Influence of nano-VC on the structural and magnetic properties of MnAlC-alloy

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Alloys of Mn55Al45C2 with additions of VC nano-particles have been synthesized and their properties evaluated. The Mn55Al45C2(VC)x (x = 0.25, 0.5 and 1) alloys have been prepared by induction melting resulting in a high content of the ferromagnetic τ-phase (> 94 wt.%). Powder X-ray diffraction indicates that nano-VC can be dissolved in the alloy matrix up to 1 at.%. On the other side, metallography investigations by scanning electron microscopy and scanning transmission electron microscope show inclusions of the nanosized additives in the microstructure. The effect of nano-VC on the grain and twin boundaries has been studied by electron backscattering diffraction. The magnetization has been measured by magnetometry up to 9 T while the domain structure has been studied using both magnetic force microscopy as well as Kerr-microscopy. For nano-VC contents above 0.25 at.%, a clear increase of the coercive force is observed, from 57 to 71 kA/m. The optimum appears to be for 0.5 at.% nano-VC which shows a 25% increase in coercive force without losing any saturation magnetization. This independent increase in coercivity is believed to originate from the nano-VC reducing the overall magnetic domain size. Overall, we observe that addition of nano-VC could be an interesting route to increase the coercive force of MnAl, without sacrificing saturation magnetization.

The mechanical-to-electrical energy conversion or vice versa is a key technology for the renewable energy and electro-mobility markets. The generators and converters used for example in electric cars or wind turbines require the use of permanent magnets. The permanent magnet market nowadays is roughly comprised of hexagonal ferrites and the rare-earth neodymium-based magnets, where Alnico and SmCo5 based magnets only play a minor role. After the rare-earth crisis in 2011, increasing attention has been placed on finding the so-called gap permanent magnets with performance exceeding those of hexagonal ferrites. One potential candidate is MnAl, which has enough anisotropy to be a permanent magnet of gap performance, i.e. to be able to supersede hexagonal ferrites but not rare-earth magnets. In the Mn–Al phase diagram, there is only a single ferromagnetic phase, τ-MnAl, which is metastable. The compound was discovered by Kono in 1958 and found to be ferromagnetic by Koch in 1960 but has still not been realized as a commercial product despite promising properties such as high saturation magnetization (MS), high Curie temperature (TC), high magnetocrystalline anisotropy (K1) and a good maximum theoretical energy product ((BH)max).

There are mainly two reasons why MnAl has not yet been commercialized. One is that before the rare-earth crisis, there was little to no incentive to develop gap magnets. The other is a more fundamental material issue on the microstructural level. The celebrated Brown’s paradox on the coercive force (Hc) of a material gives insight on why the coercive force never reaches even nearly the upper limit of the anisotropy field. According to Kronmüller’s equation,

\[ H_c = \frac{2\alpha K_1}{\mu_0 M_S} - D_{\text{eff}} M_S, \]

this is mainly due to the microstructure's ability to pin and nucleate the field. In Kronmüller’s equation \( \alpha \) is a dimensionless factor (for most materials \( \alpha < 1 \)), \( \mu_0 \) is the vacuum permeability and \( D_{\text{eff}} \) a local anisotropy constant related to the shape. In a well-controlled microstructure, \( \alpha \) takes the value of 0.1–0.3. In τ-MnAl, much struggle...
has been placed on achieving phase pure samples through a viable synthesis method. This can be done through, for instance, drop synthesis10, strip casting11, gas atomization12 and melt spinning13. It is possible to stabilize the metastable τ-phase and thus improve purity through extra alloying with for instance C14 or Ga15. For a general overview of synthesis methods see e.g. Chaturvedi et al. Two recent studies about phases in the Mn–Al–C systems were presented by Bajenova et al.17,18

In a recent study, nanoparticles of rare earths (RE) were added in MnAl (but without carbon) as (Mn0.5+Al0.5+y)RE, (0.01 ≤ x ≤ 0.09, 0.1 ≤ y ≤ 1.0)19,20, which created nanocrystallites of ~20 nm size that helped to pin domain walls and thus realizing a coercivity as large as 420 kA/m (5.43 kOe). We therefore, chose Mn55Al45C2 as base material for our investigations since it is more stable and the performance is more promising compared to the C-free τ-MnAl. The goal was to prepare Mn55Al45C2 with well dispersed nanoparticles to create pinning sites for domain walls. To achieve this, we added different amount of nano-VC to the Mn55Al45C2 base compound to elucidate its effect on the microstructure and magnetic properties.

