

[Home](http://iopscience.iop.org/) [Search](http://iopscience.iop.org/search) [Collections](http://iopscience.iop.org/collections) [Journals](http://iopscience.iop.org/journals) [About](http://iopscience.iop.org/page/aboutioppublishing) [Contact us](http://iopscience.iop.org/contact) [My IOPscience](http://iopscience.iop.org/myiopscience)

A ferromagnetic monolayer with model spin-orbit and dipole-dipole interactions

This article has been downloaded from IOPscience. Please scroll down to see the full text article. 1993 J. Phys.: Condens. Matter 5 4527

(http://iopscience.iop.org/0953-8984/5/26/023)

View [the table of contents for this issue](http://iopscience.iop.org/0953-8984/5/26), or go to the [journal homepage](http://iopscience.iop.org/0953-8984) for more

Download details: IP Address: 171.66.16.159 The article was downloaded on 12/05/2010 at 14:09

Please note that [terms and conditions apply.](http://iopscience.iop.org/page/terms)

A ferromagnetic monolayer with model spin-orbit and dipole-dipole interactions

M B Taylor† and B L Gyorffy

H H Wills Physics **Laboratory,** Royal **Fort,** 'pmdall Avenue, Bristol, **UK**

Received **19 February** 1993

Abstract. We have considered a classical spin model of a ferromagnetic monolayer with model spin-orbit and dipole-dipole interactions, which provide competing perpendicular and in-plane anisompies respectively. In particular **we** plot the **zero** lemperature phase diagram, which features, as well as the expected perpendicular and in-plane ferromagnetic phases, a series of wrexpected perpendicular antiferromagnetic phases although these are **most imponant** for anisompy parameter values higher than we would expect to **see** in real monolayers. A mean field theory at finite temperatures predicts the existence of a first-order perpendicular to in-plane reorientation transition of the magnetization for certain values of **the** parameters. **Computer** simulations of **this** system wem attempted, but were unsuccessful because of **the** large conelation lengths arising. We draw the tentative conclusion that straightforward Monte *Carlo* and molecular dynamics simulation is not a practicable way to investigate this system, and note the important technical point that data parallelization of conventional Monte Carlo methods with infinitely long-range interactions is not possible.

1. Introduction

It was first pointed out by Néel [1] that breaking the translational symmetry of a magnetically isotropic material (by introducing a surface) leads to the possibility of magnetically anisotropic terms in the Hamiltonian (a magnetic surface anisotropy). Specifically, he noted that if the anisotropy is written in a suitably general form, the leading pairwise term is zero in the bulk of a cubic crystal, but not **at** its surface.

N6el's discussion was phenomenological. More recently a particular physical basis for **the** anisotropic **terms** involved has been considered (2-41 and calculated *[5,6].* This magnetic surface anisotropy is thought to **arise** firstly from the dipole-dipole interaction, and secondly from the electron spin-orbit coupling. The dipole-dipole interaction will lead to an in-plane anisotropy, and the spin-orbit term may lead either to in-plane or to perpendicular anisotropy. Since this is a surface effect, it may best be observed in systems with a high surface-to-volume ratio. Experimentally this is likely to mean thin films or multilayers; theoretically it is convenient to examine a two-dimensional system (a monolayer). We therefore investigate a plane of spins, attempting to determine its behaviour as a function of the *two* types of anisotropy.

There **are** three important and much-studied limiting cases of this model:

(i) If the anisotropies are both removed we recover the 2D classical Heisenberg model. This has no magnetic order at non-zero temperatures since the ferromagnetic order is

t Resent address: Institute of Physics **Publishmg.** Techno **House.** Redcliffe Way, Bristol BSI **6NX. UK.**

destroyed by long-wavelength spin waves, **as** demonstrated by Mermin and Wagner **[7,81.** Various theoretical and simulation analyses have been performed on this system, see e.g. references [9-14].

(ii) If the anisotropy is such that the spins are confined to the plane they become twodimensional and we have the **2D** classical XY model which also lacks ferromagnetism, but features a Kosterlitz-Thouless transition [15,16]. Simulations of this system have been reported in references **[17-191**

(iii) In the limit of strong perpendicular anisotropy the spins become one-dimensional giving the **2D** Ising model with a finite Curie temperature *Tc.* '

To gain an idea of physically realistic values of these anisotropy strengths we use the first-principles calculations of Gay and Richter [5] and of Guo *etal* [6], and the experimental results of Gradmann **[20].**

Two previous studies motivate **us** particularly **to** examine this system: the lowtemperature mean field work of Jensen and Bennemann [2] which treats the magnetization and dipole-dipole interactions in rather simplified ways, and the renormalization group work of Pescia and Pokrovsky [4]. Both predict the existence, for some values of the interaction constants, of a reorientation temperature T_R below the Curie temperature, above which the magnetization is driven entropically from a perpendicular to an in-plane direction. Some experimental evidence of such a transition has been reported **[21-231** and the perpendicular ferromagnetic state is of great importance in the technology of high-density magnetic recording media.

