An Intro to HOQST - adiabatic master equation

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0.1 Single qubit annealing

This tutorial will recreate the single-qubit example in this paper: Decoherence in adiabatic quantum computation.

The Hamiltonian of this example is

$$H(s) = -\frac{1}{2}(1-s)\sigma_x - \frac{1}{2}s\sigma_z$$
,

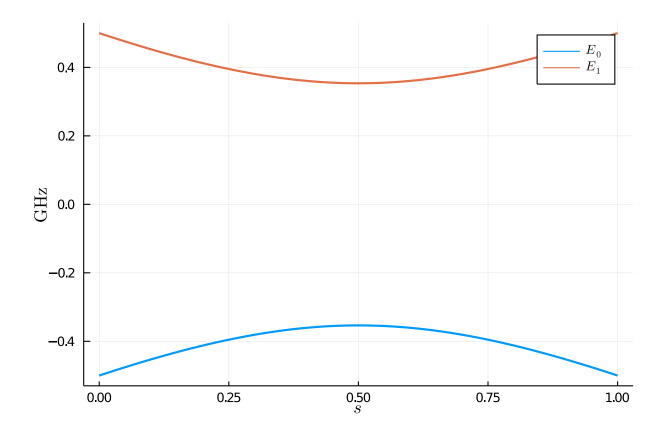
which can be constructed by the following code block

```
using OpenQuantumTools, OrdinaryDiffEq, Plots H = DenseHamiltonian([(s)->1-s, (s)->s], -[\sigma x, \sigma z]/2)
```

DenseHamiltonian with Complex{Float64}
with size: (2, 2)

This package directly interacts with Plots.jl by defining recipes. We can visualize the spectrum of the Hamiltonian by directly plotting the object:

```
# this plot recipe is for conveniently plotting the spectrum of the Hamiltonian # the first 3 arguments are: the Hamiltonian, the grid `s` and the levels to keep plot(H, 0:0.01:1, 2, linewidth=2)
```



0.1.1 Unit $(h \text{ or } \hbar)$

A keyword argument unit whose default value is :h can be provided to any Hamiltonian type's constructor. This argument specifies the unit of other input arguments. For example, setting unit to :h means the other input arguments have the unit of GHz, while setting it to : \hbar means the other input arguments have the unit of 2π GHz. To evaluate the Hamiltonian at a given time, the user should use the evaluate function instead of directly calling it. evaluate will always return the Hamiltonian value in the unit system of h = 1. The following code block shows the effects of different choices of unit:

```
 \begin{aligned} & \text{H_h} = \text{DenseHamiltonian}([(s) -> 1 - s, (s) -> s], -[\sigma x, \sigma z]/2, \text{ unit=:h}) \\ & \text{H_h} = \text{DenseHamiltonian}([(s) -> 1 - s, (s) -> s], -[\sigma x, \sigma z]/2, \text{ unit=:h}) \\ & \text{println}(\text{"Setting unit to :h"}) \\ & \text{@show evaluate}(\text{H_h}, 0.5) \\ & \text{println}(\text{"Setting unit to :h"}) \\ & \text{@show evaluate}(\text{H_h}, 0.5); \\ & \text{Setting unit to :h} \\ & \text{evaluate}(\text{H_h}, 0.5) = \text{Complex}\{\text{Float64}\}[-0.25 + 0.0\text{im} -0.25 + 0.0\text{im}; -0.25 + 0.0\text{im}] \\ & \text{Setting unit to :} \hbar @*(\text{evaluate}(\text{H}(*@\_\hbar @*(, 0.5) = Complex(*@\{\text{Float64}\}[-0.039788735772973836 + 0.0\text{im} -0.039788735772973836 + 0.0\text{im}] \\ & \text{Setting unit to :} \hbar @*(\text{evaluate}(\text{H}(*@\_\hbar @*(, 0.5) = Complex(*@\{\text{Float64}\}[-0.039788735772973836 + 0.0\text{im} -0.039788735772973836 + 0.0\text{im}] \\ & \text{H_h} = \text{DenseHamiltonian}([(s) -> 1 - s, (s) -> s], -[\sigma x, \sigma z]/2, \text{ unit=:h}) \\ & \text{Println}([s) - (s) -
```

Internally, HOQST uses a unit system of $\hbar = 1$. If we call H_h directly, its value is scaled by 2π :

```
H_h(0.5)
```

```
2\times @*(2 \text{ StaticArrays.MArray}(*@{Tuple{2,2},Complex{Float64},2,4}) with indices SOneT o(2) \times @*(SOneTo(2):-1.5708+0.0im -1.5708+0.0im-1.5708+0.0im 1.5708+0.0im
```

0.1.2 Annealing

The total Hamiltonian presented in Ref. is

$$H(s) = H_{\rm S}(s) + gS \otimes B + H_{\rm B}$$
.