Experimental details

Mn55-Al45-C2(VC) (x = 0, 0.25, 0.5, 1 and 3.5) alloys (~15 g each) were synthesized by induction melting. High purity raw materials have been used as starting materials for synthesis of the Mn55Al45C2(VC) alloys: Mn powder (Mn powder from Höganäs AB, purity 99.9%), Al (Gränges SM, purity 99.999%), C (Highways international, purity 99.999%), VC nanopowder (High purity VC nanopowder from INSCX exchange, purity > 99%, size 80–100 nm). Firstly, powders of Al and C were put together with the Mn55(VC)-pellets in an Al2O3-crucible. Afterwards the crucible was transferred to a high frequency induction furnace and vacuum pumped for 1 h. The melting was done by introducing 5 N Ar gas to have an atmosphere of 400 mbar. At the highest temperature (1300–1500 °C) the temperature was kept for 1–3 min to form a homogeneous solution. Finally, the furnace was switched off, followed by natural cooling to room temperature. The total mass loss during synthesis was less than 2 wt.%. In the event of larger losses, more Mn was added to compensate as for these alloys only Mn evaporates during melting.

All samples were characterized using a Bruker D8 powder X-ray diffractometer with a Lynx-eye position sensitive detector and Cu-Kα radiation on a zero-background single crystal Si sample holder. Phase analysis and Rietveld refinements of the X-ray data were done with the GSAS suite software21. For the Rietveld refinement of the PXRD we have used following information from literature4: the main τ-phase is a tetragonal L1₀ superstructure (P4/ mm m, AuCu-I type) with interstitial of C at [1/2 1/2 0], γ₂ phase (R3m, AlCr₂-type) and VC (Fm-3m, NaCl-type).

Microstructural evaluations were carried out using scanning electron microscopy (SEM), energy-dispersive X-ray spectrometry (EDS), electron backscattering diffraction (EBSD), magnetic force microscopy (MFM) as well as magneto-optical Kerr effect (MOKE) microscopy. Samples were cut in an Accutom-50 and subsequently mounted in Levofast resin and bakelite Phenocure®, ground and polished down to 1 μm by a standard process. The polishing was done with OPS-S non-drying colloidal silica suspension for 5 min in order to improve planeness and for relief polishing. Before etching, the samples were analyzed with a light optical microscope (LOM) in a Zeiss Axio Imager M2m microscope. They were afterwards etched in a diluted modified Keller’s reagent: with HCl 6 ml, HNO3 3 ml, HF 1 ml and H2O 290 ml. Samples were analyzed in a field emission scanning electron microscope (FE-SEM) Hitachi SU6600 equipped with an EDS system (Bruker EDX XFLASH 5010). Furthermore, the EBSD experiment was done using a Zeiss Merlin equipped with a Nordlys Max detector and evaluated by the AZtec HKL software. Characterization at high magnification was done using an FEI Titan Themis 200 scanning transmission electron microscope (STEM) equipped with SuperX EDS and spherical aberration corrector. The TEM specimen was prepared by an in-situ lift-out technique in a Zeiss Crossbeam 550 focused ion beam scanning electron microscope (FIB-SEM). The surface was protected with Pt and a final polishing step using a 5 kV ion beam was utilized to minimize beam damage from the preparation.

Magnetic force microscopy (MFM) was performed using a Bruker Dimension Icon system. Images were processed by the software GWYDION22 using background correction and flattening. Magnetic hysteresis curves were measured at 300 K using a Quantum Design PPMS with a maximum applied field of 9 T. The samples were corrected for demagnetizing effects.

An evico magnets Kerr microscope, with motion correction since the forces on the samples are rather strong, was used for magnetic domains observation. Room temperature Kerr microscopy studies were carried out in longitudinal geometry, i.e. applying an in-plane magnetic field parallel to the plane of incidence. In this configuration, out-of-plane components are dominating the magnetic contrast23.

