QEngine: An open-source C++ Library for Quantum Optimal Control of Ultracold Atoms

J. J. W. Sørensen, J. H. M. Jensen, T. Heinzel, J. F. Sherson

Department of Physics and Astronomy, Aarhus University, Ny Munkegade 120, 8000 Aarhus C Denmark

Abstract

We present the first version of the QEngine, an open-source C++ library for simulating and controlling ultracold quantum systems using optimal control theory (OCT). The most notable systems presented here are Bose-Einstein condensates, many-body systems described by Bose-Hubbard type models, and two interacting particles. These systems can all be realized experimentally using ultracold atoms in various trapping geometries including optical lattices. In addition we provide a number of optimal control algorithms including the recently introduced GROUP method. The QEngine library has a strong focus on accessibility and performance. We provide several examples of how to prepare simulations of the physical systems and apply optimal control.

Keywords: Quantum Optimal Control Theory; Bose-Einstein Condensate; Gross-Pitaevskii equation; GROUP; Bose-Hubbard; C++.

1. Introduction

In the last two decades there have been exceptional advances in the ability to engineer and understand complex quantum systems. Especially, ultracold atoms provide an excellent platform for precision measurements [1, 2], matter wave interferometry [3, 4, 5], quantum simulation [6, 7], and quantum gates [8, 9, 10]. These systems offer extensive versatility through their purity and the high level of control of both the underlying potential landscape and the interatomic interactions [11]. In order to fully utilize the potential of these quantum systems the design of efficient experimental protocols for preparing quantum states of interest poses an important challenge [12].

Many experimental control protocols rely on simple empirical or adiabatic inspiration, which are typically slow and therefore limited by decoherence and sensitivity towards experimental imperfections [12, 13]. It is often desirable...
to find fast protocols that avoid decoherence and are robust with respect to system perturbations resulting in typically highly complex controls. Such control protocols can be found within the framework of Quantum Optimal Control (QOC). In QOC improved protocols are found using optimization algorithms that seek to minimize some cost functional [14].

In the context of ultracold atomic physics, QOC has been applied to improve splitting and driving of Bose-Einstein condensates trapped on an atom chip, which can be used to realize matter wave interferometry and nonlinear atom optics [12, 15, 16]. QOC has also been applied to stabilize ultracold molecules [17, 18] and manipulate ultracold many-body systems in optical lattices [19]. In addition, it has been demonstrated that such optimal control pulses are experimentally feasible [12, 20]. There has also been fundamental studies showing that QOC can find controls saturating the fundamental quantum speed limit [21, 22, 23] where similar behavior has also been reported for ultracold atoms in a double well system [24]. QOC is a versatile tool that can be applied not only in the context of ultracold atoms but as examples also in nuclear magnetic resonance [25], control of chemical reactions [26] and nitrogen vacancies [27].

Optimal control protocols are typically designed for a certain set of experimental parameter values that may change due to modifications in the experimental setup, thereby necessitating a recalculation for new optimal controls. For instance, there is a large number of papers that discuss driving a condensate from the ground state into the first excited state with slightly different parameter values [12, 15, 16, 20, 28, 29]. A barrier for rapidly recalculating such controls is writing and rigorously testing QOC programs, which is time consuming and the programs are very slow if not properly implemented. These two requirements, performance and usability, are primary driving forces behind the design of the QEngine. The high level of usability for example enables experimentalists to readily recalculate experimental protocols.

There exist a number of alternative software packages to the QEngine for performing quantum optimal control. Many of these are implemented in MATLAB like OCTBEC [30], DYNAMO [31] and the recent WavePacket [32, 33]. The Python package QuTiP is also a widespread platform for simulation and optimal control of quantum optics [34]. Collectively these packages offer more functionality than the QEngine but they are implemented in weakly typed programming languages that are inherently less focused on performance. Especially, OCTBEC has a similar focus to the QEngine and it has been a source of inspiration for our work.

The QEngine is designed for performance. One way the QEngine achieves this is by a general reliance on templates to provide flexibility instead of virtual functions and pointers. This allows for a high number of compile-time optimizations. Templates are useful for efficiency, but they are typically a programming barrier for physicists who are not C++ professionals. In order to accommodate such users of the library, we have made considerable efforts in providing a simplified API that does not invoke any advanced language features. The auto-syntax available in modern C++ together with factory-functions and overloaded operators give a straightforward syntax close to the mathematical equations used by theoretical physicists and weakly typed languages such as MATLAB and Python. The QEngine uses the highly optimized Intel Math Kernel Library (MKL) and the C++ library Armadillo to provide efficient basic linear algebra needed for the quantum simulations and optimizations [35]. In addition, the code has also been profiled and optimized.

A comprehensive documentation for the QEngine is also available at quatomic.com and the source code is available at gitlab.com/quatomic/qengine. The library has a number of example programs that can help users get started. In this paper, we give an introduction to some of the features in QEngine but leave out several details that can be found in the online documentation. The QEngine currently supports simulation and optimal control of the Gross- Pitaevskii description of a BEC, the Bose-Hubbard model, two interacting particles, a single particle, and generic few mode models.

The paper is organized as follows. In section 2 we give a brief introduction to the physical models and optimal control theory. In section 3 we discuss how to prepare simulations in two example programs that demonstrate key functionalities in the QEngine. Quantum optimal control theory is explained in section 4 including the Group algorithm we recently introduced in Ref. [36]. Finally in section 5 we explain how to perform optimal control optimizations on the example programs from section 3. Section 6 gives a summary and outlook.

2. Overview of QEngine Features

In this section we give a brief overview of the mathematical description of the models available in the QEngine and optimal control. The starting point for modelling these ultracold atomic systems is the second quantized Hamiltonian
3.1 The simulations and optimization were dependent on the ratio $U/J$. The Hamiltonian exhibits a phase transition from a delocalized superfluid state to a Mott insulating state depending on the coupling strength $U/J$. In Eq. (1) we have used that two-particle interactions in the ultracold limit are well-described by an effective contact interaction, which in one-dimension is $V_{\text{int}}(x_1, x_2) = g_{1D} \delta(x_1 - x_2)$ where $g_{1D}$ is the system dependent coupling strength [11, 38]. Different physical models described in the QEngine emerge from different special cases of Eq. (1).

