# Precession Axis Modification to a Semi-analytical Landau-Lifshitz Solution Technique 

D. G. Porter, M. J. Donahue<br>National Institute of Standards and Technology, Gaithersburg, MD 20899


#### Abstract

A recent article [1] presents a semi-analytical method to solve the Landau-Lifshitz (LL) equation. Spin motion is computed analytically as precession about the effective field $H$, where $H$ is assumed fixed over the time step. However, the exchange field dominates at short range and varies at the time scale of neighbor spin precessions, undermining the fixed field assumption. We present an axis corrected version of this algorithm. We add a scalar multiple of $m$ to $H$ (preserving torque and hence the LL solution) to produce a more stable precession axis parallel to the cross product of the torques $m \times H$ at two closely spaced time steps. We build a predictor-corrector solver on this foundation. The second order convergence of the solver enables calculation of adjustable time steps to meet a desired error magnitude.


## I. INTRODUCTION

At the heart of many micromagnetic simulation tasks is the computation of magnetization dynamics via numerical integration of the Landau-Lifshitz equation,

$$
\begin{equation*}
\frac{d m}{d t}=\frac{\gamma}{1+\alpha^{2}} m \times H-\frac{\alpha \gamma}{1+\alpha^{2}} m \times H \times m \tag{1}
\end{equation*}
$$

where $\gamma=-221 \mathrm{kHz} /(\mathrm{A} / \mathrm{m})$ is the gyromagnetic constant, $\alpha$ is a dimensionless phenomenological damping parameter, $m$ is a unit vector in the magnetization direction, and $H$ is the effective field representing the effect of all energies included in the simulation. Solution schemes of increased efficiency are sought to permit simulations of larger objects over longer time intervals.

Some properties of (1) are noteworthy. Magnetization trajectories that solve the equation are norm-preserving. The equation computes varying magnetization direction with time, while the magnetization magnitude remains fixed. Also, the change in magnetization direction $m$ in response to the effective field $H$ is entirely a function of $m \times H$. That is, the trajectory is determined by the torque, not by the field itself. Consequently, so long as we preserve torque, we are free to modify the value of $H$ as needed to pursue other goals. Finally, we note that each of the two terms of (1) can be described by its effect on the trajectory. When the damping parameter $\alpha$ is near zero, the influence of the damping term fades, and the precession term dominates.

Many general numerical integration algorithms are not tailored to computing norm-preserving trajectories. When these algorithms are applied to (1) in micromagnetic simulation work, it is common practice to augment them with renormalization from time to time during the computation. Different renormalization schemes combined with various numerical integration algorithms give rise to different systematic computational errors.

Motivated by these factors, alternative numerical integration schemes have been proposed [1-4] that directly account for the norm-preserving nature of (1). Some of these schemes take simulation steps made up of rotations rather than straight-line increments. Such schemes
are expected to better track trajectories when precession dominates, offering the hope of acceptably accurate results even when employing larger time steps. The result is increased simulation efficiency. In this paper, we examine the scheme of [1] and offer improvements to it.

## II. ANALYSIS

The fast semi-analytical scheme presented in [1] begins with the observation that so long as $H$ remains fixed, the trajectory of $m$ can be computed analytically. Because $H$ is a function of $m$, we cannot expect it to truly remain fixed, but so long as time steps are kept small enough, the error created by this approximation can be kept acceptably low. For a typical simulation, it was reported that compared to solver schemes such as the Euler and Heun methods, full trajectory simulations to equilibrium could be completed using longer time steps, and consequently fewer total calculations.

In simulations where exchange energy plays a leading role, we expect the assumptions of the semi-analytical scheme to be undermined. With a significant portion of $H$ arising from the other spins in the simulation, and each of those spins also in motion, we expect an assumption of fixed $H$ over a time step to become invalid for shorter time steps. To illustrate this effect, we simulated a two spin system with exchange energy as the only contributor to $H$. Permalloy parameters (Exchange energy constant $A=13 \mathrm{~nJ} / \mathrm{m}$, magnetization magnitude $M=800 \mathrm{kA} / \mathrm{m}$, and $\alpha=0.01$ ) and spins a distance $\Delta=5 \mathrm{~nm}$ apart were assumed. The initial spin directions were oriented 20 deg apart. We find that the simple single step semi-analytic scheme [1] fails to converge with time steps of 0.2 ps or longer.

