

**Spirit: Multifunctional framework for atomistic spin simulations**

Gideon P. Müller,1,2,3,* Markus Hoffmann,1 Constantin Dißelkamp,1,3 Daniel Schürhoff,1,3 Stefanos Mavros,1,3 Moritz Sallermann,1 Nikolai S. Kiselev,1 Hannes Jönsson,2 and Stefan Blügel1

1Peter Grünberg Institut and Institute for Advanced Simulation, Forschungszentrum Jülich and JARA, 52425 Jülich, Germany
2Science Institute and Faculty of Physical Sciences, University of Iceland, VR-III, 107 Reykjavik, Iceland
3Department of Physics, RWTH Aachen University, 52056 Aachen, Germany

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The *Spirit* framework is designed for atomic-scale spin simulations of magnetic systems with arbitrary geometry and magnetic structure, providing a graphical user interface with powerful visualizations and an easy-to-use scripting interface. An extended Heisenberg-type spin-lattice Hamiltonian including competing exchange interactions between neighbors at arbitrary distances, higher-order exchange, Dzyaloshinskii-Moriya and dipole-dipole interactions is used to describe the energetics of a system of classical spins localized at atom positions. A variety of common simulation methods are implemented including Monte Carlo and various time evolution algorithms based on the Landau-Lifshitz-Gilbert (LLG) equation of motion. These methods can be used to determine static ground-state and metastable spin configurations, sample equilibrium and finite-temperature thermodynamical properties of magnetic materials and nanostructures, or calculate dynamical trajectories including spin torques induced by stochastic temperature or electric current. Methods for finding the mechanism and rate of thermally assisted transitions include the geodesic nudged elastic band method, which can be applied when both initial and final states are specified, and the minimum mode-following method when only the initial state is given. The lifetimes of magnetic states and rates of transitions can be evaluated within the harmonic approximation of transition-state theory. The framework offers performant central processing unit (CPU) and graphics processing unit (GPU) parallelizations. All methods are verified and applications to several systems, such as vortices, domain walls, skyrmions, and bobbers are described.

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I. INTRODUCTION

Multiscale material simulations have emerged as one of the most powerful and widespread assets in the quest for novel materials with optimal or target properties, functionalities, and performance. Simulations are employed to narrow down the design continuum of devices, to decrease the effort required in designing novel materials, to substitute experiments that seem unfeasible, and to analyze existing experiments and suggest alternatives. They can also provide understanding of the underlying physics on scales ranging from Ångström to millimeters and from femtoseconds to decades.

In this context, spintronics is a very active field where multiscale simulations [1] play an important role for the conceptualization and development of the next-generation data devices [2]. This includes nanoscale magnetic objects like domain walls or nontrivial magnetic textures such as solitons with a time dilemma of 16 orders of magnitude between writing and saving information.

The simulation approach is extremely useful as it links the desired properties to the choice of magnetic materials and their development, giving rise to a large variety of potential applications [3].

Quantum mechanics is the key to understand magnetism from a fundamental level and it is therefore common to use *ab initio* methods, such as density functional theory, to calculate interactions between atoms and various other properties.

Because of the computational complexity of such calculations, they can currently only be applied to magnetic structures in crystals with length scales in the order of 1 nm and cannot be used for time-dependent dynamics simulations on timescales relevant for spintronics.

From *ab initio* methods, one may extract parameters for more approximate, atomistic spin models, such as Heisenberg-type spin-lattice Hamiltonians. There, detailed information about the electronic structure is integrated out to effective parameters describing the interaction between pairs of classical spins, so that simulations of magnetization dynamics can be extended over the timescale of nanoseconds for systems of hundreds of nanometers.

The third level of the multiscale approach in spintronics is the well-known micromagnetic approximation [4] based on the assumption of a continuous magnetization vector field, defined at any point of the magnetic sample, that is valid when changes of the magnetization are much larger in space than the underlying atomic lattice. In contrast, the atomistic spin-lattice model covers the technologically increasingly important length scale from a few to several tens of nanometers.

Here, we present a general-purpose, open-source, i.e., publicly available, framework for atomistic spin simulations called *Spirit* [5]. There are actually a number of computational tools available for the simulation of the time- and
space-dependent magnetization evolution. Among the software packages for micromagnetic simulations, two of the most impactful and widely known ones are OOMMF [6] and MUMAX3 [7]. This software definitely revolutionized the simulation of magnetic properties of materials and the temporal behavior of devices described by the Landau-Lifshitz-Gilbert (LLG) dynamics. Based on the micromagnetic approach, these methods have well-known limitations, e.g., the description of antiferromagnets, frustrated magnets, higher order nonpairwise interactions (e.g., three-spin or four-spin interaction), stochastic spin dynamics and Monte Carlo simulations, etc. Moreover, most micromagnetic software is not interactive or provides quite limited in situ access to the parameters of the modeled system.

Among the atomistic simulation programs, UppASD [8] and VAMPIRE [9] are examples of well-tested tools that provide important functionalities beyond LLG simulations.

The functionality of the aforementioned software can be greatly extended by adding an interactive graphical user interface (GUI) that can be used to control calculations in real time to not only change parameters but also interact with the spin texture as demonstrated, for example, in Ref. [10].

In combination with such a GUI, Spirit unifies various computational methods that are commonly applied to atomistic (and to a certain extent also to micromagnetic) simulations: Monte Carlo and Landau-Lifshitz-Gilbert (LLG) dynamics [11], the geodesic nudged elastic band (GNEB) method [12], minimum mode following [13] (MMF), harmonic transition-state theory [14] (HTST), and the visualization of eigenmodes.

All of these methods are quite distinct, but complementary in nature [15]. For example, LLG dynamics can be used to simulate the time evolution of a magnetic system on a short timescale, while GNEB and/or MMF can be used to find first-order saddle points of the energy landscape, corresponding to transition states for thermally assisted transitions. These calculations can provide important information, such as the energy barrier for the transition, and can be used in HTST to calculate the lifetimes of metastable magnetic configurations over a long timescale. The integration of these methods into a single, uniform framework can lead to a significant increase in productivity.

The following section will introduce the structure of the Spirit software and subsequent sections will detail the aforementioned methods, in order of their complexity, ranked according to the derivatives of the energy required in implementing these methods. The examples provided are mainly related to magnetic skyrmions, which represents one of the most rapidly developing fields in modern nanomagnetism.

II. THE FRAMEWORK

The framework consists of modular components, as illustrated in Fig. 1: a core library for calculations and input/output; an application programming interface (API) layer to abstract the interaction with the code and provide a uniform interface across various programming languages, e.g., C/C++ and PYTHON; and a set of user interfaces to enable fast and easy interaction with simulations, powerful real-time visualization and postprocessing features, for instance, visualization of two-dimensional (2D) and three-dimensional (3D) magnetization vector fields with corresponding isosurfaces and visualization of eigenmodes.

The visualization of Spirit is available as a standalone library called VFRendering [16], which utilizes advanced features of modern OPENGL, e.g., shaders, available since version 3.2.

Note that the images of spin systems in Figs. 3 and 8 have been generated using the graphical user interface of Spirit. Other examples of the visualization features of Spirit can be found in Refs. [13,17–19]. Spirit has also been used for numerical calculations in Refs. [1,13,20,21].