**Gross-Pitaevskii Equation.** One important special case for the Bose-Einstein condensate (BEC) is in a single mode. This gives rise to the Gross-Pitaevskii Equation (GPE) describing the time evolution of a Bose-Einstein condensate (BEC) $\psi = \sqrt{N}\phi$. The corresponding Hamiltonian is

$$\hat{H}_{\text{GPE}} = \hat{H}_0 + g_{1D} |\psi(x,t)|^2,$$

where $\hat{H}_0$ is the kinetic and potential energy. The non-linear term represents the condensate self-interaction. The GPE is an important starting point for modelling the dynamics of BECs [3, 11, 12, 15].

**Bose-Hubbard.** Ultracold atoms can be loaded into a periodic optical lattice [11]. In this system it is convenient to expand $\hat{\Psi}(x)$ in terms of the localized Wannier modes on each lattice site. In a lowest band approximation the expansion reads $\hat{\Psi}(x) = \sum_i \hat{a}_i w_0(x - x_i)$. Also assuming the tight-binding approximation Eq. (1) for $L$ lattice sites becomes

$$\hat{H} = -J \sum_{i=1}^{L-1} (\hat{a}_{i+1}^\dagger \hat{a}_i + \text{h.c.}) + \frac{U}{2} \sum_{i=1}^{L} \hat{n}_i(\hat{n}_i - 1) + \sum_{i=1}^{L} V_i \hat{n}_i$$

where $J$ and $U$ are matrix elements of Eq. (1) with the lowest band Wannier functions, which describe the nearest-neighbor tunneling and on-site interaction. $V_i$ is the local external trapping potential. The ground state of this Hamiltonian exhibits a phase transition from a delocalized superfluid state to a Mott insulating state depending on the ratio $U/J$ [11]. This model is simulated in the QEngine using exact diagonalization with sparse linear algebra. The time evolution is performed using the split-step Fourier method [40].

**Two-particle.** Technological advances have enabled the preparation of single atoms in an optical lattice or tweezer arrays [11, 42, 43]. It has been proposed to use these systems as a platform for quantum computation, where the necessary two-qubit gate can be realized using controlled ultracold collisions of two atoms [8, 10, 44, 45]. It is convenient to rewrite Eq. (1) in first quantization as

$$\hat{H} = \frac{\hbar^2}{2m} \frac{\partial^2}{\partial x_1^2} + \hat{V}(x_1) - \frac{\hbar^2}{2m} \frac{\partial^2}{\partial x_2^2} + \hat{V}(x_2) + g_{1D} \delta(x_1 - x_2),$$

where $x_1$ and $x_2$ are the positions of the two atoms. The associated dynamics is also simulated using the split-step Fourier method. In a similar manner it is possible to simulate the dynamics of a single particle in the QEngine.

Figure 1: (Color Online) Snapshots of an optimized (blue) and unoptimized (red) transfer of a ground state BEC into the first excited state (green) in an anharmonic potential (black) – see section 3.1 for details. The simulations and optimization were performed using the QEngine.
Figure 2: The relative performance gain from MATLAB to C++ (higher is better) when running 100 optimization iterations in the Gross-Pitaevskii example program. The results are averaged over 15 individual runs. The inset shows the average absolute execution time for MATLAB and C++ (lower is better). The results were generated on a 2017 Macbook Pro laptop with an Intel(R) Core(TM) i7-7700HQ CPU @ 2.80GHz processor.

**Units.** In order to perform any physical simulation it is convenient to transform the Hamiltonians in Eqs. (2)-(4) into dimensionless units. A discussion of the units used in the example programs is given in the appendix.

**Quantum Optimal Control.** The QEngine enables the user to solve state transfer problems using QOC. This type of problem consists in manipulating the system dynamics in order to realize a transfer of an initial state $\psi_0$ into a target state $\psi_t$ for some fixed duration $T$. The manipulatory access to the dynamics is through one or more control fields $u(t)$ parametrizing the Hamiltonian in some way $H = H(u(t))$. In an experimental setting the control fields usually correspond to physical quantities such as the intensity or position of a laser beam. Optimization algorithms are typically used to iteratively design the control fields. An example of a state transfer problem is shown in Fig. 1 where a BEC is driven from the ground state into the first excited state using an optimized control field. In this case the control field corresponds to the position of the trap center, which is experimentally realized by adjusting magnetic fields [12, 20]. The figure shows snapshots of the transfer process before and after the optimization, illustrating that the optimization algorithm succeeds in finding an optimal control.

The QEngine offers a variety of optimization algorithms that can be applied to any of the physical models.

**Benchmark against MATLAB.** For comparison we benchmarked the QEngine against a similar implementation in MATLAB used internally in our research group. The benchmark was performed on the entire Gross-Pitaevskii example program described below, which ran for 100 optimization iterations with different grid sizes. The example program was slightly modified to give the most direct comparison between the two code bases. The results are displayed in Fig. 2, which shows that the QEngine is significantly faster.

3. Simulation Example Programs

Instructions for installing the QEngine are included in the README.md file. In order to help users get started we have included a number of example programs in the folder example_projects that are designed to illustrate many of the different features in QEngine. An overview of these programs is given in Table 1. In this paper, we discuss the two programs gpe-example.cpp and bosehubbard-example.cpp in depth, which gives a good overview of key functionalities in the QEngine. For clarity, we omit minor code details in this paper. A more detailed and complete API documentation can be found at quatomic.com, including more technical functionalities not discussed in the example programs.
### Table 1: The default example programs included in the QEngine. The table shows the file name, execution time and a short description. The results were generated on a 2017 Macbook Pro laptop with an Intel(R) Core™ i7-7700HQ CPU @ 2.80GHz processor.

| Example Program            | Execution Time | Description                                           |
|----------------------------|----------------|-------------------------------------------------------|
| gpe-example.cpp            | 582.27s        | Optimal control of driving a condensate wave function.|
| bosehubbard-example.cpp    | 234.94s        | Optimal control of a superfluid to Mott transfer.     |
| twoparticle-example.cpp    | 975.32s        | Optimal control of an ultracold atomic gate.          |
| oneparticle-example.cpp    | 0.99s          | Optimal control of a single atom in an optical tweezer.|
| twolevel-example-cpp       | 0.67s          | Optimal control of the Landau-Zener system.           |