2. **Details** of *the* model

In investigating this system we use the simplest model that *can* be expected to capture dipole-dipole interaction, and spin-orbit anisotropy: the physics of interest. We will study a Hamiltonian with **terms** representing exchange,

$$
\mathcal{H} = \mathcal{H}_{\text{ex}} + \mathcal{H}_{\text{dip}} + \mathcal{H}_{\text{so}}
$$

=
$$
-\frac{1}{2}J \sum_{ijj} \sigma_i \cdot \sigma_j + \frac{1}{2} \omega \sum_{ij} \left[\frac{\sigma_i \cdot \sigma_j}{r_{ij}^3} - 3 \frac{(\sigma_i \cdot r_{ij})(\sigma_j \cdot r_{ij})}{r_{ij}^5} \right] - \lambda \sum_i (\sigma_i^z)^2
$$
(1)

where at each site *i*, at position r_i , of a two-dimensional square lattice in the $x-y$ plane is a classical Heisenberg spin (three-dimensional unit vector) $\sigma_i = (\sigma_i^x, \sigma_i^y, \sigma_i^z)$, and where $r_{ij} = r_j - r_i$, $r_{ij} = |r_{ij}|$, and the symbol $\langle ij \rangle$ denotes summation over nearest neighbours only. The parameters of the model **are thus:**

- *J* strength *of* the pairwise short-range (nearest neighbour) exchange interaction
- ω strength of the pairwise long-range (r^{-3}) dipole-dipole interaction
- λ strength of the single site model spin-orbit term
- *T* temperature

This expression is effectively a discrete form of the Hamiltonian used by Pescia and Pokrovsky [41, and is an improvement on that used by Jensen and Bennemann **[2.3]** in that it does not simplify the dipolar interaction by representing it as a single-site term.

The limiting cases mentioned in section 1 are achieved with the following values of the parameters:

Physically, ω must be non-negative, providing an in-plane anisotropy. *J* and λ on the other hand may be positive or negative. We concem ourselves in this study only with the case $J \ge 0$, a ferromagnetic exchange, and $\lambda \ge 0$, the spin-orbit term providing a perpendicular anisotropy. These choices are motivated by our interest in the perpendicular ferromagnetic state. Note that of the limiting cases listed above this excludes the ZD *XY* model.

3. **Ground states**

By inspection of the Hamiltonian **(I)** some facts about the ground states of the system *can* be inferred.

We note first that, for the square lattice, the anisotropic terms in the Hamiltonian define no preferred direction within the plane so that states will be continuously degenerate with respect to azimuthal angle ϕ .

If *J* is large compared with the other energies in the system, then we expect a ferromagnetic state. Given a ferromagnetic system, if the in-plane anisotropy ω is large compared with the perpendicular anisotropy λ the state will be in-plane ferromagnetic, and if λ is large compared with ω it will be perpendicular ferromagnetic.

However, if λ and ω are both large compared with *J* we have a different situation: the high λ forces all the spins perpendicular to the plane causing the term $-(3/2)\omega \sum_{ij}(\sigma_i \cdot$ r_{ij})($\sigma_j \cdot r_{ij}$) r_{ij}^{-5} in the dipole-dipole interaction to vanish. The remaining part of the dipole-dipole interaction $+(1/2)\omega \sum_{ii}(\sigma_i \cdot \sigma_j)r_{ii}^{-3}$ acts like an antiferromagnetic term, and if ω is sufficiently large this will overcome the exchange term, resulting in a perpendicular antiferromagnetic state of some sort. Such a state will be highly frustrated, since the dipols dipole interaction is long-ranged, so that every spin wants to be antiparallel to every other spin.

The exact nature of such an antiferromagnetic sate is not immediately obvious. We expect two sublattices, one of spin up and one of spin down, which by symmetry will each occupy half of the lattice sites. To ascertain the geometry of these sublattices, computer simulations were performed, starting in a disordered state at high temperatures and cooling slowly to zero temperature (see section 6). The **results** of these simulations (i.e. the resulting zero-temperature states) are on their own somewhat inconclusive, since at low temperatures the system tended to settle into states which look glassy and metastable, but visual observation of graphical representations of them suggests the corresponding ordered states to which they may be close. This procedure is an alternative to the ground-state search method described by, e.g. Ducastelle $[24]$. For these large values of λ , in which the low temperature states feature entirely perpendicular spins, the observed ground states seem to fall into three categories, characterized as follows:

(i) all spins in the same direction (either up or down),

(ii) alternating stripes of up and down spins, of a characteristic width for given values of λ and ω ,

(iii) **a** 'chequerboard' pattern of up and down spins.

