

Quantum control report - Third edition

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I. INTRODUCTION

In the Shake Up challenge, one works with a state confined in a trapping potential. The state describes either a single particle or a Bose Einstein Condensate (BEC) state of matter and is in the ground state of the trapping potential. The goal is to displace the potential using a *control function* in order to excite the state to the first excited state. The problem is simulated using 2 different tools for simulating quantum mechanics. The first one is *Quantum Composer*, a visual program where a system consists of interconnected nodes with different roles e.g. potential, time, space. The second tool is the QEngine, a C++ library of high performance methods for simulating quantum mechanics. It is a goal of its own to use both tools to explore this challenge, both to test the limits of what is possible to do with the intuitive, but functionally more limited Quantum Composer, but also to find pros and cons of each tool and describe when to use what tool. Furthermore, based on those considerations it is also an objective to formulate a list of suggestions which could make Quantum Composer a more attractive tool to use as an expert.

The goal of exploring this challenge was to explore the different dynamics that arise when simulating a BEC compared to a single particle and how the dynamics depend on the nonlinearity of the BEC Hamiltonian, including how this affects the so-called Quantum Speed Limit. Adding to this is exploring if and how control functions can be clustered, how optimal solutions look compared to similar ones and how robust they are to system changes.

This report is structured as follows: In section II some background information regarding the challenge is provided. Section IV contains the results of the report and is split into 3 parts. Section ?? shows the results of exploring this challenge by simulating BECs in a number of different systems, section V explains the strategies used to reach those results and IV A discuss some of the limitations to exploring the problem using Quantum Composer that were found. The results are then discussed in section VIII.

In section IX it is concluded that the nonlinear term for the BEC drastically changes the behavior of the state compared to the single particle state for values of β . This

new behavior could, in some cases, improve the reachable fidelity compared to a single particle by up to 50%. It was also found that the value of β affects the QSL of BECs. This section also sums up the conclusions about the different approaches to creating a control function that were used in this project. Finally, a number of things for future study of the subject are suggested. The tools are compared in VII, where the strengths and weaknesses of each tool is summed up and suggestions for when to use either tool are listed.

II. BACKGROUND

The mean field of a BEC of mass m , momentum p and interatomic interaction strength β (also called g_{1D}) can be described using the Gross-Pitaevskii equation (GPE)

$$i\hbar \frac{d}{dt} \psi(x, t) = \left(\frac{\hat{p}^2}{2m} + \hat{V}(x, u(t)) + \beta |\psi(x, t)|^2 \right) \psi. \quad (1)$$

Notice how β scales the nonlinear interaction term and that for the case $\beta = 0$, we recover the time dependent Schrödinger equation for a single particle. This nonlinear term represents interaction that takes place between all of the atoms in a BEC. The factor β is both dependent on the s-wave scattering length a_{1D}^s as well as the number of atoms contained in the BEC N [1]. It is also possible to tune this interaction parameter using a Feshbach resonance [2].

Depending on the sign of β in eq. (1) the interatomic interaction will be either attractive ($\beta < 0$) or repulsive ($\beta > 0$) and because of this, we would expect the state to respectively shrink or grow in size as a result of this interaction. This is also what can be seen on Fig. 1, where the ground state of a BEC is plotted as a function of β .

We attempt to exciting the BEC from its ground state to first excited state using a control function, usually denoted $u(t)$. The control function describes how the potential evolves in time. A simple example of this is could be that $u(t)$ could describe how a quartic potential is displaced over time:

$$V(x, u(t)) = a(x - u(t))^2 + b(x - u(t))^4 \quad (2)$$

While it is not hard to write out *some* control function, it is much more difficult to find a control that successfully excites the initial state into the desired state. To quantify how well a given control function has performed, one can calculate the *fidelity* $F = |\langle \phi | \psi \rangle|^2$ to calculate the

