyrmion phase in a small temperature window of a few Kelvin below the ordering temperature (280 K). In contrast, sputter-deposited FeGe thin films show a dramatic extension of temperature stability from 280 K down to 10 K measured from topological Hall resistivity [27]. Moreover, it was shown that these same sputtered films had a finite zero-field topological Hall signal—an indication of zero field skyrmions [27,33]. These were important indications that film thickness and magnetic anisotropy could impact the skyrmion phase diagram in favorable ways for enhanced skyrmion stability.

Motivated by these experiments, Randeria and co-workers [34,35] modeled the skyrmion phase diagram for 2D systems as a function of magnetic anisotropy (K < 0 for out-of-plane hard axis, or “easy plane”; K > 0 for out-of-plane easy axis, or “easy axis”).
Dresselhaus DMI, DD \[35\]. Notably, the theoretical calculations can be tuned by layer thickness with larger DR/DD for thinner contributions to the DMI need not cancel. Because DR originates B20/TaOx), respectively (Fig. 1). Developing systems that can continuously tune skyrmions from Bloch to Néel can help realize optimal conditions for skyrmion stability and device operation (e.g. pinning potentials, skyrmion velocity, topological Hall effect). Therefore, in order to explore the predictions of Randeria and co-workers, it is crucial to develop magnetic materials where the parameters DD, DR, and K can be tuned independently.

To this end, one may consider developing heterostructures of B20 thin films. For a single interface between two materials, a non-zero Rashba DMI can develop due to the broken mirror symmetry. In a 2-component [A/B]N superlattice (with N repetitions), mirror symmetry remains intact and the Rashba DMI from the A/B interface cancels with the B/A interface. However, the magnetic anisotropy K from the interfaces is additive and does not cancel. In contrast, a 3-component superlattice [A/B/C]N can yield a non-zero overall DR because the A/B, B/C, and C/A interfacial contributions to the DMI need not cancel. Because DR originates from the interfaces while DD is from the bulk, the relative strengths can be tuned by layer thickness with larger DD/DR for thinner layers. Based on these considerations, it will be possible to independently tune the overall DD, DR, and K of the system through precisely grown B20 superlattices and heterostructures. However, to our knowledge, such artificial B20 structures have yet to be synthesized.

In this manuscript, we demonstrate the growth of B20 superlattices by molecular beam epitaxy (MBE), which establishes a route toward tunable skyrmion materials. We begin by growing FeGe (111) films on Si(111) substrates to establish a single-crystalline B20 template. We follow this growth with MnGe/FeGe and CrGe/FeGe bilayers to check for good epitaxial registry and favorable growth conditions between different B20 materials. We characterize all thin films to ensure single crystalline quality with reflection high energy electron diffraction (RHEED) and X-ray diffraction (XRD). Then, two-component [A/B]N superlattices of [MnGe/FeGe]n and [CrGe/FeGe]m are synthesized and characterized with XRD and RHEED. Finally, a three-component superlattice of [CrGe/MnGe/FeGe]n is grown and further characterized by cross-sectional scanning transmission electron microscopy (STEM) and X-ray energy dispersive spectroscopy (XEDS) to reveal chemical specificity and atomic layer resolution. This set of novel heterostructures demonstrates a fundamental materials advance for tunable skyrmions in artificial B20 systems.

2. Experimental results

The B20 thin films were grown by MBE in an ultrahigh vacuum (UHV) chamber with a base pressure of 2 × 10⁻¹¹ torr. Si(111) substrates (MTI, resistivity 10,000 Ω cm, single-side polished) were prepared by first sonicating the wafers in acetone for 5 min and isopropyl acetone for 5 min to remove any loose dirt/particles prior to treatment. The cleaned wafers were then dipped in buffered HF (Alfa Aesar) for 2 min to remove the native oxide and terminate the dangling Si bonds with H. The substrates were quickly loaded onto Ta paddles, inserted into the growth chamber, and annealed at 800 °C for 20 min to desorb the hydrogen and simultaneously obtain a 7 × 7 reconstruction. A quartz deposition monitor was utilized to determine the approximate thicknesses of the films. We used thermal effusion cells for the deposition of Fe (Alfa Aesar, 99.95%), Mn (Alfa Aesar, 99.98%), Cr (Alfa Aesar, 99.999%), and Ge (Alfa Aesar, 99.9999%) with typical cell temperatures of ~1300 °C, ~750 °C, ~1050 °C, and ~1250 °C respectively, for elemental growth rates in the range of 1.5–4 Å/min. For the growth of FeGe films and buffer layers, the substrate temperature was held at 300 °C. Growth was initiated with a ~3 Å Fe seed layer. Then elemental Fe and Ge were flux matched in a 1:1 ratio and co-deposited on the Si(111) substrate. For growth of MnGe and CrGe (flux matched 1:1) and all superlattices, the substrate temperature was maintained at 250 °C. Samples were characterized with in situ RHEED to obtain qualitative information about in-plane crystallinity. Out-of-plane lattice constants were quantified with X-ray diffraction (XRD) for all samples. For the superlattice, we performed cross-sectional STEM and XEDS for compositional mapping of individual layers.
2.1. FeGe thin films