We therefore investigate the energies of the corresponding **sets** of ordered states, which we label as indicated

ZFERRO: perpendicular ferromagnetic state: $\sigma_i^2 = +1$ **Vi** CHECK: chequered antiferromagnetic state:
 $\sigma_i^2 = \begin{cases} +1 & r_i^x + r_i^y = 2p \\ -1 & r_i^x + r_i^y = 2p + 1 \end{cases}$ CHECK: chequered antiferromagnetic state:

$$
\sigma_i^z = \begin{cases} +1 & r_i^x + r_i^y = 2p \\ -1 & r_i^x + r_i^y = 2p + 1 \end{cases}
$$

STRIPEn: striped antiferromagnetic state:

$$
\sigma_i^z = \begin{cases}\n+1 & 2pn + 1 \leq r_i^x \leq 2pn + n \\
-1 & 2pn + n + 1 \leq r_i^x \leq 2pn + 2n\n\end{cases}
$$

In the above $r_i = (r_i^x, r_i^y)$ is the lattice position of site *i* measured in units of lattice spacing, *p* represents any integer, and in all cases $\sigma_i^x \equiv \sigma_i^y \equiv 0$. Example STRIPEN phases are illustrated in figure 1. We compare these with the state we expect for small values of λ :

XFERRO: in-plane ferromagnetic state: $\sigma_i \equiv (\sigma_i^x, \sigma_i^y, \sigma_i^z) = (+1, 0, 0)$

.we have not distinguished between those states explicitly defined above and equivalent degenerate states.

The energies per site of these states **are** given **as** follows:

$$
E_{\text{XFERRO}} = -2J - \frac{1}{4}A\omega\tag{2}
$$

$$
E_{\text{ZFERRO}} = -2J + \frac{1}{2}A\omega - \lambda \tag{3}
$$

$$
E_{\text{CHECK}} = +2J + C\omega - \lambda \tag{4}
$$

$$
E_{\text{STRIPE}n} = -\left(2 - \frac{2}{n}\right)J + S_n\omega - \lambda \tag{5}
$$

where the values A , C and S_n are numerical constants depending on lattice geometry, defined using the spin states **XFERRO, CHECK** and **STRlPEn** respectively as:

$$
\text{const}^{\text{state}} = \lim_{N \to \infty} \frac{1}{2N} \sum_{i=1}^{N} \sum_{j=1}^{N} \sigma_{i}^{\text{state}} \cdot \sigma_{j}^{\text{state}} \frac{1 - \delta_{ij}}{|r_{j} - r_{i}|^{3}} \tag{6}
$$

and these are calculated numerically, giving:

$$
A = +9.0336
$$

\n
$$
C = -1.3230 = (2^{-1/2} - 1)A/2
$$

\n
$$
S_1 = -0.4677
$$

\n
$$
S_2 = +0.7908
$$

\n...

By manipulation of equations **(2)-(5)** we arrive at the zero-temperature phase diagram depicted in figure 2. The shaded rectangle near the origin represents the range of physically realistic values of ω and λ given in equation (13) and explained in section 4.

Figure Z Zem-temperature phase diagram for the monolayer. STRIPU, phases *are* **plolted** for $n \leq 100$. $a-f$: stripes 1-6 respectively. The shaded rectangle near the origin indicates the **physically realistic values for the model parameters** *(see* **section 4). All** *quantities* **are** *in* **units of** *J.*

We present also an algebraic characterization of the phase diagram by giving the following **equations** of the boundaries of **the** in-plane ferromagnetic phase with the perpendicular ferromagnetic and chequered antiferromagnetic phases respectively:

XFERRO/ZFERRO:

$$
\omega A = \frac{4}{3}\lambda \tag{8}
$$

XFERRO/CHECK:

$$
\omega A = \frac{4J}{\sqrt{2} - 1} (\lambda - 4) \tag{9}
$$

and the following expressions for the (λ, ω)

coordinates of the triple points:

XFERRO/CHECK/STRLPE I :

$$
(\lambda, \omega) = J \left(4 + \frac{(\sqrt{2} - 1)A}{(1 - \sqrt{1/2})A + S_1}, \frac{4}{(1 - \sqrt{1/2})A + S_1} \right) \tag{10}
$$

 $XFERRO/STRIPEn/STRIPE(n + 1)$:

$$
(\lambda, \omega) = J\left(\frac{2}{n+1} + \frac{A + 2S_{n+1}}{n(n+1)(S_{n+1} - S_n)}, \frac{4}{S_{n+1} - S_n}\left(\frac{1}{n} - \frac{1}{n+1}\right)\right).
$$
 (11)