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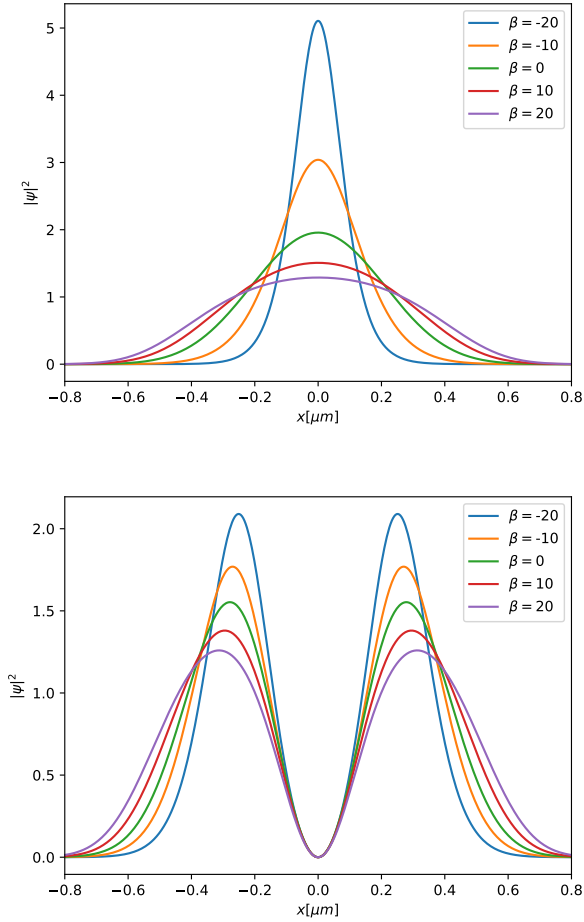


FIG. 1: **Top:** Groundstate of a BEC in a quartic trapping potential, shown as a function of the atomic interaction parameter β in 1. **Bottom:** First excited state of the same system.

overlap between the two states $|\psi\rangle$ and $|\phi\rangle$, where $|\psi\rangle$ is the current state and $|\phi\rangle$ is some desired state. Thus, the fidelity F is a number between 0 and 1, where 0 means that there is no overlap between the 2 states and 1 means that they are the same state. Usually values of 0.99 and above are set as the value that a given control function must satisfy. It is sometimes convenient to instead calculate the *infidelity* $1 - F$ to map out how well different control functions performed.

Since it can be very difficult to find a control function that is capable of reaching $F \approx 0.99$ and above, it is often useful to *optimize* an initial control function (a "seed") using different kinds of optimization algorithms. These optimization algorithms are, given enough time, usually capable of optimizing a seed into a control that can reach much higher fidelities. Regardless of which optimization algorithm is used, one must first define the optimization *problem*, which is a function that the given algorithm will

attempt to minimize. We define this problem as [3]

$$\min J(u) = \frac{1}{2}(1 - |\langle \psi_t | \psi(T) \rangle|^2) + \frac{\gamma}{2} \int_0^T \dot{u}^2 dt \quad (3)$$

where ψ_t is the target state, $\psi(T)$ is the final state evolved with the control $u(t)$ to final time T . The first term minimises infidelity and the second term penalises rapid fluctuations of the control function that can be challenging to implement experimentally. γ is called the regularization factor and is usually on the order of 10^{-4} to 10^{-6} .

The optimization algorithm GRAPE (**G**radient **A**scent **P**ulse **E**ngineering) is available to use in both Quantum Composer and the QEngine. GRAPE works by calculating the gradient of F with respect to each point of a given control function on the discretized time scale

$$\vec{\nabla}_{\vec{u}} F(\vec{u}) \quad (4)$$

where $\vec{u} = (u(t_0), u(t_1), \dots, u(T))$. When this calculation is done, GRAPE takes a step of size α_k such that the new discretized control function becomes

$$\vec{u}_{k+1} = \vec{u}_k + \alpha_k \vec{\nabla}_{\vec{u}} F(\vec{u}_k) \quad (5)$$

such that $F(\vec{u}_{k+1}) \geq F(\vec{u}_k)$. This process is continued until some convergence criterium is fulfilled, e.g. a target threshold $F(\vec{u}_k) \geq F_{target}$, a maximum number of iterations $k \leq N$ or until the step size α_k becomes too small.

Two other optimization algorithms are available to use in the QEngine: GROUP and dGROUP (dressed GROUP). Where GRAPE is a relatively simply gradient ascent algorithm, these two attempt to combine the local optimization that GRAPE does with a global search for promising regions in the fidelity landscape [4, 5]. Note, that this does not mean that an optimized seed will always be able to reach $F \approx 0.99$. One particular hindrance is the *Quantum Speed Limit* (QSL). This speed limit refers to the upper bound of how fast a state is able to evolve [6]. This speed limit is (not surprisingly) dependent on the individual system examined and in the context of the Shake Up challenge, it refers to the minimum time required to evolve from the initial state into the final state with a fidelity of at least $F \approx 0.99$.

The challenge was explored in Quantum Composer by creating different systems and trying different control functions and optimization methods. The different systems had their potentials varied and the interatomic interaction parameter β as well. The control functions themselves were varied and explored, along with the control duration T and the GRAPE optimization parameters F_{target} , N and γ .