Results and discussion

Synthesis and phase analysis. Previous investigations on the carbon stabilized τ-Mn55Al45 phase (Mn55Al45C2 made by drop synthesis)10,24,25 showed great purity and promising magnetic properties. However, despite being able to obtain close to 100% phase pure τ-MnAl the material still has challenges, mainly in terms of its microstructural properties which need to be improved to compete with ferrites. In this investigation, nano-VC additive has been selected in an attempt to achieve this. It should be taken into account that all samples in this work were synthesized by direct induction melting and all further investigations were done on the as-prepared samples. The Mn55Al45C2 alloy has been made as a reference for comparison. Investigation of the quality of Mn55Al45C2(VC) alloys has been done by powder X-ray diffraction (PXRD). Figure 1 presents a comparison of PXRD patterns for the Mn55Al45C2, Mn55Al45C2(VC)0.25, Mn55Al45C2(VC)0.5, Mn55Al45C2(VC)1, and Mn55Al45C2(VC)3 samples. Notably, peaks of VC are not visible by PXRD up to a concentration of 1 at.% of VC. A complete phase analysis of the synthesized and processed samples is presented in Table 1.
From the Rietveld refinement it was found that Mn₅₅Al₄₅C₂ has 85 wt.% of τ-phase purity while doped Mn₅₅Al₄₅C₂(VC)ₓ (x ≤ 1) alloys have greater purity (see in Table 1). From PXRD, the relative content of the τ-phase in the Mn₅₅Al₄₅C₂(VC)ₓ alloys is very close to the drop synthesized¹⁰. Using C is thus important for stabilization of the τ-phase even when introducing nano-dopants. One sample with larger content of nano-VC (3.5 at.%) has been prepared to verify the solubility limit. Additionally, one alloy without carbon (sample not presented here) but with nano-VC has been made. However, the purity of the C-free sample dropped drastically (65 wt.% of τ-phase). It should be noted that the chosen additive even promotes and amplifies in combination with carbon the stabilization/formation of the τ-phase alloys.

**Microstructural analysis.** SEM/LOM/EDX/STEM. The Mn₅₅Al₄₅C₂(VC)ₓ alloy was selected for detailed microstructural analysis. SEM observations on this etched sample (see Supplementary information Fig. S1a) show that there exist clusters (regions) of striations aligned along preferential directions. Before etching, images were collected by LOM. The polarized LOM image (Figure S1b) gives a good insight into the grain size, orientations and its distribution, which is difficult to observe by ordinary LOM/SEM even after etching. The matrix (τ-phase) that exists in all samples is very homogeneous without large agglomerates of VC.

EDX analysis cannot confirm the existence of nano-VC in the alloy, since the particles are too small in comparison with the resolution of the instrument. However, EDX mapping reveals a homogeneous distribution of V and a few Al-rich regions within the τ-phase (Figure S2). In addition, the average contents of Mn and Al measured by EDX is Mn₅₅Al₄₅ (Mn-rich region) and Mn₃₉Al₆₁ (Al-rich region) are in agreement with the τ-Mn₅₅Al₁₄₅C₂₀₀₂ and γ₂-Mn₅₅Al₄ phases, respectively. This is in good agreement with the results from PXRD.

The presence of VC particles in the τ-phase was further investigated in the TEM. Figure 2 shows a magnified image of the τ-phase, where it can be seen that particles are present within the grain. The particles are elongated in one direction, forming needle-shaped precipitates with a length of around 200 nm and width of around 20–30 nm. These are rich in V and C, indicating the presence of nanocrystalline VC. As have been described earlier, these particles are found in preferential orientations according to the LOM and SEM images. The preferential orientation could also be seen in the TEM cross section, where the red line in Fig. 2 indicates

**Table 1.** Phase analysis, magnetization at 9 T and coercivity values of the studied Mn₅₅Al₄₅C₂(VC)ₓ (x = 0, 0.25, 0.5, 1 and 3.5) alloys. Abbreviations used in the table: τ-Mn₅₅Al₄₅C₂ and γ₂-Mn₅₅Al₄₄.