As the API layer is written in the C programming language, many other languages can be used to call the corresponding functions. The core library can thus be used in many different contexts. An illustration of this flexibility is the implementation of an additional, web-based user interface [22], using JAVASCRIPT to call Spirit and WEBGL to display the simulated system.

The desktop GUI can be used to control parameters in the calculation, such as system size or interaction parameters—useful for fast testing and setup—as well as for direct inter-

![Diagram](image.png)
action with the spin textures. The latter is highly useful, for example, in setting up complex initial states \[10\] or rectifying calculations, such as GNEB paths that have diverged from their intended transition.

In order to increase productivity in repetitive or long-timescale calculations, \textit{Spirit} can be used in PYTHON scripts. Such scripts allow one to reproduce all steps which can be taken in the GUI, thus enabling the flexible and effective use of clusters and remote machines. Sample PYTHON scripts can be found in the code repository [5].

Note that the ability to use \textit{Spirit} from PYTHON also enables a straightforward integration into multiscale simulations and workflow automation frameworks, such as the atomic simulation environment (ASE) [23] or the automated interactive infrastructure and database (AiiDA) [24].

Documentation of input, features, and the APIs, as well as examples of usage, can be found online [5].

The spin-lattice Hamiltonian, as well as all implemented methods and solvers, have been abstracted from the specifics of numerical operations, allowing a generic \textit{backend}, which can optionally use OPENMP for CPU parallelization or CUDA for GPU parallelization. The performance of a simple LLG simulation over different system sizes, including dipolar interactions, is shown in Fig. 2.

The performance gain of the parallelization over the single-threaded case is obvious as 10 cores give almost an order of magnitude across a broad range of system sizes and the GPU can give another order of magnitude at larger system sizes. As expected, the speed drops with the system size. Note that when dipolar interactions are included, due to the usage of FFTs, iterations can be slowed down if a side length of the system is not a power of 2.

III. MODEL AND METHODS

A. Hamiltonian

In \textit{Spirit}, we implemented an extended Heisenberg Hamiltonian [25,26] of classical spins \( \mathbf{n}_i \) of unit length located at lattice sites \( i \), giving rise to the magnetic moment \( \mathbf{m}_i = \mu \mathbf{n}_i \). The general form

\[
\mathcal{H} = - \sum_i \mu_i \mathbf{B} \cdot \mathbf{n}_i - \sum_i \sum_j K_{ij} (\hat{\mathbf{K}}_j \cdot \mathbf{n}_i)^2
- \sum_{\langle ij \rangle} J_{ij} \mathbf{n}_i \cdot \mathbf{n}_j
- \sum_{\langle\langle ij \rangle \rangle} D_{ij} \cdot (\mathbf{n}_i \times \mathbf{n}_j)
+ \frac{1}{2} \frac{\mu_0}{4 \pi} \sum_{ij} \mathbf{n}_j \cdot \hat{r}_{ij} (\mathbf{n}_i \cdot \hat{r}_{ij}) (\mathbf{n}_j \cdot \hat{r}_{ij}) - \mathbf{n}_i \cdot \mathbf{n}_j \quad (1)
\]

includes (i) the Zeeman term describing the interaction of the spins with the external magnetic field \( \mathbf{B} \), (ii) the single-ion magnetic anisotropy, where \( \hat{\mathbf{K}}_j \) are the axes of the uniaxial anisotropies of the basis cell with the anisotropy strength \( K_j \), (iii) the symmetric exchange interaction given by \( J_{ij} \) and the antisymmetric exchange, also called Dzyaloshinskii-Moriya interaction, given by vectors \( D_{ij} \), where \( \langle ij \rangle \) denotes the unique pairs of interacting spins \( i \) and \( j \), and (iv) the dipolar interactions, where \( \hat{r}_{ij} \) denotes the unit vector of the bond connecting two spins. Quite often, the number of pairs for the exchange interactions is limited to nearest or next-nearest neighbors only.

In \textit{Spirit}, the implementation of the Hamiltonian (1) does not assume any limitation on the number of or distance between such pairs, meaning that long-ranged and nonisotropic interactions can be considered. Additionally, higher order multispin-multisite interactions [27] are implemented in \textit{Spirit} as quadruples of the form

\[
E_{\text{Quad}} = - \sum_{ijkl} K_{ijkl} (\mathbf{n}_i \cdot \mathbf{n}_j) (\mathbf{n}_k \cdot \mathbf{n}_l) \quad (2)
\]

These can represent the 4-spin–2-site [28] (also called bi-quadratic), the 4-spin–3-site [29], and the 4-spin–4-site [30] (also called “4-spin”) interactions.

Both the system geometry and the underlying lattice symmetry can be chosen arbitrarily by setting the Bravais vectors and basis cells with any given number of atoms. \textit{Spirit} also allows the pinning of individual spins or a set of spins, for instance, those belonging to the boundary layers. One can also introduce defects, such as vacancies and atoms of different types.

1. Dipolar interactions

The dipole-dipole interaction, due to its long-ranged nature, represents the most complex contribution to the Hamiltonian (1). Direct summation over all interacting spins is of complexity \( O(N^2) \), where \( N \) is the number of spins, resulting in dramatic decrease of performance of the simulations. By making use of fast Fourier transforms (FFTs) and the convolution theorem, the computational complexity can be reduced to \( O(N \log N) \). This convolution method is well known in micromagnetic simulations [31], based on a finite difference scheme. To treat arbitrary spin lattices with any given number of atoms in the basis cells, we use an adapted version of this method. In particular, we consider sublattices composed of atoms with the same index in the basis cell. One FFT is performed on each of these sublattices and additional convolutions are required to describe the
interactions between the sublattices. An efficient implementation of this scheme is achieved using high-performance, robust FFT libraries [32,33].

To verify our implementation of dipolar interactions, we compared it to the direct evaluation of the sum for random configurations with spatially nonsymmetric basis cells and checked the convergence of the stray field of a homogeneously magnetized monolayer against the analytically known result.

Here, we show that Spirit correctly reproduces the solution of typical problems, e.g., Ref. [34], by calculating the stray field-induced helicity of a ferromagnetic cube. The helicity is defined as the absolute value of the line integral over the curve $C$ which is composed of the upper edges of the cube:

$$\nu = \frac{|\int_C n \cdot ds|}{f_C|ds|}. \tag{3}$$

In the atomistic case, this is discretized into the appropriate sums.

The energy minimization was performed using a Verlet-like velocity projection method (see Appendix D). The results are shown in Fig. 3. The squared helicity is expected to approach the critical field linearly, so that a line can be fitted to the critical field of $h_c = 0.158$, which matches the expected value of $h_c = 0.159$, as shown in Ref. [34], within 1%. The two insets show illustrations of how the cube will be magnetized at $h = 0$ (left) and $h = 0.1$ (right).

FIG. 3. Helicity (3) of a ferromagnetic cube, composed of 50×50×50 spins on a simple cubic lattice with constant $a = 1\text{Å}$ and nearest-neighbor exchange of $J = 16.86$ meV. The stray-field-induced helicity $\nu$ (circles) and $\nu^2$ (triangles) are shown in dependence on the reduced external magnetic field $h$. Both $\nu$ and $h = B/(\mu_0M_s)$ are unitless parameters, with the saturation magnetization density $M_s$. The fitted curves (solid lines) show the dependence of $\nu^2$ close to the critical field is approximately linear and they give a critical field value of $h_c = 0.159$, which matches the expected value of $h_c = 0.158$, as shown in Ref. [34], within 1%. The two insets show illustrations of how the cube will be magnetized at $h = 0$ (left) and $h = 0.1$ (right).