We draw particular attention to the high- ω boundary of stability of the phase ZFERRO: we have plotted STRIPEn phase boundaries up to $n = 100$ in figure 2, but it is not clear to us what is the lowest value of ω at which a **STRIPE***n* phase is stable. On the highly plausible assumption that S_n is monotonic in *n* we are then interested in the $n \to \infty$ limit of ω in equation (11), which ought more simply to be equal to the following expression:

$$
\omega_{\max}^{\text{ZFERRO}} = \lim_{n \to \infty} \frac{4J}{n(A - S_n)}.
$$
\n(12)

Using simple graphical methods to examine the first 100 values of S_n we have been unable to determine the asymptotic behaviour of $4J/n(A - S_n)$; in particular it does not appear to be related to n by a power law. However, it may be possible analytically or numerically to find an answer to this question. This point is clearly of great interest, since if $\omega_{\text{max}}^{\text{ZFERRO}}$ is zero the perpendicular ferromagnetic phase is never a ground state of this system for finite *o.* Even if it is not zero, its value is of importance for the behaviour *of* physically realistic materials.

The general features of the perpendicular antiferromagnetic phases CHECK and STRIPEN may be understood **as** follows: where the long-ranged antiferromagnetic tendency *(o)* is very strong the system will be as antiferromagnetic as possible, resulting in the chequerboard pattern of **CHECK.** However, as the nearest-neighbour ferromagnetic exchange (J) becomes relatively more important, there is a tendency to reduce the number of upldown spin boundaries, while the long range of the dipolar interaction still **acts** to ensure that oppositely oriented spins exist at some distance away, favouring striped phases.

We have calculated energies for chequered states with squares larger than one lattice spacing, and none of these is stable, but we have **no** guarantee that there are **not** lower energy states which have not occurred to **us.** However, in view of the low temperature simulation results we think that this is unlikely.

The ground states in figure 2 except for the 2D Heisenberg case $\lambda = \omega = 0$, are not unstable to the long wavelength spin waves which destroy ferromagnetic order in that model according to the Mermin-Wagner theorem [7, 8]; the fluctuations are prevented from diverging by an energy gap of zeroth order in a wave vector proportional to λ [25] and one of first-order proportional to ω [26].

4. Physical values of the model parameters

In this section we put rough bounds on the values of the model parameters J, ω, λ which might correspond to real physical systems. As a first approximation we consider the values the parameters would take *to* model a monolayer of iron.

We can estimate the exchange interaction J by supposing that J is the same for an iron monolayer as for bulk iron, and that bulk iron is adequately approximated by a nearestneighbour classical Heisenberg ferromagnet. Then by using a series expansion value for the Curie temperature of a BCC Heisenberg ferromagnet $k_B T_C = 2.06 J$ [27] and an experimental value of Curie temperature $T_C^{\text{Fe}} = 1043 \text{ K}$ [28], we arrive at the estimate:

$$
J^{\rm Fe}\approx 40~{\rm meV}.
$$

Theoretical **15.61** and experimental **1201** wok suggest values **of** the other parameters $\lambda \approx 0.5$ meV and $\omega \approx 0.01$ meV. These are rough estimates and values will clearly vary for different materials and environments, but we expect for all available materials:

$$
\omega/J \ll 0.1 \qquad \lambda/J \ll 1 \tag{13}
$$

and this region is indicated by the shaded rectangle near the origin of the zero-temperature phase diagram figure 2.

Most systems will fall well inside **this** region so it can be seen that of the phases plotted the moa impoltant are **XFERRO** and **ZFERRO.** It can therefore be seen that most of the phase diagram plotted corresponds to values of ω and λ which we do not expect to see in real monolayers, but we feel that the statistical mechanical interest of these unphysical regions is sufficient to justify depicting them.

We note also that cases of physical interest are far from the $\lambda = \infty$ 2D Ising limiting case and near to the $\lambda = 0$ 2D Heisenberg limiting case of the system, and we expect to see behaviour which is like a modified Heisenberg model, and unlike an Ising model.

