Efficient optimization method for finding minimum energy paths of magnetic transitions

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Abstract

Efficient algorithms for the calculation of minimum energy paths of magnetic transitions are implemented within the geodesic nudged elastic band (GNEB) approach. While an objective function is not available for GNEB and a traditional line search cannot be performed, the use of limited memory Broyden–Fletcher–Goldfarb–Shanno (LBFGS) and conjugate gradient algorithms in conjunction with orthogonal spin optimization (OSO) approach is shown to greatly outperform the previously used velocity projection and dissipative Landau–Lifschitz dynamics optimization methods. The implementation makes use of energy weighted springs for the distribution of the discretization points along the path and this is found to improve performance significantly. The various methods are applied to several test problems using a Heisenberg-type Hamiltonian, extended in some cases to include Dzyaloshinskii–Moriya and exchange interactions beyond nearest neighbours. Minimum energy paths are found for magnetization reversals in a nano-island, collapse of skyrmions in two-dimensional layers and annihilation of a chiral bobber near the surface of a three-dimensional magnet. The LBFGS-OSO method is found to outperform the dynamics based approaches by up to a factor of 8 in some cases.

Keywords: minimum energy paths, geodesic nudged elastic band method, magnetism, skyrmions

(Some figures may appear in colour only in the online journal)

1. Introduction

An estimation of the thermal stability of a magnetic state can be of critical importance, for example in the context of various magnetic devices. Each (meta) stable magnetic state corresponds to a local minimum on the energy surface characterizing the system. When a magnetic element is small enough and the temperature high enough, thermal fluctuations due to the coupling to a heat bath can induce magnetic transitions. The lifetime of a state at a given temperature can be estimated using harmonic transition state theory \cite{1–3}. In addition to the local minimum characterizing the initial state, the relevant first order saddle points on the energy surface need to be found in order to estimate the activation energy and entropy for each transition mechanism and thereby estimate parameters in the Arrhenius rate expression. This can be accomplished using the geodesic nudged elastic band (GNEB) method \cite{4} for finding
minimum energy paths (MEPs) between the given initial state and each one of the final states. A maximum in energy along an MEP corresponds to a saddle point on the energy surface. The MEP may have more than one maximum and can reveal unexpected metastable states as intermediate minima, but for the purpose of lifetime estimation the highest maximum is the most relevant one. Once the MEP has been estimated closely enough to reveal its shape and the location of the highest maximum has been established well enough, it can be efficacious enough to refine the MEP there [5] or use a saddle point search method to converge more thoroughly on the maximum [6].

Previous implementations of the GNEB method have mainly used dissipative spin dynamics or velocity projection algorithm based on spherical coordinates [4, 7–11]. The calculations can require substantial computational effort, especially when long range interactions are included or when each energy evaluation requires self-consistent-field iterations such as in density functional theory (DFT) [12] or non-collinear Alexander-Anderson calculations [13]. It is, therefore, important to develop an implementation of GNEB that requires as few energy and energy gradient evaluations as possible to obtain convergence to an MEP. For rearrangements of atoms, for example in chemical reactions and diffusion events, well-documented approaches for efficient optimization of MEPS have been developed [14–16], but for calculations of magnetic systems less work has been done. The difficulty arising for magnetic systems is an additional constraint on the length of the magnetic moments. As a result, conventional optimization algorithm cannot be applied without significant modification.

A recently proposed approach to the problem of finding local minima, the orthogonal spin optimization (OSO) [17], corresponds to rotations of the magnetic vectors on a sphere and can be combined with algorithms developed for unconstrained optimization. The rotations are performed in Euclidean space and do not require the introduction of variables confined to a sphere. Here, we apply the OSO approach to GNEB calculations of MEPS and compare the performance of various optimization methods. The combination of OSO with the limited-memory Broyden–Fletcher–Goldfarb–Shanno (LBFGS) optimization method is found to be most efficient in that it requires the fewest energy and gradient evaluations to reach convergence to MEPS apart from the case where the transition path levels off from an upward trajectory, and a first order saddle point is undefined. In the latter case, the conjugate gradient algorithms appeared to be most efficient. These methods offer a significant improvement over previously used methods for GNEB calculations. The algorithms have been implemented in the LAMMPS software which has been used in the present calculations [18].