| No | Sample                  | Phase analysis | Magnetometry data |
|----|-------------------------|----------------|------------------|
|    |                         | τ-phase parameters | Phase abundance (wt.%) | τ-phase parameters |                |
|    |                         | V(Å³) | c/a | Phase abundance (wt.%) | M(kA/m) | Hc(kA/m) |
| 1  | Mn₅₅Al₄₅C₂              | 27.599(3) | 1.3039 | 85.0 τ + 15.0 γ₂ | 541   | 57       |
| 2  | Mn₅₅Al₄₅C₂(VC)₀.₂₅       | 27.617(3) | 1.3055 | 97.5 τ + 2.5 γ₂ | 565   | 57       |
| 3  | Mn₅₅Al₄₅C₂(VC)₀.₅        | 27.620(4) | 1.3049 | 94.0 τ + 6.0 γ₂ | 544   | 71       |
| 4  | Mn₅₅Al₄₅C₂(VC)₁          | 27.630(1) | 1.3040 | 96.3 τ + 3.7 γ₂ | 499   | 69       |
| 5  | Mn₅₅Al₄₅C₂(VC)₃.₅       | 27.816(1) | 1.3060 | 86 τ + 5 γ₂ + 9 VC | -     | -       |

**Figure 1.** Comparison of the PXRD patterns for the as synthesized Mn₅₅Al₄₅C₂(VC)ₓ (x = 0, 0.25, 0.5, 1 and 3.5) alloys.

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the [103] direction. From these results, it is suggested that the particles have a preferred orientation in regard to the τ-phase, but the same orientation relationship was seen across twin boundaries in the TEM (Figure S3). Due to the high melting point of VC, it is expected that the added nanoparticles are distributed in the liquid phase, but remain as solid particles due to the high melting point and acting as nucleation sites for the τ-phase. Further investigation of this phenomenon is required to understand the solidification behavior of the alloy with VC, but is outside the scope of this study.

**EBSD.** Figure 3 shows the EBSD results for the Mn₅₅Al₄₅C₂, Mn₅₅Al₄₅C₂(VC)₀.₅ and Mn₅₅Al₄₅C(VC)₁ samples. It can be seen in Fig. 3a,d,g that all samples contain a combination of the τ-phase (shown in red) with some γ₂-phase (shown in blue) randomly distributed within the samples. Figure 3b,e,h,c,f,i show EBSD orientation maps of the τ-phase and the γ₂-phase for the three samples. The τ-phase is randomly oriented, but all individual grains of the γ₂-phase appear to be oriented in similar directions, which could originate from solidification if the γ₂-phase solidifies with a preferred orientation in regard to the thermal gradient. In the last few years, much work has been done to elucidate certain features of the microstructure and its effect on the performance of τ-MnAl such as anti-phase defects, dislocations, interfaces and twins. As many of these features are coexisting, often together with a difference in tetragonality (c/a ratio) of the τ phase, it is still not fully resolved which of these features have a negative impact on magnetic properties like magnetization, remanence and coercivity. Very recently, two studies were presented which show that for instance the coercivity can be negatively affected by twinning and that dislocations instead improve the coercivity.

Figure 4 a-c shows the distribution of grain boundaries in the Mn₅₅Al₄₅C₂, Mn₅₅Al₄₅C₂(VC)₀.₅ and Mn₅₅Al₄₅C(VC)₁ samples, respectively. It can be seen that a large amount of low angle grain boundaries (almost 50%) are formed in the sample without nano-VC, as well as a large fraction of grain boundaries with angles of 75˚ and 85˚ corresponding to <100> and <110> twins. With the addition of nano-VC, the amount of small angle grain boundaries decreases significantly to 17% and 14% for 0.₅ at.% and 1 at.% respectively. Figure 4d–f show the distribution of the <100> 75˚ and <110> 85˚ twin boundaries in blue and red, respectively. Figure 4g–i show the local misorientation for the Mn₅₅Al₄₅C₂, Mn₅₅Al₄₅C₂(VC)₀.₅ and Mn₅₅Al₄₅C(VC)₁ samples, respectively. Furthermore, the coherent boundaries in the twins are illustrated in Fig. 4j,k for the <100> 75° and <110> 85° twins, respectively.

**Magnetic characterization.** Magnetic measurements. Hysteresis measurements were performed up to 7200 kA/m (9 T) on samples of Mn₅₅Al₄₅C₂, Mn₅₅Al₄₅C₂(VC)₀.₂₅, Mn₅₅Al₄₅C₂(VC)₀.₅ and Mn₅₅Al₄₅C₂(VC)₁ to investigate the magnetic properties. All curves were corrected for demagnetizing effects. The results can be seen in Fig. 5 and Table 1.

