Jumping the Gap: Can Tetrataenite become a "Hard" Permanent Magnet?

C. D. Woodgate¹, J. B. Staunton¹, L. H. Lewis²

¹University of Warwick, Coventry, UK ²Northeastern University, Boston, USA



Motivation

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- Consists entirely of two abundant 3d transition metals.
- Dispute in the literature as to the value of its uniaxial anisotropy energy.



Context 0●0

Existing Dat

Theoretical Insight

Conclusions

Magnetocrystalline Anisotropy Energy (MAE)

• Tetragonal sample with magnetisation $\mathbf{M} = |\mathbf{M}|(\cos\theta\cos\phi, \cos\theta\sin\phi, \sin\theta).$



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Magnetocrystalline Anisotropy Energy (MAE)

- Tetragonal sample with magnetisation $\mathbf{M} = |\mathbf{M}|(\cos\theta\cos\phi, \cos\theta\sin\phi, \sin\theta).$
- Magnetocrystalline anisotropy energy written

$$K(\theta,\phi) = K_1 \sin^2 \theta + K_2 \sin^4 \theta + K_3 \sin^4 \theta \cos 4\phi + \dots$$
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Conventional to measure/calculate uniaxial MAE:

$$K_U = K(\hat{\mathbf{z}}) - K(\hat{\mathbf{x}}) = K_1 + K_2 + K_3$$



History of L10 FeNi

▶ L1₀ phase first reported in 1960s by Néel and co-workers¹².

¹Néel *et al.*, J. Appl. Phys. **35** 873 (1964)

²Paulevé et al., J. Appl. Phys. 55 989 (1968)

³M. Werwiński, W. Marciniak, J. Phys. D: Appl. Phys. **50** 495008 (2017)

History of $L1_0$ FeNi

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- Also found to occur naturally in metallic meteorites, mineral name tetrataenite.

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- Also found to occur naturally in metallic meteorites, mineral name tetrataenite.
- Challenging to synthesise. Recent experimental efforts have focussed on manufacturing thin films or on targeted processing.
- Extensively studied using density functional theory calculations, *e.g.* Ref. 3. Primarily for perfectly ordered structures at T = 0 K, though.

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• Néel *et al.*:
$$K_1 = 0.32$$
, $K_2 = 0.23$.

Paulevé et al.: $K_1 = 0.3$, $K_2 = 0.17$, $K_3 = 0.08$.

JOURNAL OF APPLIED PHYSICS VOLUME 35, NO. 3 (TWO PARTS-PART 2) MARCH 1964

Magnetic Properties of an Iron-Nickel Single Crystal Ordered by Neutron Bombardment

L. NÉEL, J. PAULEVE, R. PAUTHENET, J. LAUGIER, AND D. DAUTREPPE Centre d'Etudes Nucléaires et Laboratoire d'Electrostatique et de Physique du Métal, Grenoble, France

JOURNAL OF APPLIED PHYSICS VOLUME 39, NUMBER 2 1 FEBRUARY 1968

Magnetization Curves of Fe-Ni (50-50) Single Crystals Ordered by Neutron Irradiation with an Applied Magnetic Field

J. PAULEVÉ, A. CHAMBEROD, K. KREBS, AND A. BOURRET

Centre d'Etudes Nucléaires de Grenoble, France

Theoretical Insight

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Potential Origins of Large K_2 and K_3 ?

In our DFT calculations, we find negligible K₂, K₃.



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Conclusions

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- In our DFT calculations, we find negligible K₂, K₃.
- Néel proposes bulk sample made up of individual L1₀ nanocrystals, layering directions not all aligned¹.
- Exchange interaction between nanoscale domains yields large, macroscopic K₂ and K₃.



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Degree of Atomic Order

Even if I have a single-variant nanocrystal, is is perfectly ordered?

⁴C. D. Woodgate, C. E. Patrick, L. H. Lewis, J. B. Staunton, arXiv:2307.15470

Degree of Atomic Order

- Even if I have a single-variant nanocrystal, is is perfectly ordered?
- Define atomic order parameter, S to quantify how well-ordered sample is⁴.



⁴C. D. Woodgate, C. E. Patrick, L. H. Lewis, J. B. Staunton, arXiv:2307.15470

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Degree of Atomic Order



Clear disagreement between theory and experiment once effects of imperfect atomic order are considered⁴.

⁴C. D. Woodgate, C. E. Patrick, L. H. Lewis, J. B. Staunton, arXiv:2307.15470

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Theoretical Insight 000●

Effects of Finite Temperature



 Robust computationally predicted finite temperature performance⁴.

⁴C. D. Woodgate, C. E. Patrick, L. H. Lewis, J. B. Staunton, arXiv:2307.15470

Summary: Aspects to Consider

Aspect 1: Polycrystalline, Multivariant Samples

Experimentally FeNi has significant K_2 , K_3 ; these are *not* detected in DFT calculations. Nanoscale origins?

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Aspect 2: Imperfect Atomic Order

Decreasing atomic order decreases computationally predicted MAE, makes for large discrepancy between theory and experiment.

Aspect 3: Finite Temperature

Tetrataenite has good finite temperature performance, but increasing temperature decreases MAE compared to T = 0.

Context	Existing Data	Theoretical Insight	Conclusions
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Acknowledgements People

University of Warwick, UK

- Christopher D. Woodgate
- Julie B. Staunton

Northeastern University, USA

Laura H. Lewis

University of Oxford, UK

Christopher E. Patrick

Funding

- C.D.W. supported by a studentship within EPSRC-funded CDT: warwick.ac.uk/hetsys
- EPSRC (UK)
- NSF (US)
- DOE (US)