2. Methodology

Two types of optimization methods are tested here. The first type is based on equations of motion, either dissipative Landau–Lifshitz (dis-LL) dynamics or effective particle dynamics using velocity projection optimization (VPO) [4]. The other type involves the OSO approach [17] in combination with either conjugate gradient (CG) or LBFGS optimization [22]. The presentation of these methods is brief here as they have been presented in detail elsewhere in the context of finding local minima on the energy surface. References are given to publications that provide more detail. The GNEB method is also reviewed briefly for completeness.

2.1. Dissipative spin dynamics

Minimization of the energy of magnetic systems is often carried out using the Landau–Lifshitz equation of motion with only the dissipation term included

\[
\frac{d\tilde{s}_i}{dt} = -\alpha \tilde{\delta}_i \times \tilde{s}_i \times \tilde{\omega}_i, \tag{1}
\]

where \(\alpha\) is a damping parameter and

\[
\tilde{\omega}_i = -\frac{1}{\hbar} \frac{\partial E}{\partial \tilde{s}_i}, \tag{2}
\]

with \(\hbar\) being a reduced Planck constant. This is referred to as dis-LL method and it generates a steepest descent path on the energy surface [17]. The implementation in the LAMMPS software [18, 20] is based on a symplectic integrator with adaptive time step [20]. At each iteration the time step is

\[
\Delta t = \frac{2\pi}{\kappa |\tilde{\omega}_{max}|}, \tag{3}
\]

where \(|\tilde{\omega}_{max}|\) is the maximum precession frequency in the system, and \(\kappa\) is a parameter. Results are presented here for two values of \(\kappa\), 10 and 20. The smaller value gives larger time step and faster convergence but can lead to instability. An even smaller value, \(\kappa = 5\), leads to convergence problems.

2.2. Velocity projection optimisation

Another way to generate a steepest descent path to a minimum on the energy surface is to use effective equations of motion that mimic those of a particle. A pseudo mass is given to the spin and the orientation and velocity calculated using a discretization of particle equations of motion, such as the velocity Verlet algorithm [21]. In order to reach the minimum, a projection is applied to the velocity associated with spin \(i\) in such a way that at each step only the component of the velocity in the direction of the force, the negative of the gradient of the energy with respect to the \(i\)th spin projected on the tangent space, \(\tilde{f}_i = -\mathcal{P} \left( \frac{\partial E}{\partial \tilde{s}_i} \right)\), is kept when the inner product of velocity and force is positive, while the velocity is zeroed if the inner product is negative

\[
\beta = \sum_i \tilde{f}_i \cdot \tilde{v}_i / \sum_i \tilde{f}_i \cdot \tilde{f}_i \tag{4}
\]

if \(\beta < 0\) then set \(\beta = 0\) \(\tag{5}\)

\(\tilde{v}_i \leftarrow \beta \tilde{f}_i\) \(\tag{6}\)

In this way, the pseudo-particle can accelerate when the gradient keeps pointing in a similar direction, but the velocity is zeroed when the particle has gone past the minimum.
The algorithm is analogous to a variable time step algorithm tracing out the steepest descent path. The VPO method was used in the original implementation of the GNEB method and more detail is provided in reference [4]. Here, the VPO has been used in conjunction with OSO as described in reference [17] and briefly discussed below.

2.3. Orthogonal spin optimization

In orthogonal spin optimization (OSO) [17], the spin directions are updated according to the following iteration process

$$\dot{s}_i \leftarrow e^{-P_i^T} s_i$$

where $P_i$ is a skew-symmetric matrix corresponding to rotation towards the energy minimum. For a gradient descent algorithm, $P_i$ is

$$P_i = \begin{pmatrix} 0 & -g_{ic} & g_{iy} \\ g_{ic} & 0 & -g_{ix} \\ -g_{iy} & g_{ix} & 0 \end{pmatrix}, \quad \vec{g}_i = \dot{s}_i \times \frac{\partial E}{\partial \vec{s}_i}$$