FeGe is a magnetic material with B20 crystal structure and is known to host the skyrmion phase [23,27–29,31,32]. The helical pitch length is approximately 70 nm, and the skyrmions form an incommensurate hexagonal lattice. It has an ordering temperature of 280 K, which is the highest known value for a B20 skyrmion material. Its bulk cubic lattice constant [30] is 4.679 Å. For FeGe films with (111) orientation, this corresponds to an out-of-plane lattice constant of 8.10 Å. A quadrupole-layer (QL) period of 2.701 Å, as shown in Fig. 2a (a QL consists of four atomic layers: dense Fe, dense Ge, sparse Fe, sparse Ge). Of the three B20 materials presented in this manuscript, FeGe is the most well-known and studied.

We first established the growth of FeGe thin films on Si(111) substrates. RHEED patterns of the Si(111) surface taken along the [1 1 2] (Fig. 2c) and [1 1 0] (Fig. 2d) in-plane directions show a clear 7 × 7 reconstruction pattern after the 800 °C anneal. This indicates a clean and well-ordered surface for subsequent growth of B20 materials. For growth of FeGe in the B20 phase, the substrate temperature was reduced to 300 °C. After depositing the ~3 Å Fe seed layer (Fig. 2c inset), elemental sources of Fe and Ge were co-deposited with a flux ratio of 1:1. The pressure during growth was 1 × 10^-5 torr. Fig. 2e and 2f show RHEED patterns for a ~40 nm FeGe film. Notably, the FeGe lattice is rotated relative to the Si(111) lattice in order to minimize the lattice mismatch, thereby having the <1 1 0> axes of FeGe align with the <1 1 2> axes of Si(111), and the <1 1 2> axes of FeGe align with the <1 1 0> axes of Si(111). This epitaxial alignment is illustrated in the crystal structure diagrams in Fig. 2b. The sharp streaks in the RHEED pattern are indicative of flat terraces, and the distinct patterns along the [1 1 0] (Fig. 2e) and [1 1 2] (Fig. 2f) directions of FeGe indicate in-plane single crystal order.

To further investigate the crystallinity and orientation of the FeGe films, we performed XRD in the 0–2θ geometry. Fig. 2g shows an XRD scan for the ~40 nm FeGe/Si(111) sample. The Si(111) and Si(2 2 2) peaks are present from the substrate. Additionally, an FeGe (1 1 1) peak appears at 33.10°, and no other phases of FeGe are present, showing that single phase FeGe was synthesized. From this, we calculated the QL period to be 2.703 Å, which is larger than the bulk value by ~0.065%. This difference can be attributed to in-plane compressive strain on the FeGe film due to the 0.05% lattice mismatch between FeGe(1 1 1) and Si(1 1 1).

Having established the single crystal growth of FeGe films on Si(1 1 1), we utilized this as a template for subsequent growth of B20 overlayers and superlattices.

2.2. MnGe thin films and superlattices

MnGe is another skyrmion B20 crystal material. The helical pitch length is ~3 nm and forms a square lattice of skyrmions [42]. It has an ordering temperature of 170 K, and shows the largest topological Hall effect of the B20 compounds [11]. The bulk lattice constant [43] is 4.795 Å, which corresponds to a QL period of 2.768 Å.