5. Mean field theory

By making the substitutions:

$$
\sigma_i = \langle \sigma_i \rangle + \delta \sigma_i \tag{14}
$$

in the Hamiltonian (1) and discarding terms in $\delta\sigma_i \cdot \delta\sigma_i$, we can obtain a mean field Hamiltonian. Since we expect the ordered state to be ferromagnetic in the physical (small λ , small ω) region of the phase diagram we further assume homogeneity

$$
\langle \sigma_i \rangle \equiv m = (m^x, m^y, m^z) \tag{15}
$$

and thereby arrive at the following expression for the mean field free energy:

$$
F = -k_{\text{B}}T \ln \text{Tr} \ e^{-\beta H}
$$

= $\frac{N}{2} \left[\left(4J + \frac{\omega A}{2} \right) (m^x)^2 + \left(4J + \frac{\omega A}{2} \right) (m^y)^2 + (4J - \omega A)(m^z)^2 \right]$
- $k_{\text{B}}T N \ln \int_0^{2\pi} d\phi \int_0^{\pi} d\theta \sin \theta \exp \beta \left[\left(4J + \frac{\omega A}{2} \right) m^x \sin \theta \cos \phi + \left(4J + \frac{\omega A}{2} \right) m^y \sin \theta \sin \phi + (4J - \omega A) m^z \cos \theta + \lambda \cos^2 \theta \right]$ (16)

where N is the number of sites in the system, β is the reciprocal temperature $(k_BT)^{-1}$, and A is the constant defined in equation (7).

Given this free energy $F = F(m, J, T, \omega, \lambda)$, which can be evaluated and manipulated numerically, the behaviour of the **system** can be determined. In particular the Curie temperature T_c and the temperature of the first-order reorientation transition T_R can be located iteratively using the equations:

$$
\left(\frac{\partial^2 F(T,m)}{\partial m^2}\right)_{m=0, T=T_{\rm C}} = 0\tag{17}
$$

$$
\left(F_{\perp}^0 - F_{\parallel}^0\right)_{T=T_R} = 0\tag{18}
$$

where F_{\perp}^0 and F_{\parallel}^0 are the values of the free energy when it is minimized with respect to m under the constraint of perpendicular and in-plane magnetization respectively.

Sample free energy surfaces in the regions of these **transitions** are illustrated in figure **3.**

magnetization m : T_C is that temperature at which the two minima coalesce, and (b) as a function **of temperature and angle** θ **of the magnetization from the vertical;** T_R **is that temperature at which** the minimum switches from $\theta = 0$ to $\theta = \pi/2$.

The finite temperature mean field phase diagram determined using equations (17) and (18) is plotted in figure **4.** We expect this **to** be qualitatively correct for values of *w* and A sufficiently small that the perpendicular antifemmagnetic phases **are** not important and sufficiently large that Mermin-Wagner fluctuations do not destroy the ferromagnetism at temperatures significantly above zero. Rather small values of ω and λ , even within the range given in equation **(13). are** plotted **so** that the former of these conditions is most likely to be satisfied (for indications that the latter is likely to be satisfied **see** figure *5* and its discussion) and because most real materials will probably lie in this range.

Solution of a mean field theory accommodating the large *n* **STRIPEn** phases (the only physical ones) would require considerable computational effort, and has not been attempted.

Figure 4. Mean field *T* against λ phase diagrams for small λ , ω monolayers. (a) shows the phases for $\omega = 0.01J$, and (b) shows the variation of the transition temperatures for some **physically likely values of** ω **. All quantities are in units of** J **.**

6. Computer simulation

The most powerful computing hardware available to us was a massively parallel (32×32) **[8+ 11-bit** processors) **AMT** DAP *5* 1OC. The architecture of this machine. is Single Instruction Multiple Data **(SIMD),** allowing efficient performance of an algorithm which exhibits data parallelism (the **same** operation may be performed **on** many *data* simultaneously), but not one which only exhibits algorithmic parallelism (several qualitatively different operations may be **performed** simultaneously).

For computer simulation of an idealized spin system such **as** this, the obvious method is Metropolis Monte Carlo **[29,301,** but a problem **arises** with data parallelization of the Monte Carlo algorithm.

Conventionally in Monte Carlo simulation, each step (modification of the spin state) **requires** making an **alteration** to **a** single spin and assessing the difference *611 this makes* in

the energy of the system. Providing that the change in energy associated with a spin σ_i at site *i* is not affected by the state of a spin σ_i at site *j*, then attempted updates can be made to the states of spins σ_i and σ_j simultaneously as required. However, if spins σ_i and σ_j are neighbours in terms of interaction range this is no longer possible, since the change in energy associated with the modification of spin σ_i is undefined if spin σ_i is simultaneously being modified so that it has no determinate state. Formally, spins σ_i and σ_j can be updated simultaneously only if:

$$
\frac{\partial^2 \mathcal{H}}{\partial \sigma_i \partial \sigma_j} \equiv 0. \tag{19}
$$