Any localized solutions with an identical integer index $Q$ belong to the same homotopy class. It means that such vector fields can be continuously transformed into one another—without the appearance of singularities. An example of a large variety of topologically nontrivial solutions with $|Q| \geq 1$ are the skyrmions in the conventional model of chiral magnets, which were recently discussed in Ref. [10]. Topologically trivial magnetic configurations have $Q = 0$ and irrespective of morphology can always be continuously transformed into the collinear state, $\mathbf{n}(r) = \mathbf{n}_0$. Note that the homotopy classification is not applicable to vector field configurations with noninteger $Q$.

To avoid ambiguity, in the calculation of the topological index with Eq. (4) one should follow the sign convention, assuming that the polarity of the solutions always obeys the condition $\mathbf{n}_0 = (0,0,1)$ for $|r| \to \infty$ [35].

2. Topological charge

When the order parameter of the system represents a smooth unit vector field $\mathbf{n}(r)$, defined at any point of 2D space, $r \in \mathbb{R}^2$, the field configurations corresponding to localized solutions $\{\mathbf{n}(r) \to \mathbf{n}_0\}$ for $|r| \to \infty$ can be classified using the topological concept of homotopy.

The whole domain of definition of the order parameter can be mapped to an $S^2$ sphere: $\mathbb{R}^2 \cup \{\infty\} \leftrightarrow S^2$. Note that the space of the order parameter $\mathbf{n}(r)$, which in magnetic systems represents a magnetization field $\mathbf{n}(r) = \mathbf{m}(r)/\mu$, is also a sphere $S^2$. The map $S^2 \to S^2$ leads to a homotopy classification of localized solutions in 2D with the topological invariant

$$Q = \frac{1}{4\pi} \int_{\mathbb{R}^2} \mathbf{n} \cdot (\partial_r \mathbf{n} \times \partial_s \mathbf{n}) \, dr. \tag{4}$$

B. Monte Carlo

The Monte Carlo method is well known in physics and has a broad range of applications [36]. We have implemented a basic METROPOLIS algorithm with a cone angle for the displacement of the spins [11,37]. This requires only the calculation of the energy, making it the most straightforward method of those implemented in Spirit. While it is a useful tool to calculate equilibrium properties, the drawback is that it cannot resolve time-dependent processes.

One iteration of the METROPOLIS algorithm will sequentially—but in random order—pick each spin in the system once and perform a trial step. Trial steps are preformed by defining a relative basis in which the current spin is the $z$ axis and choosing a new spin direction by uniformly distributed random variables $\phi \in [0, 2\pi]$ and $\cos(\theta) \in [0, \cos(\theta_{cone})]$, where $\theta_{cone}$ is the opening angle of the cone.

The trial step is accepted with a probability

$$P = e^{-\Delta E/k_B T}, \tag{5}$$

where $\Delta E$ is the energy difference between the previous spin configuration and the trial step. The cone angle can be set by an adaptive feedback algorithm according to a desired acceptance-rejection ratio.

Using this method, one can, for example, calculate the critical temperature of a spin system. It is known that the isothermal susceptibility is related to the magnetization...
The Landau-Lifshitz-Gilbert (LLG) equation [45,46] is the well-established equation of motion for the dynamical propagation of classical spins. In its explicit form and including spin torque and temperature contributions [47,48], it can be written as

\[
\frac{\partial \mathbf{n}_i}{\partial t} = -\frac{\gamma}{(1 + \alpha^2)\mu_i} \mathbf{n}_i \times \mathbf{B}^{\text{eff}}_i - \frac{\gamma'\alpha}{(1 + \alpha^2)\mu_i} \mathbf{n}_i \times (\mathbf{n}_i \times \mathbf{B}^{\text{eff}}_i) - \frac{\alpha - \beta}{(1 + \alpha^2)} \mathbf{u}_i \times (\hat{\mathbf{j}}_e \cdot \nabla) \mathbf{n}_i + \frac{1 + \beta\alpha}{(1 + \alpha^2)} \mathbf{u}_i \times [\mathbf{n}_i \times (\hat{\mathbf{j}}_e \cdot \nabla)\mathbf{n}_i],
\]

in which the terms correspond to (i) precession, (ii) damping, (iii) precession-like current-induced spin torque, and (iv) damping-like current-induced spin torque. \( \gamma \) is the electron gyromagnetic ratio, \( \alpha \) is the damping parameter, \( \mathbf{B}^{\text{eff}}_i \) is the effective field, \( \beta \) is a nonadiabatic parameter, \( u = J_s P \hat{g} M_s / (2eM_s) \) with \( J_s \) the current density, \( P \) is the polarization of the current, \( e \) is the electron charge, \( M_s \) is the saturation magnetization, \( \hat{\mathbf{j}}_e \) is the electron current normal vector, and \( \nabla \) is \( \partial / \partial r \) the spatial gradient acting on the spin orientation.

The effective field always contains a component related to the energy gradient \( \mathbf{B}^{\text{th}}_i = -\partial H / \partial \mathbf{n}_i \), but in this notation for the LLG equation, the effective field may contain also a stochastic thermal field, i.e., \( \mathbf{B}_i \rightarrow \mathbf{B}^{\text{th}}_i + \mathbf{B}_i^{\text{th}} \), given by

\[
\mathbf{B}^{\text{th}}_i(t) = \sqrt{2D_i \eta_i(t)} = \sqrt{\frac{2\kappa_B T \xi_i}{\gamma}} \eta_i(t),
\]

where the magnitude is given by the fluctuation-dissipation theorem and \( \eta_i \) is white noise, such that the ensemble average and variance of the thermal field fulfill \( \langle \mathbf{B}^{\text{th}}_i(t) \rangle = 0 \) and \( \langle \mathbf{B}^{\text{th}}_i(t) \mathbf{B}^{\text{th}}_j(t') \rangle = 2D_i \delta_{ij} \delta(t - t') \) respectively. To achieve these properties in an implementation, the vectors \( \eta_i(t) \) can each be created from three independent standard normally distributed random values at every time step. Note also that in time-integration schemes, to fulfill the fluctuation-dissipation relation, the thermal field needs to be normalized by the time step with a factor \( 1 / \sqrt{\delta t} \).

For more details on the implementation of the stochastic LLG equation, see, for example, Refs. [49–51] and references therein.

Sampling of the stochastic LLG for the same parameters as shown in Fig. 4 is presented in Appendix E, verifying the implementation and the equivalence of the stochastic LLG and Monte Carlo methods.

In order to evolve a spin system in time according to this equation, quite a few well-known solvers can be applied. In Spirit, currently Heun’s method [11], a fourth-order Runge-Kutta solver, Depondt’s Heun-like method [52], and Mentink’s semi-implicit method B (SIB) [49] are implemented (see Appendixes B and C for details).

These methods can also be used for energy minimization by considering only the damping part of the LLG equation. However, experience has shown that a Verlet-like velocity...
spin in an external magnetic field, as shown in Fig. 5. The error is the difference between the numerically calculated value and the analytical solution (11). Note that the error may depend strongly on the time step and damping. While the Heun method matches well with results shown in Ref. [9], giving an error within $10^{-4}$, the Depondt method shows a lower error of around $3 \times 10^{-7}$ with respect to the analytical solution.