We first established the growth of MnGe thin films on FeGe/Si(1 1 1) to determine suitable growth conditions for synthesizing MnGe/FeGe superlattices. Starting with a ~5 nm FeGe base layer grown on Si(1 1 1), a ~40 nm MnGe overlayer was co-deposited from elemental sources of Mn and Ge. The flux ratio was held at 1:1 and the substrate temperature was maintained at 250 °C during growth. Fig. 3a and 3b show the RHEED images for the MnGe film along the [1 1 0] and [1 1 2] directions, respectively. The RHEED images are streaky, indicative of flat terrace growth on the FeGe base layer. To quantify the out-of-plane lattice constant, Fig. 3c shows XRD for the MnGe/FeGe/Si(111) bilayer. First, there are no other phases of MnGe, present other than MnGe. Second, a MnGe(1 1 1) peak appears at a 2θ of 32.41°. This corresponds to a QL period of 2.759 Å, which is smaller than the bulk value by 0.33%. Comparing this value to the 2.5% lattice mismatch between MnGe and FeGe suggests that the ~40 nm MnGe film has relaxed to its bulk-like lattice structure. There is an additional shoulder peak which can be attributed to the ~5 nm FeGe base layer. The RHEED combined with XRD data shows that MnGe grows single crystalline on the FeGe(1 1 1) template.

Having found suitable growth conditions for MnGe/FeGe/Si(111) bilayers, we subsequently demonstrated the ability to synthesize a two-component superlattice comprised of MnGe (~2 nm) and FeGe (~2 nm). The first layer grown was FeGe/Si(111) where the substrate temperature was held at 300 °C. After we obtained a good RHEED image of FeGe, the substrate was cooled to 250 °C for the remainder of the superlattice growth. Fig. 3e and 3f show the RHEED for the final FeGe and MnGe layers, respectively. Both images are qualitatively very similar, and during growth, there was very little change in the RHEED images for the different layers. Fig. 3d shows the XRD scan of the MnGe/FeGe/Si(111) superlattice. Instead of two peaks for MnGe and FeGe, a single peak is present at a 2θ of 32.72°, which corresponds to a QL period of 2.734 Å, which lies between the measured QL periods of thick films of FeGe (2.703 Å) and MnGe (2.759 Å). Looking more closely near this peak reveals the presence of a weak satellite peak as indicated by the arrow in the inset of Fig. 3d.

2.3. CrGe thin films and superlattices

In addition to magnetic B20 materials, we are also interested in incorporating nonmagnetic materials. CrGe is a B20 paramagnetic metal that does not have a magnetic ordering temperature and, consequently, is not known to host a skyrmion phase [44–47]. Its bulk lattice parameter is 4.790 Å (QL period of 2.766 Å) and is very closely lattice matched to MnGe. To the best of our knowledge, the growth of CrGe thin films has yet to be demonstrated.

Prior to growth of CrGe, a ~5 nm FeGe base layer was grown on a 7 × 7 reconstructed Si(111) substrate at 300 °C. Elemental sources of Cr and Ge were co-deposited on the FeGe base layer at 250 °C. Fig. 4a and 4b show the CrGe RHEED images for a ~30 nm thick film. The RHEED images are qualitatively similar to MnGe and FeGe, but the CrGe streaks have bright spots and arrowhead-like features which indicate the onset of faceting and 3D growth [48]. In spite of these features, XRD was performed to obtain the out-of-plane CrGe lattice constant (Fig. 4c). In addition to the FeGe(111) peak from the buffer layer, a CrGe(111) peak appears at a 2θ of 32.38°, corresponding to a QL period of 2.761 Å. This is smaller than the bulk value by 0.15%, which suggests that the ~30 nm CrGe film has relaxed to its bulk-like lattice structure.

