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### 1 (1) Overview

### 2 Title

Spinsim: a GPU optimized simulator of spin-half and spin-one quantum systems

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### 5 Abstract

The *Spinsim python* package simulates spin-half and spin-one quantum mechanical systems following a time dependent Schroedinger equation. It makes use of `numba.cuda`, which is an *LLVM* (Low Level Virtual Machine) compiler for *Nvidia Cuda* compatible systems using GPU parallelisation. Along with other optimisations, this allows for speed improvements of up to four orders of magnitude while keeping staying just as accurate. *Spinsim* is available for installation on *PyPI*, and the source code is available on *github*. The initial use-case for the package will be to simulate quantum sensing-based Bose Einstein condensate (BEC) experiments for the Monash University School of Physics and Astronomy spinor BEC lab,

but we anticipate it will be useful in simulating any range of spin-half or spin-one quantum systems with time dependent Hamiltonians that cannot be solved analytically. These appear in the fields of nuclear magnetic resonance (NMR), nuclear quadrupole resonance (NQR) and magnetic resonance imaging (MRI) experiments and quantum sensing, and with the spin-one systems of nitrogen vacancy centres (NVCs) and BECs.

## 6 Keywords

Time dependent Schrödinger equation; Quantum; Physics; Spin one; Spin half; Integrator; Exponentiator; GPU; Parallel computing; Solver; python; numba;

## 7 Introduction

### 7.1 Motivation

Ultracold rubidium atoms have proven their effectiveness in state of the art technologies in quantum sensing [1], the use of quantum mechanics to make precise measurements of small signals. The rotation of these atoms can be modelled as quantum spin systems, which is the quantum mechanical model for objects with angular momentum. The simplest spin system, spin-half (ie spin quantum number of  $\frac{1}{2}$ , also referred to as a qubit), is quantized into just two quantum spin levels, and this describes the motion of some fundamental particles such as electrons. However, systems more practical for sensing, such as ultracold rubidium atoms, are more accurately described as a spin-one quantum system (ie spin quantum number of 1, also referred to as a qutrit), which is quantized into three quantum spin levels.

The design of sensing protocols requires many steps of verification, including simulation. This is especially important, since running real experiments can be expensive and time consuming, and thus it is more practical to debug such protocols quickly and cheaply on a computer. In general, any design of experiments using spin systems could benefit from a fast, accurate method of simulation.

In the past, the spinor Bose Einstein condensate (spinor BEC) lab at Monash University used an in-house, *cython* based script, on which this package is based, and standard differential equation solvers (such as *Mathematica*'s function `NDSolve`) to solve the Schroedinger equation for quantum sensing spin systems. Spin one systems are sometimes approximated to spin-half for a faster execution time, at the cost of modelling all effects of the system. However, these methods are not completely optimized for our use-case, and therefore come with some issues.

First, while the execution time for these solvers is acceptable for running a small number of experiments, for certain experiments involving large arrays of independent atom clouds (which require many thousands of simulations to be run), this time accumulates to the order of many hours, or even multiple days.

Second, the Schroedinger equation has the geometric property of being norm preserving. In other words, the time-evolution operator for a system between two points in time must be unitary. As such, numerical solutions to the Schroedinger equation should also preserve this property. For many numerical methods like those in the Runge Kutta family, the approximations used might not be norm preserving, and the evaluated quantum state may diverge towards an infinite norm, or converge to zero if run for many iterations.

Third, our system (and similar spin systems) can be very oscillatory. In standard conditions for our application, the expected spin projection of a system that we want to solve for can rotate in physical space (alternatively viewed as a point rotating around an abstract object known as a *Bloch sphere*) at a rate of 700kHz. Standard integration methods require very small time-steps in order to accurately depict these oscillations.

Given the recent boom in machine learning, many research computers are now equipped with advanced graphics processing units (GPUs). Their many cores, which can range from hundreds to tens of thousands in number, allows them to run some highly parallel algorithms much faster than a central processing unit (CPU). Examples include calculating colours for many pixels on a screen in the titular graphics processing, or the weights in a large neural network. By finding ways to parallelize the problem of solving quantum spin systems, we can use the many cores of a GPU to our advantage in this context as well.

## 8 Implementation and architecture

### 8.1 Quantum mechanics background

The *Spinsim* package solves the time-dependent Schrödinger equation

$$i\hbar \frac{d\psi(t)}{dt} = H(t)\psi(t), \quad (1)$$

where the quantum state  $\psi(t) \in \mathbb{C}^N$  is assumed normalized, and the Hamiltonian  $H(t) \in \mathbb{C}^{N \times N}$  is Hermitian. Here  $N$  is the number of levels in the quantum system. Often we are considering systems with spin of half or one. Spin-half is the  $N = 2$  case, and spin-one is  $N = 3$ . We set  $\hbar = 1$ , so that the Hamiltonian has physical dimension of angular frequency.

Rather than parametrizing the problem in terms of the matrix elements of  $H(t)$ , we consider time varying real coefficients in a linear combination of fixed operators,

$$H(t) = \sum_{j=1}^{N^2-1} \omega_j(t) \mathcal{A}_j. \quad (2)$$

Here we exclude the identity so that the Hamiltonian is traceless, which corresponds to choosing a physically meaningless energy zero point. It is well known that a charged spin-half system with magnetic moment  $\vec{\mu}$ , and gyromagnetic ratio  $\gamma$ , in a magnetic field  $\vec{B}(t)$ , has Hamiltonian

$$H(t) = -\vec{B}(t) \cdot \vec{\mu} \quad (3)$$

$$= -\gamma (B_x(t)J_x + B_y(t)J_y + B_z(t)J_z) \quad (4)$$

$$= \omega_x(t)J_x + \omega_y(t)J_y + \omega_z(t)J_z. \quad (5)$$

Here  $J_x$ ,  $J_y$  and  $J_z$  are spin operators, equal to the Pauli matrices [2, (p169)] halved. Because of the equivalence of the magnetic field components  $B_j(t)$  and the  $\omega_j(t)$ , we henceforth refer to the latter as field functions.