To start with, there is no strong correlation between the magnetization measured in an applied magnetic field of 9 T and the amount of τ-phase. Compare for instance Mn₅₅Al₄₅C₂(VC)₀.₅ and Mn₅₅Al₄₅C₂(VC)₁, where first one has slightly less concentration of τ-phase compared to the last one (94.0 vs. 96.3 wt%, respectively). However, measured magnetization values show opposite trend (544 kA/m and 499 kA/m, cf. Table 1). Increasing the C content of (Mn₀.₅₄Al₀.₄₆)₁₀₀₋ₓCₓ increases saturation magnetization slightly from carbon free to some carbon
added, while too much decreases the saturation magnetization. This is however further complicated by the fact that the $c/a$ ratio varies between the samples, which is also known to affect magnetization. The $\tau$-phase in all $\text{Mn}_{55}\text{Al}_{45}\text{C}_x$ ($x = 0.25, 0.50$ and $1.0$) alloys has higher $c/a$ ratio than in $\text{Mn}_{55}\text{Al}_{45}\text{C}_2$, which also has been seen to affect both $H_A$ and $T_C$. This increase in $c/a$ ratio is most probably due to differences in the C content of the samples.

The vanadium carbide seems to have more pronounced effect on the coercive force, which is varying between $57–71$ kA/m. The optimum appears to be for $0.5$ at.% nano-VC which shows a $25\%$ increase in coercive force (from 57 to 71 kA/m) without losing any saturation magnetization.

**MFM and Ker microscopy.** We performed magnetic force microscopy (MFM) studies on the as-cast samples for $\text{Mn}_{55}\text{Al}_{45}\text{C}_2$, $\text{Mn}_{55}\text{Al}_{45}\text{C}_2(\text{VC})_{0.5}$, and $\text{Mn}_{55}\text{Al}_{45}\text{C}_2(\text{VC})_1$, to investigate the influence of added precipitates on the magnetic microstructure. The as-cast samples were polished in the same way as it had been done for the SEM studies. The measurements were performed on a Dimension Icon system from Bruker. Each sample was investigated at different positions and one representative result is shown in Fig. 6.

Qualitatively, smaller domains can be seen for the samples with VC precipitates, to investigate the domain size quantitatively we use the stereological procedure. For the stereological procedure, arbitrary line profiles are drawn through the domain image and the changes of the domain state along each line is recorded. The domain width is calculated as follows:

$$\text{domain width} = \frac{2 \times \text{line length}}{\pi \times \text{number of domain changes}}$$

Applying this procedure to each domain image and averaging the results for one sample yield the domain width distributions presented in Fig. 7.
Comparing the distributions of Mn$_{55}$Al$_{45}$C$_2$, Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.5}$ and Mn$_{55}$Al$_{45}$C$_2$(VC)$_1$ samples and (d–f) show the corresponding <100> 75° and <110> 85° twins in red and blue. (g–i) show the local misorientation for the Mn$_{55}$Al$_{45}$C$_2$, Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.5}$ and Mn$_{55}$Al$_{45}$C$_2$(VC)$_1$ samples. (j,k) illustrate the <100> 75° and <110> 85° twin boundaries, respectively. For simplicity, an atomic ratio of Al:Mn = 1:1 was assumed; red atoms (corners) are Al while blue atoms (center) are Mn.

Figure 4. (a–c) show the grain boundary distribution for the Mn$_{55}$Al$_{45}$C$_2$, Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.5}$ and Mn$_{55}$Al$_{45}$C$_2$(VC)$_1$ samples and (d–f) show the corresponding <100> 75° and <110> 85° twins in red and blue. (g–i) show the local misorientation for the Mn$_{55}$Al$_{45}$C$_2$, Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.5}$ and Mn$_{55}$Al$_{45}$C$_2$(VC)$_1$ samples. (j,k) illustrate the <100> 75° and <110> 85° twin boundaries, respectively. For simplicity, an atomic ratio of Al:Mn = 1:1 was assumed; red atoms (corners) are Al while blue atoms (center) are Mn.

Comparing the distributions of Mn$_{55}$Al$_{45}$C$_2$, Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.5}$ and Mn$_{55}$Al$_{45}$C$_2$(VC)$_1$, it can be seen that the samples with VC precipitates have a reduced domain width, verifying the more qualitative observations based on Fig. 6.

Investigating the Mn$_{55}$Al$_{45}$C$_2$(VC)$_1$ sample in more detail, we noticed at different locations a correlation between the domain appearance and precipitates visible in the topography. Figure 6 shows such a correlation between striations of precipitates in the topography measurement and the smaller domains in the magnetic domain image. The STEM study (Fig. 2) has shown that the lines of particles are indeed the VC precipitates.

