An Introduction to Python and Qutip

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Setting up

• Files at https://github.com/peterkirton/qutipdemo

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- Files at https://github.com/peterkirton/qutipdemo
- Run ipython (or python) to start a python session
- Scripts can be run either using
 - run script_name from ipython (better)
 - python script_name from bash
- import numpy as np
- from scipy.foo import bar

Numpy/Scipy

- Provide large set of library functions for scientific applications
- Linear algebra
- Differential equations
- Optimization
- Fourier transforms etc, etc.

Quick examples 1: Linear algebra

- A = np.array([[1, 2], [3, 4]])
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- A = np.array([[1, 2], [3, 4]])
- val, vec = linalg.eig(A)
- B = np.array([[1,2,3],[4,5,6]])
- U,s,Vh = linalg.svd(B)
- Also routines for sparse matrices etc.

Quick examples 2: ODE

- Use ode class from scipy.integrate
- need to specify function of differentials
- Solve:

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Split into 1st order:

$$\frac{dv}{dx} = -5y + J_0(x) \qquad \qquad \frac{dy}{dx} = v$$

Qutip

- A Python toolbox for simulating open quantum systems
- (Relatively) simple to use
- Wraps up useful Numpy/Scipy routines
- Efficient
- Easy to set up complex Hilbert/Liouville spaces
- from qutip import *
 - Brings all qutip functions into the current namespace

States

- fock(N, m) N number of basis states, m Fock state
- $fock_dm(N, m)$
- coherent(N, α) α displacement
- coherent_dm(N, α)
- thermal_dm(N, n) n thermal occupation number

Operators

- qeye(N)
- create(N)
- num(N)
- displace(N, α)
- ullet squeeze(N,sp) sp squeezing parameter
- sigmax(), sigmap() etc
- sigmap() \neq create(2)
- Spin states defined so that fock(2,1)= $|\downarrow\rangle=|1\rangle$

Functions

- Append to an object of class Qobj e.g. create(5).dag()=destroy(5)
- .dag()
- .eigenstates()
- .groundstate()
- .tr()

Some things try

- vac = fock(5,0)
- a = create(5)
- a*vac
- (a**4)*vac
- a.dag()*vac

Two spin model

Two coupled spins

$$H = \Omega_1 \sigma_x^1 + \Omega_2 \sigma_x^2 + g(\sigma_+^1 \sigma_-^2 + \sigma_-^1 \sigma_+^2)$$
$$\dot{\rho} = -i[H, \rho] + \gamma_1 \mathcal{L}[\sigma_-^1] + \gamma_2 \mathcal{L}[\sigma_-^2]$$

- All operators need to be in tensor product space
- σ_x^1 =tensor(sigmax(), Is)
- $\sigma_+^1 \sigma_-^2 = \text{tensor(sigmap(), sigmam())}$
- decay is a list

Rabi model

Coupled spin-photon

$$H = \Omega_0 \sigma_z + \Omega_c a^\dagger a + g \sigma_x (a + a^\dagger)$$

 $\dot{
ho} = -i[H,
ho] + \kappa \mathcal{L}[a]$

- Need to truncate photon Hilbert space Nphot = 10
- Check convergence with this

Waveguide

• Hamiltonian:

$$H = J \sum_{i} a_{i}^{\dagger} a_{i+1} + h.c.$$

$$\dot{\rho} = -i[H, \rho] + \Gamma \sum_{i} \mathcal{L}[a_{i}]$$

- Restrict to certain excitation number subspace
- Use enr_destroy to create a; operators

Other Useful Things

- steadystate steady state density matrix (or expectation values) of superoperator (or Hamiltonian, decay pair)
- correlation_ss ss two-time correlation function
- spectrum_ss
- correlation_2op_2t non-steady two-time correlation function
- bloch_redfield_tensor, bloch_redfield_solve
- etc., etc.

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- Python is not C
- Use the features:
 - Iterables: enumerate rather than range
 - in is really useful
 - Collections module has really useful datatypes
 - Avoid temp variables a,b =b,a
 - try: Except blocks
- import this



Links

- Web:
 - http://qutip.org/
 - http://python.org/
 - https://www.scipy.org/
- Books
 - Learn Python the hard way
 - A Primer on Scientific Programming with Python