#### 3.1. Gross-Pitaevskii Example Program

As a first example, we describe optimal control of a BEC trapped on an atom-chip. We focus on the control problem described in Refs. [12, 16, 46] where a BEC is transferred from the ground state into the first excited state as shown in Fig. 1. The physical motivation for this problem is to create a source of twin-atom beams, which is the matter-wave analogue to twin-photon beams [46]. The mechanism behind twin beam emission is binary collisions of two excited atoms. The collision may cause atoms to de-excite into the radial ground state mode while simultaneously populating twin momentum states $|±k_0\rangle$ along the axial $z$-direction due to conservation of momentum and energy. The characteristic timescale for the collision induced decay is a few milliseconds ($\approx 3\text{ms}$), and as a consequence the duration of the preparation stage into the excited state must be well below this threshold [12]. The atom-chip experiment has two tightly confined transverse directions ($x$ and $y$) and a weakly confined axial direction ($z$). One of the transverse directions (say $y$) has a tighter confinement freezing out excitations. The dynamics along the axial direction is slow compared to the transverse directions, and we may only consider a one-dimensional GPE along the $x$-direction. This requires an appropriate effective coupling constant [12, 47].

The potential along the $x$-direction is parameterized by a single control field $u(t)$, which is well-approximated by the anharmonic potential

$$V(x,u(t)) = p_2(x - u(t))^2 + p_4(x - u(t))^4 + p_6(x - u(t))^6,$$

where the $p_i$’s are constants obtained experimentally [12, 46]. The initial state is taken to be the ground state of $V(x,0)$, and in the optimal control part we take the target state to be the first excited state of $V(x,0)$ as shown in Fig. 1. Measuring length in units of micrometers and time in units of milliseconds, the effective mean field interaction strength for 700 atoms is $g_{1D} = 1.8299$ (see Appendix A).

To use the QEngine in a program, we need to include the QEngine header file

```cpp
#include <qengine/qengine.h>
#include <iostream>

using namespace qengine;
```

The `qengine.h` header exposes the different simulation models and optimal control algorithms. At the highest level the QEngine library defines the namespace `qengine`. The `qengine` namespace contains most API-functionality across the different types of physics and optimal control. We will make use of the `DataContainer` class defined in the QEngine, which can be used to save data to a .json file format or optionally to a MATLAB .mat file format. This makes it easy to export data for visualization and post-processing.

First we set up the control field $u(t)$

```cpp
const auto dt = 0.002;
const auto duration = 1.25; // corresponds to 1.25ms
const auto n_steps = floor(duration/dt) + 1;
const auto ts = makeTimeControl(n_steps, dt);
const auto initialAmplitude = 0.55;
const auto u = initialAmplitude*sin(PI/duration*ts); // control field
```

The `makeTimeControl` function returns a single control field with linearly spaced values, which can be used to compose more complicated control fields. In this case the control field $u$ is half a sine period with amplitude 0.55. This will also act as our initial guess in the optimal control algorithms. It is possible to access the control field values
at time index \( i \) by calling \( \text{u.get}(i) \), returning an \( \text{RVec} \) whose entries are the values for each control field at that time index. The control field values at the first and last time index can be easily accessed with \( \text{u.getFront()} \) and \( \text{u.getBack()} \), respectively. In the present case we only have a single control field.

The concept of a Hilbert space is mimicked in the QEngine for each type of physical model.

```cpp
const auto kinFactor = 0.36537; // T = -kinFactor*d^2/dx^2
const auto s = gpe::makeHilbertSpace(-2,+2,256,kinFactor);
const auto x = s.x(); // FunctionOfX of x=grid values
```

Having defined both \( x \) and \( u(t) \) we can create the control-dependent potential \( V(x,u(t)) \) Eq. (5)

```cpp
const auto p2 = 65.8392;
const auto p4 = 97.6349;
const auto p6 = -15.3850;

const auto V_func = [&x,p2,p4,p6]( const real u)
{  // By saving intermediate calculations we reduce overall computation time
    const auto x_u = x - u;
    const auto x_uPow2 = x_u * x_u;
    const auto x_uPow4 = x_uPow2 * x_uPow2;
    const auto x_uPow6 = x_uPow2 * x_uPow4;
    return p2*x_uPow2 + p4*x_uPow4 + p6*x_uPow6;
};
```

```cpp
const auto u_initial = u.getFront().front(); // first entry in first time index
const auto V = makePotentialFunction(V_func, u_initial);
```

The lambda function \( V_{\text{func}} \) takes a \textit{real} number and returns a \textit{FunctionOfX} evaluated with the given control value. To create a potential object the lambda function and an initial control field value are combined in \textit{makePotentialFunction}. The \( V \) object encapsulates the idea of a potential, and calling \( V(\text{newControlValue}) \) evaluates the \( V_{\text{func}} \) lambda with \textit{newControlValue} and returns a potential operator. In the present case \textit{newControlValue} is of type \textit{real}. The kinetic energy operator can simply be extracted from the Hilbert space. It is represented by the 5-diagonal approximation to the second derivative with non-periodic boundary conditions. The mean field interaction is equally succinctly handled. Assembling the Hamiltonian operator is then straightforward,

```cpp
const auto T = s.T();
const auto g1D = 1.8299;
const auto meanfield = makeGpeTerm(g1D);

const auto H = T + V + meanfield;
```

The \( H \) object can be called in the same way as the underlying potential by \( H(\text{newControlValue}) \). Note that the type of \( H \) is auto deduced to be a GPE Hamiltonian by the compiler. Omitting the \textit{meanfield} term would change the type deduction to a single particle Hamiltonian. This would still validate code since the GPE Hilbert space is the same as the single particle Hilbert space. The QEngine defines a convenient syntax for creating general linear combinations of eigenstates for operators. Let \( A \) be an operator and let \( \{ \phi_i \} \) be the corresponding eigenstates and suppose we wanted to create the linear combination \( \psi = \phi_0 - 2i\phi_1 \). This is readily achieved with the lines,

```cpp
const auto comb = A[0] - 2.0*i*A[1];  // syntax object
const auto psi = makeWavefunction(comb);  // evaluate syntax
```

The states we need for the example are individual eigenstates

```cpp
const auto psi_0 = makeWavefunction(H(u.getFront())[0]);
const auto psi_1 = makeWavefunction(H(u.getBack())[1]);
```

The QEngine currently only supports calculation of the ground state and first excited state for GPE type physics, but will not raise an error if higher excited states are queried. There are no such restrictions for the other types of physics.

