# Time-Dependent Schrödinger Equation with Magnetic Field

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## 1 Introduction

#### 1.1 Mathematic model

In this report we consider a spinless particle in  $\mathbb{R}^d$  with mass  $m \in \mathbb{R}_{\geq 0}$  and charge  $e \in \mathbb{R}$  in a homogeneous magnetic field B(t). We follow the notation introduced in [1] and quickly recap the important elements. For a full derivation please consult [1]. In quantum mechanics, the time evolution of a particle subject to a magnetic field is governed by the Pauli equation

$$i\hbar \partial_t \Psi(x,t) = H_P(t)\Psi(x,t) \tag{1}$$

$$H_P(t) := \frac{1}{2m} \sum_{k=1}^{d} (p_k - eA_k(x,t))^2 + e\phi(x,t) + \tilde{V}(x,t)$$
 (2)

where  $\tilde{V}(x,t)$  is some external potential. The magnetic field 2-form dA associated with B(t) is independent of x because of the homogeneity of B(t) and we can thus rewrite the magnetic vector potential to

$$A(x,t) := \frac{1}{2} B_{jk}(t) x^j \mathrm{d}x^k, \tag{3}$$

where  $B(t) = (B_{jk}(t))_{i,k=1}^d$  is a real, skew-symmetric matrix. Using the operators

$$L_{jk} := x_j p_k - x_k p_j \tag{4}$$

$$H_B(t) := -\sum_{j,k=1}^{d} B_{jk}(t) L_{jk}$$
 (5)

the Pauli-Hamiltonian takes the form

$$H_P(t) = \frac{1}{2m} \left( \hbar^2(-\Delta) - e \sum_{1 \le j < k \le d} B_{jk}(t) L_{jk} + \frac{e^2}{4} \|B(t)x\|_{\mathbb{R}^d}^2 \right) + e\phi(x,t) + \tilde{V}(x,t). \tag{6}$$

### 1.2 Numerical model

We introduce  $\epsilon^2 := \hbar$  and redefine t, x and B to find the simplified form

$$H_P(t) = -\Delta + H_B(t) + V(x,t) \tag{7}$$

where  $V(x,t) := \frac{1}{2m} \frac{e^2}{4} ||B(t)x||_{\mathbb{R}^d}^2 + e\phi(x,t) + \tilde{V}(x,t)$  can be considered to be a total effective potential. The Schrödinger equation

$$i\epsilon^2 \partial_t \Psi(x,t) = H_P(t)\Psi(x,t)$$
 (H)

is then split up into three separate parts that can be solved numerically.

$$i\epsilon^2 \partial_t \Psi = -\Delta \Psi \tag{K}$$

$$i\epsilon^2 \partial_t \Psi = H_B(t)\Psi$$
 (M)

$$i\epsilon^2 \partial_t \Psi = V(x, t)\Psi$$
 (P)

(K) can be solved discretely in Fourier-space and (P) by pointwise multiplication with  $e^{-i/\epsilon^2 \int_{t_0}^t dt V(x,t)}$ .

(M) is reduced to the linear differential equation

$$\frac{d}{dt}y(t) = B(t)y(t) \tag{B}$$

[2] proves the existence of a flow map  $U(t,t_0)$  which is a solution to (B). The unitary representation

$$\rho: SO(d) \longrightarrow U(L^2(\mathbb{R}^d)) \tag{8}$$

$$R \longmapsto (\rho(R)\Psi)(x) = \Psi(R^{-1}x) \tag{9}$$

maps the solution  $U(t,t_0)$  of (B) to a solution of (M). The proof of this statement can be found in [1]. The exact flow map  $U(t,t_0)$  is approximated through a Magnus expansion proposed by [3]. Direct calculation shows  $[-\Delta, H_B(t)] = 0$ . Thus the flow maps solving (K) and (M) yield a solution to the differential equation

$$i\epsilon^2 \partial_t \Psi = (-\Delta + H_B(t))\Psi$$
 (K+M)

in the following way: Denote the solutions to (K) and (M) by  $\Phi_{-\Delta}$  and  $\Phi_{H_B}$  respectively. This means that the flow map to (K) satisfies

$$i\frac{d}{dt}\Phi_{-\Delta}(t,t_0) = -\Delta\Phi_{-\Delta}(t,t_0), \quad \Phi_{-\Delta}(t_0,t_0) = id, \tag{10}$$

and analogously for the flow map to (M). The flow map  $\Phi_{-\Delta+H_B}$  which is a solution to (K+M) then follows as a simple multiplication

$$\Phi_{-\Delta + H_B}(t, t_0) = \Phi_{H_B}(t, t_0)\Phi_{-\Delta}(t, t_0) = \Phi_{-\Delta}(t, t_0)\Phi_{H_B}(t, t_0). \tag{11}$$

Finally, combining the solutions to (K+M) and (P) using a splitting scheme leads to a solution of H.