Thus, except in a (rather uninteresting) non-interacting spin system, it is not possible to update all spins in parallel. Some parallelization can be performed, however, by splitting the set **S** of all sites *i* into subsets \mathbf{R}_i such that:

$$
\forall i, j \in \mathbf{R}_l, \ i \neq j \ \colon \ \frac{\partial^2 \mathcal{H}}{\partial \sigma_i \partial \sigma_j} \equiv 0 \tag{20}
$$

and then for each *l*, the spins at all the elements of \mathbf{R}_l can be updated simultaneously. If this splitting can be done in such a way that the size of all the subsets \mathbf{R}_l are multiples of the number of processors (in our case **1024)** then the problem can be parallelized with maximal efficiency (i.e. without leaving any processors idle). Typically it is convenient to perform this decomposition into subsets so that all \mathbf{R}_l are equivalent (mappable onto each other by translation) and **as** large **as** possible. **A** simple example *of* such a procedure is the case of nearest-neighbour interactions on a two-dimensional square lattice, in which case the lattice may be decomposed into two square sublattices \mathbf{R}_1 and \mathbf{R}_2 like the black and white squares on a chess board.

As the range of interactions becomes longer this lattice decomposition procedure becomes more difficult. An interaction of infinite range (one in which there is no cut*off)* may be defined as follows: α , be decomposed into two some a chess board,

ange of interactions becomes difficult. An interaction of fined as follows:
 $\forall i, j \in S, \frac{\partial^2 \mathcal{H}}{\partial \sigma_i \partial \sigma_j} \neq 0$

$$
\forall i, j \in \mathbf{S}, \ \frac{\partial^2 \mathcal{H}}{\partial \sigma_i \partial \sigma_j} \neq 0 \tag{21}
$$

and in this case the largest subset \mathbf{R}_l that can be selected is one site. We therefore conclude that data parallelization of conventional Monte Carlo simulation of systems with infinitely long-ranged forces is not possible.

This is an important fact which we have not seen pointed out elsewhere, and it will become more important **as** parallelization becomes a more common way of increasing computer power **[31].**

The only solution appears to be an 'unconventional' Monte Carlo in which each trial modification in the state of the system would consist *of* a small change in each *of* the spins rather than a large change in a single spin. Although this achieves data parallelism so that no processors need lie idle it will not allow a high rate of lattice updates per unit time, so it **seems** unlikely that it will be. efficient. It may however deserve further investigation **as** a method of introducing spin waves rather than spin flips **as** trial spin state modifications.

For this reason we simulated using the molecular dynamics method **[30]** instead. Molecular dynamics is an inherently parallel method and does not present the same problems as Monte Carlo methods. For spin systems such **as** ours this involves introducing artificial dynamics and is probably slightly less efficient than Monte Carlo methods. but seems otherwise satisfactory. Some test Monte Carlo simulations were performed (on a scalar processor) for comparison, and good agreement between the Monte Carlo and molecular dynamics results was found.

The simulation was performed on a square $L \times L$ cell with periodic boundary conditions, and an Ewald sum **[32]** was used to take account of its periodic images for evaluation of the long-ranged dipolar interaction energy.

Unfortunately, the simulations produced no conclusive **results.** The reason **for** this was that the correlation length ξ became larger than the linear dimension L of the simulation box at temperatures significantly above the Curie temperature. Simulation results at lower temperatures could then not be trusted, so that neither *of* the transitions of interest was represented reliably by the calculations. That this divergence could not be assumed to be in the immediate vicinity of T_c was demonstrated by the fact that ξ became comparable with *L* at $T \approx 0.75$ for the system $\omega = \lambda = 0$, the 2D Heisenberg model, for which T_c is known to be zero **[7,8].** It is therefore clear that accurate results are not available near enough the critical region for the usual finite size scaling methods to be of use.

Test simulations, not reported here (but see **[33]),** were performed in various regions of the phase diagram and in no case did they lead to reliable conclusions concerning this system. By careful analysis of the results, particularly of the spin-spin correlation function for the case $\lambda = \omega = 0$, it seemed likely that the simulation box size used was too small by several orders of magnitude. We therefore regretfully conclude that a computer simulation of this type is unhelpful in investigating this system at finite temperatures.

However, **as** mentioned in section 3 simulation results provided important indications about the ground states.

7. **Discussion** and comparison with other results

We **are** not familiar with any other work on this or similar systems which has identified the striped or chequered perpendicular antiferromagnetic phases shown in figure 2, although our simple energy considerations leave us in no doubt that they are more stable at low temperatures than the ferromagnetic phases usually assumed to dominate the phase diagram, and our simulations suggest that they are indeed ground states of the system. The lack of consideration of these phases in previous studies is no doubt a consequence of the fact that for values of the anisotropy parameters corresponding to real physical materials these phases are not obvious, but we reiterate from section **3** that it seems possible to *us* that the limit of large-striped phases extends down to the limit of small ω , in which case the perpendicular ferromagnetic state would not **be** stable for any finite value of *w.* The fact that perpendicular ferromagnetism is seen experimentally could be compatible with this as a consequence of finite sample size, or of external fields applied for magnetization measurements which could act to stabilize the ferromagnetic phase.