Initializing the container and filling in time independent data is done by

```cpp
DataContainer dc; // empty data container
dc["dt"] = dt;
dc["duration"] = duration;
```
dc["x"] = x.vec();
dc["psi_t"] = psi_t.vec();

Once all data has been collected, calling dc.save("<path/to/dest>.<file-extension>") creates a file in either the .json or .mat file format, where e.g. the variable x is stored with the field name "x" and corresponding field values.

To perform time evolution we initialize a stepper with fixed stepping interval length \( dt \). We then loop over the entire control and append a few quantities of interest at each instant of time for saving.

```c++
auto stepper = makeFixedTimeStepper(H,psi_0,dt);
for(auto i = 0; i < n_steps; i++)
{
    const auto& psi = stepper.state();
    dc["V"].append(V(u.get(i)).vec());
    dc["psis"].append(psi.vec());
    dc["overlap"].append(overlap(psi, psi_t));
    dc["fidelity"].append(fidelity(psi, psi_t));
    if(i < n_steps -1) stepper.step(u.get(i+1));
}
```

The propagation from \( \psi(t) \) to \( \psi(t + dt) \) is performed by `stepper.step(u.get(i+1))` using the midpoint rule. Only \( u(t + dt) \) is needed since \( u(t) \) is stored internally from the previous step.

In the program we subsequently take additional steps with the final Hamiltonian held constant by using the `stepper.cstep()` function in an otherwise identical loop to the one above.

The result of the simulation is illustrated in Fig. 3 where the density of the condensate is plotted as a function of time. In the atom-chip experiment the objective is to transfer the initial state into the first excited state, which is not accomplished in the unoptimized transfer Fig. 3.

### 3.2. Bose-Hubbard Example Program

In this example, we discuss the simulation of bosons in an optical lattice described by the Bose-Hubbard model Eq. (3). Creating a Mott state with one particle on each site is important for many experimental applications such as quantum logic gate operations [48, 49, 50, 51, 52], quantum simulation [53], and single atom transistors [54]. Experimentally the system is initialized in the superfluid state and must be dynamically transferred into the Mott state [55, 56]. However, near the phase transition the gap between the ground state and the first excited state closes in an infinite system. This implies diverging transfer times for adiabatic solutions. There have been both experimental and numerical attempts to find improved transfer protocols [19, 55, 57, 58].

Here we consider a transfer from the ground state at \( U = 4 \) into a Mott like ground state at \( U = 30 \) with a weak harmonic external potential in units of \( J \). It is necessary to impose a minimal \( U_{\text{min}} = 2 \) since the Bose-Hubbard model assumes a sufficiently deep lattice [11]. In a similar manner, it is not experimentally feasible to have arbitrarily large values of \( U \). We take the upper bound to be \( U_{\text{max}} = 40 \). Later we will apply QOC to find optimized solutions, which must also satisfy these experimental and modeling constraints on \( U \). The constraints can be accommodated by introducing a nonlinear transformation \( U(u) = A(\tanh(u) + B) \), where \( u \) is a non-physical but unbounded control field. Here \( A = U_{\text{max}}/(1 + B) \) and \( B = (1 + U_{\text{min}}/U_{\text{max}})/(1 - U_{\text{min}}/U_{\text{max}}) \) restricts the physical control \( U_{\text{min}} < U < U_{\text{max}} \).

```c++
const auto Umin = 2.0;
const auto Umax = 40.0;

const auto B = (1+Umin/Umax)/(1-Umin/Umax); // transformation params
const auto A = Umax/(1+B);

const auto UFromu = [A,B](auto u){ return A*(tanh(u) + B);}; // control to physical U
const auto uFromU = [A,B](auto U){ return atanh(U/A-B );}; // physical U to control
```

```c++
const auto dt = 0.002;
const auto duration = 2.2;
const auto n_steps = floor(duration / dt) + 1;

const auto ts = makeTimeControl(n_steps, dt);
const auto u = uFromU(Umin + 0.5*exp(log((30-Umin)/0.5)*ts/duration));
```
Here we use an exponential ramp for $U$, which will later be used as the starting point for the QOC algorithms [19].

As in the previous example we begin by creating the underlying Hilbert space and subsequently initialize the terms in the Hamiltonian. We also demonstrate how to add a weak confinement potential.

```cpp
class auto space = bosehubbard::makeHilbertSpace(5,5);
class auto periodicBoundaries = false;
class auto hoppingOperator = space.makeHoppingOperator(periodicBoundaries);
class auto onSiteOperator = space.makeOnSiteOperator();
class auto sitePositions = linspace(-1.0, +1.0, space.nSites());
class auto potential = 0.1* pow(sitePositions,2);
class auto V = space.transformPotential(potential); // transform to site indices

const auto H_J = -1.0* hoppingOperator; // J = 1.0
const auto H_const = H_J + V; // Constant parts of Hamiltonian is added

// Transform to site indices

const auto H_func = [&](const real u) {
    return H_const + 0.5* UFromu(u)* onSiteOperator;
};

class auto H = makeOperatorFunction(H_func, u.getFront().front());
```

As in the previous example the full Hamiltonian $H$ is assembled using a lambda function $H_{func}$ and an initial control value $u.getFront().front()$. After initializing the Hamiltonian we set up the superfluid state and the Mott like state. We then initialize the time stepper. The default stepper is a Lanczos propagator, which uses a user supplied Krylov order [41].

```cpp

const auto psi_0 = makeState(H(u.getFront())[0]);
class auto psi_t = makeState(H(u.getBack())[0]);
class auto krylovOrder = 4;
class auto stepper = makeFixedTimeStepper(H, psi_0, krylovOrder, dt);
```

These lines of code complete the necessary steps to set up a Bose-Hubbard simulation. Exactly as in the previous example we propagate over the control and in this case save the single-particle density matrix by

```
dc["rho1"].append(space.singleParticleDensityMatrix(state));
```

`state` is the instantaneous state from the `stepper` when propagated over the control $u$. The result of the Bose-Hubbard simulation is illustrated in Fig. 4 where the on-site density is plotted as a function of time. In the superfluid-Mott transfer the objective is to reach the Mott insulator type state, which is not accomplished in the unoptimized transfer Fig. 4.