#### 1.3 Splitting

Consider a splitting scheme with the coefficients  $(a_i, b_i)$  for  $1 \le i \le n$ . Consider the time grids

$$t_i = t_0 + (t - t_0) \sum_{j=1}^{i} b_j$$
 and  $s_i = t_0 + (t - t_0) \sum_{j=1}^{i} a_j$  (12)

to an initial time  $t_0$ . [1] derives an explicit expression of the solution to H as follows:

$$\Phi_{H}\left(t_{0}+Nh,t_{0}\right) \approx \left(\prod_{j=0}^{N-1}\prod_{i=0}^{n-1}\Phi_{-\Delta}\left(t_{i+1},t_{i}\right)\Phi_{\rho(U(t_{0}+Nh,t_{i}+jh))V}\left(s_{i+1}+jh,s_{i}+jh\right)\right)\Phi_{H_{B}}\left(t_{0}+Nh,t_{0}\right)$$
(13)

where h is the splitting time step,  $\rho$  the representation defined in eq. 8 and  $U(t, t_0)$  the exact flow map to (B).

## 2 Implementation

#### 2.1 WaveBlocksND

//something about waveblocks

#### 2.2 Code

```
r"""The WaveBlocks Project

This file contains the Fourier Magnetic Propagator class. The wavefunction

imath: '\Psi' is propagated in time with a splitting of the
exponential :math: '\exp(-\frac{i}{\varepsilon^2}\\tau H)'.

dauthor: R. Bourquin
copyright: Copyright (C) 2012, 2016 R. Bourquin
clicense: Modified BSD License
"""

from numpy import array, complexfloating, dot, exp, eye, zeros, shape
from numpy.fft import fftn, ifftn
from scipy.linalg import expm

from WaveBlocksND.BlockFactory import BlockFactory
from WaveBlocksND.Fropagator import Propagator
from WaveBlocksND.KineticOperator import KineticOperator
from WaveBlocksND.MagneticField import MagneticField
```

```
20 from WaveBlocksND.SplittingParameters import SplittingParameters
21
__all__ = ["FourierMagneticPropagator"]
23
24
25 class FourierMagneticPropagator(Propagator, SplittingParameters):
      r"""This class can numerically propagate given initial values :math: `\Psi(x_0, t_0
26
      a potential hyper surface :math: `V(x)`, in presence of a magnetic field. The
27
      propagation is done with a splitting
      of the time propagation operator :math:'\exp(-\frac{i}{\varepsilon^2} \tau H)'.
28
       Available splitting schemes are implemented in :py:class:'SplittingParameters'.
29
30
31
32
      def __init__(self, parameters, potential, initial_values):
33
          r"""Initialize a new :py:class:'FourierMagneticPropagator' instance.
      Precalculate the
           the kinetic operator :math: 'T_e' and the potential operator :math: 'V_e'
34
          used for time propagation.
35
36
37
          :param parameters: The set of simulation parameters. It must contain at least
                               the semi-classical parameter :math: '\varepsilon' and the
38
                               time step size :math: '\tau'.
39
          :param potential: The potential :math: {}^{\circ}V(x) {}^{\circ} governing the time evolution.
40
           :type potential: A :py:class:'MatrixPotential' instance.
41
          :param initial_values: The initial values :math:'\Psi(\Gamma, t_0)' given
42
                                  in the canonical basis.
43
          :type initial_values: A :py:class:'WaveFunction' instance.
44
45
46
           :raise: :py:class:'ValueError' If the number of components of :math:'\Psi'
      does not match the
                               number of energy surfaces :math:'\lambda_i(x)' of the
47
      potential.
48
          :raise: :py:class:'ValueError' If the number of components of :math:'\Psi'
      does not match the dimension of the magnetic field :math: \vec{B}(x).
50
          :raise: :py:class:'ValueError' If the dimensions of the splitting scheme
51
      parameters :math:'a' and :math:'b' are not equal.
52
          # The embedded 'MatrixPotential' instance representing the potential 'V'.
53
          self._potential = potential
54
55
56
           # The initial values of the components '\psi_i' sampled at the given grid.
57
          self._psi = initial_values
58
          if self._potential.get_number_components() != self._psi.get_number_components
59
       ():
               raise ValueError("Potential dimension and number of components do not
60
      match.")
61
          # The time step size.
          self._dt = parameters["dt"]
63
64
65
           # Final time.
           self._T = parameters["T"]
66
67
          # The model parameter '\varepsilon'.
68
```

```
69
           self._eps = parameters["eps"]
70
           # Spacial dimension d
71
           self._dimension = parameters["dimension"]
72
73