In the spin-one case there is no standard basis of operators  $\mathcal{A}_j$ . Choices include the Gell-Mann matrices [3], and multiple dipole-quadrupole bases [4, 5]. In general, we

can choose any basis from the 8-dimensional Lie algebra  $\mathfrak{su}(3)$ , which is the vector space of traceless Hermitian operators that can generate transformations (from the corresponding Lie group  $SU(3)$ ) in the spin-one system. We focus on a particular subfamily of spin-one systems for which the Hamiltonian is a linear combination of matrices from a 4-dimensional subspace of  $\mathfrak{su}(3)$ , consisting of the spin matrices  $J_x$ ,  $J_y$  and  $J_z$ , (labelled  $L_x$ ,  $L_y$  and  $L_z$  in Reference [4]) and a single quadrupole operator  $Q = \text{diag}(1, -2, 1)/3$ ,

$$H(t) = \omega_x(t)J_x + \omega_y(t)J_y + \omega_z(t)J_z + \omega_q(t)Q. \quad (6)$$

Note that  $Q$  is proportional to  $Q_{zz}$  [4] and  $Q_0$  [5] from alternative quadrupole bases. The  $J_x$ ,  $J_y$ ,  $J_z$  and  $Q$  are the only operators necessary to simulate many spin-one quantum systems in arbitrary bias fields, but with single-photon coupling. This includes quadratic Zeeman splitting described by the Breit-Rabi formula [6] important to experiments in our lab. The *Spinsim* simulator can also be configured to solve a general spin-one system by setting the Hamiltonian to an arbitrary point in  $\mathfrak{su}(3)$  using the full quadrupole basis, which extends the possible Hamiltonian to

$$H(t) = \omega_x(t)J_x + \omega_y(t)J_y + \omega_z(t)J_z + \omega_q(t)Q \\ + \omega_{u1}(t)U_1 + \omega_{u2}(t)U_2 + \omega_{v1}(t)V_1 + \omega_{v2}(t)V_2. \quad (7)$$

Here the additional operators are those defined in [5]. Note that this is included for completeness, but has not been thoroughly tested, as our lab has no physical context for modelling the  $U_1, U_2, V_1$  and  $V_2$  operators.

As well as integrating the Schrödinger equation, *Spinsim* also has the functionality to calculate the expected spin projection  $\langle \vec{J} \rangle(t)$  of a system from its state. In an experimental setting, one cannot measure the value of the state itself, and must instead measure observables such as spin projection. If this is done for an ensemble of systems, the average spin projection over all systems will converge to the expected value as calculated here. As an alternative to solving the Schrödinger equation, if one is only interested in the dynamics of the expected spin projection, one can instead solve for  $\langle \vec{J} \rangle(t)$  directly using the Bloch equations [7]. These are three coupled real equations as compared to three coupled complex equations for the spin-one Schrödinger equation. However, the Bloch equations cannot be used to compute  $\psi(t)$ , and can only fully model spin-half systems (and spin-one systems approximated to spin-half). Thus, with *Spinsim* we choose to solve the full Schrödinger equation instead.

## 8.2 Unitary time evolution and the Magnus expansion

Since  $H(t)$  is Hermitian, it follows from Equation (1) that  $\|\psi(t)\|^2 = \|\psi(t_0)\|^2$  for all  $t$ . Therefore it is possible to write  $\psi(t)$  in terms of a unitary transformation  $\mathcal{U}(t, t_0)$  of the state  $\psi(t_0)$ , for any time  $t_0$ , that is,  $\psi(t) = \mathcal{U}(t, t_0)\psi(t_0)$ . It then follows from Equation (1) that  $\mathcal{U}(t, t_0)$  also follows a Schrödinger equation,

$$i\hbar \frac{d\mathcal{U}(t, t_0)}{dt} = H(t)\mathcal{U}(t, t_0). \quad (8)$$

Thus, to solve Equation (1) for  $\psi(t)$  given a particular  $\psi(t_0)$ , one only needs to solve Equation (8) for  $\mathcal{U}(t, t_0)$ . Since  $\mathcal{U}(t, t_0)$  is unitary, it can be written as a matrix exponential of a anti-Hermitian (also termed skew-Hermitian) matrix. If the Hamiltonian is a of constant value  $H(t) = \mathcal{H}$ , then this exponential is

$$\mathcal{U}(t, t_0) = \exp(-i(t - t_0)\mathcal{H}). \quad (9)$$

For a time varying Hamiltonian, the general solution for  $\mathcal{U}(t, t_0)$  is much more complex, because it encapsulates the full solution to the time-dependent Schrödinger equation. The well known Dyson series [8] gives an explicit expression for  $\mathcal{U}(t, t_0)$  in terms of multiple time integrals over nested time commutators of  $H(t')$ . While the Dyson series has recently been used numerically [8], it is challenging to work with because once truncated, it is in general no longer unitary. The Magnus series, in contrast, gives  $\mathcal{U}(t, t_0)$  in terms of an exponential of a series of anti-Hermitian operators, specifically

$$\mathcal{U}(t, t_0) = \exp(\Omega(t, t_0)) = \exp\left(\sum_{m=1}^{\infty} \Omega_m(t, t_0)\right). \quad (10)$$

As such, the Magnus series explicitly preserves unitarity when truncated [9]. The terms in the Magnus series are integrals over nested time commutators of the Hermitian matrix  $A(t')$ , where, in the case of quantum mechanics,  $A(t') = -iH(t')$ . For instance, the first three terms are [10]

$$\Omega_1(t, t_0) = \int_0^t dt_1 A(t_1) \quad (11)$$

$$\Omega_2(t, t_0) = \frac{1}{2} \int_{t_0}^t dt_1 \int_{t_0}^{t_1} dt_2 [A(t_1), A(t_2)] \quad (12)$$

$$\Omega_3(t, t_0) = \frac{1}{6} \int_{t_0}^t dt_1 \int_{t_0}^{t_1} dt_2 \int_{t_0}^{t_2} dt_3 ([A(t_1), [A(t_2), A(t_3)]] + [A(t_3), [A(t_2), A(t_1)]]) , \quad (13)$$

where  $[X, Y] = XY - YX$  is the commutator. Note that, when truncated to first order, the Magnus expansion  $\mathcal{U}(t, t_0) = \exp\left(\int_{t_0}^t dt_1 (-iH(t_1))\right)$  reduces to Equation (9), with  $H(t)$  approximated by its average value over the interval  $[t_0, t]$ . Convergence of the Magnus series for  $\mathcal{U}(t, t_0)$  is not in general guaranteed, but it is under the condition that  $\int_{t_0}^t dt_1 \|H(t_1)\| < \xi \approx 1.08686870 \dots$  [10]. Furthermore, each subsequent term in the expansion increases in complexity rapidly. For these reasons, the Magnus expansion is generally used as a time-stepping method rather than a single step to solve the complete system. Designing a Magnus-based solver is a trade-off between the number of terms used in each expansion, and the number of time-steps used. Time stepping is based on the fact that time evolution operators can be split into a product via

$$\mathcal{U}(t, t_0) = \mathcal{U}(t, t_{n-1})\mathcal{U}(t_{n-1}, t_{n-2}) \cdots \mathcal{U}(t_2, t_1)\mathcal{U}(t_1, t_0). \quad (14)$$

Each of the time-evolution operators can be approximated by a Magnus expansion.

Many unitary time-stepping techniques have been developed based on the Magnus expansion [11]. Some of these techniques use Gauss-Legendre quadrature sampling and the Baker–Campbell–Hausdorff formula to respectively avoid the integration and time commutators of the Hamiltonian, producing simple expressions that can be used for unitary time-stepping [12].

