





RadicalPy



Radical pair spin dynamics for the masses

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Motivation



- The radical pair mechanism has gained popularity in recent years:
 - Biological magnetoreception
- Spin chemistry is a relatively niche area of science and can be daunting for people trained in different fields.
- Current toolboxes, such as EasySpin¹ and Spinach² focus on EPR and NMR spin dynamics simulations that require the proprietary commercial³ software MATLAB.
- We aim to build an intuitive (object-oriented) open-source framework in the Python programming language (which is freely available) specific to radical pair spin dynamics.
- Designed for specialists and non-specialists alike to learn and be able perform a plethora of spin dynamics simulations with relative ease.

^{1.} S. Stoll, A. Schweiger, *J. Magn. Reson.*, 178(1), 42-55 (2006)

^{2.} H. J. Hogben, M. Krzystyniak, G. T. P. Charnock, P. J. Hore, I. Kuprov, J. Magn. Reson., 208, 179-194 (2011)

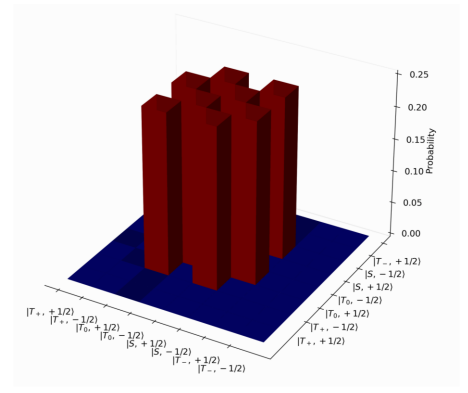
^{3.} Non-negligible price \$\$\$\$\$

RadicalPy

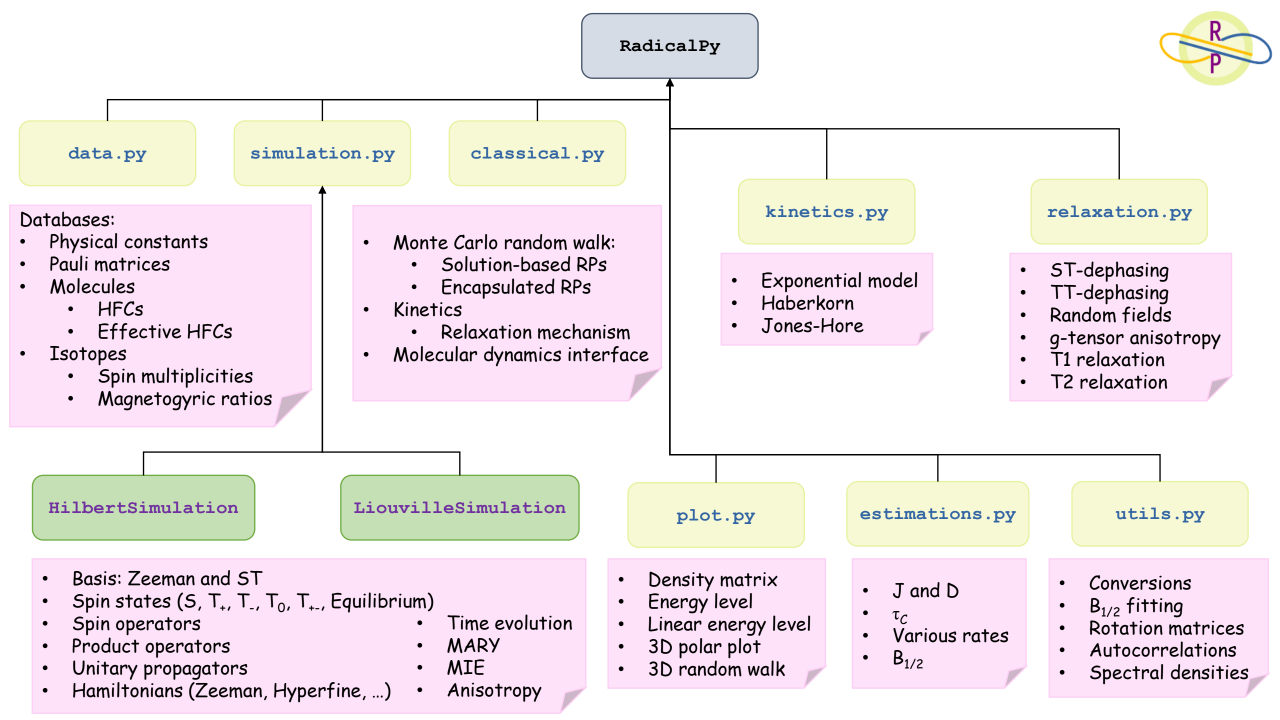


- Open-source, written in Python, available on GitHub:
 - https://github.com/Spin-Chemistry-Labs/radicalpy
 - python -m pip install radicalpy
 - https://radicalpy.readthedocs.io/
- Tutorials
 - Jupyter Notebook (coming soon)
- CI/CD (automated tests, automated documentation)
- License: MIT License
- Standard Python packages:
 - NumPy (http://numpy.scipy.org/)
 - SciPy (http://www.scipy.org/)
 - Matpotlib (<u>http://matplotlib.sourceforge.net/</u>)
- 4,236 lines of code (so far)
- Still in development