           # The position space grid nodes '\Gamma'.
74
           self._grid = initial_values.get_grid()
75
76
           # The kinetic operator 'T' defined in momentum space.
77
           self._KO = KineticOperator(self._grid, self._eps)
78
79
           # Exponential '\exp(-i/2*eps^2*dt*T)' used in the Strang splitting.
80
81
           # not used
           self._KO.calculate_exponential(-0.5j * self._dt * self._eps**2)
82
83
           self._TE = self._KO.evaluate_exponential_at()
84
           # Exponential '\exp(-i/eps^2*dt*V)' used in the Strang splitting.
85
           # not used
86
           self._potential.calculate_exponential(-0.5j * self._dt / self._eps**2)
87
           VE = self._potential.evaluate_exponential_at(self._grid)
88
           self._VE = tuple([ve.reshape(self._grid.get_number_nodes()) for ve in VE])
89
90
91
           # The magnetic field
           self._B = MagneticField(parameters["B"])
92
           # check if magnetic field and potential are of same dimension
93
           if self._B.get_dimension() != self._dimension:
94
               raise ValueError("Spacial dimension of potential and magnetic field must
95
       be the same")
96
97
           #precalculate the splitting needed
           self._a, self._b = self.build(parameters["splitting_method"])
98
           if shape(self._a) != shape(self._b):
99
                raise ValueError("Splitting scheme shapes must be the same")
           # Get inital data as function
           packet_descr = parameters["initvals"][0]
           self._initalpacket = BlockFactory().create_wavepacket(packet_descr)
104
105
106
107
       # TODO: Consider removing this, duplicate
       def get_number_components(self):
108
           r"""Get the number :math:'N' of components of :math:'\Psi'.
109
110
           :return: The number :math:'N'.
           return self._potential.get_number_components()
113
114
115
       def get_wavefunction(self):
116
           {\tt r"""}{\tt Get \ the \ wavefunction \ that \ stores \ the \ current \ data \ :math:`{\tt Psi(\Gamma)'}.
117
118
           :return: The :py:class:'WaveFunction' instance.
119
120
           return self._psi
121
123
124
     def get_operators(self):
```

```
r"""Get the kinetic and potential operators :math:'T(\Omega)' and :math:'V(\
       Gamma) '.
126
           :return: A tuple :math: '(T, V)' containing two ''ndarrays''.
127
128
           # TODO: What kind of object exactly do we want to return?
           self._KO.calculate_operator()
130
           T = self._KO.evaluate_at()
132
           V = self._potential.evaluate_at(self._grid)
           V = tuple([v.reshape(self._grid.get_number_nodes()) for v in V])
134
           return (T, V)
135
136
       Ostaticmethod
138
       def _Magnus_CF4(tspan, B, N, *args):
139
           r"""Returns the Fourth Order Magnus integrator :math: '\Omega(A)' according to
        [#]_.
140
           :param tspan: Full timespan of expansion.
141
142
           :param B: Magnetic field matrix :math: (B(t) = (B_{j,k}(t))_{1 \leq j,k} (t)
143
       14.
144
           :param N: Number of timesteps for the expansion.
145
146
           :param *args: Additional arguments for the magnetic field :math:'B(t, *args)'
147
148
           .. [#] S. Blanes and P.C. Moan. "Fourth- and sixth-order commutator-free
149
       Magnus integrators for linear and non-linear dynamical systems". Applied Numerical
       Mathematics, 56(12):1519 - 1537, 2006.
150
           # Magnus constants
151
           c1 = 0.5*(1.0 - 0.5773502691896258)
152
           c2 = 0.5*(1.0 + 0.5773502691896258)
           a1 = 0.5*(0.5 - 0.5773502691896258)
154
           a2 = 0.5*(0.5 + 0.5773502691896258)
156
           R = 1.*eye( len( B(1.*tspan[0], *args) ) )
157
           h = (tspan[1] - tspan[0]) / (1.*N)
158
159
           for k in range(N):
               t0 = k*h + tspan[0]
160
               t1 = t0 + c1*h
161
               t2 = t0 + c2*h
162
               B1 = B(t1, *args)
               B2 = B(t2, *args)
               R = dot(expm(a1*h*B1+a2*h*B2), dot(expm(a2*h*B1+a1*h*B2), R))
165
           return R
167
168
169
       def post_propagate(self, tspan):
170
           r"""Given an initial wavepacket :math:'\Psi_0' at time :math:'t=0', calculate
       the propagated wavepacket :math: '\Psi' at time :math: 'tspan \[ 0 \]'. We perform :