### 4. Theory of Quantum Optimal Control

In the previous section we have described how to set up simulations in the QEngine. In this section we briefly review QOC and related algorithms. In the next section we describe how to apply QOC to the example programs.

Consider the problem of engineering a single control field $u(t)$ realizing the state transfer from $\psi_0$ to $\psi_t$ in duration $T$ constrained by the equation of motion $i\hbar \dot{\psi} = \hat{H}(u)\psi$ for all $t$. We may consider $\hat{H} = \hat{H}_0$ to be the general case, as taking $\beta = 0$ produces the usual Schrödinger equation. In QOC this problem is posed as a minimization of the cost functional [36]

$$J[\psi, \chi, u] = J_F[\psi] + J_s[u] + J_{\text{sp}}[\psi, \chi, u] = \frac{1}{2} (1 - |\langle \psi|\psi(T) \rangle|^2) + \gamma \int_0^T u^2 dt + \Re \int_0^T \langle \chi | (i\hbar \partial_t - \hat{H}_0(u) - \beta |\psi|^2) |\psi \rangle dt,$$

where the time dependence of most quantities has been suppressed for readability. The first term is minimal for perfect transfers up to a global phase i.e. when the fidelity $F = |\langle \psi|\psi(T) \rangle|^2$ is 1. The second term penalizes rapid temporal fluctuations in the control field, which are typically not experimentally feasible. The relative importance between the first and second term is determined by a regularization hyperparameter $\gamma \geq 0$ where higher values shifts preference towards smoother controls. Usually $\gamma \sim 10^{-7} - 10^{-5}$. The last term containing the Lagrange multiplier $\chi(t)$ ensures the equation of motion is obeyed at all times.
Figure 3: (a) the density of the condensate $|\psi(x,t)|^2$ when propagated along the unoptimized control (b) from the Gross-Pitaevskii example program. The initial control gives $F = 0.23$. After the vertical dashed line the control is held constant.

Figure 4: (a) the on-site density $\langle n_i \rangle$ when propagated along the unoptimized control (b) from the Bose-Hubbard example program. The initial control gives $F = 0.81$. After the vertical dashed line the control is held constant.

4.1. grape

Setting the first Gateaux variations of $J$ wrt. the functions $\{\psi(t), \chi(t), u(t)\}$ to zero

\[ D_{\delta \psi}J = D_{\delta \chi}J = D_{\delta u}J = 0, \]

and assuming the variations of the control vanish at the boundaries ($t = 0$ and $t = T$) lead to the first order optimality conditions

\[ i\hbar \dot{\psi} = (\bar{H}_0(u) + \beta |\psi|^2) \psi, \quad \psi(0) = \psi_0, \]
\[ i\hbar \dot{\chi} = \left(\bar{H}_0(u) + 2\beta |\psi|^2\right) \chi + \beta \psi^2 \chi^*, \quad \chi(T) = i \langle \bar{\psi} \bar{\psi}(T) \rangle \psi_i, \]
\[ \gamma \ddot{u} = -\Re \left\{ \chi \left[ \frac{d\bar{H}_0(u)}{du} \right] \psi \right\}, \quad u(0) = u_0, \quad u(T) = u_T. \]

At this point we may think of $J$ as a functional of only $u$, $J = J[u]$, with the corresponding dynamics of $\psi$ and $\chi$ determined by the equations of motion above. An analytical approach to solving this set of equations is not generally feasible. However, we may define the gradient of $J$ wrt. the $X$ norm as the element $\nabla_X J$ fulfilling the relation

\[ D_{\delta u}J = \langle \nabla_X J, \delta u \rangle_X, \]

where $\delta u$ is an arbitrary variation. The common choices of the norm are $X = L^2$ and $X = H^1$ defined as $(f, g)_{L^2} = \int_0^T f(t)g(t)dt$ and $(f, g)_{H^1} = \int_0^T \tilde{f}(t)\tilde{g}(t)dt$, respectively [59]. Eq. (12) establishes an indirect way of calculating the gradient and for the norms above we obtain

\[ \nabla_{L^2} J(t) = -\Re \left\{ \chi(t) \left[ \frac{dH_0(u(t))}{du} \right] \psi(t) \right\} - \gamma \ddot{u}(t), \]
\[ \nabla_{H^1} J(t) = -\nabla_{L^2} J(t). \]

The $L^2$ gradient may not vanish at the boundaries so we must artificially enforce $\nabla_{L^2} J(0) = \nabla_{L^2} J(T) = 0$ to respect Eq. (11). In solving the Poisson Eq. (14) for the $H^1$ gradient we may conveniently choose Dirichlet boundary conditions $\nabla_{H^1} J(0) = \nabla_{H^1} J(T) = 0$ directly so Eq. (11) is always fulfilled.
The control is iterated towards a local minimum of the cost functional by the update rule

\[ u^{(k+1)} = u^{(k)} + \alpha^{(k)} p^{(k)}, \]  

where \( \alpha^{(k)} \) is a suitable step size along the search direction \( p^{(k)} \) for the \( k \)’th iteration. Typically the search direction is based on gradient information \[60]. The simplest choice is searching in the direction of steepest descent \( p^{(k)} = -\nabla_X J[u^{(k)}] \) where we again are free to choose either \( X = L^2 \) or \( X = H^1 \). Another common search direction is the Newton direction \( p_k^{(k)} = (\nabla^2_X J[u^{(k)}])^{-1} \nabla_X J[u^{(k)}] \), which takes into account the local curvature of the functional. This requires an expensive calculation of the Hessian \( \nabla^2_X J \), while also having no guarantee of invertibility far from the critical points of \( J \). In practice one uses methods like \( \text{l-bfgs} \) to build an approximation \( p^{(k)} \approx p_k^{(k)} \) to the search direction at iteration \( k \) based on the gradients calculated in iterations 1, 2, …, \( k \) \[59]. In passing we note that our numerical experiments suggest that restarting the \( \text{l-bfgs} \) algorithm family, the dimensionality \( M \) and performing gradient-based optimization in the \( M \)-dimensional space of real coefficients \( c = [c_1, c_2, \ldots, c_M]^T \), which is usually on the order of tens, \( M \ll N \) \[36]. This gives GROUP a much smaller optimization dimensionality than GRAPE and is also independent of the duration and size of the time steps. In Eq. (18) \( u_0(t) \) is a reference control and \( S(t) \) is any shape-function that goes to zero for \( t = 0 \) and \( t = T \), together enforcing appropriate boundary conditions Eq. (11). Since we are now optimizing the expansion coefficients \( c \), the first Gâteaux variations of \( J \) with \( \psi(t), \chi(t), c \) are set to zero

\[ D_{\chi} J = D_{\psi} J = D_{\delta c} J = 0. \]  