Time evolution of a 3D density matrix of a 1-proton singlet born radical pair (with automated spin state labels)



github://examples/density_matrix_3d.py



Building molecules: Interface to molecule and isotope databases



from radicalpy.simulation import Molecule

```
Helpful error messages:

Molecule("adenine_cation", ["buz"])
    Traceback (most recent call last):
    ...
    ValueError: Available nuclei below.
    N6-H2 (hfc = -0.66)
    N6-H1 (hfc = -0.63)
    C8-H (hfc = -0.55)
```

Using the isotope database:

Custom molecules:

Building simulators: Laying the groundwork for simulations



```
import radicalpy as rp
flavin = rp.simulation.Molecule("flavin anion", ["H25"])
Z = rp.simulation.Molecule("Z")
sim = rp.simulation.HilbertSimulation([flavin, Z])
print(sim)
Number of electrons: 2
Number of nuclei: 1
Number of particles: 3
Multiplicities: [2, 2, 2]
Magnetogyric ratios (mT): [-176085963.023, -176085963.023, 267522.18744]
Nuclei: ['H25']
Couplings: [0]
HFCs (mT): [array([[ 0.47570582,  0.04192216, -0.00921916], 
       [0.04192216, 0.4626044, -0.00827001],
       [-0.00921916, -0.00827001, 0.42560999]])]
                      H = sim.total hamiltonian(B=0, D=0, J=0)
                      plt.spy(H)
                      plt.show()
```

$$H_Z = \gamma_e B_0 [\hat{S}_{Lx} \sin \theta \cos \phi + \hat{S}_{Ly} \sin \theta \sin \phi + \hat{S}_{Lz} \cos \theta]$$

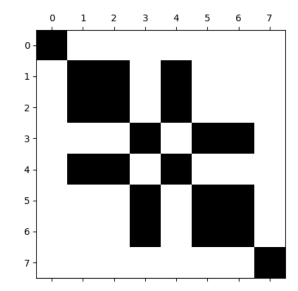
$$H_{H} = \sum_{k=1}^{N_{L}} [\hat{S}_{L} \cdot A_{Lk} \cdot \hat{I}_{Lk}]$$

$$L = \{A, B\}$$

$$H_J = -J\left(2\hat{S}_A \cdot \hat{S}_B + \frac{1}{2}\hat{E}\right) \qquad \qquad \hat{E} = \hat{P}^S + \hat{P}^T$$

