Solve the GPE in a 1D parabolic trap

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April 12, 2019

1 Introduction

In this simple example we find a ground state of the Gross-Pitaevskii equation in a harmonic trap.

The mean field order parameter evolves according to

$$i\hbar \frac{\partial \psi(x,t)}{\partial t} = \left(-\frac{\hbar^2 \partial_x^2}{2m} + V(x,t) + g|\psi(x,t)|^2\right)\psi(x,t)$$

2 Loading the package

```
First, we load some useful packages.
```

```
using Plots, LaTeXStrings, Pkg, Revise gr(legend=false,titlefontsize=12,size=(500,300),transpose=true,colorbar=false);
```

Now load FourierGPE

using FourierGPE

Let's define a convenient plot function

```
function showpsi(x, \( \psi \)
    p1 = plot(x,abs2.(\( \psi \)))
    xlabel!(L"x/a_x");ylabel!(L"\\psi|^2")
    p2 = plot(x,angle.(\( \psi \)))
    xlabel!(L"x/a_x");ylabel!(L"\\textrm{\phase}(\\psi)")
    p = plot(\( \psi \), \( \psi \) zize=(600,400))
    return p
end

showpsi (generic function with 1 method)
```

3 User parameters

We reserve a place for user parameters.

3.1 Declaring the potential

Let's define the trapping potential.

```
import FourierGPE.V
V(x,t) = 0.5*x^2
V (generic function with 3 methods)
```

We only require that it is a scalar function because alter we will evaluate it using a broad-casted dot-call.

4 Initial condition

Let's define a useful Thomas-Fermi wavefunction

```
\psi 0(x,\mu,g) = \operatorname{sqrt}(\mu/g) * \operatorname{sqrt}(\max(1.0-V(x,0.0)/\mu,0.0) + \operatorname{im}*0.0)  x = X[1]; The initial state is now created as \psi i = \psi 0.(x,\mu,g) \psi i : += (\operatorname{randn}(N...) \mid > \operatorname{complex}) \phi i = \operatorname{kspace}(\psi i, \operatorname{sim}) \operatorname{Opack!} \sin = \phi i;
```

5 Evolution in k-space

```
The FFTW library is used to evolve the Gross-Pitaevskii equation in k-space sol = runsim(sim.\phi i, sim);
0.481494 seconds (1.42 M allocations: 68.921 MiB, 5.51% gc time)
```

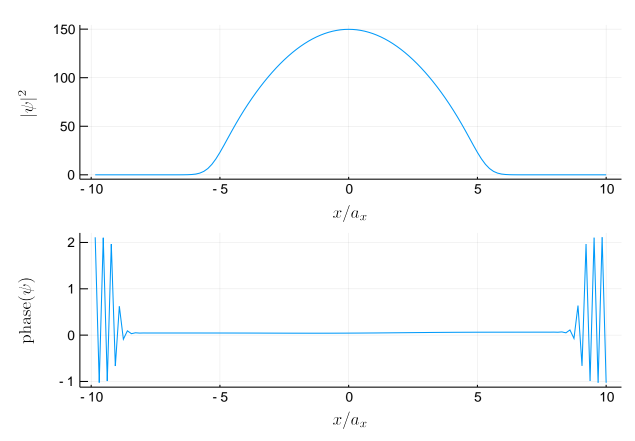
Here we save the entire solution as a single variable sol.

Let's have a look at the final state and verify we have a ground state

```
\phig = sol[end]

\psig = xspace(\phig,sim)

showpsi(x,\psig)
```



The initial Thomas-Fermi state has been evolved for a default time $t=2/\gamma$ which is a characteristic damping time for the dissipative system with dimensionless damping γ . The solution will approach the ground state satisfying $L\psi_0 = \mu\psi_0$ on a timescale of order $1/\gamma$. The figure shows a smooth density profile and a completely homogeneous phase profile over the region of finite atomic density, as required for the ground state. The indeterminate phase evident at large |x| is unimportant.