In particular, we use the commutator free, fourth order method (CF4) from Reference [11]. Suppose we wish to evaluate a time-step of  $\mathcal{U}(t + dt, t)$  using the CF4 method. The Hamiltonians are sampled at times

$$t_1 = t + \frac{1}{2} \left( 1 - \frac{1}{\sqrt{3}} \right) dt \quad \text{and} \quad (15)$$

$$t_2 = t + \frac{1}{2} \left( 1 + \frac{1}{\sqrt{3}} \right) dt, \quad (16)$$

based on the second order Gauss-Legendre quadrature

$$\overline{H}_1 = \frac{3 + 2\sqrt{3}}{12} H(t_1) + \frac{3 - 2\sqrt{3}}{12} H(t_2) \quad \text{and} \quad (17)$$

$$\overline{H}_2 = \frac{3 - 2\sqrt{3}}{12} H(t_1) + \frac{3 + 2\sqrt{3}}{12} H(t_2). \quad (18)$$

Then

$$\mathcal{U}(t + dt, t) \approx \exp(-i\overline{H}_2 dt) \exp(-i\overline{H}_1 dt) \quad (19)$$

is used to approximate the time-evolution operator.

### 8.3 Lie-Trotter based Exponentiator

Evaluating matrix exponentials is a core part of the integration algorithm. Rather than exponentiating the Hamiltonian directly as in Equation (19), *Spinsim* works with the field functions  $\omega_j(t)$ .

For spin-half, the exponentiator is in an analytic form in  $\omega_x, \omega_y$  and  $\omega_z$ . For spin-one, an exponentiator based on the Lie-Trotter product formula [13]

$$\exp(X + Y) = \lim_{n \rightarrow \infty} \left( \exp\left(\frac{X}{n}\right) \exp\left(\frac{Y}{n}\right) \right)^n, \quad (20)$$

is used. An advantage of the Lie-Trotter approach is that  $\exp(-iA_k/n)$  has known analytic forms for the Lie algebra basis elements  $A_k$ . Hence the unitary time-evolution operator is approximated by  $U = \exp(-iHt) \approx T^n$  where

$$T = \exp\left(\frac{-i\omega_x J_x}{n}\right) \exp\left(\frac{-i\omega_y J_y}{n}\right) \exp\left(\frac{-i\omega_z J_z}{n}\right) \exp\left(\frac{-i\omega_q Q}{n}\right). \quad (21)$$

In fact, commutation relations between the Lie basis operators and the leapfrog splitting method [14] allow us to write

$$T = \exp\left(-i\frac{1}{2}D_{z,q}\right) \exp(-i\Phi J_\phi) \exp\left(-i\frac{1}{2}D_{z,q}\right). \quad (22)$$

Here  $z = \omega_z/n$ ,  $q = \omega_q/n$ ,  $\Phi = \sqrt{\omega_x^2 + \omega_y^2}/n$ ,  $\phi = \arctan2(\omega_y, \omega_x)$  (where  $\arctan2(y, x)$  is the two argument arctangent [15]),  $D_{z,q} = zJ_z + qQ$  and  $J_\phi = \cos(\phi)J_x + \sin(\phi)J_y$ . The element-wise analytic form of this is

$$T = \begin{pmatrix} (\cos(\frac{\Phi}{2})e^{-iz/2}e^{-iq/6})^2 & \frac{-i}{\sqrt{2}}\sin(\Phi)e^{iq/6}e^{-iz/2}e^{-i\phi} & -(\sin(\frac{\Phi}{2})e^{iq/6}e^{-i\phi})^2 \\ \frac{-i}{\sqrt{2}}\sin(\Phi)e^{iq/6}e^{-iz/2}e^{i\phi} & \cos(\Phi)e^{i4q} & \frac{-i}{\sqrt{2}}\sin(\Phi)e^{iq/6}e^{iz/2}e^{-i\phi} \\ -(\sin(\frac{\Phi}{2})e^{-iq/6}e^{i\phi})^2 & \frac{-i}{\sqrt{2}}\sin(\Phi)e^{iq/6}e^{iz/2}e^{i\phi} & (\cos(\frac{\Phi}{2})e^{iz/2}e^{-iq/6})^2 \end{pmatrix}. \quad (23)$$

Matrix exponentiation is completed by raising  $T$  to an integer  $n$ . To approximate the limit in Equation (20),  $n$  must be a large number. We choose  $n$  to be of the form  $n = 2^\tau$ , as raising matrices to powers of two can be done efficiently by iterative squaring. This can be done just by iterative squaring, which requires a small number of operations. While there are some loose upper bounds for the  $n$  required to reach a desired accuracy for the matrix exponential [16], using these to choose a value of  $n$  can, in practice, cause floating-point errors from over-squaring. Instead, we tested the formula on  $10^5$  matrices for different values of  $\tau$ , and compared them to results given by `scipy.linalg.expm()`, from the popular python library *SciPy* [17]. We found that the error minimized after a number of squares of  $\tau = 24$ . Thus, by default,  $n = 2^{24}$ .

In addition to those introduced by the Lie-Trotter approximation, we find errors when manipulating matrices close to the identity. Because  $n$  is very large,  $T$  is very close to the identity. In particular, the limited precision of floating-point numbers is unable to represent the diagonal of  $T$  and its squares accurately. Errors appear in two forms. One form is that iterative squaring of matrices close to the identity produces floating-point cancellation errors. It can be avoided by working with the difference of the matrices from the identity when iteratively squaring. That is, if  $A = I + a$ , then instead of calculating the square  $S = I + s = A^2$ , we find  $s = (a + 2I)a$ . The other form of error is that if we use this method, we must initially calculate  $T - I$  accurately, rather than  $T$ . This cannot be done by simply subtracting the identity from Equation (23), as doing so would introduce cancellation error. It can be avoided by replacing the exponentials and trigonometric functions along the diagonal with specialized implementations of functions such as  $\text{expm1}(x) = \exp(x) - 1$ , designed for this purpose [18]. The identity is added to the result of the iterative residual squaring and the full exponential is returned.

This spin-one exponentiator evolves Hamiltonians spanned by  $J_x$ ,  $J_y$ ,  $J_z$  and  $Q$  which is sufficient for three level systems in arbitrary bias fields, but with single-photon coupling. An exponentiator capable of evolving an arbitrary spin-one Hamiltonian, for example with different coupling between the lower and upper pairs of states, or with two-photon coupling, is included in the package. When assessing accuracy for spin-one problems, we have used the single-photon exponentiator.

Note that, the methods for both spin-half and spin-one use analytic forms of matrix exponentials so that  $T$  is unitary by construction;  $\mathcal{U} = T^n$  is then also unitary. Simulations in *Spinsim* thus maintain unitarity and so conserve probability even over very large numbers of time-steps.