$$H_{D} = \sum_{j=x,y,z} \sum_{k=x,y,z} D_{jk} \hat{S}_{Aj} \hat{S}_{Ak}$$

$$H = H_Z + H_H + H_J + H_D$$



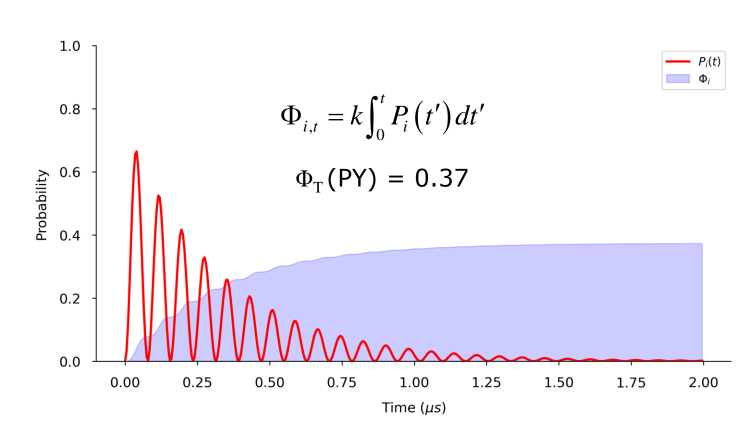
Time evolution and product yield



Hilbert space simulation:

Time evolution and product (triplet) yield for a singlet born flavin-Z radical pair

```
time = np.arange(0, 2e-6, 5e-9)
rhos = sim.time evolution(State.SINGLET, time, H)
k = 3e6
kinetics=[rp.kinetics.Exponential(k)]
time evol = sim.product probability(State.TRIPLET,
                                    rhosl
sim.apply hilbert kinetics(time, time evol, kinetics)
product yield, product yield sum =
          sim.product yield(time evol, time, k)
x = time * 1e6
plt.plot(x, time evol, color="red", linewidth=2)
plt.fill between(x, product yield,
                 color="blue", alpha=0.2)
plt.xlabel("Time ($\mu s$)")
plt.ylabel("Probability"); plt.ylim([0, 1])
plt.legend([r"$P i(t)$", r"$\Phi i$"])
print(f"PY = {product yield sum}")
```



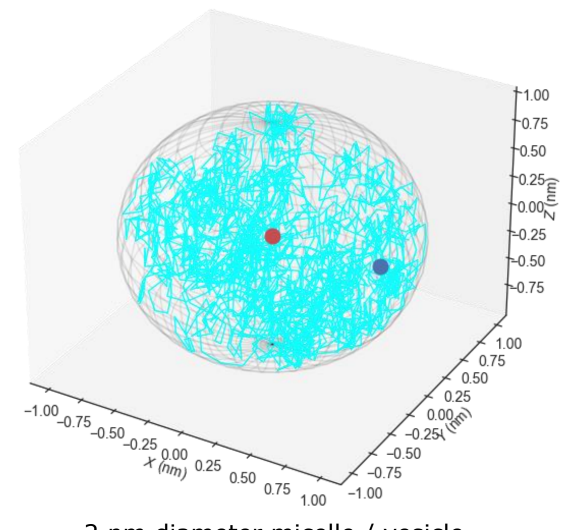
Monte Carlo random walk: Encapsulated radical pair motion



Diffusional motion of the radical pair can be considered as random jumps between two points separated by distance Δr^1 ,

$$\Delta r = \sqrt{6D_{AB}\Delta t}$$

 Δt = time interval D_{AB} = mutual diffusion coefficient of the radical pair

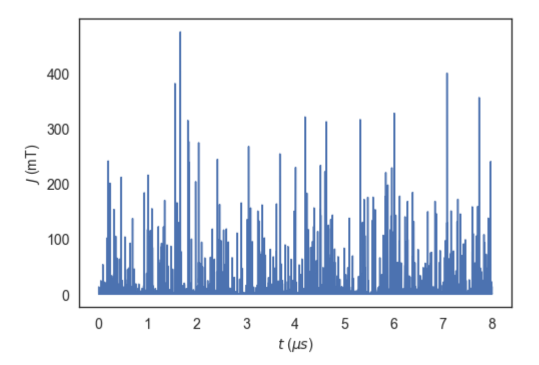


2 nm diameter micelle / vesicle

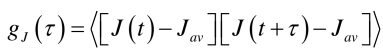
1. T. Miura and H. Murai, J. Phys. Chem. B, 119, 5534-5544 (2015)