We denote S the coupling and $\{gB, H_B\}$ the bath.

Coupling For constant coupling operators, we can use the constructor **ConstantCouplings**. Like Hamiltonian's case, there is a keyword argument unit to specify the input unit.

```
coupling = ConstantCouplings(["Z"])
```

```
ConstantCouplings with AbstractArray\{T,2\} where T and string representation: ["Z"]
```

Bath A bath instance can be any object which implements the following three methods:

- 1. Correlation function: correlation(τ , bath)
- 2. Spectrum density: $\gamma(\omega, \text{ bath})$
- 3. Lamb shift: $S(\omega, bath)$

Redfield/Adiabatic ME solvers require those three methods. Currently, we have built-in support for the Ohmic bath. An Ohmic bath object can be created by:

```
η = 1e-4
fc = 4
T = 16
bath = Ohmic(η, fc, T)

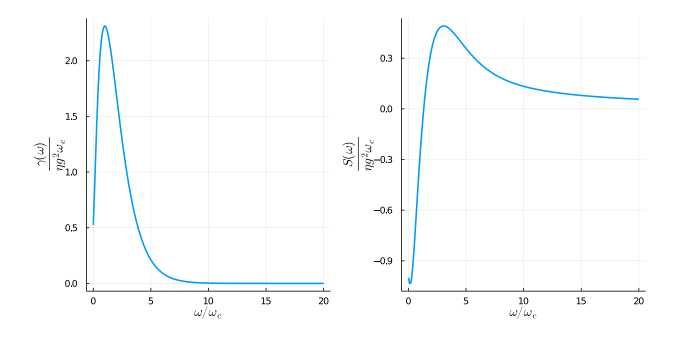
Ohmic bath instance:
η@*( (unitless): 0.0001(*@ω@*(c (GHz): 4.0T (mK): 16.0)

info_freq is a convenient function to convert each quantity into the same unit.

info_freq(bath)

ω@*(c (GHz): 4.0T (GHz): 0.33338579560200365

We can also directly plot the spectrum density of Ohmic bath:
p1 = plot(bath, :γ, range(0,20,length=200), label="", size=(800, 400), linewidth=2)
p2 = plot(bath, :S, range(0,20,length=200), label="", size=(800, 400), linewidth=2)
plot(p1, p2, layout=(1,2), left_margin=3Plots.Measures.mm)
```



Annealing object Finally, we can assemble the annealing object by

```
# Hamiltonian
H = DenseHamiltonian([(s)->1-s, (s)->s], -[\sigma x, \sigma z]/2, unit=:\hbar )
# initial state
u0 = PauliVec[1][1]
# coupling
coupling = ConstantCouplings(["Z"], unit=:\hbar )
# bath
bath = Ohmic(1e-4, 4, 16)
annealing = Annealing(H, u0; coupling=coupling, bath=bath)

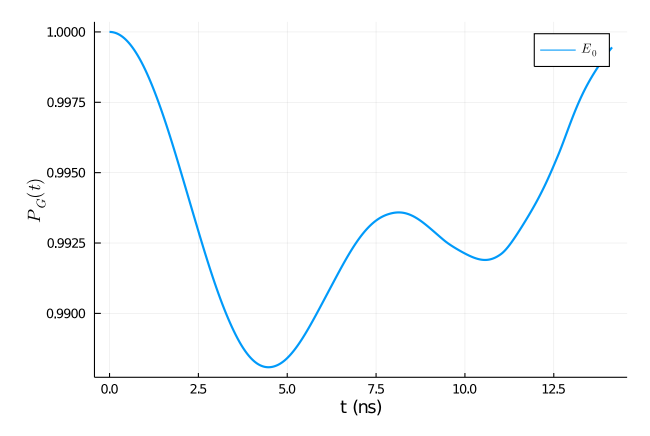
Annealing with hType QTBase.DenseHamiltonian{Complex{Float64}} and uType Ar
ray{Complex{Float64},1}
u0 with size: (2,)
```

Because we want to compare our results to the Ref., we need to set the unit to $\hbar = 1$.