For this simple two-spin, exchange-only system, we can analytically demonstrate an alternative scheme that performs better. The exchange field at spin $1, H_{1}$, due to the magnetization of $\operatorname{spin} 2, m_{2}$, is conventionally expressed,

$$
\begin{equation*}
H_{1}=\frac{2 A}{\mu_{0} M \Delta^{2}} m_{2} \tag{2}
\end{equation*}
$$

Recall that the dynamics of spin 1 are determined only
by the torque $m_{1} \times H_{1}$ however. This means we may add any scalar multiple of $m_{1}$ to $H_{1}$ without changing the dynamics. With this in mind, let

$$
\begin{equation*}
\tilde{H}=H_{1}=H_{2}=\frac{2 A}{\mu_{0} M \Delta^{2}}\left(m_{1}+m_{2}\right) \tag{3}
\end{equation*}
$$

and we see that both spins in the system may have their dynamics computed as in response to a common field value. We may further examine how a common field value defined in this manner evolves in time.

$$
\begin{align*}
\frac{d \tilde{H}}{d t} & =\frac{2 A}{\mu_{0} M \Delta^{2}}\left(\frac{d m_{1}}{d t}+\frac{d m_{2}}{d t}\right)  \tag{4}\\
& =\frac{4 A^{2} \alpha|\gamma|}{\left(\mu_{0} M \Delta^{2}\right)^{2}} \sin (\theta) \tan \left(\frac{\theta}{2}\right) \frac{m_{1}+m_{2}}{2} \tag{5}
\end{align*}
$$

where $\theta$ is the angle between $m_{1}$ and $m_{2}$. The derivation comes from substituting (1) for the $d m / d t$ terms. Note that both $\tilde{H}$ and $d \tilde{H} / d t$ are in the direction of $m_{1}+$ $m_{2}$, so we conclude the value $\tilde{H}$ increases in magnitude, but has fixed direction. Its maximum value is reached as $\theta$ reaches 0 , corresponding to a maximum precession frequency of

$$
\begin{equation*}
f_{\max }=\frac{2 A|\gamma|}{\pi \mu_{0} M \Delta^{2}} \tag{6}
\end{equation*}
$$

Computing dynamics according to the common field value $\tilde{H}$ more closely fulfills the fixed field assumption, and permits larger time steps without divergence, or loss of accuracy beyond specified bounds. In effect, we have replaced the effective field with a different one directed along a corrected axis, for which the efficiency gains are greater. We find that when we apply our axis-corrected semi-analytic scheme to our example problem, solutions continue to converge to the proper equilibrium state for time steps up to 6 ps , a significant increase over the 0.2 ps achieved by the uncorrected scheme.

For other two-spin simulations with other energies in addition to exchange, such a common field value does not arise, but the same strategy of adding a scalar multiple of $m$ to $H$ to get an effective field that changes direction less rapidly is still effective. At each time step, we are computing the torque $m \times H$. Given the value of $m \times H$ at time $t=-\tau$ and at time $t=0$, an axis is determined by

$$
\begin{equation*}
a=(m \times H)(-\tau) \times(m \times H)(0) \tag{7}
\end{equation*}
$$

At $t=0$, we add the appropriate scalar multiple of $m$ to the effective field $H$ to produce a corrected $\tilde{H}$ that is parallel to that axis. That is, we solve

$$
\begin{equation*}
\tilde{H}=(H+\lambda m)(0)=\beta a \tag{8}
\end{equation*}
$$

for suitable scalar values of $\lambda$ and $\beta$. Consider the inner product

$$
\begin{equation*}
\tilde{H} \cdot(m \times H \times m) \tag{9}
\end{equation*}
$$



FIG. 1: Comparison of relative error vs time step length for the original semi-analytic predictor-corrector solver, and the proposed axis-corrected alternative.

Make the substitutions $\tilde{H}=H+\lambda m$ and $\tilde{H}=\beta a$ in turn, simplify and compare to see that

$$
\begin{equation*}
H \cdot(m \times H \times m)=\beta a \cdot(m \times H \times m) \tag{10}
\end{equation*}
$$

Solving for $\beta$,

$$
\begin{equation*}
\beta=\frac{H \cdot(m \times H \times m)}{a \cdot(m \times H \times m)} \tag{11}
\end{equation*}
$$

we have all we need to determine $\lambda$ and $\tilde{H}$. The analytic LL solution assuming the fixed value $\tilde{H}$ over the next time interval then yields the value of $m$ at time $t=\tau$. When we employ this scheme to the exchange only case, we compute the common $\tilde{H}$ field detailed above. This strategy applies when other energies are represented as well.