Comparing the location of the smaller and larger domains in the magnetic contrast image in Fig. 8 with the location of the precipitates in the topography, a clear correlation between smaller domain sizes and the location of precipitates can be drawn. This further corroborates the beneficial effect of the VC precipitates on the magnetic properties.

To investigate the domain structure under applied field we performed Kerr microscopy studies on the polished surface of the as-cast pieces of Mn$_{55}$Al$_{45}$C$_2$, Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.5}$ and Mn$_{55}$Al$_{45}$C$_2$(VC)$_1$. The measurements were performed using the polar and longitudinal magneto-optical Kerr effect (MOKE); therefore, we are sensitive to...
magnetization components both out-of-plane and parallel to the applied in-plane magnetic field. The investigated as-cast samples have various orientations of the grains. Hence, a preferred anisotropy axis could not be identified and each grain had an almost unique response to the applied magnetic field. Figure 9 shows representative locations for each of the three samples. While the first row illustrates the MOKE images without any applied field, the second row shows the same location under the maximum applied field of 269 kA/m. The microstructure of the samples is dominating these images, but in several grains the domain structure can be identified. In each

Figure 5. Magnetic hysteresis curves for Mn$_{55}$Al$_{45}$C$_2$, Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.25}$, Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.5}$ and Mn$_{55}$Al$_{45}$C$_2$(VC)$_{1.0}$. Inset shows behavior near the coercive field.

Figure 6. Representative results from MFM measurements of Mn$_{55}$Al$_{45}$C$_2$, Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.5}$ and Mn$_{55}$Al$_{45}$C$_2$(VC)$_{1}$. First row shows the topography of the measured sample, while the second row represents the magnetic contrast illustrating the magnetic domain structure at the measured position.
of the three samples, grains could be identified which saturated with the applied field of 269 kA/m. Examples are marked with a green border around that specific grain. Additionally, there are also grains which could not completely be saturated, which are marked by yellow borders. A plausible reason for this vastly different behavior is the random orientation of the grains in the as-cast samples. Therefore, the anisotropy axis within saturated grains is more or less aligned with the applied field direction.

Another observation based on the microstructure is that the grain size seems to be reduced for the samples Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.5}$ and Mn$_{55}$Al$_{45}$C$_2$(VC)$_1$ in comparison with Mn$_{55}$Al$_{45}$C$_2$. But this observation cannot be quantified using the Kerr microscopy studies, as the domain structure visible in the images will interfere in any grain size analysis.

The lower two rows in Fig. 9 are typical domain contrast images recorded with a Kerr microscope. To receive a clear domain contrast, the Kerr image at a certain applied magnetic field is subtracted from the image at a different field. The resulting image illustrates the changes in the domain structure between the two applied fields.

**Figure 7.** Domain width distributions for Mn$_{55}$Al$_{45}$C$_2$, Mn$_{55}$Al$_{45}$C$_2$(VC)$_{0.5}$ and Mn$_{55}$Al$_{45}$C$_2$(VC)$_1$. A clear peak shift to lower domain width is visible in both the samples with 0.5% and 1% VC precipitates.

**Figure 8.** Magnetic force microscopy measurement of the Mn$_{55}$Al$_{45}$C$_2$(VC)$_1$ sample. Left image shows the topography with clear visible lines of precipitates. The right image illustrates the magnetic domains and a correlation between small domains and the location of the precipitates can be seen.
Figure 9. Kerr microscopy investigation of as-cast Mn_{55}Al_{45}C_{2}, Mn_{55}Al_{45}C_{2}(VC)_{0.5}, and Mn_{55}Al_{45}C_{2}(VC)_{1} alloys. The first row are measurements taken at 0 kA/m, while the second row shows images taken at an applied field of 269 kA/m. In all samples, grains can be identified which can be saturated (marked green) and which cannot be saturated (marked yellow). Row three and four illustrate the domain changes coming from 269 kA/m to 0 kA/m (third row) and coming from -269 kA/m to 0 kA/m (fourth row). As expected the domains in the grains are reversing their orientation (black to white and vice versa), but no obvious pinning sites can be identified. The images are cut-outs of the bigger measurements and represent 90 × 90 µm², i.e. the same size as the MFM images in Fig. 8; the red scale bar illustrates 10 µm.
Therefore, we can investigate the domain dynamics by applying the magnetic field stepwise. While increasing it from 0 kA/m up to 269 kA/m, we notice in all three samples that the domains change similarly. After reaching the maximum applied field at 269 kA/m we reduce the field stepwise, first down to remanence (0 kA/m) and then continuing to -269 kA/m, and ending the hysteresis loop again at 0 kA/m. Videos of these measurements can be found in the Supplementary materials. The third row in Fig. 9 shows the corresponding domain changes going from 269 kA/m to 0 kA/m, while the fourth raw illustrates the domain changes going from -269 kA/m to 0 kA/m. It is obvious that the black and white domains in row 3 are inverted in row 4. But at a closer examination of the domain change structure, it becomes clear that there is no evidence of strong pinning in the investigated samples, as there is no complete inversion of the domains. Only the expected reversing of the domain preference is visible, revealing more white than black domains.