An easy test for the validity of the implemented dynamics solvers is the Larmor precession and the damping of a single spin in an external magnetic field of $B = 1$ T with a damping of $\alpha = 0.1$ and a time step of $dt = 10$ fs, using the Depondt method. The error is the difference between the numerically calculated value and the analytical solution (11). Note that the error may depend strongly on the time step and damping. While the Heun method matches well with results shown in Ref. [9], giving an error within $10^{-4}$, the Depondt method shows a lower error of around $3 \times 10^{-7}$ with respect to the analytical solution.

The errors of the Depondt solver, shown in Fig. 5, match those of an equivalent calculation given in Ref. [9].

In order to verify our implementation of spin-current-induced torques, the results from Ref. [48] on the velocity of a domain wall in a head-to-head spin chain were reproduced for various nonadiabatic parameters $\beta$. The chain is oriented along the $x$ axis and the first and last spins are fixed in $+x$ and $-x$ directions, respectively.

As a subset of the general Hamiltonian (1), the Hamiltonian for this example can be written as follows:

$$\mathcal{H} = -\sum_i K_{ix} n_i^2 + K_{iy} n_{iy} - J \sum_{(ij)} n_i \cdot n_j . \quad (12)$$

The reference provides analytical equations against which the numerical results were checked. In Fig. 6, we show the data for the average domain wall velocity $\langle v \rangle$ over applied current $u$ in normalized units.

The approximate prediction [53] $\langle v \rangle = \sqrt{u^2 - u_w^2} / (1 + \alpha^2)$ fits the results well, as shown in Fig. 6. As expected, we observe the Walker breakdown [54] and a critical effective velocity of $u_c \approx 0.0414$, which is in close agreement with the reported value of $u_c \approx 0.0416$. Note for $\beta = 0.1$ and currents larger than $u_w$, as well as for $\beta = 0$ and currents larger than $u_c$, the wall starts rotating around the $x$ axis.

D. Geodesic nudged elastic band method

When determining the rates of some rare transition events or the lifetimes of metastable magnetic states, LLG dynamics simulations typically are typically unfeasible due to the disparity between the timescales of the simulation and the transition events. An approach to this problem is given by a set of rate theory methods, namely the geodesic nudged elastic band [12] (GNEB) and minimum mode following [13] (MMF) methods together with harmonic transition-state theory [14] (HTST). The latter two are higher order methods,
requiring knowledge of the second derivatives of the energy—the Hessian matrix—and will be described in the following sections.

The GNEB method is a way of calculating minimum energy transition paths between two predetermined configurations. The path is discretized by a number of spin configurations, which in the following are called images.

In order to converge from an initial guess to a stable, energy-minimized path, spring forces are applied along the path tangents, while energy gradient forces are applied orthogonal to the path tangents. The total force therefore reads

\[ F_v^{\text{tot}} = F_v^S + F_v^E, \]

where \( v \) is the image index along the chain, \( F_v^S \) is a spring force, and \( F_v^E \) is an energy gradient force. The forces in this section are \( 3N \)-dimensional vectors.

A simple definition of the spring force, which gives an equidistant distribution of images in phase space, is given by

\[ F_v^S = (l_{v-1,v} - l_{v,v+1}) \tau_v, \]

where \( l_{v,\mu} \) is a measure of distance between images \( v \) and \( \mu \) and \( \tau_v \) is the (normalized) path tangent at image \( v \).

The \( F_v^E \) should pull each image toward the minimum energy path, while leaving the distance to other images unchanged. They can be defined to be orthogonal to the path by orthogonalizing with respect to the tangents

\[ F_v^E = -\nabla E_v + (\nabla E_v \cdot \tau_v) \tau_v, \]

where \( \nabla_v = \partial / \partial n_v \).

The path tangents can be easily approximated by finite differences between spin configurations, but in order to avoid the formation of kinks in the path the definitions given in Ref. [55] should be used.

In order to precisely find the point of highest energy along the minimum energy path, a first-order saddle point of the energy landscape, one can use a so-called climbing image (CI) [56]. Convergence onto the saddle point is achieved by homogeneous interpolation between the initial and final images and a chain with two climbing and one falling images. The climbing images converge onto the saddle points and the falling image converges onto an additional local minimum, so that the energy barriers are known exactly.

The implementation of the GNEB method can be further tested using a conceptually simple process, which has enough degrees of freedom to pose a challenge for convergence: the destruction of a skyrmion tube in a chiral magnetic thin film.

The parameter set is chosen in accordance with a calculation presented in Ref. [57], where a novel particle-like state is shown to emerge along the minimum energy path—the chiral bobber.

The nucleation of a pair of Bloch points, cutting the skyrmion tube in half, is reported, resulting in the formation of one chiral bobber at each surface of the film. In fact, as we show in Fig. 8, also a single Bloch point can be nucleated at one of the film’s free surfaces. For these calculations, the specific parameters are \( J = 1 \) meV and \( D = 0.45J \), meaning that the incommensurate spin spiral has a period of \( L_D = 13.96 \) \( a \).

We note that the conical phase background—corresponding to the ground state of the system—introduces additional modes with little energy cost associated and this can slow the convergence to the minimum energy path.

The climbing-image method [56] was used to converge nearby images onto the maxima along the path. Analogous to a suggestion in Ref. [56], the spring forces were modified to distribute the images evenly along the energy curve. The latter improves the convergence onto the maxima, as the resolution for the finite-difference calculation of the tangents.

![Figure 7](image-url) FIG. 7. An illustration of the GNEB method for a single-spin system (the Hamiltonian and corresponding parameters are given in Appendix G). The two-dimensional energy landscape is shown superimposed on a unit sphere. The initial guess (green), relaxed path (blue), and final path using climbing and falling images (red) are shown.
at the saddle points is increased. As is common to calculate cubic polynomials to interpolate between the discrete points, the segment length of these polynomials can be used for the spring forces between the images. In *Spirit*, an additional parameter is implemented, with which one can set the weighting of energy versus reaction coordinate.

Without the climbing image method, energy barrier calculations may be quite imprecise, especially when the resolution near the maximum is low. This is illustrated by the fact that we observe a ratio of the energy barriers between the collapse of the bobber and the Bloch point nucleation of only 3.3, while Ref. [57]—not using climbing images—reports a ratio of 4.3.

The GNEB calculations reveal a crossover between the two Bloch point nucleation mechanisms, where at increasing field it becomes favorable to nucleate just one Bloch point at the surface. It can further be seen that the energy barrier for the collapse of the bobber goes to zero right below the critical field $H_D$, meaning that—in the frame of this model—it can only be stabilized in the conical phase.

In order to give additional quantitative reference results for this parameter set, the dependence of the energy barrier on the external magnetic field is also presented in Fig. 9.

### E. Harmonic transition-state theory

As certain processes may be too rare or the desired timescale to be simulated too large to allow for dynamical simulations, other approaches are essential in estimating stability and the calculation of lifetimes of metastable magnetic states.

One can employ the well-known transition-state theory [58], which has been used extensively, e.g., in chemical reaction and diffusion calculations [59]. The rate of transitions can be estimated from the probability of finding the system in the most restrictive and least likely region separating the initial state from possible final states—the transition state, sometimes also called dividing surface.