## III. PREDICTOR-CORRECTOR SOLVERS

In [1] the semi-analytic step was used as a foundation to construct a predictor-corrector solver. The axiscorrected semi-analytic scheme also supports a predictorcorrector extension. To determine the effectiveness of our axis correction, we simulated a two spin system with exchange, demagnetization, and cubic anisotropy energies. First we used several runs of a $5 / 4$ Runge-KuttaFehlberg solver [5] at various time steps to compute a converged baseline solution over a 10 ps interval. Then the predictor-corrector solver from [1] and our axis corrected predictor-corrector solver were used to compute solutions over the same interval, using a variety of time steps for both. The error at $t=10 \mathrm{ps}$ relative to the baseline solution was taken as a figure of merit. Figure 1 displays the results. The axis-corrected solver yields about an order of magnitude less error compared to the referenced predictor-corrector scheme. Equivalently, the


FIG. 2: Comparison of effectiveness of adjustable time step determination applied to the original semi-analytic predictorcorrector solver, and to the proposed axis-corrected alternative. Magnetization components $m_{x}$ and $m_{y}$ as a function of time are plotted against the scale on the left. Time step lengths as a function of time are plotted against the scale on the right. Note the scaling of the third curve for sake of visibility makes values appear one hundred times greater than they are.
axis-corrected solver achieves the same magnitude of error with three times smaller time steps.

## IV. ADJUSTABLE TIME STEP SOLVERS

Figure 1 clearly demonstrates that both solution algorithms exhibit second order convergence. This means they are suitable foundations for the construction of adjustable time step algorithms that dynamically grow and shrink the time step duration to keep the overall calculation within a desired error magnitude.

Figure 2 illustrates the results of the adjustable time step solvers based on the two predictor-corrector solvers.

Again a two-spin system is simulated, this time with exchange, demagnetization, cubic anisotropy, and Zeeman energies in the simulation. The $x$ and $y$ components of $m$ for one of the spins is displayed, showing its precession and approach to convergence to an equilibrium direction after 5 ns of simulated time. The same system was simulated using the Runge-Kutta-Fehlberg solver with a fixed time step of 1 fs to produce a baseline solution. In Fig 2 both solvers compute results with errors less than $2 \times 10^{-6}$ relative to the baseline solution. However, the required time steps to achieve that error level are quite distinct.

Note that the time step duration as a function of time for the original semi-analytic predictor-corrector solver is displayed with a magnification of one hundred times. In this case, the time step adjustments never produce a time step even as long as 2 fs . In contrast the time step adjustments applied to the axis-corrected version of the semi-analytic predictor-corrector solver are able to reach time steps of more than 200 fs by the end of the 5 ns simulation interval, and the time steps appear to still be lengthening at that point.

Because the axis corrected solver over time takes longer and longer time steps while maintaining the same error level, it is able to achieve the same computational results with thirty times fewer calculations.

## V. SUMMARY

In this article, three new developments have been presented. First, we have described an axis correction that improves on a previously published Landau-Lifshitz solution technique, and demonstrated its benefits. Second, we have analyzed the convergence of these solution techniques and demonstrated that they exhibit second order convergence. Third, we have taken advantage of the second order convergence property of these solution techniques to implement adjustable time step algorithms that permit even more striking demonstration of the advantage of axis correction.
[1] Ben Van de Wiele, Femke Olyslager, and Luc Dupré, "Fast semianalytical time integration schemes for the LandauLifshitz equation", IEEE Trans. Magn., vol. 43, no. 6, pp. 2917-2919, June 2007.
[2] P. B. Visscher and Xuebing Feng, "Quaternion-based algorithm for micromagnetics", Phys. Rev. B, vol. 65, no. 10, pp. 104412, February 2002.
[3] P. S. Krishnaprasad and Xiaobo Tan, "Cayley transforms in micromagnetics", Physica B, vol. 306, pp. 195-199, 2001.
[4] Massimiliano d'Aquino, Claudio Serpico, and Giovanni Miano, "Geometrical integration of Landau-LifshitzGilbert equation based on the mid-point rule", Journal of Computational Physics, vol. 209, pp. 730-753, 2005.
[5] Michael J. Donahue and Donald Gene Porter, "OOMMF user's guide, version 1.0", NISTIR 6376, National Institute of Standards and Technology, 100 Bureau Drive, Gaithersburg, MD 20899, 1999.