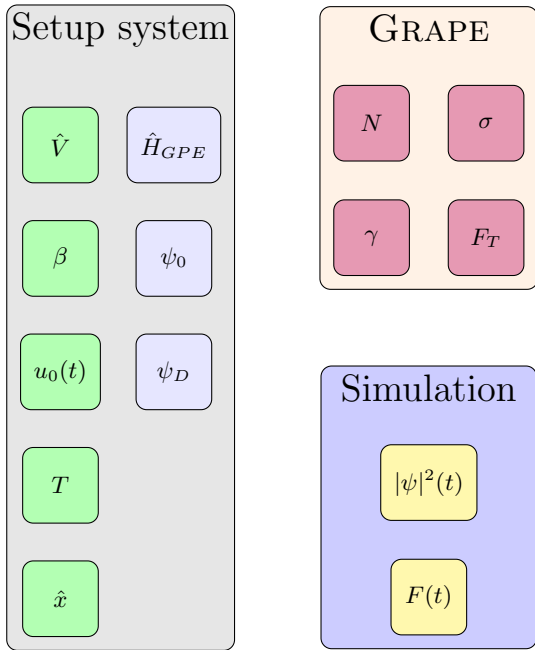


FIG. 2: Work flow in Quantum Composer

III. QUANTUM COMPOSER

Quantum Composer is a tool for visualizing the quantum mechanics of a system, either statically or as the system evolves in time. To do this, one combines building blocks (nodes) containing the various components (e.g. a potential \hat{V}) to build new nodes (e.g. a wave function Ψ). Some of these nodes can be visualized on graphs to show f.x. a wave function norm squared and can be evolved over time using a time node. The use of Composer in this report can be grouped into three overall groups: System setup, GRAPE optimization and simulation. These groups and their components have been visualized on Fig. 2.

To simulate a quantum control problem, one first has to set up a system that describes the problem. This includes choosing boundaries of the Hilbert space and discretizing the space into a number of points. It also includes things like describing the potential of the system, the time interval (t_0, T) . Specifically for quantum control problems it also involves defining how the potential depends on one or multiple control functions and how these initially look. From these steps, it is then possible to build the Hamiltonian of the system and from that we can create wave functions, in particular ψ_0 and ψ_D that are relevant for a quantum control problem.

It is then possible to optimize the initial control function using GRAPE. The optimization procedure requires a number of arguments, some optional (γ and σ), others required (N and F_T). The control function will then be updated until a convergence criteria is reached, after which GRAPE outputs the optimized control function. In

Composer it is possible to map the fidelity of the current and previous control iterations to show how fidelity improves as GRAPE iterates. Likewise, it is also possible to plot the initial and current control function to see how they differ. A screenshot from Composer that shows how this optimization procedure is set up can be seen on Fig. 3.

When the system is simulated, it is possible to analyze the wave function at the current point in time. Thus we are able to calculate and visualize different variables in our system as they are being simulated. This includes calculating the fidelity F as a function of time, but we can also plot how our wave function evolves in time by mapping out $|\psi|^2(x, t)$. When the simulation is finished, the final fidelity $F(T)$ can be recorded.

The data presented in Composer can in most cases not be saved directly through the program, but must instead be written down in e.g. a textfile separately. This file containing data can then be further analyzed and plotted. In this case it is done using Python and its plotting module `Matplotlib.pyplot`.

IV. RESULTS

A. Numerical limitations

1. Spatial resolution & performance

One obvious source of numerical inaccuracies is the resolution of the quantized Hilbert space one uses. Throughout the report, the spatial grid was divided into 256 points. We can compare this resolution to other resolutions by calculating the difference between a high resolution wave function with 1024 points and lower resolution wave functions. This difference between the different resolutions for the ground state and first excited state can be seen on the Fig. 4. As can be seen on the figure we find that the lower the resolution of a wave function is, the bigger the difference between that wave function and a higher resolution one.