This produces the same equations of motion as Eqs. (9)-(10). Effectively now \( J = J[c] \). Choosing the inner product to be the usual vector dot product for \( X = \mathbb{R}^M \), the corresponding gradient of \( J \) with \( c \) is then defined as the element \( \nabla_{\mathbb{R}^M} J \) fulfilling the relation

\[ D_{\delta c} J = \langle \nabla_{\mathbb{R}^M} J, \delta c \rangle_{\mathbb{R}^M} = \nabla_{\mathbb{R}^M} J \cdot \delta c = \sum_{m=1}^{M} \frac{\partial J}{\partial c_m} \delta c_m, \]
Table 2: Simplest API to instantiate different control algorithms in the QEngine. The grape algorithms (left) only require a state transfer problem object, whereas the group algorithms (right) additionally require a basis specification. The basisMaker object creates a new basis in each superiteration according to prescription (23).

\[ \frac{\partial J}{\partial c_m} = \int_0^T \left( -\Re \left\{ \chi \frac{d\hat{H}_0(u)}{du} \right\} \psi - \gamma \ddot{u} \right) S(t)f_m(t)dt = \int_0^T \nabla_{L^2} J(t)f_m(t)dt, \]  

where we identified the term in parenthesis to simply be the $L^2$ grape gradient from Eq. (13). Calculating the group gradient amounts to first calculating the usual $L^2$ grape gradient and subsequently performing $M$ inexpensive one-dimensional integrals \[36\]. The coefficients are then iterated according to

\[ c^{(k+1)} = c^{(k)} + \alpha^{(k)} p^{(k)}, \]  

where $p^{(k)}$ is either the steepest descent direction or the l-bfgs direction both utilizing the gradient Eq. (21).

4.3. Dressed GROUP

A caveat of the parametrization Eq. (18) is that we may induce local trap minima not inherent to the control problem, but rather due to the parametrization itself. These types of traps are known as artificial traps \[61\]. A method to escape such traps was proposed in Ref. \[61\]. We may let $f_m = f_m(t; \theta_m)$ where $\theta_m$ is a set of values that is usually drawn at random. For example we may take $f_m(t, \theta_m) = \sin((m + \theta_m)\pi t/T)$ where $-0.5 \leq \theta_m \leq 0.5$ is drawn from a uniform distribution. Then, if the algorithm gets trapped at (possibly) an artificial minimum, we set $u_0(t) \leftarrow u(t)$, re-initialize the algorithm with coefficients $c = 0$, and draw a new set of basis functions $f_m$ defined by a new set of values $\theta_m$,

\[ u_0(t) \leftarrow u(t), \quad c \leftarrow 0, \quad f_m(t; \theta_m) \leftarrow f_m(t; \theta^*_m), \]  

where $\theta^*_m$ are new random values. This changes the topology of the optimization landscape and the artificial trap may have been eliminated. Effectively, this corresponds to restarting the group algorithm with a new parametrization basis. These restarts are known as superiterations \[61\]. This modification is referred to as dressed group, or dGROUP for short \[36\].

5. Optimal Control Example Programs

All algorithms described in the previous section are readily available in the QEngine. These algorithms can be instantiated by the simple API listed in Table 2. Additional options such as stopping criteria, data collection, and step size finding method can be set independently of the physical model. If these are not specified, default options are used. As a result, the code in the following sections is valid for all example programs.

We now extend the example programs in section 3 by performing optimal control on the systems. Initially we will use the simplest API to perform QOC on the Bose-Hubbard example program. Afterwards we will show a more advanced API for the GPE example program.
5.1. Control of Bose-Hubbard Program – Simple API

Here we continue the example program from section 3.2. In this example we prepare a state transfer problem and solve it using GRAPE, GROUP, and dGROUP. A state transfer problem is encapsulated by a problem object,

```
auto problem = makeStateTransferProblem(H, psi_0, psi_t, u, krylovOrder);
```

Here H is the Hamiltonian, psi_0 is the initial state, psi_t is the target state, and u is the initial control field, which in this case is an exponential ramp. The krylovOrder is the order used internally in the timestep. This problem object maintains a control field and is responsible for calculating the corresponding cost functional and gradient. These quantities are used internally by the optimization algorithm to update the control according to Eq. (15) or (22). Having set up the problem, we can apply the different algorithms using the simple API listed in Table 2.

```
// GRAPE
auto GRAPE = makeGrape_bfgs_L2(problem);
GRAPE.optimize();
const auto u_grape = GRAPE.problem().control();

// GROUP
const auto M = 60; // basis size
const auto shapeFunction = makeSigmoidShapeFunction(ts, 0.999);
const auto maxRand = 0.0; // -maxRand < theta_m < maxRand
const auto basis = shapeFunction* makeSineBasis(M, u.metaData(), maxRand);
auto GROUP = makeGroup_bfgs(problem, basis);
GROUP.optimize(); // begin optimization
const auto u_group = GROUP.problem().control(); // extract GROUP optimized control

// dGROUP
auto basisMaker = makeBasisMaker([M, maxRand, &u, &shapeFunction](){
    return shapeFunction*makeSineBasis(M, u.metaData(), maxRand);
});
auto dGROUP = makeDGroup_bfgs(problem, basisMaker);
dGROUP.optimize(); // begin optimization
const auto u_dgroup = dGROUP.problem().control(); // extract dGROUP optimized control
```

After construction, calling .optimize() begins the optimization algorithm. Once the optimization is completed, the optimized control fields are extracted by .problem().control(). GRAPE optimizes the control field directly whereas GROUP uses a reduced basis, which must be supplied by the user – see Table 2. GROUP also uses a shape function to enforce boundary conditions on the control field Eq. (18). The default shape function makeShapeFunction is a symmetric sigmoid function depending on parameters like dt and the number of controls, which are conveniently supplied through u.metaData(). GROUP uses the same basis for the entire optimization whereas dGROUP uses a new basis in each superiteration through the prescription (23). The new basis is constructed from the basisMaker object.