Having found suitable growth conditions for CrGe thin films, we applied these similar growth parameters to the fabrication of a two-component superlattice. A [CrGe/FeGe]10 superlattice was grown on Si(1 1 1) at 250 °C with each layer having a thickness of approximately 2 nm. The RHEED patterns during superlattice growth are shown in Fig. 4e and 4f. In contrast to the growth of the ~30 nm CrGe film, the RHEED is streaky and absent of any arrowhead features. Lastly, Fig. 4d shows an XRD scan of the superlattice structure, and a [CrGe/FeGe]10(1 1 1) peak is located at 2θ of 32.68°, corresponding to a QL period of 2.737 Å. A closer look near this peak clearly shows the presence of additional satellite peaks due to the superlattice structure (Fig. 4d inset). The stronger satellite peaks for CrGe compared to MnGe suggest sharper interfaces. The spacing of the satellite peaks corresponds to a superlattice period of ~5 nm, which is consistent with the designed period of
Fig. 2. (a) Cross-sectional diagram of FeGe showing the (111) lattice constant of 8.104 Å and the quadruple layer (QL) period of 2.701 Å. (b) Diagram of the epitaxial alignment of FeGe(111) on Si(111) with 30° lattice rotation. (c, d) RHEED images of a 7 × 7 reconstructed Si(111) surface shown for in-plane directions of ½ 11/2 2 and ½ 11/2 10/138, respectively. Inset of (c) RHEED after deposition of the Fe seed layer, showing a characteristic asymmetric pattern. (e, f) RHEED patterns measured along the ½ 11/2 10/138 and ½ 11/2 2 directions of the ~40 nm FeGe film, respectively. (g) XRD scan of the FeGe film. Single-phase, single crystal FeGe is shown amongst the Si substrate peaks.
With this, we have established that B20 CrGe layers are suitable for artificial B20 heterostructures.

2.4. [CrGe/MnGe/FeGe] trilayer superlattice

With suitable growth conditions obtained for CrGe, MnGe, and FeGe, we proceeded to the synthesis of a three-component B20 superlattice. A [CrGe/MnGe/FeGe]₈ superlattice structure was grown at 250 °C on a 7 × 7 reconstructed Si(111) surface. Every layer had a target thickness of approximately 2 nm. Fig. 5a, 5b, and 5c show RHEED patterns in the [1 1 0] direction for the topmost FeGe, MnGe, and CrGe layers in the superlattice, respectively. The RHEED is streaky indicating smooth 2D surfaces, and all three layers have similar patterns. Fig. 5d shows the XRD scan, which exhibits a [CrGe/MnGe/FeGe]₈(111) peak at 2θ of 32.65° corresponding to a QL period of 2.740 Å. The inset of Fig. 5d shows the presence of satellite peaks corresponding to a superlattice period of ~7 nm, which is consistent with the designed period of ~6 nm.

To gain further insight into the structural and compositional quality of the three-component superlattice structure, cross-sectional STEM and XEDS were performed on an aberration corrected FEI Titan³ G2 60-300 S/TEM equipped with a four quadrant Super-X XEDS detection system. A STEM image and XEDS compositional maps acquired from a cross-section with the viewing direction parallel to the FeGe [1 1 0] axis are shown in Fig. 6. The contrast in the high-angle annular dark field (HAADF) STEM image (Fig. 6a) arises from differences in atomic number (Z), where heavier elements have higher intensity. The compositional maps were extracted from the hyperspectral XEDS dataset. The uniform intensity in the Ge map (Fig. 6b) demonstrates the high degree of Ge homogeneity, which is consistent with the stoichiometry of the B20 materials grown. Fig. 6d, 6e, and 6f are the elemental maps for Fe, Mn, and Cr, respectively, which show the quality of the

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**Fig. 3.** Thin film: RHEED for MnGe thin film grown on FeGe buffer layers is shown along the (a) [1 1 0] and (b) [1 1 2] directions. (c) XRD for a ~40 nm thick MnGe film. The MnGe(111) peak is shown alongside the ~5 nm FeGe buffer layer. Superlattice: (d) XRD for the superlattice structure shows a single (1 1 1) peak. Inset: a weak satellite peak (arrow) due to superlattice structure is observed. (e, f) RHEED patterns are shown for the final topmost layers in a [MnGe/FeGe]₈ superlattice for FeGe [1 1 0](111) and CrGe [1 1 0](111), respectively.
MBE growth process with consistent, high quality layers with relatively low interdiffusion. The composite elemental XEDS map of Fe, Mn, and Cr (Fig. 6c) shows the relative positions of the layers and provides further confirmation of the quality of the multilayer film.