Within the harmonic approximation to transition-state theory [14] (HTST), one can make simplifications allowing the analytical calculation of the transition rate. The rate is then given by an Arrhenius-type law with an exponential dependence on the inverse temperature $T$ and the energy barrier of the transition $\Delta E$, as well as more complicated pre-exponential factors:

$$\Gamma^{\text{HTST}} = \frac{v}{2\pi} \Omega_0 e^{-\Delta E/\kappa_B T},$$

(17)

with

$$\Omega_0 = \sqrt{\frac{\text{det}^{'H^M}}{\text{det}^{'H^S}}} = \sqrt{\frac{\prod_i \lambda_i^M}{\prod_i \lambda_i^S}},$$

(18)

$$v = \sqrt{2\pi \kappa_B T N_0^M - N_0^S V^S} \sqrt{\frac{1}{VM} \sum_i \frac{a_i^2}{\lambda_i^S}},$$

(19)

where the $M$ and $S$ superscripts indicate the minimum and first-order saddle point of the transition. The $\lambda_i$ are eigenvalues of the Hessian matrix (see Appendix H), $V$ are the phase-space volumes of zero modes (if present, otherwise $V = 1$), $N_0$ are the number of zero modes—modes with zero eigenvalue—and $a_i$ are coefficients in the expansion of the velocity along the unstable mode. The primes next to determinants, products, and sums denote that only positive eigenvalues are taken into account.
The factors $a_i$ are in fact velocities: the first row of the dynamical matrix $V$ transformed into the eigenbasis of the Hessian according to
\[ V|^{2N} = \Lambda^T T^T V|^{3N} T \Lambda, \]
where $T$ is a $3N \times 2N$ basis matrix of the tangent space and $\Lambda_{ij}$ denote the matrix of the Hessians eigenvectors (in $2N$ representation, i.e., in the basis $T$). See Appendix H for more information.

The implementation has been verified against UppASD [8], and we additionally present an example for the calculation of the lifetime $\tau$ of an isolated skyrmion in a two-dimensional system. As parameters, we chose $J = 1$ meV and $D = 0.6$ meV and only the radial collapse mechanism is considered, making for a simple structure of the dependence on external field and temperature. Note that this example is purely illustrative and while larger skyrmions would exhibit longer lifetimes, the parameters are chosen to produce a small skyrmion in order to reduce the computational effort. Figure 10 shows the results for an external field varied between 3.5 and 5 T and temperature between 2 and 5 K.

HTST as well as Langers theory [60], which is closely related, have recently both been used to calculate skyrmion lifetimes [61–63]. These calculations show that energy barriers are in general not sufficient for the estimation of the stability of metastable magnetic states and the entropic contributions in the pre-exponential factors can have a significant impact.

There are two translational zero modes at the initial state minimum, while—due to the lattice discretization and the defectlike shape of the skyrmion at the saddle point—there are no zero modes at the saddle point. Consequently, the transition rate prefactor has a linear temperature dependence.

**F. Minimum mode following method**

To find the first-order saddle points on the energy surface, without prior knowledge of the possible final states, the minimum mode following method [13] can be used. The effective force acting on a spin configuration is defined as
\[ F^{\text{eff}} = F - 2(F \cdot \hat{\lambda})\hat{\lambda}, \]
where $F = -\nabla \mathcal{H}$ is the negative gradient of the energy and $\hat{\lambda}$ is the normalized eigenvector corresponding to the lowest, negative eigenvalue of the Hessian matrix of second derivatives. Note that these vectors and the dot product are $3N$ dimensional for a system with $N$ spins.

The calculation of second derivatives requires further attention, as the requirement of constant length effectively constrains the spins $\mathbf{n}_i$ to a submanifold $\mathcal{M}^{\text{phys}} \subset \mathcal{E}$ of an embedding space $\mathcal{E} = \mathbb{R}^{3N}$. As is shown in Ref. [13], the covariant second derivatives, valid at all points of the phase space, can be calculated using a projector-based approach [64]. The corresponding $2N \times 2N$ Hessian matrix can be represented as
\[ H_{ij} = T_j^T \mathcal{H}_i T_j - T_j^T I (n_j \cdot \nabla \mathcal{H}) T_j, \]
where $i$ and $j$ are spin indices, $\mathcal{H}$ is the smooth continuation of the Hamiltonian to the embedding space, $\mathcal{H}_i = \partial^2 \mathcal{H}$, $I$ is the $3 \times 3$ unit matrix, and $T_j$ is a $3 \times 2$ matrix that transforms into a tangent space basis of spin $i$. As the Hessian matrix (22) is represented in the $2N$-dimensional tangent basis, the evaluation of an eigenmode in the $3N$ representation of the embedding space $\mathcal{E}$ requires a transformation back, i.e., $\lambda^{|^{2N}} = T \lambda^{|^{3N}}$.

Further details on the above mathematical concepts and notations can be found in Appendix H.
For a single spin, the energy landscape and force vectors can be visualized easily as the phase space is two dimensional. An illustration of the method is shown in Fig. 11 for a system consisting of one movable spin interacting with a second, pinned spin. The parameters of the Hamiltonian are, relative to the exchange constant $J$,

$$K = 4J, \quad D = (0, 0, 1J),$$

(23)

where the anisotropy $K$ is used to reduce the symmetry of the energy landscape.

The figure illustrates how the minimum mode can be used to invert the right part of the gradient force in order to obtain a force that directs the system to a first-order saddle point.

The test of a larger and far more complex system has been given in Ref. [13], where the minimum mode-following method revealed the existence of a skyrmion duplication mechanism. By defining the force field in the above way, previously unknown transition mechanisms can be found and subsequently used in the calculation of lifetimes.

Applying this saddle-point search method to three-dimensional systems will likely identify an even larger variety of mechanisms, as the additional dimension can significantly increase the amount of possible transitions.

**IV. CONCLUSIONS**

The functionality of a comprehensive simulation framework, *Spirit*, for studies of atomic-scale magnetic systems is presented and various example applications described. It is an open-source software written in the C++ programming language and is available for free under the so-called MIT license (see Ref. [5]). *Spirit* is a very flexible, high-performance, and interactive tool, able to simulate, for example, ferromagnets, antiferromagnets, synthetic antiferromagnets, ferrimagnets, noncollinear magnetic structures, vortices, or skyrmions. Arbitrary geometries and interactions can be described, such as bulk systems, thin films, exchange bias, multilayers, nanotubes, or core-shell nanoparticles. The computational domain can be treated by open and periodic boundary conditions and can be subjected to external magnetic fields, temperature, and spin-current-induced torques. Because it can be used with the PYTHON programming language, *Spirit* can integrate perfectly into multiscale simulations and workflow automation frameworks, such as ASE [23] or AiiDA [24]. It can be used on most common architectures, such as desktop and laptop computers, clusters or supercomputers, and even mobile devices. The calculations can be parallelized both on CPUs and GPUs.

Various simulation methods have been implemented, including Monte Carlo, Landau-Lifshitz-Gilbert dynamics, Langevin dynamics, geodesic nudged elastic band, and minimum mode-following methods as well as the calculations of transition rates and lifetimes within the harmonic approximation to transition-state theory. The basic algorithms of these methods have been outlined, their implementation verified, and applications to several systems, such as vortices, domain walls, skyrmions, and boobers, are described. The parameters of the simulation can be set and modified in real time through a graphical user interface and the output of the simulations can be visualized easily.

We note that a micromagnetic description of the energetics could easily be implemented in *Spirit* and the micromagnetic calculations would then be able to make use of the various simulation methods and visualization features.