## 8.4 Discretisation and parallelisation

Frequently we wish to sample the state at times  $t_k$  spaced more coarsely than the integration time-step. Consider discrete times  $t_k = t_0 + Dt \cdot k$  where  $Dt$  is the time-step of the time-series, in contrast to  $dt$ , the time-step of the integration. Denoting  $\psi_k = \psi(t_k)$  and  $\mathcal{U}_k = \mathcal{U}(t_k, t_{k-1})$ , the time-series of states  $\psi_k$  and time-evolution operators  $\mathcal{U}_k$  satisfy

$$\psi_k = \mathcal{U}_k \psi_{k-1}. \quad (24)$$

This presents an opportunity for parallelism. While  $\psi_k$  depends on  $\psi_{k-1}$ , the operators  $\mathcal{U}_k$  depend only on the field functions, and not on previous states  $\psi_{k'}$  or previous operators  $\mathcal{U}_{k'}$ . Hence, the time-evolution operators  $\mathcal{U}_k$  can be calculated in parallel.

*Spinsim* splits the full simulation into time intervals  $[t_{k-1}, t_k]$ , and calculates time-evolution operators  $\mathcal{U}_k$  for these intervals in parallel on a GPU. When all the  $\mathcal{U}_k$  are calculated, the CPU then multiplies them together (a comparatively less demanding job than calculating them) using Equation (24) to determine the output samples  $\psi_k$  of the evolving state.

Beyond discretisation for parallelisation, we discretize our time-evolution operator into individual integration time-steps. Each of the  $\mathcal{U}_k$  is further split into products  $u_{L-1}^k \cdots u_0^k$  of  $L$  time-evolution operators, now separated by the integration time-step  $dt = Dt/L$ .

To choose an appropriate value of  $dt$ , users can refer to the accuracy plots in Figures 2a and 2c under *Evaluation of accuracy*. To choose an appropriate value of  $Dt$ , users must consider three factors. For the most benefit out of GPU parallelisation, users should choose  $Dt$  so that the number of time samples  $K$  is at least as large as the number of GPU cores available for *Spinsim*. Users should also make sure that  $Dt$  represents a large enough sample rate for the state  $\psi_k$  or expected spin projection  $\langle \vec{J} \rangle_k$  to be used in their further analysis. Finally, if  $K$  is too large, then the GPU will run out of memory on compilation, which will raise an exception. We found that the 10GiB *GeForce RTX 3080* could run at full memory capacity with 60 million sample points for spin-one mode and 120 million sample points for spin-half.

## 8.5 Dynamically rotating frame

Hamiltonians which have dominant and slowly-varying terms induce rotations around a primary axis, which is usually chosen to be the quantisation axis (which is the  $z$  axis when representing spin operators with the Pauli matrices). Transforming from the lab (standard) frame of reference into one rotating around this axis by rotation operator  $R_{\omega_r}(t) = \exp(i\omega_r J_z t)$ , the resulting system follows a new Hamiltonian

$$H_r(t) = R_{\omega_r}(t)(\omega_x(t)J_x + \omega_y(t)J_y)R_{\omega_r}(-t) + (\omega_z(t) - \omega_r)J_z + \omega_q(t)Q \quad (25)$$

It is a common technique in solving quantum mechanical problems to enter rotating frames, and their more abstract counterparts of interaction pictures [2, (p336,338)]. This is almost always done to enable the Rotating Wave Approximation (RWA),



which is an assumption that the oscillatory components of  $H_r(t)$  on average make minor contributions to time evolution, and can be ignored. In some cases, this allows one to obtain analytic, but approximate, solutions to the quantum system. Note that the RWA is never invoked by *Spinsim*, as doing this would reduce the accuracy of simulation results; although a problem already expressed in a rotating frame can of course be input by the user if desired. In fact, because of the speed increase of *Spinsim* compared to alternatives, users who would have previously simulated their systems using the RWA due to speed concerns may now wish to use *Spinsim* to simulate beyond RWA effects that would previously have been missed.

Instead we take advantage of rotating frames to increase numerical accuracy. By our assumption that  $|\omega_r| \approx |\omega_z(t)| \geq |\omega_x(t)|, |\omega_y(t)|, |\omega_q(t)|$  for the duration we have entered the frame, then the rotating frame Hamiltonian will have a smaller norm than the lab frame Hamiltonian. This will yield more accurate results when making discrete time-steps by the integrator, as the distances stepped are smaller.

A different rotating frame is entered by *Spinsim* for each time interval  $[t_{k-1}, t_k]$ . This dynamic application allows for the rotating frame to be used even during sweeps of the dominant term  $\omega_z(t)$  where it is only approximately constant locally. Here we choose  $\omega_r = \omega_z(t_{k-1} + Dt/2)$  as an approximate average value of  $\omega_z(t)$  over the duration. The calculated time-evolution operator  $\mathcal{U}_k^r$  is then transformed back to the lab frame as

$$\mathcal{U}_k = R_{\omega_r}(-Dt)\mathcal{U}_k^r. \quad (26)$$

This functionality can be disabled for systems that do not fit these criteria.

## 8.6 Integrator architecture

A visual summary of the structure of *Spinsim* is shown in Figure 1. The integrator in the *Spinsim* package calls a kernel (multithreaded function) to be run on a *Cuda* capable *Nvidia* GPU in parallel, with a different thread being allocated to each of the  $\mathcal{U}_k$ . Alternatively, the kernel can be run in parallel on a multicore CPU if a compatible GPU is not available. The kernel returns when each of the  $\mathcal{U}_k$  have been evaluated.

Each thread begins with initialisation of variables, which includes moving into the rotating frame. It then starts a loop to find each of the time-stepping operators  $u_l^k$ . Within the loop, the integrator enters a device function to sample the user-provided field functions  $\omega_j(t)$  (device functions are GPU subroutines, and in practice all device functions here are compiled inline for speed). After this, it applies the rotating frame transformation  $R_{\omega_r}(t)$ , before calculating the Gauss-Legendre weightings  $\bar{H}_1$  and  $\bar{H}_2$ . Next, the matrix exponentials are taken within another device function. Finally,  $u_l^k$  is premultiplied to  $\mathcal{U}_k^r$  (which is initialized to the identity), and the loop continues.

When the loop has finished,  $\mathcal{U}_k^r$  is transformed to  $\mathcal{U}_k$  as in Equation (26), and this is returned. Once all threads have executed, using Equation (24), the state  $\psi_k$  is calculated in a (CPU) `numba.jit()`ed function from the  $\mathcal{U}_k$  and an initial condition  $\psi_{\text{init}} = \psi(t_0)$ .

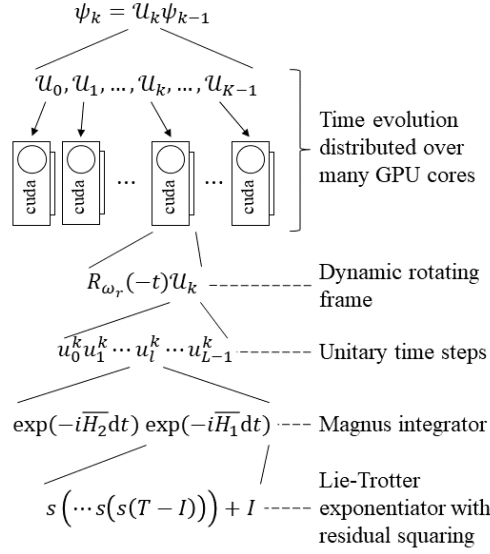


Figure 1: The mathematical and computational breakdown of *Spinsim*. The procedural problem of evaluating the state time series  $\psi_k$  is broken into the parallel problem of evaluating time-evolution operators  $\mathcal{U}_k$ . Each of the  $\mathcal{U}_k$  are found independently on a separate GPU thread. The problem is brought into a dynamic rotating frame to increase numerical accuracy. The  $\mathcal{U}_k$  are broken further into time-stepping operators  $u_k$ , each of which are evaluated using the Magnus CF4 method. Matrix exponentials are evaluated using a Lie-Trotter based method, including a residual squaring technique to avoid over-squaring.