0.1.3 Closed system

There are several interfaces in HOQST that might be handy. The first one is the Schrodinger equation solver:

```
tf = 10*sqrt(2)
@time sol = solve_schrodinger(annealing, tf, alg=Tsit5(), retol=1e-4)
# The following line of code is a convenient recipe to plot the instantaneous population
during the evolution.
# It currently only supports Hamiltonian with annealing parameter s = t/tf from 0 to 1.
# The third argument can be either a list or a number. When it is a list, it specifies
the energy levels to plot (starting from 0); when it is a number, it specifies the total
number of levels to plot.
plot(sol, H, [0], 0:0.01:tf, linewidth=2, xlabel = "t (ns)", ylabel="\$P_G(t)\$")
0.007976 seconds (2.42 k allocations: 147.051 KiB)
```



The solution is an ODESolution object in DifferentialEquations.jl package. More details for the interface can be found here. The state vector's value at a given time can be obtained by directly calling the ODESolution object.

```
sol(0.5)
```

```
2-element Array{Complex{Float64},1}:
0.6856253144209079 + 0.1750041214939618im
0.6861430138705714 + 0.1688172359560306im
```

Other interfaces include

```
# You need to solve the unitary first before trying to solve Redfield equation
@time U = solve_unitary(annealing, tf, alg=Tsit5(), abstol=1e-8, retol=1e-8);
@time solve_von_neumann(annealing, tf, alg=Tsit5(), abstol=1e-8, retol=1e-8);
```

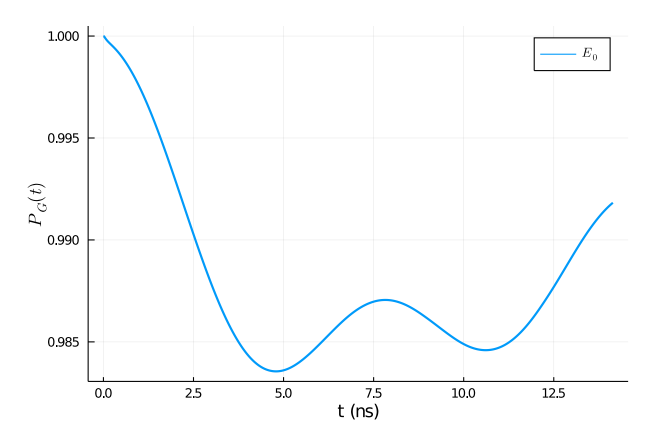
0.1.4 Open System

Time-dependent Redfield equation The time-dependent Redfield equation solver needs

- 1. Annealing object
- 2. Total annealing time
- 3. Pre-calculated unitary

to work. The following code block illustrates how to supply the above three objects to the Redfield solver. Besides, all the other keyword arguments in DifferentialEquations.jl are supported.

```
tf = 10*sqrt(2)
U = solve_unitary(annealing, tf, alg=Tsit5(), abstol=1e-8, retol=1e-8);
sol = solve_redfield(annealing, tf, U; alg=Tsit5(), abstol=1e-8, retol=1e-8)
plot(sol, H, [0], 0:0.01:tf, linewidth=2, xlabel="t (ns)", ylabel="\$P_G(t)\$")
```