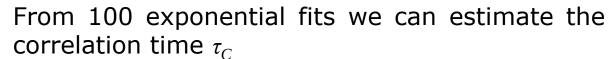
Correlation time estimation

Exchange interaction^{1,2}: $J(r) = J_0 e^{-\alpha(r(t)-r_{\min})}$



Autocorrelation function²:





```
acf j = rp.utils.autocorrelation(J, factor=2)
acf j popt, acf j pcov = curve fit(
    rp.utils.multiexponential fit, x2, y2, p0=a + taus)
acf_j_As, acf_j_taus, tauc_j =
rp.estimations.correlation_time_from_fit(*acf_j_popt)
     0.35
     0.30
     0.25
                            J\tau_{C} = 0.41 \text{ ns}
E 0.15 )
     0.20
     0.10
     0.05
     0.00
```

10⁻⁷

τ (s)

10⁻⁶

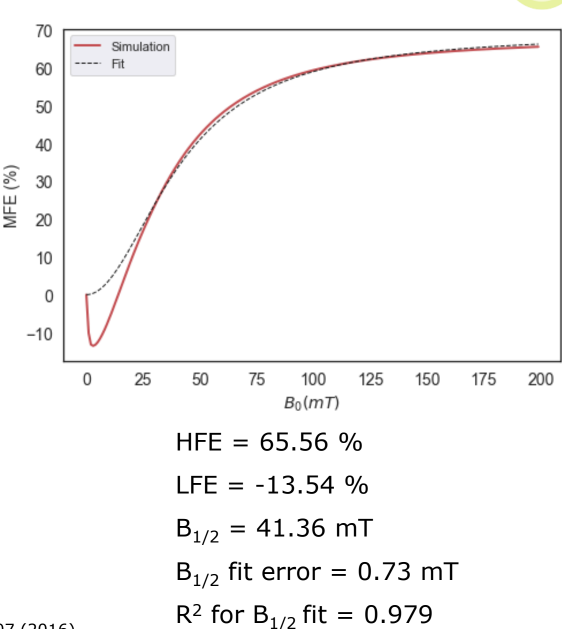
- 1. T. Miura and H. Murai, J. Phys. Chem. B, 119, 5534-5544 (2015)
- 2. D. R. Kattnig, J. K. Sowa, I. A. Solov'yov, and P. J. Hore, New J. Phys. 18, 063007 (2016)

ST-dephasing rate estimation and MARY



Using Redfield theory one can obtain¹:

```
k_{STD} = 4 \left\langle \left[ \left( J(r(t)) - J_{av} \right)^{2} \right] \right\rangle \tau_{C} = 1.33 \times 10^{9} \, s^{-1}
kSTD = rp.estimations.k STD(J, tauc j)
rftb = rp.simulation.Molecule("flavin anion", ["N5", "H25"])
indole = rp.simulation.Molecule("tryptophan cation",
                                                  ["H7", "Hbeta1"])
sim = rp.simulation.LiouvilleSimulation([rftb, indole])
time = np.arange(0, 15e-6, 5e-9)
Bs = np.arange(0, 200, 1)
results = sim.MARY(
    init state=State.TRIPLET,
    obs state=State.TRIPLET,
    time=time,
    B=Bs,
    D=0,
    J=0,
    kinetics=[
         kinetics.Haberkorn(8.4e6, State.SINGLET),
         kinetics.HaberkornFree(1.6e6),
    relaxations=[relaxation.SingletTripletDephasing(kSTD)],
Bhalf, x model MARY, y model MARY, MARY fit error, R2 =
                                       rp.utils.Bhalf fit(Bs, MARY)
```