**ACKNOWLEDGMENTS**

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**APPENDIX A: DETERMINATION OF TOPOLOGICAL CHARGE FOR SPIN DENSITY ON A LATTICE**

For the proper definition of the topological charge of a discrete lattice of spins $n(x_i, y_i)$, where $i$ runs over all lattice sites, we follow the definition given by Berg and Lüscher [65], and arrive at the following expression:

$$Q = \frac{1}{4\pi} \sum_i A_{ij},$$

(A1)

with

$$\cos \left( \frac{A_{ij}}{2} \right) = \frac{1 + n_i \cdot n_j + n_i \cdot n_k + n_j \cdot n_k}{\sqrt{2(1 + n_i \cdot n_j)(1 + n_i \cdot n_k)(1 + n_j \cdot n_k)}}$$

(A2)

where $l$ runs over all elementary triangles of any triangulated regular lattice and $A_{ij}$ is the solid angle, i.e., the area of the spherical triangle with vertices $n_i, n_j, n_k$; see Fig. 12. The sign of $A_{ij}$ is determined as sign ($A_{ij}$) = sign ($n_i \cdot (n_j \times n_k)$).

The sites $i$, $j$, and $k$ of each elementary triangle are numbered in a counterclockwise sense relative to the surface normal vector $\hat{r}_\perp$ pointing in positive direction of the $z$ axis. The latter means that the numbering should satisfy the condition $\hat{r}_\perp \cdot (r_{ij} \times r_{ik}) > 0$, where $r_{ij}$ is a connection vector directed from lattice site $i$ to $j$.

The parameter $A_{ij}$ can be thought of as local topological charge density, which takes values in the range of $[-\pi/4, \pi/4]$. We illustrate this in Fig. 12, where the definition of the topological charge on a discrete lattice as given in the main text. $A_{ij}$ is the area of a spherical triangle defined by vectors $n_i$, $n_j$, and $n_k$ located at the vertices of a triangle of lattice points (indicated shaded).
$-2\pi < A_i < +2\pi$. According to Berg and Lüscher [65], there is a set of exceptional spin configurations for which $Q$ is not defined but still measurable as $A_i$ in (A2) is defined for all possible spin configurations. The exceptional spin configurations correspond to the case when a spherical triangle degenerates to a great circle $A_i = 2\pi$. In this case, the orientation of $A_i$ becomes ambiguous and the position of these elementary triangles $*i$ are considered as exceptional configurations or topological defects of a two-dimensional magnetic structure.

These topological defects satisfy the following conditions:

$$n_i \cdot (n_j \times n_k) = 0, \quad \text{and} \quad |n_i + n_j + n_k| \leq 1. \quad (A3)$$

The elementary triangle $*i$, for which the condition (A3) is satisfied, can be considered as the position at which the localization of a topological defect takes place. It is important to note that the definition of the topological charge given above remains correct only for spatially extended two-dimensional systems. This means that a topological analysis of the spin structure on a finite-size domain is only defined if periodical boundary conditions are present. In the case of open boundary conditions, strictly speaking, the topological charge is not defined.

**APPENDIX B: HEUN’S SOLVER**

To simplify the following discussion, we write the LLG equation (9) as

$$\frac{\partial n_i(t)}{\partial t} = n_i(t) \times A_i(t, \{n_j(t)\}), \quad (B1)$$

where $\{n_j\}$ is the set of all spins and we keep the explicit time dependence of $A_i$, as the Hamiltonian can be time dependent, for example, when an AC magnetic field is used.

Heun’s method is a common and illustrative way to solve ordinary differential equations (ODEs) by first calculating an intermediate prediction step and then “averaging” to obtain the final approximation. When we denote the time step $\delta t$, for an ODE of the form

$$\frac{\partial y(t)}{\partial t} = f(t, y(t)), \quad y(t_0) = y_0, \quad (B2)$$

the predicted value $y^p$ is first calculated as

$$y^p(t + \delta t) = y(t) + \delta t f(t, y(t)) \quad (B3)$$

and then the approximation for the next step as

$$y(t + \delta t) = y(t) + \frac{\delta t}{2} \left[ f(t, y) + f(t + \delta t, y^p(t + \delta t)) \right]. \quad (B4)$$

When applied to the LLG equation, where $f \equiv n \times A$, this integration scheme obviously does not intrinsically preserve the spin length, requiring the renormalization of the vectors $n_i$ after a given number of iterations, depending on the required precision. Note that Heun’s method falls into the category of Runge-Kutta methods, which function analogously and therefore all have this property.

In order to improve on this, Ref. [52] proposes to make use of the fact that the spins are only allowed to rotate, by writing an appropriate rotation matrix $R_i$, which is calculated directly from the field $A_i$. Applied to Heun’s method, the prediction step (B3) reads

$$n_i^p(t + \delta t) = R_i(A_i(t, \{n_j(t)\})) \cdot n_i(t). \quad (B5)$$

To perform the correction step (B4), one needs the correction field $A^c$, which is calculated from the average of the initial and predicted fields:

$$A^c_i = \frac{A_i(t, \{n_j(t)\}) + A^p_i(t + \delta t, \{n_j^p(t + \delta t)\})}{2} \quad (B6)$$

From this, in turn, the rotation matrix for the correction step $R_i^c(A^c_i)$ is obtained and the final step of the scheme reads

$$n_i(t + \delta t) = R_i^c(A^c_i) \cdot n_i(t). \quad (B7)$$

Higher order Runge-Kutta schemes could apply this approach analogously.

**APPENDIX C: SEMI-IMPLICIT MIDPOINT SOLVER**

The semi-implicit scheme B (SIB), described in Ref. [49], makes use of an implicit midpoint (IMP) structure. In contrast to Runge-Kutta-type schemes, such as described in Appendix B, this intrinsically preserves the spin length.

IMP schemes solve differential equations of the form $y(t) = f(t, y(t))$, $y(t_0) = y_0$ [see Eq. (9) of the main text] and an iteration step is defined as

$$y(t + \delta t) = y(t) + \frac{\delta t}{2} \left[ f(t + \frac{\delta t}{2}, y(t) + \frac{\delta t}{2} f(t, y(t))) \right]. \quad (C1)$$

For the LLG equation (B1) and a time step $\delta t$, this leads us to

$$n_i(t + \delta t) = n_i(t) + \frac{\delta t}{2} \left\{ n_i(t) + \frac{n_i(t) + n_i(t + \delta t)}{2} \times A_i(t + \frac{\delta t}{2}, \frac{n_i(t) + n_i(t + \delta t)}{2}) \right\}. \quad (C2)$$

The SIB scheme uses a predictor $n_i^p$ to reduce the implicitness of the equation above by removing the dependence of $A_i$ on $n_i(t) + \delta t)$. To preserve the spin length, the predictor is obtained with the IMP structure:

$$n_i^p(t + \delta t) = n_i(t) + \frac{\delta t}{2} \left\{ n_i(t) + n_i^p(t + \delta t) \frac{A_i(t, \{n_j(t)\})}{2} \right\}. \quad (C3)$$

Equation (C3) can be rewritten as

$$M \cdot n_i^p(t + \delta t) = M^T \cdot n_i(t) \quad (C4)$$

with the matrix

$$M = I + \text{skew}(A) = \begin{pmatrix} 1 & -A_z & A_y \\ A_z & 1 & -A_x \\ -A_y & A_x & 1 \end{pmatrix}. \quad (C5)$$