5.1 Default simulation parameters

The default parameters are given in the declaration of Sim, which allows parameter interdependence. The struct Sim is declared as:

```
@with_kw mutable struct Sim{D} <: Simulation{D} @deftype Float64
    L::NTuple{D,Float64}
    N::NTuple{D,Int64}
    \mu = 15.0
    g = 0.1
    \gamma = 0.5; @assert \gamma >= 0.0
    ti = 0.0
    tf = 2/\gamma
    Nt::Int64 = 200
    t::LinRange{Float64} = LinRange(ti,tf,Nt)
    \phii::Array{Complex{Float64},D} = zeros(N...) |> complex
```

```
params::UserParams # optional parameters
X::NTuple{D,Array{Float64,1}} = xvecs(L...,N...)
K::NTuple{D,Array{Float64,1}} = kvecs(L...,N...)
espec::Array{Complex{Float64},D} = k2(L...,N...)
T::TransformLibrary{D} = Transforms{D}()
```

where we see a set of default parameters, and then some useful transform fields built using the parameters. Note that the transforms have to be built after building X,K.

6 Dark soliton in harmonically trapped system

We found a ground state by imaginary time propagation. Now we can impose a phase and density imprint consistent with a dark soliton. We will use the solution for the homogeneous system, which will be a reasonable approximation if we impose it on a smooth background solution.

6.1 Imprinting a dark soliton

```
\begin{split} &\psi \mathbf{f} = \mathtt{sol}[\mathtt{end}] \\ &\mathbf{c} = \mathtt{sqrt}(\mu) \\ &\xi = 1/\mathtt{sqrt}(\mathtt{g*abs2.}(\psi \mathtt{f}[\mathtt{Int}(\mathtt{end/2})])) \\ &\mathbf{v} = 0.3*\mathtt{c}; \end{split} Soliton speed is determined by depth and local healing length. Start at xs = 0.0 xs = 0. \mathbf{v} = 0.5*\mathtt{c} \mathbf{f} = \mathtt{sqrt}(1-(\mathtt{v/c})^2) \psi \mathbf{s} = \mathbf{0}. \psi \mathbf{f} * (\mathtt{f*tanh}(\mathtt{f*(x - xs)/\xi}) + \mathtt{im*v/c});
```

6.2 Initilize Simulation

We can recycle our earlier parameter choices, modifying the damping and simulation timescale

```
γ = 0.0
tf = 8*pi/sqrt(2);
simSoliton = Sim(γ=γ,tf=tf)

Error: Field 'L' has no default, supply it with keyword.

@pack! simSoliton = T,X,K,espec

Error: UndefVarError: simSoliton not defined
initsim!(simSoliton)

Error: UndefVarError: simSoliton not defined

@unpack_Sim simSoliton;

Error: UndefVarError: simSoliton not defined
```

In doing so, we have to specify the dimension of the simulation in this case (an improved constructor needed).

6.3 Solve equation of motion

```
As before, we specify the initial condition in momentum space, and evolve \phii = kspace(\psis,simSoliton)

Error: UndefVarError: simSoliton not defined

Opack! simSoliton = \phii;

Error: UndefVarError: simSoliton not defined

sols = runsim(simSoliton.\phii,simSoliton);

Error: UndefVarError: simSoliton not defined
```

6.4 View the solution using Plots

```
y = g*abs2.(sols[Nt-4])
Error: UndefVarError: sols not defined
plot(x,y,fill=(y,0.4),size=(400,200),legend=false,grid=false)
Error: UndefVarError: y not defined
xlims!(-10,10);ylims!(0,1.2*\mu)
xlabel!(L"x/a_x")
ylabel!(L"\mu(x)/\hbar\omega_x")
title!(L"\textrm{local}\;\mu",titlefontsize=12)
```