1. D. R. Kattnig, J. K. Sowa, I. A. Solov'yov, and P. J. Hore, New J. Phys. 18, 063007 (2016)

Radical pair anisotropy



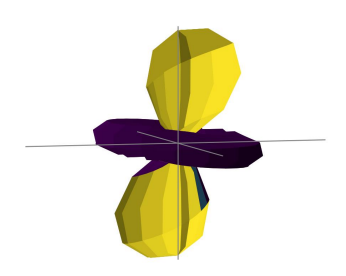
```
theta = np.linspace(0, np.pi, 17)
phi = np.linspace(0, 2 * np.pi, 32)
flavin = rp.simulation.Molecule("flavin anion", ["N5", "N10"])
Z = rp.simulation.Molecule("zorro", [])
sim = rp.simulation.HilbertSimulation([flavin, Z])
time = np.arange(0, 5e-6, 5e-9)
B0 = 0.05
k = 1e6
results = sim.anisotropy(
    init state=State.SINGLET,
    obs state=State.SINGLET,
    time=time,
    theta=theta,
    phi=phi,
    B=B0,
    D=0,
    J=0,
    kinetics=[rp.kinetics.Exponential(k)],
Y = results["product yield sums"]
delta phi s, gamma s = rp.utils.yield anisotropy(Y,
                                  theta, phi)
Y = Y - rp.utils.spherical average(Y, theta, phi)
rp.plot.anisotropy surface(theta, phi, Y)
```

D. R. Kattnig, I. A. Solov'yov, and P. J. Hore, *Phys. Chem. Chem. Phys.*, 18, 12443 (2016)

O. Efimova and P. J. Hore, *Biophysical Journal*, 94, 1565–1574 (2008)

$$\Phi_{S}^{anis}(\theta,\phi) = \Phi_{S}(\theta,\phi) - \langle \Phi_{S} \rangle$$

$$\langle \Phi_S \rangle = \frac{1}{4\pi} \int_0^{2\pi} \int_0^{\pi} \Phi_S(\theta, \phi) \sin\theta d\theta d\phi$$



$$\Delta\Phi_S = \max \Phi_S - \min \Phi_S = 0.262$$

$$\Gamma = \frac{\Delta \Phi_S}{\langle \Phi_S \rangle} = 0.702$$

Outlook



simulation.py

- Spherical tensor basis
- Two-site model
- RYDMR
- Field switching (SEMF)
- Time-resolved MARY
- Semi-classical
- FPR SCRP
- Coherent control

relaxation.py

Rotationally modulated relaxation (Bloch-Redfield-Wangsness theory)

utils.py

- Wigner functions
- Irreducible spherical tensors
- Gaussian / Orca parser

kinetics.py

Diffusion model (Noyes)

Democratising radical pair spin dynamics will require a community:

- Comments
- Remarks
- New molecules (json files)
- Pull requests (expanding the code base)

Future work:

 Large scale (more accurate) spin dynamics simulations on supercomputers (e.g., Fugaku)

Acknowledgements



Supervisors
Jonathan R Woodward (Tokyo)
Kiminori Maeda (Saitama)

Collaborator
Peter J Hore (Oxford)

Help with correlation time estimation

Daniel Kattnig

Funding





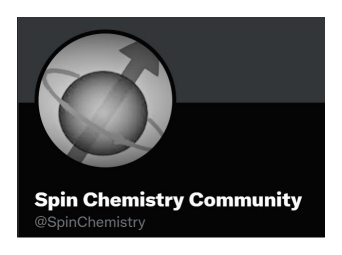


Spin Chemistry Community

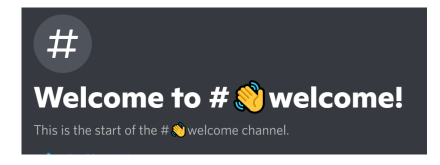




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