In order to improve the performance of the gradient descent algorithms (dis-LL and VPO) as well as the CG algorithm (without line search) in GNEB calculations, the gradient is scaled to make it dimensionless and the magnitude adjusted according to the adaptive time step given by equation (3)

$$\vec{g}_i \leftarrow \Delta t \frac{\vec{g}_i}{\hat{h}}$$

while for the LBFGS algorithm a maximum-rotation parameter is used [17]

$$P_i \leftarrow P_i \frac{\theta_{\text{max}}}{\theta_{\text{rms}}} \quad \text{if} \quad \theta_{\text{max}} < 1,$$

where $\theta_{\text{rms}} = \sqrt{-\sum_i \text{Tr} \left[ P_i^T \right] / 2N}$, $N$ is the number of spins and $\theta_{\text{max}}$ is a parameter, chosen to be $J\tau/300$. This makes the LBFGS algorithm stable and can be particularly important in the beginning of a minimization since the line search procedure is not performed. A description of OSO combined with CG and LBFGS optimization for finding local energy minima is given in reference [17]. Here, we extend that work to calculations of MEPs. The Fletcher–Reeves CG algorithm has been used.

2.4. Geodesic nudged elastic band

For completeness, we briefly review the GNEB approach. It is an extension of the nudged elastic band that is frequently used in calculations of MEPs for atomic rearrangements, such as chemical reactions and diffusion events [23, 24]. There, a path connecting the initial state minimum to the final state minimum is represented with a set of discrete replicas of the system, referred to as images. This provides a discretized representation of a path that is initially started by some interpolation between the initial and final states and then converges to an MEP using some iterative optimization algorithm.

The distribution of the images, i.e. the discretization points, is controlled with a spring force that acts only along the path and only between adjacent images. GNEB takes into account that the variables represent orientation of vectors such as magnetic momenta rather than Cartesian coordinates of particles.

The effective gradient acting on spin $i$ of image $\nu$ is

$$\vec{h}_{\nu} = \vec{h}_{\nu}^\perp + \vec{h}_{\nu}^\parallel$$

where $\vec{h}_{\nu}^\perp = \frac{\partial E}{\partial \vec{s}_{\nu}} \cdot \bar{\mathcal{P}} \hat{t}_{\nu}$

and $\vec{h}_{\nu}^\parallel = [k_\nu D(\nu + 1, \nu) - k_{\nu-1} D(\nu, \nu - 1)] \hat{t}_{\nu}$.

Here, $\mathcal{P}$ is a projection operator, $\bar{\mathcal{P}} = \mathcal{P} / |\mathcal{P}|$, $D(\nu + 1, \nu)$ the geodesic distance between images $\nu + 1$ and $\nu$, $k_\nu$ a spring constant and $\hat{t}$ the unit tangent to the path estimated from adjacent higher energy image [27]. For more detail see reference [4].

In previous implementations of GNEB the spring constant has been taken to be the same for all pairs of images. In the present implementation, the spring constants are chosen to be energy weighted in order to obtain better resolution of the path in high energy regions since the main goal is to find the energy maximum along the path.

The spring constant is evaluated from

$$k_\nu = \begin{cases} k_{\text{max}} \Delta k \left( \frac{E_{\text{max}} - X_\nu}{E_{\text{max}} - E_{\text{ref}}} \right) & \text{if} \quad X_\nu > E_{\text{ref}}, \\ k_{\text{max}} - \Delta k & \text{if} \quad X_\nu \leq E_{\text{ref}}. \end{cases}$$

Here, $X_\nu = \max(E_\nu, E_{\nu+1})$ and $E_{\text{ref}}$ is the energy of the higher end-point of the path. The magnitude of the spring constants increases linearly towards the current maximum along the path. This is found to significantly improve the rate of convergence especially in the LBFGS calculations. Without this scaling, a larger number of images are needed to represent the path in order to obtain better resolution of the path. The spring forces should be of similar magnitude as the energy gradient in order to obtain smooth convergence. Here, we chose the minimum value of $k$ to be one tenth of the exchange parameter, $J$, and it increases linearly by up to a factor of 2 according to equation (14).

In GNEB calculations, the gradient vector used in the minimization algorithms should be replaced by the effective GNEB gradient, equation (11),

$$\frac{\partial E}{\partial \vec{s}_{\nu}} \leftarrow \vec{h}_{\nu}$$

and all scalar products in the minimization algorithms should involve summation with respect to both spin and image indices. In this way the optimization algorithms are applied to the whole chain of images simultaneously.