Considering this, one would think that it would always be better to use as high spatial resolution of a given system as possible. This would of course be true if we had access to unlimited computational power. We instead have to balance the desire for a high spatial resolution and more accurate wave functions against the increased time it takes to do calculations for a higher dimension Hilbert space. As an example of this increased calculation time, a system was set up with the following parameters: $\beta = 4$, $V(x, u(t)) = a(x - u(t))^2 + b(x - u(t))^4$ with $a = 50$, $b = 128$, $u(t) = 0.5 \cos(2\pi t/T)$ and $T = 0.8\tau \approx 1.09$ ms. This system was then optimized for 100 iterations using GRAPE before the system was simulated with the optimized control function. The results can be seen on Fig. 5. From this result we can reassure ourselves that choosing a grid resolution of 256 points gives us a high accuracy

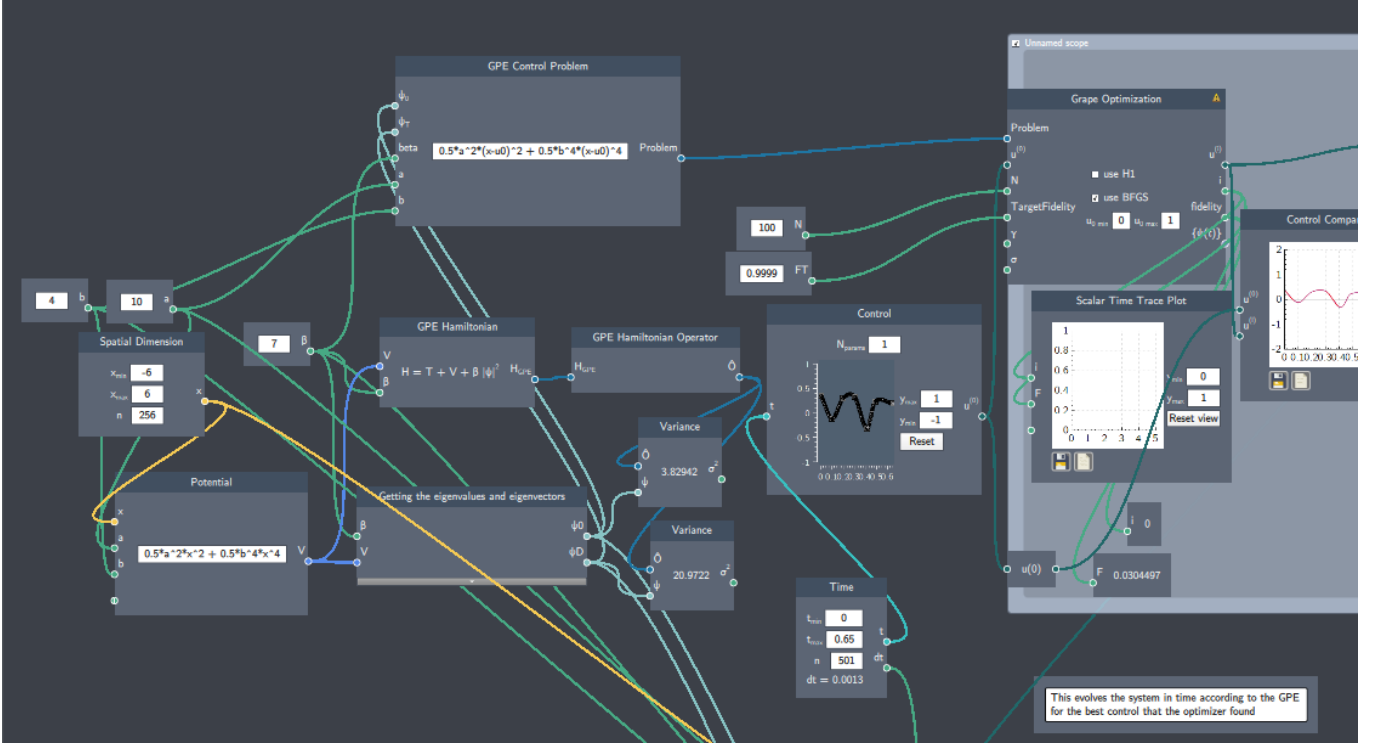


FIG. 3: Screenshot from Quantum Composer partially showing a flowfile that optimizes a control function using GRAPE. The optimized solution is then simulated in a scope not shown.

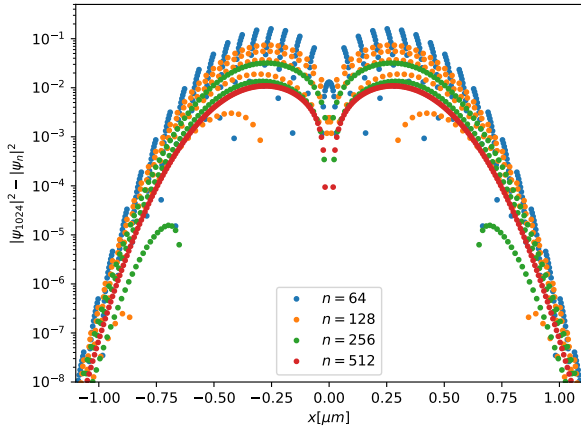


FIG. 4: Difference between different resolutions of the initial wave function compared to a high resolution wave function.