Conceming finite temperature behaviour *of* the system, however, since simulations have failed to provide positive results, we have only the mean field ones of section 5. The mean field values for $T_C(\lambda, \omega)$ and $T_R(\lambda, \omega)$ shown in figure 4 can be compared with those predicted by other theories.

We consider first the work of Jensen and Benneman **[2,3].** Using a rather simplified mean field approach, they find a reorientation transition in agreement with our expectations, but they predict that the transition will, under certain circumstances, be continuous, proceeding via a canted magnetic state of which we see no evidence. They also predict

Figure *5.* **Polyakov renormalization goup phase diagram for monolayer from [41 (labelled solid** lines), compared with the mean field phase diagram from figure $4(b)$ (unlabelled broken curves). **All quantities are in unitr of** *J,*

that the magnetic order will decay in a range of 20–30 K around T_R . However, their model treats the dipolar and uniaxial anisotropies in the same way, effectively as single-site terms, merely reversing the sign between the tWo **[3],** and this is quite inadequate since the dipolar interaction only stabilizes the ferromagnetic state in virtue of its long-range nature. **Our** mean field theory treats the dipolar interaction correctly **as** pairwise and long-ranged and is in agreement with their prediction *of* a reorientation transition, although not with the other details mentioned above.

Pescia and Pokrovsky **(41** have applied an adaptation of a renormalization group method developed by Polyakov **[34].** This, unlike the simple mean field theory, correctly predicts the absence of finite temperature magnetization for the *20* Heisenberg model, so the crossover between this case and the small but finite ω , λ case is of particular interest to us. Expressions for the transition temperatures from this theory **are as** follows:

$$
T_{\rm C} = \frac{4\pi J}{8 + \ln J - \ln \lambda} \tag{22}
$$

$$
T_{\rm R} = \frac{\pi J (4\lambda - 3A\omega)}{\lambda (\ln J - \ln \lambda)}
$$
(23)

A phase diagram corresponding to these expressions is plotted in figure *5* where it is also compared with the mean field **mult** already given in figure **4@).** Concerning the crossover between isotropic and anisotropic behaviour we see that it is very rapid: only very small values of λ are needed to stabilize the ferromagnetism even at high temperatures and to bring *TC* to near its mean field value of the order *of* J. Thus we might suppose that the mean field theory which is unable to describe the instability of the pure isotwpic case **is** a reasonably good approximation even for the rather small ω , λ of interest to us.

This **is** encouraging, but certainly there **are** doubts **as** to the validity of equations **(22)** and **(23).** It is quite clear that within this theory the dipolar anisotropy is not dealt with

correctly, since T_c is independent of ω , and in particular $(T_c)_{\alpha=0} \equiv 0$ for all values of ω , i.e. the dipolar interaction is not capable of stabilizing the ferromagnetic state. This is certainly not the case **[261.**

Furthermore, the recent work of Levanyuk and Garcia 1351 claims that the Polyakov renormalization procedure is inappropriate for anisotropic spins at low temperatures. Levanyuk and Garcia investigated this model themselves using an ingenious low temperature expansion, and found no evidence for a reorientation transition. This is contrary to our expectations, based on the persuasive entropic arguments emphasized by Jensen and Benneman as well **as** our own mean field results. Since the approach of Levanyuk and Garcia is a low temperature one, the status of the mean field theory at high temperatures is somewhat uncertain. It may also be relevant for the comparison of results that the model explored here is classical in contrast to the quantum mechanical system of [351.

8. Conclusions

We have modelled a magnetic monolayer with a nearest-neighbour ferromagnetic exchange interaction, a long-range dipolar interaction, and a single site spin-orbit-type anisotropy.

We have mapped exactly most of the zero-temperature phase diagram and we have discovered, rather unexpectedly, a series of perpendicular antiferromagnetic phases which to our knowledge were not previously known *to* exist. These phases **are** most pronounced for values of the interaction parameters much larger than those we would expect for a **real** monolayer, but depending on the behaviour in the limit of large modulation length of the antiferromagnetic phase boundaries, which we have been unable to determine, these phases may replace the ferromagnetic perpendicular state **as** the ground states of some real monolayers.

We have failed to perform useful simulations of this system, and draw the tentative conclusion that straightforward Monte Carlo or molecular dynamics simulations of practicable sizes cannot be used to investigate it. We note also the important technical point that data parallelization of conventional Monte Carlo simulation of systems with infinitely long-range interactions is not possible.