For the highest energy image, the spring forces are not included, only the effective gradient along the path. It pulls
the image towards the point of maximum energy and this is referred to as the climbing image [25]. The density of images can become different on the two sides of the climbing image. In some cases another image can during the iterative optimization of the path become higher in energy than the climbing image. Then, the assignment a climbing image is changed to the higher energy one. In such cases, the LBFGS algorithm is reset. In the calculations presented here, a climbing image has been used from the beginning of the iterative optimization of the paths.

The initial path between the two local minima is generated by a linear interpolation using Rodrigues’ formula as discussed in reference [4]. Iterative optimization of the path then brings it to the closest MEP. In some cases, two or more MEPs connect the end-point minima. Then the path obtained from an initial linear interpolation may not be the one corresponding to the lowest energy barrier and thereby not represent the most likely mechanism. For complex systems, genetic algorithms can be used to sample the MEPs in order to find the optimal one [26].

In some cases, the initial path is stationary in that the gradient of the energy is zero at the images even though the path does not correspond to an MEP. This occurs for one of the systems used here in the performance tests (system A, see below). The initial path corresponds in this case to an energy ridge and the maximum along the path is a higher order saddle point on the energy surface. In order to push the initial path off the ridge so the iterative optimization can bring the images to an MEP, random noise is added to the spin configurations in the initial path. The unit vector describing the axis of rotation for spin $i$, defined as in equation (16) of reference [4], is then tilted in a random way by

$$\hat{k}_i \leftarrow \frac{\hat{k}_i + \vec{\xi}_i}{|\hat{k}_i + \vec{\xi}_i|},$$

where $\vec{\xi}_i$ is a three-dimensional vector with components generated from random numbers distributed uniformly over the interval $(-\epsilon, \epsilon)$.

### 3. Simulated systems

The energy of the systems simulated in the tests presented here is given by an extended Heisenberg Hamiltonian

$$E = -\sum_{i>j} J_{ij} \hat{s}_i \cdot \hat{s}_j + \sum_{i>j} \vec{D}_{ij} \cdot \hat{s}_i \times \hat{s}_j - \sum_i \mu_i \hat{B} - \sum_m K^m \sum_i (\hat{s}_i \cdot \hat{k}_m)^2$$

where $\hat{s}$ denotes unit vector in the direction of magnetic moment, $J_{ij}$ is exchange parameter for interaction between spins $i$ and $j$, $\vec{D}_{ij}$ a Dzyaloshinskii–Moriya vector, $\mu_i$ the magnitude of spin $i$ in the unit of $\mu_B$ the Bohr magneton, and $K^m$ an uniaxial anisotropy associated with direction $\hat{k}^m$. In the last term sum with index $m$ runs over all anisotropies in the system, for example, easy axis and easy plane. $(i > j)$ denotes sum among first neighbouring spins and each interaction between pair of spins is taken once. The values of the parameters in the Hamiltonian are taken from previous simulations of these systems, as specified below.

MEPs of transitions in five different systems are calculated with the various optimization methods to compare performance. The systems are briefly presented below, and references given to more complete descriptions.

#### 3.1. System A: Fe nanoisland on W(110) surface

A rectangular island consisting of $30 \times 10$ Fe atoms sitting in a commensurate way on a W(110) substrate is simulated in the same way as in reference [3]. Insets of figure 1 show a schematic of the island. The parameters of the Hamiltonian are chosen to be $J = 25.6$ meV (only nearest neighbour interaction), $K_1 = 1.2$ meV corresponding to $k = (0,1,0)^T$, $K_2 = -0.5$ meV corresponding to $k = (0,0,1)^T$, and $D = |D_{ij}| = |\vec{B}| = 0$. The remagnetization takes place by formation of a temporary domain wall in the direction of the narrower island width, as shown in figure 1 (the energy curve is obtained using cubic polynomial interpolation between the GNEB images based on both energy and gradient along the path [4]). A study of the effect of island shape and size on the mechanism and rate of remagnetization has previously been presented and comparison made with experimental data [3].
Figure 2. Minimum energy paths for the collapse of Néel skyrmions in 2D systems and formation ferromagnetic final states. Red stars correspond to GNEB images. (a) Co/Pt (system B). Results of a calculation using 12 images are shown, in order to better describe the path, but the benchmark calculations were carried out with 8 images. (b) PdFe/Ir (system C) where (c), (d) and (e) show the corresponding initial state minimum, saddle point, and final state minimum, respectively. Color indicates the out of plane spin component, blue corresponding to 1, white 0 and red $-1$.