The result of the dGROUP optimization is shown in Fig. 6a where the on-site density is plotted as a function of time for the optimized control. After the vertical dashed line the density and the control is constant since the target state is an eigenstate. In Fig. 6 the controls from the other optimization algorithms are plotted. In this example the algorithms find very similar solutions.

5.2. Control of Gross-Pitaevskii Program – Advanced API

In this section we continue the example program from section 3.1. Here we perform GRAPE, GROUP, and dGROUP optimizations using the more advanced API.

Calculating the gradient Eq. (13) requires the derivative of the Hamiltonian wrt. the control. The default behavior in the simple API is to calculate the derivative numerically using finite differences. It is more efficient and accurate to manually supply the analytic derivative

```
const auto dHdu_func = [&x,p2,p4,p6](const real u) {
    auto x_u = x-u;
    auto x_u_Pow2 = x_u*x_u;
    auto x_u_Pow3 = x_u*x_u_Pow2;
}
```
Figure 5: (a) the density of the condensate $|\psi(x,t)|^2$ when propagated along the optimized dGROUP control from the Gross-Pitaevskii example program. (b) the initial control $u_0$ and the optimized controls from GRAPE $F = 0.992$, GROUP $F = 0.992$ and dGROUP $F = 0.994$ – see legend.

Figure 6: (a) the on-site density $\langle \hat{n} \rangle$ when propagated along the optimized dGROUP control from the Bose-Hubbard example program. (b) the initial control $u_0$ and the optimized controls from GRAPE, GROUP and dGROUP – see legend. All algorithms have fidelity $F = 0.99$ and find almost identical controls.

```cpp
class dHdu: public dHduBase2
{
public:
    dHdu(const PotentialFunction &potential) : super(potential) {}

    //...Implementation...

    const auto dHdu = makeAnalyticDiffPotential(makePotentialFunction(dHdu_func, u_initial));

    auto problem = makeStateTransferProblem(H, dHdu, psi_0, psi_t, u) + 1e-5*Regularization(u) + 2e3*Boundaries(u, RVec{-1}, RVec{+1});

    Note the resemblance to the cost functional Eq. (7) when initializing the state transfer problem. The last term adds soft bounds to the optimization problem as in Eq. (16) that penalizes control values outside $|u(t)| \geq 1 \mu$m, which is set by experimental constraints on the control problem [12].

    The QEngine allows for arbitrary stopping conditions within its optimization algorithms. After each iteration the optimizer calls a stopper object to check if the optimization should be stopped. This object is instantiated by the makeStopper function. Any callable taking the optimizer type as its argument and returning a boolean can be passed in to this function. In this example we pass an inline lambda into makeStopper using auto& to deduce the optimizer type.

    // Stopper object
    const auto stopper = makeStopper([](auto & optimizer) -> bool
    {
        bool stop = false;
        if (optimizer.problem().fidelity() > 0.999)
        { std::cout << "Fidelity criterion satisfied" << std::endl; stop = true; }
        if (optimizer.previousStepSize() < 1e-7)
        { std::cout << "Step size too small" << std::endl; stop = true; }
        if (optimizer.iteration() == 2000)
        { std::cout << "Max iterations exceeded" << std::endl; stop = true; }
        if (stop)
        { std::cout << "STOPPING" << std::endl; }
        return stop;
    });

    auto x_u_Pow5 = x_u_Pow2 * x_u_Pow3;
    return -(2*p2*x_u + 4*p4*x_u_Pow3 + 6*p6*x_u_Pow5);
};
```
After each iteration the optimizer also calls a collector object and defines what should be displayed, saved, and so on in each iteration. The collector works similarly to the stopper and is instantiated by the makeCollector function. In this example the fidelity of the current control is saved to the DataContainer, and a status message is printed to console.

```cpp
// Collector object
const auto collector = makeCollector([&dc, n_steps] (auto & optimizer) {
  dc["fidelityHistory"].append(optimizer.problem().fidelity());
  std::cout <<
  "ITER " << optimizer.iteration() << " | " <<
  "fidelity : " << optimizer.problem().fidelity() << "\t " <<
  "stepsize : " << optimizer.stepSize() << "\t " <<
  "fpp : " << round(optimizer.problem().nPropagationSteps()/n_steps) << "\t " <<
  std::endl;
});
```

Next we define a line search algorithm that calculates a suitable step size $\alpha^k$ along the search direction $p^k$ in Eqs. (15) and (22). The QEngine supplies an interpolating line search algorithm that can be used out of the box by calling the makeInterpolatingStepSizeFinder function. It is also possible to create custom line search algorithms. As a simple example, a constant step size strategy is commented out to show the interface for custom algorithms.

```cpp
// Stepsize finder object
const auto maxStepSize = 5.0;
const auto maxInitGuess = 1.0;
const auto stepSizeFinder = makeInterpolatingStepSizeFinder (maxStepSize, maxInitGuess);
```

Restarting the L-BFGS algorithm is sometimes beneficial as noted in section 4. In this example we simply use the default option. The optimizer of choice is then created by calling the corresponding make function with the additional objects defined above.

```cpp
/// GRAPE
auto GRAPE = makeGrape_bfgs_L2 (problem, stopper, collector, stepSizeFinder);
```