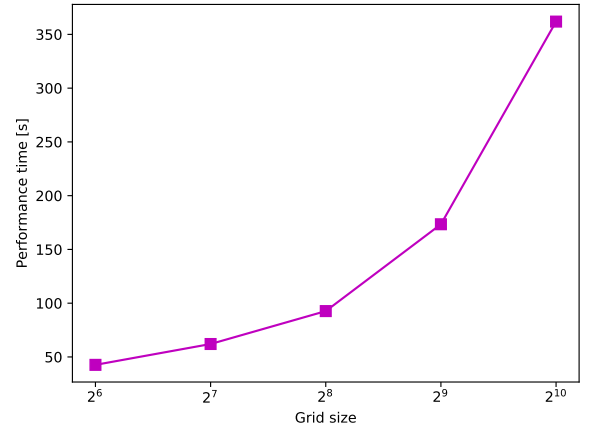


FIG. 5: The time it takes for Quantum Composer to solve the same problem as a function of the spatial grid resolution.

2. Eigenvectors & eigenvalues

compared to lower resolutions while also keeping performance time lower than for higher resolutions, where the performance time increases more rapidly.

In order to reach accurate results when making these computations, one has to ensure that the states that are analyzed are described within acceptable numerical accuracy. We can ensure that we are working with the ground state as our initial state in Composer by calculating the

variance of our initial state wrt. the Hamiltonian

$$\text{Var}(\psi)_H = \langle \psi | \hat{H}^2 | \psi \rangle - (\langle \psi | \hat{H} | \psi \rangle)^2 \quad (6)$$

This gives us a numerical estimate of how "close" our initial state is to the actual ground state of \hat{H}_{GPE} since

$$\psi \text{ is an eigenstate of } \hat{H}_{GPE} \iff \text{Var}(\psi)_H = 0 \quad (7)$$

The way the energy spectrum of \hat{H}_{GPE} is calculated is not trivial [5], but the variance can generally be reduced by adjusting different parameters of the system or Composer file, e.g. spatial dimension x or initial conditions for the H_{GPE} spectrum node. Despite these possible fixes, the numerical limitations of Quantum Composer become apparent as the "Shake Up" problem is explored in increasingly greater details and limits and accuracy are pushed further and further. The following paragraphs describe areas where the numerical limitations are significantly limiting the exploration of certain aspects of this problem.

As the work with the QSL shown in figure 11 was explored, it was desirable to examine how the energy levels of the state changed as a function of β . The idea was that β could change the spacing between the different levels of the potential well and perhaps make the spacing more even for some high or low values of β . As was shown in figure 10, the even spacing of a harmonic quadratic potential makes it difficult to reach the high fidelities desired. To investigate this, the energy difference between the first excited state and the ground state as well as the difference between the second and first excited state was calculated in Composer.

Despite not being able to calculate the second excited state of a BEC, it was also desired to explore how the ground and first excited state energies changed as a function of β . These energies are plotted on figure 6 and as can be seen, the uncertainty regarding the first excited state increases for values of β below -10 . As mentioned when estimating the optimizable limit of negative β 's, this could explain why $\beta = -10$ is the optimizable limit found.

Even when working within the low-variance range of this problem, one still has to be careful of numerical errors caused by the control function used. These problems with the control function seem to be caused by a combination of different factors: 1. Using 'high' frequency control functions. 2. Initial control functions having large displacements. 3. Larger control durations. As an example, one can see how the $k = 4$ seed from figure 13 performed for a duration of $T \approx 0.85\text{ms}$ compared to a $k = 5$ with longer duration $T \approx 1.0\text{ms}$ with a smaller amplitude. The comparison can be seen in figure 7 where the resulting state evolution of each of these optimized seeds are shown. The consequence of this error is that the control space that is possible to explore in Composer

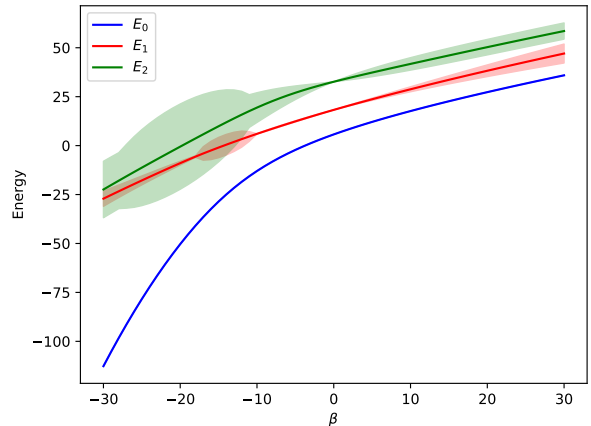


FIG. 6: Calculated energies of the first 3 energy eigenstates of a BEC in a quartic potential as a function of β .