       math:'n = \lceil tspan\[ 0 \] /dt \rceil' steps of size :math:'dt'.
           :param tspan: :py class:'ndarray' consisting of end time at position 0, other
173
       positions are irrelevant.
174
```

```
175
           # (ignoriere tspan[0])
176
           nsteps = int(tspan[0] / self._dt + 0.5)
177
           print("Perform " + str(nsteps) + " steps from t = 0.0 to t = " + str(tspan[0])
178
179
180
           # Magnetfeld Matrix B(t)
181
182
           B = lambda t: self._B(t)
183
           #how many components does Psi have
184
           N = self._psi.get_number_components()
185
186
           #start time t_0 = 0?
187
           t0 = 0
188
189
           t_a = t0
           t_b = t0
190
           #calculate R = U(t0 + N*h, t0)
192
           #Use N = n_steps to account for large time difference
193
194
           t_interval = array([t0, tspan[0]])
           R = FourierMagneticPropagator._Magnus_CF4(t_interval, B, nsteps)
195
196
           \# rotate the grid by the transpose of R
197
           self._grid.rotate(R.T)
198
199
           # Compute rotated initial data
200
201
           X = self._grid.get_nodes(flat=True)
           values = self._initalpacket.evaluate_at(X, prefactor=True)
202
203
           values = tuple([val.reshape(self._grid.get_number_nodes()) for val in values])
           self._psi.set_values(values)
204
205
           self._grid.rotate(R)
206
207
           #calculate the necessary timesteps
208
           for j in range(nsteps):
209
                for i in range(len(self._a)):
210
                    # Integral -\int_{tspan}[0]^{tspan}[1]B^2(s)ds und zugehörige
211
       Propagation
212
                    # (siehe Paper, Remark 3.1)
                    minus_B_squared = lambda t: (-1.0) * dot(B(t), B(t))
213
                    A = 1.0 / 8.0 * MagneticField.matrix_quad(minus_B_squared, t_a, t_a +
214
       self._a[i]*self._dt)
215
216
                    X = self._grid.get_nodes(flat=True)
                    VB = sum(X * dot(A, X))
217
                    VB = VB.reshape(self._grid.get_number_nodes())
218
                    prop = exp(-1.0j / self._eps**2 * VB) # ev. -0.5j durch -1j ersetzen
219
220
                    values = self._psi.get_values()
221
                    values = [prop * component for component in values]
222
223
                    self._potential.calculate_exponential(-1.0j * self._a[i]*self._dt /
224
       self._eps**2)
225
226
                    self._grid.rotate(R.T)
                    VE = self._potential.evaluate_exponential_at(self._grid)
227
```

```
self._VE = tuple([ve.reshape(self._grid.get_number_nodes()) for ve in
       VE])
229
230
                    #apply it
                    values = [self._VE * component for component in values]
231
                    self._grid.rotate(R)
232
                    t_interval[0] = t_b
234
                    t_interval[1] = t_b + self._b[i]*self._dt
235
236
                    U = (FourierMagneticPropagator._Magnus_CF4(t_interval, B, 1)).T
237
                    R = dot(R, U)
238
239
                    if(R.shape != U.shape):
                        raise ValueError("Shapes of R and U do not match")
240
241
242
                    #check for obsolete splitting steps
                    if(self._b[i] != 0):
243
                        values = [fftn(component) for component in values]
244
245
                        # Apply the kinetic operator
246
247
                        self._KO = KineticOperator(self._grid, self._eps)
                        self._KO.calculate_exponential(-0.5j * self._eps**2 * self._b[i]*
248
       self._dt)
249
                        TE = self._KO.evaluate_exponential_at()
250
                        values = [TE * component for component in values]
251
252
253
                        # Go back to real space
                        values = [ifftn(component) for component in values]
254
255
                    #Apply
256
                    self._psi.set_values(values)
257
258
                    #update t_a and t_b
259
                    t_a = t_a + self._a[i]*self._dt
                    t_b = t_b + self._b[i]*self._dt
261
262
           return tspan[0]
263
264
265
       def propagate(self, tspan):
266
           r"""This method does nothing.
267
268
```

### 3 Results

# 4 Summary and Conclusion

## References

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