These results from the domain dynamics are expected, as there are no big differences between the samples in the magnetization measurements for the available field range of the Kerr microscope (Fig. 5 and Table 1).

Conclusions
The effects of the nano-VC on the microstructure and magnetic properties of Mn55Al45C2 have been investigated by various techniques. It was shown that the nano-VC containing Mn55Al45C2(VC)x (x = 0.25, 0.50 and 1.0) samples were purer in τ-phase in comparison to the reference (≥ 94 wt.% vs. 85 wt.% of α-phase, respectively). Metallography studies demonstrate a microstructure with nanometer sized inclusions of carbides. The addition of nano-VC significantly decreases the amount of small angle grain boundaries as compared to Mn55Al45C2. In relation to this, the presence of twin boundaries with angles of 75˚ and 85˚ corresponding to <100> and <110> twins, respectively, increases with the addition of nano-VC.

For nano-VC contents above 0.25 at.%, a clear increase of the coercive force is observed, from 57 to 71 kA/m. The optimum appears to be for 0.5 at.% nano-VC which shows a 25% increase in coercive force without losing any saturation magnetization. This independent increase in coercivity is believed to originate from the nano-VC reducing the overall magnetic domain size. Also, as there is a weaker than usual correlation of τ-phase purity and magnetization in these alloys, it is concluded that saturation magnetization is here not only determined by the amount of τ-phase, but also the amount of carbon, which has been reported to have effects on saturation magnetization. Comparing carbon free τ-phase MnAl to carbon substituted τ-phase MnAl show that some amount of carbon is beneficial for saturation magnetization, while too much lowers the saturation magnetization. This has also been reported to decrease Kc and Tc. However, all of this is further complicated by the fact that atomic order, carbon content and c/a ratio (which are all interacting) varies in these samples. Separating these effects from each other is however outside of the scope of this work.

From magnetic force microscopy, we can conclude that nano-VC precipitates reduce the domain size in Mn55Al45C2. This effect can be caused by nucleation of magnetic domain walls by the precipitates or being an indirect effect of a reduced grain size, but most likely it is a combined effect of both.

Kerr microscopy studies revealed that there are no measurable differences in the dynamic response to the applied magnetic field for the investigated Mn55Al45C2, Mn55Al45C2(VC)0.5 and Mn55Al45C2(VC)1 samples. This finding is in good agreement with the magnetization measurements. Additionally, we can see a large variety of grain orientations leading to a vastly different response of the magnetic domains to the applied field. However, we cannot identify traces of strong pinning within a single grain in neither the Mn55Al45C2(VC)0.5 or Mn55Al45C2(VC)1 sample.

Overall, we can conclude that addition of nano-VC could be an interesting route to increase the coercive force of MnAl, without sacrificing saturation magnetization. However, before this can be realized, much deeper investigations on the effects of inclusions of nanoparticles in MnAl are needed.

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Author contributions
V.S. (synthesis, PXRD, Rietveld refinement and directed the project), H.S. (MFM and Ker microscopy), D.H. (magnetic measurements), D.K. (STEM/EDX and EBSD), R.P. (Ker microscopy) and F.O. (LOM and SEM/EDX) performed the experiments, analyzed the data and wrote the paper. B.S., H.V., G.A., P.S. and M.S.: organized the research, supervision, resources, project administration and writing—review & editing. All authors took part at joint discussion of the results and commented on the manuscript.

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