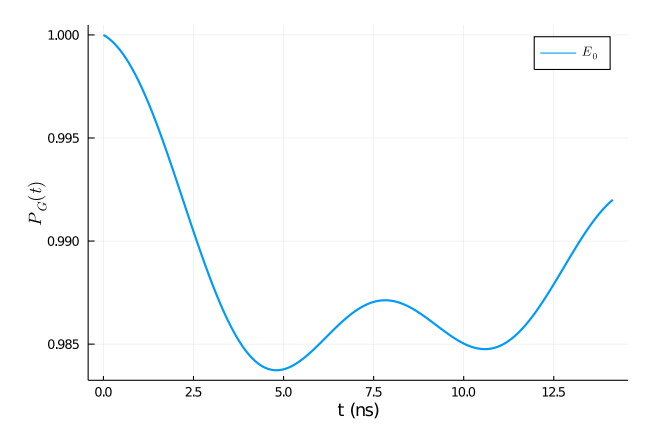


Adiabatic master equation The adiabatic master equation solver needs

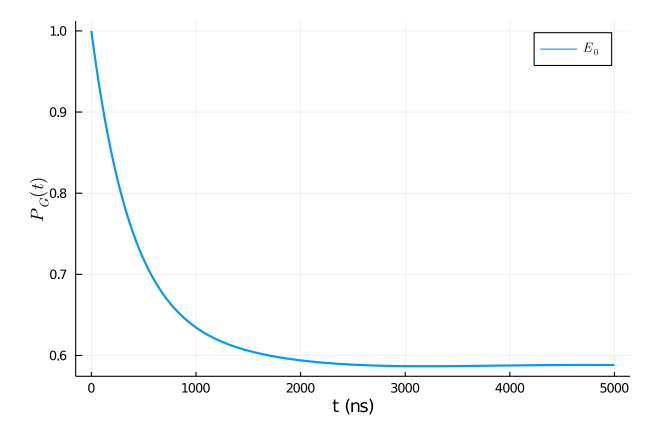
- 1. Annealing object
- 2. Total Annealing time

Besides other keyword arguments supported in DifferentialEquations.jl, it is highly recommended to add the ω _hint keyword argument. By doing this, the solver will pre-compute the quantity $S(\omega)$ in Lambshift within the range specified by ω _hint to speed up the computation.

```
tf = 10*sqrt(2)  
@time sol = solve_ame(annealing, tf; alg=Tsit5(), \omega_{\rm hint=range}(-6, 6, length=100), reltol=1e-4)  
plot(sol, H, [0], 0:0.01:tf, linewidth=2, xlabel="t (ns)", ylabel="\$P_G(t)\$")  
0.010107 seconds (193.16 k allocations: 4.270 MiB)
```



We can also solve the AME for a longer annealing time:



The above results agree with Fig 2 of the reference paper.

Trajectory method for adiabatic master equation The package also supports the trajectory method for AME. More details of this method can be found in this paper. The basic workflow is to create an ODE EnsembleProblem via build_ensembles interface. Then, the resulting EnsembleProblem object can be solved by the native Parallel Ensemble Simulations interface of DifferentialEquations.jl. The following code block solves the same annealing process described above($t_f = 5000(ns)$) using multithreading. To keep the running time reasonably short, we simulate only 3000 trajectories in this example. The result may not converge to the true solution yet. The user is encouraged to try more trajectories and see how the result converges.

The codes can also be deployed on high-performance clusters using Julia's native distributed computing module.

```
tf = 5000
# total number of trajectories
num_trajectories = 3000
# construct the `EnsembleProblem`
# `safetycopy` needs to be true because the current trajectories implementation is not thread-safe.
prob = build_ensembles(annealing, tf, :ame, ω_hint=range(-6, 6, length=100),
safetycopy=true)
# to use multi-threads, you need to start Julia kernel with multiple threads
# julia --threads 8
sol = solve(prob, Tsit5(), EnsembleThreads(), trajectories=num_trajectories,
reltol=1e-6, saveat=range(0,tf,length=100))
t_axis = range(0,tf,length=100)
dataset = []
```

```
for t in t_axis
   w, v = eigen_decomp(H, t/tf)
   push!(dataset, [abs2(normalize(so(t))' * v[:, 1]) for so in sol])
end
# the following codes calculate the instantaneous ground state population and its error
bar by averaging over all the trajectories
pop_mean = []
pop_sem = []
for data in dataset
   p_mean = sum(data) / num_trajectories
   p_sem = sqrt(sum((x)->(x-p_mean)^2, data)) / num_trajectories
   push!(pop_mean, p_mean)
   push! (pop_sem, p_sem)
end
scatter(t_axis, pop_mean, marker=:d, yerror=2*pop_sem, label="Trajectory", markersize=6)
plot!(sol_ame, H, [0], t_axis, linewidth=2, label="Non-trajectory")
xlabel!("t (ns)")
ylabel!("\P_G(s)\")
    1.0
                                                                       Trajectory
                                                                       Non-trajectory
    0.9
 P_G(s) \\ {\rm gc}
    0.7
    0.6
```

2000

t (ns)

3000

1000

0

4000

5000