Figure 3. Collapse of a Néel skyrmion in a system of spins with up to fifth nearest neighbor exchange interaction (system D). Left axis and solid line: energy along the path. Right axis and dashed line: projection of the gradient vector along the path tangent. Results of a GNEB calculation with 16 images is shown in order to better resolve the path, but the benchmark calculations were carried out with eight images. The gradient undergoes rapid change near the saddle point, so the larger number of images is needed to obtain convergence with the tighter tolerance of $1 \times 10^{-6}$ J.
3.2. System B: Skyrmion in 2D Co overlayer on Pt

The simulation cell contains $200 \times 200$ atoms subject to periodic boundary conditions in the $x$- and $y$-direction representing a Co overlayer on a Pt(111) surface. The values of the parameters in the Hamiltonian are $J = 29$ meV (only nearest neighbour interaction), $D = 1.5$ meV, $B = 0$, $K = 0.293$ meV, and $k = (0, 0, 1)^T$. This system size and set of parameters have been used in previous studies of the stability of skyrmions in this system [28]. See also reference [29].

3.3. System C: Skyrmion in 2D PdFe/Ir with nearest neighbour exchange

The simulation cell contains $200 \times 200$ atoms subject to periodic boundary conditions in the $x$- and $y$-direction representing a face centered cubic PdFe overlayer on Ir(111). The parameter values are $J = 7.36$ meV (only nearest neighbour interaction), $D = -2.78$ meV, $B = 1.5$ T, $\mu_i = 3$, $K = 0.7$ meV, $k = (0, 0, 1)^T$. This system size and set of parameters have been used in previous studies of the stability of Néel skyrmions in this system [30]. Figure 2(b) shows the MEP between the skyrmion minimum and the ferromagnetic minimum, and insets show the spin configurations at these points as well as the saddle point. The saddle point has significantly higher energy than the skyrmion state minimum in this case.

3.4. System D: Skyrmion in 2D PdFe/Ir with long range exchange

This is the same as system C except that the exchange interaction includes up to 5th neighbours rather than just nearest neighbours, and $D = -3.0$ meV, $B = 0.5$ T. The values of the exchange coupling constants are taken from estimates obtained using density functional theory calculations [30]. A representation of $J$ as a function of distance between the atoms was used as input in the LAMMPS simulation software, see the appendix. The long range exchange interaction together with other parameters used turns out to bring in a challenging feature as the gradient varies rapidly near the saddle point for skyrmion collapse, see figure 3.

3.5. System E: Chiral bobber in 3D system

The simulation cell contains $30 \times 30 \times 30$ atoms subject to periodic boundary conditions in the $x$–$y$ plane representing a chiral bobber located near one of the boundaries, see figure 4. The final state of the transition is the conical state. The parameter values are $J = 1$ meV (only nearest neighbour interaction), $D = 0.45$ meV, $B = 2.8$ T, $\mu_i = 1$. No anisotropy is included for
this system. The direction of the Dzyaloshinskii–Moriya vector is chosen to give Bloch type chirality. This system size and set of parameters is the same as in reported simulation studies in references [19, 31].

4. Results

The results of the performance tests are summarized in table 1. The number of iterations needed to reach convergence using a total of eight images (i.e. six movable images) to represent the GNEB path is reported. For all four optimization methods, there is one energy and gradient evaluation per movable image for each iteration. The OSO-LBFGS method outperforms the other methods for problem B–E, especially those based on equations of motion, namely the dis-LL and VPO methods. In some cases the OSO-LBFGS method is an order of magnitude more efficient than the frequently used dis-LL method. For problem A, however, the OSO-CG algorithm requires fewer iterations than OSO-LBFGS.