In Fig. 7, an XEDS line scan profile taken parallel to the growth direction and averaged horizontally from the maps in Fig. 6 shows that the layers are uniform in their compositions, with consistent stoichiometric Ge throughout all layers. Right at the substrate interface, there is an increase in the Fe:Ge ratio with significant amounts of Si, indicating the presence of an interfacial FeSi layer with possible Ge substitution. Additionally, there is a small amount (<2 at%) of Mn right at the interface, due to residual Mn in the chamber and its relatively high vapor pressure compared to Fe and Cr. The varying height of the Cr, Mn, and Fe curves for each layer, as well as the partial overlap of each peak are likely due to both the spatial resolution of the technique, as well as the fact that each layer is not perfectly parallel, leading to a further reduction in spatial resolution during averaging.

Regarding the initial FeGe growth with the presence of Mn impurities and formation of FeSi, we make a few comments. First, good quality FeGe films have been grown before and after the Mn cell was introduced to the MBE system, so the presence of small amounts of Mn is not essential for achieving good FeGe growth (nor does it inhibit good growth). Second, we have routinely achieved good FeGe growth with and without the Fe seed layer, so we do not view it as a required step. However, we believe it might reduce the diffusion of Fe out from the FeGe overlayer. In both cases, we observe the same type of RHEED patterns as shown in the inset of Fig. 2c. This pattern typically persists for the first ~2 nm of growth before it transitions to the final streaky pattern (i.e. Fig. 2e) at ~5 nm. This is consistent with the Si and Ge depth profiles in Fig. 7.

![Fig. 4. Thin film: RHEED for CrGe thin film grown on FeGe buffer layers is shown along the (a) [1 1 0] and (b) [1 1 2] directions. Arrowhead features indicate the occurrence of faceting. (c) XRD for a ~30 nm thick CrGe film. The CrGe(1 1 1) peak is shown alongside the ~5 nm FeGe buffer layer. Superlattice: (d) XRD for the superlattice structure shows a single (1 1 1) peak. Inset: satellite peaks (arrows) are observed due to the superlattice structure. (e, f) RHEED patterns are shown for the final topmost layers in a CrGe/FeGe superlattice for FeGe [1 1 0](1 1 1) and CrGe [1 1 0](1 1 1), respectively.](image)
Fig. 5. RHEED and XRD characterization of a three-component superlattice [CrGe/MnGe/FeGe]$_8$. (a–c) RHEED patterns are shown for the topmost layers FeGe, MnGe, and CrGe layers, respectively. The patterns are measured along the $\frac{1}{2}10$ in-plane direction of the films. Qualitatively, the RHEED images are very similar between each layer. (d) XRD scan of the trilayer superlattice shows a single (111) peak with satellite peaks from the superlattice structure (arrows in the inset).

Fig. 6. Cross-sectional STEM XEDS for the [CrGe/MnGe/FeGe]$_8$ superlattice. (a) HAADF image shows initial contrast for superlattice structure. (b) An elemental map for Ge. (c) A composite elemental map showing Fe (red), Cr (green), Si (light blue), and Mn (blue). (d–f) Separate elemental maps for Fe, Mn, and Cr, respectively. All scale bars are 20 nm. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Fig. 7. Horizontally averaged STEM XEDS line profile normal to the interface shown in the maps of Fig. 6. The presence of an interfacial FeSi layer can be seen in the first few nanometers, as well as a small amount of residual Mn.

The high-resolution HAADF-STEM image in Fig. 8 demonstrates the high-quality single crystalline layers of the superlattice, as evidenced by the coherent interfaces and block-like, zig-zag structure characteristic of B20 crystals along [1 1 0] (Fig. 2b). Additionally, the contrast in the HAADF-STEM image, especially at the FeGe/MnGe interface where Z-contrast is highest, shows that the interfaces are nearly atomically sharp, further demonstrating the quality of the growth.

3. Conclusion

We have synthesized the first B20 superlattices by MBE, including two-component superlattices ([MnGe/FeGe] and [CrGe/FeGe]) and a three-component superlattice ([CrGe/MnGe/FeGe]). Characterization by RHEED, XRD, and cross-sectional STEM indicate high crystalline quality and relatively low elemental interdiffusion at the interfaces. These studies establish the growth of a new class of artificial materials that offer a path toward skyrmion systems with tunable $D_0$, $D_a$, and $K$ for fundamental scientific studies and applications in magnetic memory and logic.

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References