The right-hand side of Eq. (C4) can be easily calculated as

$$M^T n_i = n_i + n_i \times A_i =: a. \quad (C6)$$

To solve Eq. (C4), we use Cramer’s rule. The components $n_{i,\alpha}$ with $\alpha = x, y, z$ of $n_i^p$ are calculated with

$$n_{i,\alpha}^p = \frac{\text{det}(M_{\alpha})}{\text{det}(M)}, \quad (C7)$$

where $M_{\alpha}$ is the matrix obtained by removing the $\alpha$-th column.
where $M^\alpha$ is the same matrix as $M$ but column $\alpha$ is replaced with the vector $a$, for example,

$$M^\alpha = \begin{pmatrix} a_x & -A_z & A_y \\ a_y & 1 & -A_x \\ a_z & A_x & 1 \end{pmatrix}.$$  

(C8)

We now use the predictor $n_i^n$ in the IMP step (C2) to calculate $n_i(t + \delta t)$:

$$n_i(t + \delta t) = n_i(t) + \delta t \frac{n_i(t) + n_i(t + \delta t)}{2} \times A_i \left( t + \frac{\delta t}{2} \left\{ n_j(t) + n_j(t + \delta t) \right\} \right).$$  

(C9)

The correction step is analogous to the prediction step [compare Eqs. (C3) and (C9)], meaning that the scheme (C7) can be applied to obtain $n_i(t + \delta t)$, too.

APPENDIX D: VELOCITY PROJECTION SOLVER

This description is derived from Ref. [12]. Verlet-like methods generally find application in solving second-order differential equations of the form $\ddot{x}(t) = F(t, x(t), v(t))$, $x(t_0) = x_0$, and $\dot{x}(t_0) = v_0$, such as Newton’s equation of motion.

One formulation of this method is to increment both the position and the velocity at each time step,

$$x(t + \delta t) = x(t) + \delta t v(t) + \frac{1}{2m} \delta t^2 F(t),$$  

(D1)

$$v(t + \delta t) = v(t) + \frac{1}{2m} \delta t (F(t + F(t + \delta t)).$$  

(D2)

The velocity projection is used to accelerate convergence toward local minima and to avoid overstepping due to momentum. The velocity at each time step is damped by projecting it on the force

$$v \rightarrow \begin{cases} (v \cdot F)/|F|^2, & (v \cdot F) > 0 \\ 0 & \text{else} \end{cases}.$$  

(D3)

Note that the dot product and norm in this equation denote those of $3N$-dimensional vectors.

To apply this scheme to the energy minimization of a spin system, we therefore no longer solve the LLG equation but instead pretend that the spins are massive particles moving on the surfaces of spheres. The force is then simply

$$F_i = -\frac{\partial H}{\partial n_i}.$$  

(D4)

As the method does not conserve the length of the spins, they should be renormalized after each iteration:

$$n_i(t + \delta t) \rightarrow n_i(t + \delta t) / |n_i(t + \delta t)|.$$  

(D5)

Note that this scheme, too, would most likely benefit from the usage of rotations instead of displacements.

APPENDIX E: STOCHASTIC LLG

Instead of Monte Carlo, one can also sample the stochastic LLG equation over time. We present here the results of such sampling for the same system and parameters, as the example shown in Fig. 4. Recall the expected critical temperature $T_c = 1.44 J/k_B \approx 16.71$ K. The energy per spin $E$ and normalized values of the total magnetization $M$, susceptibility $\chi$, specific heat $c_V$, and fourth-order Binder cumulant $U_4$ are shown. The value obtained from the simulation is $T_c \approx 16.92$ K, an agreement with the expectation of 1.2%. The exponent is fitted with $b \approx 0.33$. At each temperature, $2 \times 10^5$ thermalization steps were made before taking $10^5$ samples.

APPENDIX F: GNEB TANGENTS AND FORCES

For spin systems, special care has to be taken due to the fact that the phase space is curved (the spins are restricted to unit spheres; see also Appendix H).

The expression for $l_{\nu,\mu}$ should not be the Euclidean distance norm, but the geodesic (here, the great circle) distance. Further, the tangents $\tau_\nu$ need to lie in the space tangent to their corresponding image. One may correct the tangents for example by a simple projection, orthogonalizing the corresponding three-component subvectors with respect to the spins

$$\tau_{\nu,i} \rightarrow \tau_{\nu,i} - (\nu_{\nu,i} \cdot \nu_{\nu,i}) \nu_{\nu,i}.$$  

(F1)

After this, the tangent needs to be renormalized $\tau_\nu \rightarrow \tau_\nu / |\tau_\nu|$. This tangent projection is illustrated for a single spin in Fig. 14. As the spring forces are constructed from tangent vectors, they are by definition in the tangent space.

Finally, for the energy gradient force, the same scheme as for the tangents can be applied, and we write for each spin

$$F_{\nu,i}^E \rightarrow F_{\nu,i}^E - (F_{\nu,i}^E \cdot \nu_{\nu,i}) \nu_{\nu,i}.$$  

(F2)
This tangent is denoted to the spin configuration so that it correctly points along the path. In a difference calculation, it needs to be projected onto the tangent plane $\tau^\nu$ for a single-spin system. After a tangent $\tau_i^{FD}$ is determined by finite difference calculation, it needs to be normalized, which for a multispin system needs to be performed in $3N$ dimensions.

**APPENDIX G: GNEB PARAMETERS OF THE SINGLE-SPIN SYSTEM**

The energy surface of the single-spin system, shown in Fig. 7 in the main text to illustrate the geodesic nudged elastic band method, is defined for a single spin as a sum of Gaussians of the form

$$ \mathcal{H} = \sum_i \mathcal{H}_i = \sum_i a_i \exp \left[ - \frac{(1 - n \cdot c_i)^2}{2 \sigma_i^2} \right], $$

with parameters given in Table I.

**APPENDIX H: DETAILS ON THE CURVED MANIFOLD**

The following has been detailed in the supplementary material of Ref. [13], but the key ideas are reproduced here. Both the HTST and MMF methods require the calculation and diagonalization of the Hessian matrix. However, when treating Riemannian manifolds, the second derivatives do not have an intrinsic geometrical meaning and therefore need to be treated with special care [66].

In a spin system where the spin length is fixed, the manifold $\mathcal{M}_{\text{phys}}$ of physical states is composed of the direct product of $N$ spheres:

$$ \mathcal{M}_{\text{phys}} = \bigotimes_{i=1}^{N} \mathbb{S}^2 \subset \mathbb{R}^{3N}. $$

Hence, $\mathcal{M}_{\text{phys}}$ is a submanifold of the embedding Euclidean space $\mathcal{E} = \mathbb{R}^{3N}$.

It turns out to be convenient to treat the spins and derivatives with respect to their orientations in a $3N$-dimensional Cartesian representation. This also avoids problems of other representations, such as the singularities which arise at the poles of spherical coordinates. The derivatives in the embedding space $\mathcal{E}$ are readily calculated by extending the Hamiltonian $\mathcal{H}$, which is defined on $\mathcal{M}_{\text{phys}}$ to a function $\mathcal{H}$ on $\mathcal{E}$.

While we denote the gradient taken in the embedding space $\mathcal{E}$ as $\partial \mathcal{H}$, the gradient taken on the manifold $\mathcal{M}_{\text{phys}}$ has to lie in the tangent space to the manifold, which we write as a projection $P_\perp \partial \mathcal{H}$. The Hessian matrix of second derivatives in the embedding space $\mathcal{E}$ is denoted $\partial^2 \mathcal{H}$.