The value of the $\kappa$ parameter in the variable time step expression equation (3) has significant effect on the number of iterations needed to obtain convergence with the OSO-GC and dis-LL methods. If the value of $\kappa$ is increased from 10 to 20, the time steps are half as large and the number of iterations correspondingly double. This has, however, only a minor effect on the number of iterations needed for the VPO method. There, the system accelerates when the gradient keeps pointing in a similar direction, as will occur more often when a smaller time step is used. As a result, the algorithm effectively adjusts the magnitude of the displacement. For system A, the number of iterations needed for the VPO algorithm even decreases when a larger value of $\kappa$ is used.

The energy weighted spring forces make it possible to reach convergence with fewer images as the density of images becomes higher near the saddle point. A certain density is needed in order for the tangent estimate in the discrete GNEB representation of the path to be accurate enough. For example, in calculations with OSO-LBFGS for system C, convergence is not reached with equal spring constants and 6 movable images when the tolerance is set to $1 \times 10^{-6}$ J. However, when the number of movable images is increased to 12, convergence can be reached. At the same time the number of iterations required increases to 267 (from 101). The use of energy weighted spring constants, therefore, decreases the computational effort significantly—by a factor of four in this case.

A special issue arises for the magnetization reversal of the Fe nano-island, system A. The initial path is a stationary path, i.e. with zero energy gradient, because of symmetry and it lies along an energy ridge going through a higher order saddle point on the energy surface. It is necessary to break the symmetry in some way for the iterative GNEB calculations to converge to an MEP. The optimal reversal mechanism corresponds to a temporary domain wall in the direction of the narrower width of the island and starting from either one of the two ends of the island. As the domain wall propagates through the island, the energy is practically constant, as can be seen from figure 1. Another MEP corresponds to a temporary domain wall in the direction of the wider width of the island but it corresponds to higher energy barrier. By adding large enough random rotations of the spins in the initial path as described in section 2, equation (16), the iterative optimization of the path converges to either one of the two optimal and symmetrically equivalent MEPs.

Table 1. Number of iterations required in order to converge on minimum energy path for the five systems studied, A–E. The GNEB calculation is considered to be converged when the effective torques evaluated with GNEB gradients given by equation (11) with respect to each one of the spins in all the images has a magnitude below a tolerance of $1 \times 10^{-9}$ J, except for system D where a tolerance of $5 \times 10^{-5}$ J is used, as none of the optimization methods give convergence for the smaller value when 8 images (6 movable) are used. This is because of the sharp variations in the gradient near the saddle point as the skyrmion collapses when long range exchange interactions are included in the Hamiltonian. The results for OSO-CG, VPO and dis-LL were obtained with $\kappa = 10$ in equation (3). Results obtained with $\kappa = 20$ are shown in brackets.

| System | OSO-LBFGS | OSO-CG | VPO | dis-LL |
|--------|-----------|--------|-----|--------|
| A      | 814       | 505 (976) | 2425 (1629) | 1408 (2813) |
| B      | 102       | 313 (623)  | 590 (685)   | 888 (1776)  |
| C      | 101       | 164 (323)  | 610 (643)   | 455 (911)   |
| D      | 89        | 253 (518)  | 676 (747)   | 725 (1448)  |
| E      | 180       | 455 (893)  | 968 (1086)  | 1299 (2610) |

Table 2. Number of iterations needed for convergence in the GNEB calculations of remagnetization in the $30 \times 10$ Fe island on a W(110) surface (system A). The tolerance for convergence is $1 \times 10^{-6}$ J. Results are given for various values of the magnitude of the random rotations of spins in the initial path, $\epsilon$ in equation (16). For the largest value, $\epsilon = 0.1$, results of calculations using two different random seeds are given (one being the same as in table 1). Those calculations both converge on the minimum energy path with a maximum of 4.097 J corresponding to the activation energy, $E_a$. Calculations started from paths with $\epsilon = 0$ or 0.01 converge on the uniform rotation path with a maximum of 14.062 J. The results for OSO-CG, VPO and dis-LL, were obtained with $\kappa = 10$ in equation (3). Results obtained with $\kappa = 20$ are shown in brackets.