## 8.7 Compilation of integrator

The *Spinsim* integrator is constructed and compiled just in time, using the *Numba* [19] python package. This is achieved via the `numba.cuda.jit()` decorator, which compiles python functions into *Nvidia Cuda* [20] kernels using an LLVM [21] (Low Level Virtual Machine) compiler. The particular device functions used are not predetermined, but are instead chosen based on user input to decide on a closure. This technique has multiple advantages. First, the field functions  $\omega_j(t)$  are provided by the user as a plain python method. Note that this method must be `numba.cuda.jit()` compatible. This allows users to define  $\omega_j(t)$  in a way that compiles and executes fast, does not put many restrictions on the form of the function, and returns the accurate results of analytic functions (compared to the errors seen in interpolation). Compiling the simulator also allows the user to set meta-parameters, and choose the features they want to use in a way that does not require experience with the `numba.cuda` library. This was especially useful for running benchmarks comparing integration methods from previous versions of the software to the current one, CF4. The default settings should be optimal for most users, although tuning the values of *Cuda* meta-parameters `max_registers` and `threads_per_block` could improve performance for different GPUs. Third, just in

time compilation also allows the user to select a target device other than *Cuda* for compilation, so the simulator can run, using the same algorithm, on a multicore CPU in parallel instead of a GPU. In particular, this allows for compatibility with products running *Apple's MacOS 10.14 Mojave* and later, which are incompatible with *Cuda* devices and software.

This functionality is interfaced through an object of class `spinsim.Simulator`. The *Cuda* kernel is defined as per the user's instructions on construction of the instance, and it is used by calling the method `spinsim.Simulator.evaluate()`. Users are returned an instance of `spinsim.Results` including the time, state, time-evolution operator, and expected spin projection. Note that the expected spin projection is calculated as a lazy parameter if needed, rather than returned by the simulator object.

The *Spinsim* package is designed so that a single simulator instance can be used to execute many simulations, sweeping through parameters while not needing to be recompiled. This is done through the `sweep_parameters[]` argument to the user-provided field functions  $\omega_j(t)$ . First the user must instantiate a `spinsim.Simulator` object. They can then set the value for `sweep_parameters[]` for a particular simulation each time `spinsim.Simulator.evaluate()` is called. As `sweep_parameters[]` is a `numpy.ndarray` [22], one can use this functionality to sweep many parameters using the same simulator object. For a use-case example, an MRI experimenter might want to simulate many spin systems at different locations  $(x, y)$ , in a magnetic field gradient  $\omega_z(t) = x - 2y$ . To do this they could choose to set `sweep_parameters = [x, y]`, and define `field_sample[2] = sweep_parameters[0] - 2*sweep_parameters[1]`. This feature is demonstrated with examples in the documentation.

## 9 Quality control

### 9.1 Evaluation of accuracy

All accuracy benchmarks were run in *Cuda* mode, on the desktop computer with the *Ryzen 7 5800X* and *GeForce RTX 3080*, which from Figure 4 are the fastest CPU and GPU from the devices tested. Note that these benchmarks do not take into account the amount of time required to JIT compile simulation code for the first time (order of seconds), as we want to look at this in the limit of running many simulations while sweeping through a parameter that differs in each one.

Benchmarks were performed using `neural-sense.sim.benchmark` (where `neural-sense` [23] is the quantum sensing package that *Spinsim* was written for). This simulation involves continuously driving transitions in the system for 100 ms, while exposing it to a 1 ms pulsed signal that the system should be able to sense. The system has the hamiltonian

$$H(t) = \omega J_z + 2\Omega \cos(\omega t) J_x + \Omega_p p_\Omega(t - t_p) J_z, \quad (27)$$

where  $\omega = 2\pi \cdot 700$  kHz,  $\Omega = 2\pi \cdot 1$  kHz,  $\Omega_p = 2\pi \cdot 70$  Hz,  $t_p = 233$  ms and  $p_\Omega(t)$  is a single cycle of a sine wave with frequency  $\Omega$ .

We first wanted to test the accuracy of the different integration techniques for various integration time-steps. Here we wanted to test the advantages of using a

Magnus-based integration method. Accuracy was calculated by taking the quantum state simulation evaluations of a typical quantum sensing experiment and finding the root mean squared (RMS) error from a baseline simulation run by `scipy.integrate.ivan_solve()` as part of the *SciPy* python package, via

$$\epsilon = \frac{1}{K} \sqrt{\sum_{k=0}^{K-1} \sum_{m_j=-j}^j |\psi_{k,(m_j)} - \psi_{k,(m_j)}^{\text{baseline}}|^2}, \quad (28)$$

where  $j \in \{\frac{1}{2}, 1\}$  is the spin quantum number of the system. This baseline was computed in 2.4 hours (in comparison to order of 100 ms of a second these *Spinsim* tests were executed in), and was also used for comparisons to other software packages.

Error vs time-step plots are shown in Figures 2a and 2c. In order to find how long a simulation method takes to complete for a given accuracy, we also measured the execution time for each of the simulations. These plots are shown in Figures 2b and 2d. The former sets of plots will be of interest to users choosing an appropriate time-step for simulations, and the latter for estimating the execution time of such simulations. In all of these comparisons, errors above  $10^{-3}$  were counted as a failed simulation, as the maximum possible error for a quantum state saturates given that it is a point on a unit complex sphere. Errors below  $10^{-11}$  were also excluded, as this was the order of magnitude of the errors in the reference simulation.

The integration techniques tested were the Magnus-based CF4, as well as two Euler-based sampling methods. A midpoint Euler method was chosen as the simplest (and fastest for a given integration time-step) possible sampling method, whereas a Heun-Euler sampling method was used as a comparison to previous versions of this code. These are respectively labelled as the modified Euler method and the improved Euler method in Reference [24, p328]. We benchmarked these methods both while using and not using the rotating frame option. This was done separately for spin-one and spin-half systems, to ensure they both yield accurate results.

From Figure 2, we find that overall, the results that *Spinsim* gives are accurate to those of *SciPy*. Figures 2a, and 2c show that using the Magnus-based integration method is up to 3 orders of magnitude more accurate when compared the Euler-based methods. Also, using the rotating frame increased the accuracy here by 4 orders of magnitude for any individual integration method. When comparing speed vs accuracy in Figures 2b, and 2d, the advantage that CF4 gives in terms of accuracy far outweighs its slower execution speed when compared to midpoint Euler methods.