In this extrinsic view onto the spin manifold, the covariant second derivatives can be extracted from a projector approach [64], where for any scalar function $f$ on the manifold $\mathcal{M}_{\text{phys}}$, the covariant Hessian is defined as

$$ \text{Hess}(f(x)[z]) = P_\perp \partial^2 f(x)[z] + W_x(z, P_\perp \partial f(x)[z]). $$

$W_x$ denotes the Weingarten map, which, for a spherical manifold, for any vector $v$ at a point $x$ is given by

$$ W_x(z, v) = -z x^T v, $$

where $z$ is a tangent vector to the sphere at $x$.

To calculate the Hessian, we insert $v = P_\perp \perp \partial f(x)[z]$ and retrieve

$$ W_x(z, P_\perp \perp \partial f(x)[z]) = -z x^T P_\perp \partial f(x)[z] = -z x^T x^T \partial f(x)[z] = -z x^T \partial f(x)[z], $$

where $x^T \partial f(x)[z]$ is the scalar product of the spin with the gradient.

To illustrate the implementation in Spirit [5], we switch notation to matrix representation and drop the subscript $x$. For spin indices $i$ and $j$, the gradient $\partial \mathcal{H}$ can be written as a three-dimensional object $\nabla_i \mathcal{H}$ and the second derivative $\partial^2 \mathcal{H}$ as a matrix $\mathcal{H}$.

In Euclidean representation, the Hessian of Eq. (H2) becomes as a $3N \times 3N$ matrix,

$$ H_j^{3N} = (H_{ij})^{3N} = \begin{pmatrix} H_{11}^{3N} & H_{12}^{3N} & \cdots \\ H_{21}^{3N} & H_{22}^{3N} & \cdots \\ \vdots & \vdots & \ddots \end{pmatrix}. $$

consisting of $N^2$ blocks, each corresponding to a different spin-spin subspace. It is obtained by acting with Eq. (H2) on the Euclidean basis vectors of the embedding space $\mathcal{E}$.

These subspace matrices of size $3 \times 3$ are given by

$$ H_{ij}^{3N} = P_i \mathcal{H}_{ij} - \delta_{ij} I n_j \cdot \nabla_j \mathcal{H}, $$

where $I$ denotes the $3 \times 3$ unit matrix.

The matrix $H_j^{3N}$, of course, describes $3N$ degrees of freedom, while there can only be $2N$ physical eigenmodes of the

---

**TABLE I.** Parameters of the Gaussians in the energy surface of the single-spin system shown in Fig. 7 in the main text.

<table>
<thead>
<tr>
<th>$a$</th>
<th>$\sigma$</th>
<th>$c_x$</th>
<th>$c_y$</th>
<th>$c_z$</th>
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<tbody>
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<td>-0.20</td>
<td>0.00</td>
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<td>-1.00</td>
<td>0.20</td>
<td>-0.20</td>
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<tr>
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<td>1.00</td>
<td>-0.20</td>
<td>-0.10</td>
</tr>
<tr>
<td>009</td>
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<td>0.50</td>
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</tr>
<tr>
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<td>0.80</td>
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<tr>
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<td>0.10</td>
<td>0.50</td>
<td>1.20</td>
<td>-0.40</td>
</tr>
<tr>
<td>-090</td>
<td>0.10</td>
<td>0.20</td>
<td>-0.90</td>
<td>-0.40</td>
</tr>
</tbody>
</table>
spins, spanning the tangent space to the spin configuration. In order to remove the unphysical degrees of freedom in the embedding space $\mathcal{E}$, it is sufficient to transform the matrix into a tangent space basis, which we can write as $H_{ij} = T_j^T H_{ij}^{SN} T_i$, where $T_i$ is the basis transformation matrix of spin $i$ fulfilling $T_j^T P = T_j$ and $T_j^T T = I^{2N}$.

The true Hessian $H = (H_{ij})$ of Eq. (H2) in the $2N \times 2N$ matrix representation, containing only the physical degrees of freedom, is therefore defined as

$$H_{ij} = T_j^T \tilde{H}_{ij} T_i - T_j^T I (n_j \cdot \nabla \tilde{H}) T_i.$$  \hspace{1cm} (H7)

Note that this reduction of dimensionality also improves the numerical efficiency of the diagonalization.

As the eigenmodes $\lambda^{2N}_i$ are represented in the tangent basis, the $3N$ representation needs to be calculated by $\lambda_i^{3N} = T^{\dagger} \lambda_i^{2N}$. While the $3 \times 2$ basis matrix $T_i$ can be calculated quite arbitrarily by choice of two orthonormal vectors, tangent to the spin $n_i$, we found it convenient to use the unit vectors of spherical coordinates $\theta$ and $\phi$:

$$T = \{e_0, e_\varphi\} = \begin{pmatrix} \cos \theta \cos \varphi & -\sin \varphi \\ \cos \theta \sin \varphi & \cos \varphi \\ -\sin \theta & 0 \end{pmatrix},$$  \hspace{1cm} (H8)

where $r_{xy} = \sin \theta = \sqrt{1-z^2}$.

Note that the poles need to be excluded, but since the basis does not need to be continuous over the manifold, one may, e.g., orthogonalize $e_0$ and $e_\varphi$ with respect to the spin vector to obtain suitable tangent vectors.

Finally, the Hessian matrix in the embedding space $\mathcal{E} = \mathbb{R}^{3N}$ is needed, denoted $\tilde{H}_{ij}^{SN}$. The atomistic Hamiltonian can generally be written in matrix form,

$$\mathcal{H} = -\sum_j A_{ij} n_j - \sum_{(ij)} n_i B_{ij} n_j,$$  \hspace{1cm} (H9)

where $A_{ij}$ are matrices of size $3 \times 3$ describing the linear contributions, such as the Zeeman term, and $B_{ij}$ are matrices describing the quadratic contributions, such as anisotropy, exchange, DMI, and dipolar interactions. The Hessian matrix is then naturally given by

$$\tilde{H}_{ij} = \partial^2 \mathcal{H} = -2B_{ij},$$  \hspace{1cm} (H10)

**APPENDIX I: MINIMUM MODES IN THE INTERACTING SPIN SYSTEM**

Recall Eq. (21) of the main text, which can be written

$$F^{\text{eff}} = -\nabla \mathcal{H} + 2(\hat{\lambda} \cdot \nabla \mathcal{H}) \hat{\lambda},$$  \hspace{1cm} (I1)

where $(\cdot)$ denotes the dot product of $3N$-dimensional vectors. Figure 15 illustrates how the minimum eigenmode axis $\hat{\lambda}$ is oriented and in which direction, and therefore, the gradient force is inverted. Figures 11 and 15 can be compared to equivalent Cartesian-coordinate figures from molecular calculations in Ref. [67].

Equation (21) describes the reflection of the energy gradient’s component along the mode. Following this force, the system climbs in energy along the mode, while minimizing the energy in the orthogonal degrees of freedom. As can be seen in Fig. 15, the minimum mode reflects the symmetry of the energy landscape, as it is pointed approximately between the two minima. This is especially obvious in the area close to the saddle point. Consequently, the MMF force acts analogous to the climbing image method [see Eq. (16)], but using the lowest eigenmode instead of a transition path tangent.