| $\epsilon$ | OSO-LBFGS | OSO-CG | VPO | dis-LL |
|------------|-----------|--------|-----|--------|
| 0          | 90        | 84 (160) | 278 (426) | 233 (466) |
| 0.01       | 76        | 92 (173) | 274 (402) | 235 (470) |
| 0.1        | 1044      | 436 (850) | 1847 (1697) | 1230 (2459) |
| 0.1        | 814       | 505 (976) | 2425 (1629) | 1408 (2813) |
When small random displacements are added to the initial path with $\epsilon = 0.01$, the iterative optimizations still converge on the energy ridge. Points on the ridge are at a maximum along the direction of some of the vibrational modes orthogonal to the path, but along all other modes they are at a minimum. With larger random displacements, $\epsilon = 0.1$, the symmetry is broken sufficiently strongly for the optimization to converge on an MEP where the rise in energy is only 4.097 J, corresponding to the activation energy for thermal transitions. The uniform rotation is, therefore, far from being the preferred mechanism for thermally activated magnetization reversals. For small enough islands with less than 14 atoms along each side, the uniform rotation mechanism becomes preferred [3].

The number of iterations needed for convergence is different for different sets of random numbers, generated from a different random number seed, as can be seen from the last two lines in table 2. This is to be expected since the initial paths are different.

5. Conclusion

The results of the calculations presented here show that the OSO-LBFGS optimization method gives the best performance of the four methods tested here, apart from system A where the transition mechanism involves formation and propagation of a domain wall through an island. There, the OSO-CG method turned out to require fewer iterations when the smaller value of $\kappa$ is used. The number of energy and gradient evaluations needed for convergence of the OSO-LBFGS method is in some cases an order of magnitude smaller than for the commonly used dis-LL method. This performance difference is particularly important when calculations are carried out using sophisticated models of the magnetic system, such as long range interactions (such as dipole–dipole interaction) and self-consistent field methods. In such cases the evaluation of the energy and gradients of the energy account for most of the computational effort.

The present implementation of the GNEB method makes use of energy weighted spring constants. While this was proposed two decades ago in the context of atomic rearrangements [25], it has not become commonly used and has not been implemented previously for magnetic systems. It is found to enhance the performance of the optimization methods, especially the OSO-LBFGS method. Partly this is the result of an improved tangent estimate when the density of images is higher near the climbing image. The magnitude of the spring forces can also influence the convergence rate through the approximations of the Hessian in the LBFGS method. There, the spring force and energy gradient are coupled together.

In addition to the information about performance of the various optimization methods, the calculations present here reveal an interesting effect that be present when the long range exchange interaction is included in the Hamiltonian. Near the saddle point for annihilation of a skyrmion, the gradient of the energy changes abruptly, as shown in figure 4. As a result, the climbing image may not converge on the saddle point unless the resolution of the path is high enough and the displacement steps in the optimization algorithm are small enough. The density of images near the saddle point needs to be high enough to resolve the oscillations in the gradient in order for the GNEB calculation to converge.

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Appendix. Long range exchange interaction in system D

In system D representing face centered cubic PdFe overlayer on Ir(111) the exchange interaction includes up to 5th neighbours. The values of the exchange coefficients have been estimated using DFT calculations and are presented in reference [30]. But, in the input for the LAMMPS software [18] the exchange interaction needs to be specified as a continuous function of the distance between the atoms. The DFT values are, therefore, interpolated using the functional form (figure A1)

$$J(r) = 2a \sin(b r) + 2c \sin^3(d r) \over r^2$$

with parameter values

$$a = 11.5852$$
$$b = 2.4973$$
$$c = 8.0713$$
$$d = 1.3404.$$

From this interpolation function, the following set of $J$ values is obtained (the target DFT values from reference are shown in brackets)

$$J_1 = 28.81 (28.8)$$
$$J_2 = -5.05 (-4.96)$$
$$J_3 = -5.21 (-5.38)$$
$$J_4 = 0.91 (1.04)$$
$$J_5 = 1.59 (1.48)$$

$$\eta_{\text{tev}}$$
An interpolation of the values of the exchange coefficient, $J$, calculated using DFT and presented in reference [30] to provide the exchange coefficient as a continuous function of the distance between the two atoms.

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