We have formulated and solved, for physically realistic values of the anisotropy parameters, a finite temperature mean field theory which treats the dipolar interaction correctly **as** pairwise and long-ranged, and this exhibits an entropically driven first order reorientation transition at a temperature T_R from the perpendicular to the in-plane ferromagnetic state, and a Curie temperature *Tc.* In predicting a reorientation transition the mean field theory is in agreement with the previous work of Pescia and Pokrovsky **[4]** and of Jensen and Benneman **[2,3],** but not with that of Levanyuk and Garcia [35]. Our disagreement with [35] may be a consequence of our use of classical, rather than quantum mechanical spins, although it should be remembered that the treatment of Levanyuk and Garcia is essentially a low temperature one. We feel on entropic grounds that the reorientation transition ought to exist **121,** but have no proof that the mean field theory predicts the correct behaviour in this respect.

Acknowledgments

We would like to acknowledge several useful conversations with **Dr** M P Allen. MBT was financially supported by a **SERC** studentship and the calculations were canied out on the departmental **DAF'** provided by the Computational Science Initiative.

References

- [I] N&l **L** 1953 Comptes *Rendvr* **237** ¹⁴⁶⁸
- 121 lensen PI and Bennemann **K** H 1990 Phys. *Rev.* B **42** 849
- [3] lensen P **J** and Bennemann **K H** 1992 Preprinr
- [4] Pescia D and Pokrovsky V L **1990** Phys. Rev. *Lett.* **65 20**
- [SI Gay I G and **Richler** R 1987 *1.* Appl. Phys. **61** 3362
- (61 **Guo G Y.** Temmerman W M **and Ekn** H 1991 *J.* Phys: Cnndens. *Mailer* 3 **8205**
- [7] Mermin N D and Wagner H 1966 Phys. Rev. Lert. **17** 1133
- [E] Memin N D 1968 Phys. Rev. **176** 250
- [9] Shenker **S H** and Tobochnik **J** 1980 Phys. Rev. **B 22** *4462*
- [IO] Kawabata C and Bishop A R 1980 *Snlid Srure* Cnmn. 33 453
- [I 11 Gunman A 1 1978 *J.* Phys. A: *Murh* Gen. **11** 545
- 1121 Watson R E. Blume **M** and Vineyard G H 1970 Phys. Rev. B **2** 684
- 1131 Takahashi M 1987 Phys. Rev. B **36** 3791
- [I41 Fukugita M and Oyanagi **Y** 1983 Phys. *Lett.* **123** 871
- **[IS]** Koslerlilz M and Thouless D I 1973 *J.* Phys. *C: Solid* Stare Phys. **6** 1181
- [16] Kosterlitz M 1974 *J. Phys. C: Solid State Phys.* 7 1046
- [I71 Miyashita **S.** Nishimori H, **Kumda A and Swki** M 1978 *Prop. Theor.* Phys. **60** 1669
- [I81 Tobochnik J **and** Chester **G** V 1979 Phys. Rev. **B** *M* 3761
- [I91 Van Himbergen J **E** and Chakmvarty **S** 1981 Phys. Rev. B 23 359
- **[201** Gradmann U 1986 *J. Mugn. Magn.* Muter. **54-51** 733
- [211 Pappas D P. **Kmmper** K P and Hopsrer H 1990 Phys. Rev. *Leii.* **64** 3179
- [221 **Krebs** J 1, **lonker** B T and *Prim* **G A** 1988 *J.* Appl. Phys. **63** 3467
- [23] Stampanoni M. Vaterlaus A. Aeschlimann M and Meier F 1987 Phys. Rev. Lett. 59 2483
- [24] Ducastelle F 1991 Order and Stability in Alloys (New York: North-Holland)
- 1251 Herring *C* and **Killcl** C 1951 Phys. *Rev.* 81 869
- E61 Stamps **R** Land Hillebrands B 1991 Phys. Rev. B **44** 12417
- [271 Rushbrooke B **S.** Baker G A and Wood P J 1974 *Phase* Transitions and Criticul Phenomew vol 111, *ed* C **Domb** and M **S Green (London:** Academic)
- [28] Massalski T B 1986 *Binary Alloy Phase Diagrams* (Menlo Park, OH: American Society for Metals)
- [291 Metropolis **M.** Rosenbluth A **W,** Rosenbluth M N, Teller A H and Teller E 1953 *J.* Chem. Phys. **21** 1087
- [301 Allen M P and Tildesley D J 1987 Cnmpurer Simulation *of* Liquids *(Oxford* Oxford University **Press)**
- [31] Charles D 1992 *NwSci.* **135** 1837 26
- 1321 Heyes D **M** 1981 *Surf Sci. Len.* **110** MI9
- (331 Taylor **M B** 1992 PhD Thesis University of Bristol
- [34] Polyakov A M 1975 Phys. *kil.* **59B** 79
- [35] Levanyuk A P and Garcia N 1992 J. Phys: Condens. Matter 4 10277