## 9.2 Comparison to alternatives

We ran the same error and execution time benchmarks on some alternative packages, listed in Table 1, to compare *Spinsim*'s performance to theirs. To obtain simulation results of different accuracies, the step sizes of the alternative integrators were limited to a maximum value. In some cases, the maximum number of steps was modified in some cases to allow for the smaller step sizes. Apart from that, the

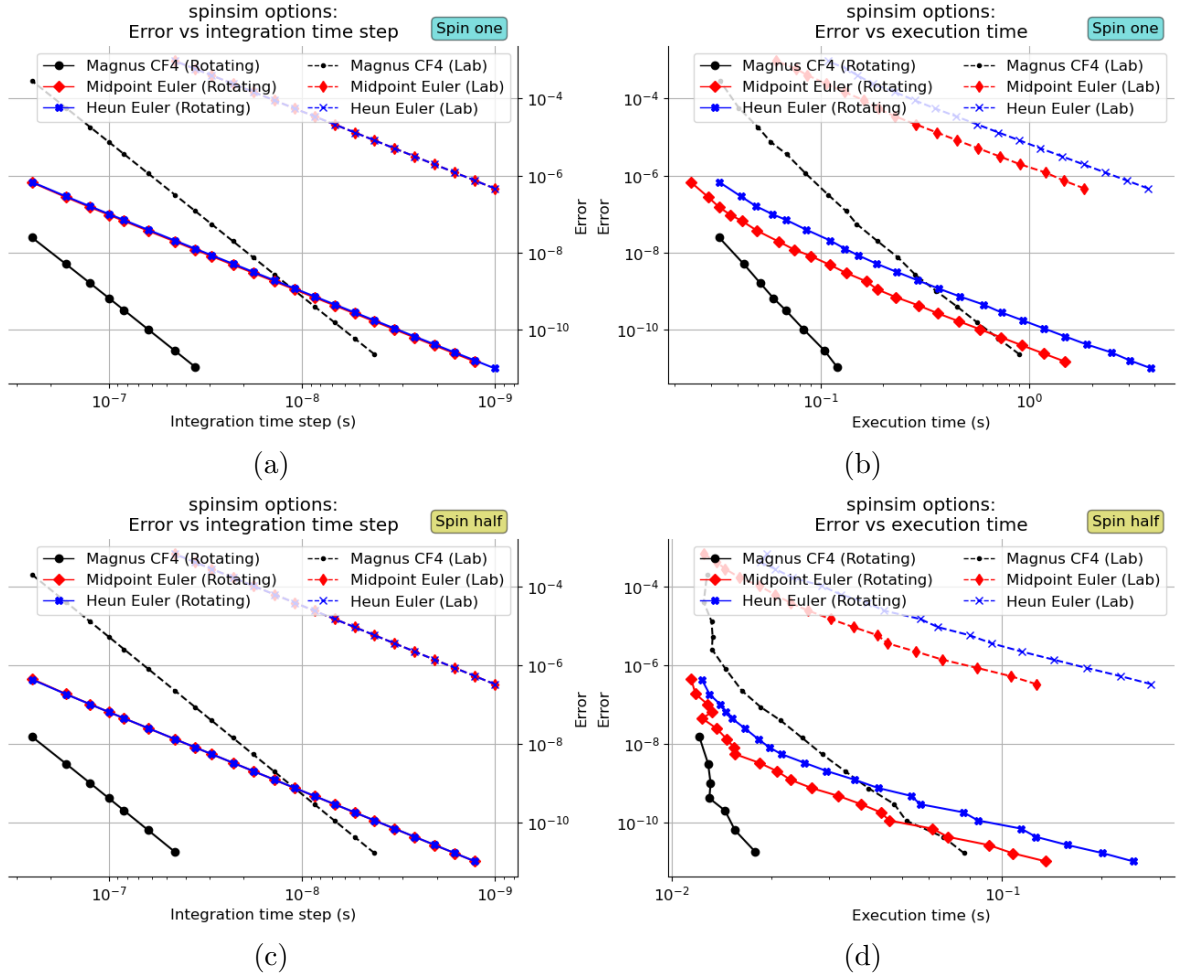


Figure 2: Speed and accuracy of the spin-one and spin-half options of *Spinsim*. A simulation of a typical neural sensing experiment was run for every integration time-step, for each of the possible integration techniques (Magnus-based commutator free 4, and two Euler methods). In the simulation, transitions are continuously driven in the spin system for a duration of 100ms, and an additional small signal is injected for 1ms. See Equation (27). Each technique was tested while both using and not using a transformation into a rotating frame. Both execution time and error were recorded for each of the simulations. Error is RMS error compared to a long running *SciPy* baseline.

integrator settings were left untouched from the default values, as a representation of what a user would experience using a generic solver for spin system problems. Similarly to the internal *Spinsim* benchmarks (see *Evaluation of accuracy*), the expected spin projection was evaluated in each case, but only the states were compared to calculate a relative error via Equation (28). Again, we used the longest running *SciPy* simulation as a baseline for comparison, as the accuracy of *Mathematica* plateaus at small time-steps. Functions used for sampling were compiled to *LLVM* and *cython* in *Spinsim* and *QuTip* respectively. In both cases, the time taken to complete a simulation was measured using a second simulation using the

Table 1: The software packages used for and verification of *Spinsim*.

Software package	Function	Details
Spinsim	<code>Simulator()</code>	The best performing <i>Spinsim</i> configuration, using the CF4 integrator and the rotating frame mode. This was run both on CPU and GPU.
QuTip [25]	<code>sesolve()</code>	The Schrödinger equation solver from the popular <i>python</i> quantum mechanics library, <i>QuTip</i> . This was chosen as a comparison to a specially designed solver used within the physics community for this application. Some quantum mechanics simulation packages such as the recent <i>scqubits</i> [26] use <i>QuTip</i> for calculating time evolution. Like <i>Spinsim</i> , <i>QuTip</i> allows users to sample from compiled functions, and uses parallelisation.
Mathematica [27]	<code>NDSolve()</code>	A generic ODE solver from the <i>Mathematica</i> software. This was chosen as it is popular with our lab group for simulating magnetometry experiments.
SciPy [17]	<code>integrate.ivp_solve()</code>	A generic ODE solver from the popular <i>python</i> scientific computing library. This was chosen as a comparison to a generic solver from within the python ecosystem.

already compiled functions, to represent the use-case of sweeping through many simulations.