To perform the optimization and retrieve the optimized control we invoke .optimize() on each optimizer. The optimization runs until the stopper function returns true. For the GROUP optimizations we use a sine basis Eq. (18) with no randomization. On construction the reference control and coefficients in Eq. (18) are set to $u$ and $c = [0, 0, 0, ... , 0]^T$. We can manually update them to e.g. $c = [0.55, 0, 0, ... , 0]^T$ where 0.55 is the initial amplitude as illustrated.

```cpp
auto u_grape = GRAPE.problem().control(); // extract GRAPE optimized control
auto GROUP = makeGroup_bfgs (problem, basis, stopper, collector, stepSizeFinder);
```

```cpp
collector (GROUP);
```
For dGROUP we first prepare a shape-function and the BasisMaker as in the Bose Hubbard example program. In dGROUP there is also the possibility to supply a user-defined dressedRestarter object that in each iteration checks if the algorithm should re-initialize with a new random basis as in Eq. (23), which requires a non-zero bound `maxRand` on the random values \( \theta_m \). As a simple example we re-initialize the algorithm every 100 iterations or if there is only a small decrease in the cost

```cpp
// dGROUP
maxRand = 0.1;
const auto basisMaker = makeRandSineBasisMaker(basisSize, shapeFunction, maxRand);

auto restart_func = [ tol { 1e-6 }](const auto & dGROUP) mutable
{
    auto stepSize = dGROUP.stepSize();
    if (stepSize < tol)
    {
        std::cout << "New superiteration in dGROUP algorithm." << std::endl;
        return true;
    }
    return false;
};
const auto dRestarter = makeDressedRestarter(restart_func);

problem.update(0* u);
dGROUP = makeDGroup_bfgs(problem, basisMaker, stopper, collector, stepSizeFinder, dRestarter);
dGROUP.problem().update(cs); // set initial coefficient vector \( \vec{c} = [0.55, 0, 0, \ldots, 0] \)

collector(dGROUP); // collect iteration 0
dGROUP.optimize(); // begin optimization

const auto u_dgroup = dGROUP.problem().control(); // extract dGROUP optimized control
```

Finally the data from all the optimizations is saved into a json file by calling `dc.save("gpe-example.json")` or alternatively a mat file with `dc.save("gpe-example.mat")` if matio has been configured.

The result of the dGROUP optimization is shown in Fig. 5a where the condensate density is plotted as a function of time. The control and the density is constant after the vertical line since the target state is an eigenstate. The final controls from all the optimization algorithms are displayed in Fig. 5b.

6. Summary and Outlook

We have introduced and described how to use the QEngine. In the example programs we showed how the auto-syntax combined with factory-functions allows the user to straightforwardly set up optimal control simulations of ultracold atomic systems. However, the QEngine is limited in the number of physical models it supports e.g. we only simulate the dynamics of one-dimensional systems in the current version. We plan to release future versions of the QEngine that can simulate a wider range of dynamics. We also plan to include more sophisticated optimal control for instance the quality of the gradients can be improved by also taking the temporal discretization into account.

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Appendix A. Units and Nondimensionalization

The SI-unit system usually results in very small numerical values for quantum mechanical simulations, which makes simulations impractical or infeasible. For this reason it is beneficial to rescale physical quantities into characteristic scales of the problem such that most values are of order unity. This is achieved using a process known as nondimensionalization where quantities in SI-units are written in product form e.g. \( a \) becomes \( a = \alpha \tilde{a} \) where \( \alpha \) carries both the dimension of \( a \) and a magnitude while \( \tilde{a} \) is a non-dimensional scaling value. This is done for all quantities and substituted into the equations of motion, which leaves a new set of working equations involving only
the dimensionless scaling values. As an example consider the GPE example program discussed in section 3.1 where the relevant quantities are

\[
x = \chi \tilde{x}, \quad t = \tau \tilde{t}, \quad V = \epsilon \tilde{V}, \quad \psi = \xi \tilde{\psi}, \quad g_{1D} = \gamma \tilde{g}_{1D}
\]

We may a priori take length to be measured in micrometer and time to be measured in milliseconds.

\[
\chi = 1 \mu m, \quad \tau = 1 ms
\]

The three remaining units are chosen conveniently as

\[
\epsilon = \frac{\hbar^2}{2m\chi^2}, \quad \xi = \sqrt{\frac{\hbar}{\chi}}, \quad \gamma = \frac{\epsilon}{\xi^2},
\]

where \(\kappa\) is the kinetic factor. Substituting into the GPE

\[
i \frac{\hbar}{\tau} \xi \frac{\partial \tilde{\psi}(\tilde{x}, \tilde{t})}{\partial \tilde{t}} = \left( -\kappa \left( \frac{\hbar^2}{\epsilon \chi^2} \right) \frac{\partial^2 \tilde{\psi}(\tilde{x})}{\partial \tilde{x}^2} + \epsilon \tilde{V}(\tilde{x}) + \gamma \xi^2 \tilde{g}_{1D} |\tilde{\psi}(\tilde{x}, \tilde{t})|^2 \right) \xi \tilde{\psi}(\tilde{x}, \tilde{t})
\]

Dividing by \(\epsilon\) and requiring \(\hbar/\tau \epsilon = 1\) or equivalently \(\kappa = \tau \hbar/2m\chi^2\) gives

\[
i \frac{\partial \tilde{\psi}(\tilde{x}, \tilde{t})}{\partial \tilde{t}} = \left( -\kappa \frac{\partial^2 \tilde{\psi}(\tilde{x})}{\partial \tilde{x}^2} + \tilde{V}(\tilde{x}) + \tilde{g}_{1D} |\tilde{\psi}(\tilde{x}, \tilde{t})|^2 \right) \tilde{\psi}(\tilde{x}, \tilde{t}),
\]

which is the dimensionless form the GPE solved in the example programs. For the potential we find \(\tilde{p}_i\) by comparing

\[
\tilde{V} = \tilde{V} = \sum_{i=2,4,6} \tilde{p}_i (x - u)^i = \sum_{i=2,4,6} \left( \frac{p_i x^i}{\epsilon} \right) (\tilde{x} - \tilde{u})^i = \sum_{i=2,4,6} \tilde{p}_i (\tilde{x} - \tilde{u})^i
\]

In these units the nondimensionalized scaling values used in the simulation for \(N = 700\) are

\[
\kappa = 0.36537, \quad \tilde{p}_2 = 65.8392, \quad \tilde{p}_4 = 97.6349, \quad \tilde{p}_6 = -15.3850, \quad \tilde{g}_{1D} = 1.8299.
\]

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