The speed of only one simulation was measured for each benchmark. However, it might be possible to increase the average speed of many benchmarks from *Mathematica* and *SciPy* packages by using multithreading to run multiple simulations at a time. When this was attempted using *Mathematica*, the kernels crashed as the 32GiB of RAM was not enough to run them all at once. Multithreading was not attempted using *SciPy*, due to the fact that running the full set of benchmarks of only a single simulation per integration time-step already consumes over 11 hours of computational time. Regardless, for a fair comparison, both *Mathematica* and *SciPy* results are plotted with an artificial reduction in execution time by a factor of 8 (dotted line in Figures 2b and 2d), which is an upper bound for the speed increase that could be obtained by running them parallel on an 8 core processor. Simulations from *Spinsim* and *QuTip* automatically run multithreaded, so this comparison is not plotted for these packages.

From Figure 3, for any given error tolerance, *Spinsim* is over 3 orders of magnitude faster than *Mathematica* and *QuTip*, and 4 orders of magnitude more accurate than *SciPy*. In practice, this means that a 25 minute *SciPy* simulation is reduced to 50ms, and a three week long *SciPy* batch simulation of 1000 separate systems (a realistic requirement for testing quantum sensing protocols) would take less than one minute in *Spinsim*.

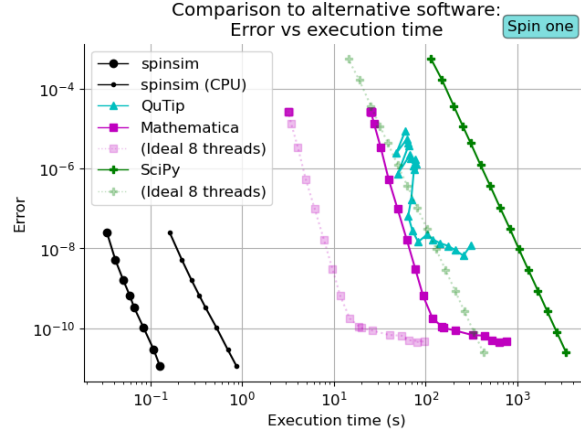


Figure 3: Speed vs accuracy of two alternative integration packages. A simulation of a typical neural sensing experiment was run for every integration time-step, for each of alternative packages (`qutip.sesolve()` from *QuTip*, `NDSolve()` from *Mathematica*, and `scipy.integrate.ivp_solve()` from *SciPy*). In the simulation, transitions are continuously driven in the spin system for a duration of 100ms, and an additional small signal is injected for 1ms. See Equation (27). Both execution time and error were recorded for each of the simulations. Error is RMS error compared to a long running *SciPy* baseline. The *Mathematica* and *SciPy* results are also shown with a speed up by a factor of 8 to represent the upper bound of hypothetical parallelisation across an 8 core CPU.

### 9.3 Parallelisation performance

Once the algorithm behind *Spinsim* was developed, we wanted to check its execution speed while running on various devices. The main reason for this test was to quantify the speed increase of parallelisation by comparing execution speeds on highly parallel devices (being GPUs), and highly procedural devices (being CPUs). Speed benchmarks were performed using `neural-sense.sim.benchmark`, by comparing the evaluation speed of typical spin-one sensing experiments on different devices. Results of these benchmarks are shown in Figure 4. The integration code was compiled by `numba` for multicore CPUs, CPUs running single threaded, and *Nvidia Cuda* compatible GPUs. The compiled code was then run on different models of each of these devices. These test devices are given in Table 2.

The results in Figure 4 show the benefit to using parallelisation when solving a spin system problem. Moving from the 6 core *Core i7-8750H* CPU to the 12 core *Ryzen 9 5900X* CPU doubles the execution speed, as does moving from the

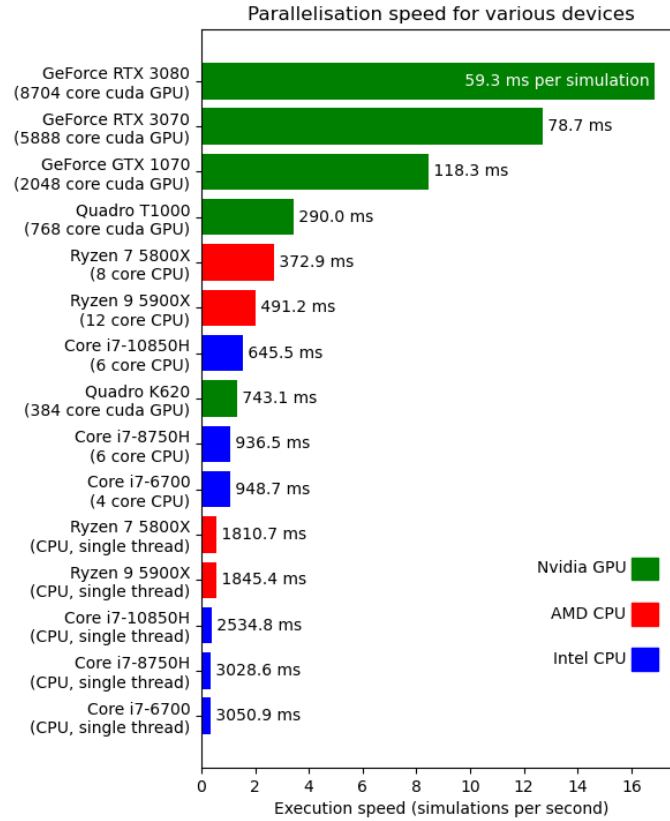


Figure 4: Evaluation speed of a simulation of a typical spin-one sensing experiment on both CPUs and GPUs. Integration time-step is set to 100ns. Transitions are continuously driven in the spin system for a duration of 100ms, and an additional small signal is injected for 1ms. See Equation (27). Evaluation time is determined by an average of 100 similar simulations for each device, where each individual simulation varies in dressing amplitude (transition frequency).



Table 2: Devices used in the parallelisation speed test. These devices are part of individual computers, which are separated here by horizontal lines.

Name	Device	RAM (GiB)	Cores	Cooling
Core i7-6700	Intel CPU	16	4	Air
Quadro K620	Nvidia GPU	2	384	Air
Core i7-8750H	Intel CPU	16	6	Air
GeForce GTX 1070	Nvidia GPU	8	2048	Air
Core i7-10850H	Intel CPU	32	6	Air
Quadro T1000	Nvidia GPU	4	768	Air
Ryzen 9 5900X	AMD CPU	32	12	Air
GeForce RTX 3070	Nvidia GPU	8	5888	Air
Ryzen 7 5800X	AMD CPU	32	8	Liquid
GeForce RTX 3080	Nvidia GPU	10	8704	Air

384 core *Quadro K620* GPU to the 768 core *Quadro T1000* GPU. So, in these cases, performance scales in proportion to thread count. Moving from a single core processor to a high end GPU increases performance by well over an order of magnitude on three of the five computers used for testing. Even the low end *Quadro K620* was an improvement over the *Core i7-6700* used by the same computer. Execution speed vs number of cuda cores starts to plateau as the number of cores increases. This happens because the time it takes to transfer memory from RAM to VRAM (dedicated graphics memory), which is independent on the number of cores of the GPU, becomes comparable to the execution time of the simulator logic. However, there is still a large improvement from using the high end *GeForce RTX 3070* to the *GeForce RTX 3080*, with the latter simulating the experiment almost twice as fast as it would take to run the simulated experiment in the real world.

Surprisingly, we found that the *Ryzen 7 5800X* 8 core CPU was able to execute the benchmark faster than the *Ryzen 9 5900X* 12 core CPU. This can be explained by the fact that the *Ryzen 7 5800X* was cooled by liquid rather than air, meaning it was likely able to boost to a higher core clock, and resist thermal throttling.

Users can view 4 to decide on how much their current or future hardware will be able to take advantage of *Spinsim*'s parallelism. Another factor not shown in the plot is that, in practice, running highly code parallel code on a CPU on a personal computer will severely limit the responsiveness of other applications, as it can utilize the entire CPU (as it should). In contrast, this does not happen when running a GPU based program, as it requires very little CPU utilisation to function. This can be convenient when running simulations on a personal laptop or desktop, as other work on the computer does not have to halt while simulations are being run.

## 9.4 Testing

During the accuracy tests, it was confirmed that all possible modes of *Spinsim* agree with a baseline *SciPy* simulation, as close as the user wants up to the error of  $10^{-11}$  of that baseline. Therefore, *Spinsim* can be trusted for accuracy. The Lie Trotter matrix exponentiator was tested separately from the full system, as well

as benchmarked separately against `scipy.linalg.expm()` from *SciPy*. These tests and benchmarks were run as part of the `neural_sense` package. The simulator has also been adopted by members of our lab, who have given advice on user experience. The kernel execution was profiled thoroughly, and changes were made to optimize VRAM and register usage and transfer. This was done specifically for the development hardware of the *GeForce GTX 1070*, so one may get some performance increases by changing some GPU specific meta parameters when instantiating the `spinsim.Simulator` object.

A good way to confirm that *Spinsim* is functioning properly after an installation is to run the tutorial code provided and compare the outputs. Otherwise, one can reproduce the benchmarks shown here using `neural_sense.sim.benchmark`.

## 10 (2) Availability

### 11 Operating system

Developed and tested on *Windows 10*. CPU functionality tested on *MacOS 10.16 Big Sur* (note that *MacOS 10.14 Mojave* and higher is not compatible with *Cuda* hardware and software). The package (including *Cuda* functionality) is in principle compatible with Linux, but functionality has not been tested.

### 12 Programming language

Python (3.7 or greater)

### 13 Additional system requirements

To use the (default) *Nvidia Cuda* GPU parallelisation, one needs to have a *Cuda* compatible *Nvidia* GPU [28]. For *Cuda* mode to function, one also needs to install the *Nvidia Cuda* toolkit [29]. If *Cuda* is not available on the system, the simulator will automatically parallelize over multicore CPUs instead.

### 14 Dependencies

numba (0.50.1 or greater)

numpy (1.19.3)

matplotlib (for example code, 3.2)

neuralsense (for benchmark code)

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Original conception of released version of algorithm.

## 16 Software location:

### Archive

**Name:** Monash Bridges

**Persistent identifier:** 10.26180/13285460

**Licence:** Apache 2.0

**Publisher:** Alex Tritt

**Version published:** 1.0.0

**Date published:** dd/mm/yy

### Code repository

**Name:** GitHub

**Persistent identifier:** <https://github.com/alexander-tritt-monash/spinsim>

**Licence:** BSD 3 Clause

**Date published:** 18/11/20

## 17 Language

English.

## 18 (3) Reuse potential

### 18.1 Use potential and limitations

*Spinsim* will be useful for any research group needing quick, accurate, and / or large numbers of simulations involving spin-half or spin-one systems. This is immediately relevant to developing new quantum sensing protocols with spin-half and spin-one systems. This package is being used in the context of Bose Einstein Condensate (BEC) magnetic sensing protocol design by our lab.

This project is to be able to measure neural signals using BECs. The electrical pulses made by neurons are currently measured using electrical probes, which is intrusive and damages the cells. We instead propose to sense the small magnetic fields that these electrical currents produce. Rubidium BECs can potentially be made sensitive enough to these tiny magnetic fields that they can be measured by them. *Spinsim* was written to simulate possible measurement protocols for this, showing the behaviour of the array of spin-one atoms interacting with the magnetic fields of the neurons, control signals, and noise. The package is also now being used to simulate other BEC magnetometry experiments by the lab group.

Another example of spin based magnetic field sensing is the use of Nitrogen Vacancy Centres (NVCs). These are spin-one structures found in diamond doped with Nitrogen atoms. This leaves a vacancy in a position adjacent to the Nitrogen atom, which pairs of electrons occupy to obtain the spin-one properties. Similar to BECs, NVCs can be placed and addressed in 2D arrays in order to take many samples in one measurement. A paper was only recently released covering simulation experiments of magnetic neural pulse sensing using NVCs [30], which is something that *Spinsim* could be useful for.

*Spinsim* is designed to simulate small dimensional quantum systems, including large arrays of non-interacting spin systems. This means that it would not be able to integrate large arrays of entangled states or interacting particles. As a result, despite being fast at simulating qubits, it is inappropriate for the package to be used for quantum computing. In addition, *Spinsim* is currently designed to integrate the time evolution of pure states only. This means that it may not be adequate for use in some Nuclear Magnetic Resonance (NMR) applications where relaxation [31] is important (or other kinds of simulations involving decoherence).

With these restrictions in mind, *Spinsim* could be used for some simplified simulations in various areas of NMR. There are many atomic nuclei with spins of half (eg protons, Carbon 13) and, and fewer that have spins of one (eg Lithium 6, Nitrogen 14) [32], which, if relaxation and interactions between systems are not important for the application, *Spinsim* could be used to simulate for spectroscopy experiments, for example. The inclusion of a quadrupole operator means that, with the same level of simplifications, *Spinsim* should be able to simulate Nuclear Quadrupole Resonance (NQR) spectroscopy for spin-one nuclei [33], such as Nitrogen 14, provided a suitable coordinate system is chosen. This technique measures energy level differences between levels split by electric field gradients, rather than static magnetic bias fields. Another possible use-case could be for Magnetic Resonance Imaging (MRI) simulation and pulse sequence design. MRI uses measures the response of spins of an array of spin-half protons to a spatially varying pulse sequence [34], which essentially just corresponds to many separate *Spinsim* simulations of spins at different positions in space. While this package offers some advantages over state of the art simulators in the field [35], with its use of quantum mechanics over classical mechanics, and its absence of rotating wave approximations, its parametrized pulse sequence definitions and geometric integrator, again, the lack of interacting particles and decoherence features are may limit its use in this area.

## 18.2 Support

Documentation for *Spinsim* is available on *Read the Docs*. This documentation contains a thorough tutorial on how to use the package, and installation instructions. For direct support with the *Spinsim* package, one can open an issue in the *github* repository. One can also use this contact to suggest extensions to the package. *Spinsim* is planned to be maintained by the Monash University spinor BEC lab into the future.

## 19 Acknowledgements

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## 21 Competing interests

The authors declare that they have no competing interests.

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