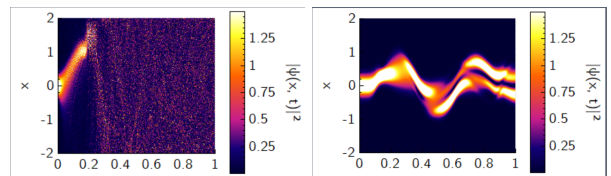


FIG. 7: Comparison of the state evolution for a numerically broken state (left) and a numerically working state (right). On the first axis is normalized control time and on the second axis position in μm .

is limited to what can be handled without breaking the state due to inaccuracy. Thus, high frequency controls can be explored, but if sufficiently large durations are desired, then the amplitude of this oscillation must be scaled to accommodate this.

B. Challenge exploration

Through simulations it seems that BECs have very different dynamics compared to single particles. Without using GRAPE, one can study how the similarity between single particles and BECs changes depending on the value of β . Keeping the control function and control duration the same, we find that what was a good solution for a single particle reaches a lower and lower fidelity as we change β away from 0, but that it also reaches revivals with a higher fidelity compared to neighboring values. This behavior is shown in figure 8 where the non-optimized control

$$u_{\text{non-opt}}(t) = 0.12 \cdot \sin(0.86 \cdot \omega_{01}t) \quad (8)$$

with final time $T = 3.49$ was simulated for different values of β . We define ω_{01} to be the energy frequency between the ground and first excited state. This varying

should not come as a surprise. The function in eq. (8) was chosen because of its high fidelity score around $\beta = 1$, but expecting this control function to perform well for the wide range of other systems (i.e. values of β) it is tested on would not be expected. While the behavior of similar systems is not vastly different, this small change to the system dynamics does mean that some systems will not reach a high fidelity at the final time, while others will. This can also be seen on Fig. 8, where the fidelity as a function of time has been plotted for 2 different interaction strengths. So when one does not optimize the control function it does not seem viable to use it on a broad range of systems as it then becomes a matter of 'timing' the control duration T to reach a high fidelity instead of examining the performance of the control and making changes according to what was observed.

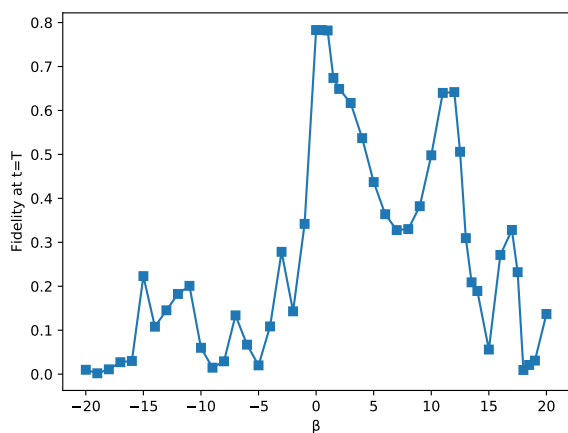


FIG. 8: Fidelity of a non-optimized solution for different values of β in a quartic potential.

Due to the same spaced energy levels of a quadratic potential, it is difficult to excite the state exclusively to the first excited state. It is likely that some part of the state is driven into higher excited states and so it is challenging to reach high fidelities with this kind of potential. Nonetheless, the nonlinear contribution to the GPE Hamiltonian seems to make it possible to improve the fidelity of control functions compared to the levels reachable for a single particle. In a quadratic potential different controls were optimized to estimate the highest reachable fidelity for this kind of potential as a function of the nonlinear interaction strength β . These controls were optimized with GRAPE for 100 iterations and the highest reachable fidelity was then noted, the results can be seen on Fig. 10. As can be seen on the figure, it is actually possible to reach higher levels of fidelity for all measured values of $\beta \neq 0$ with control duration $T \approx 1.36$. The highest result shows that it is possible to improve the fidelity score with about 50% compared to the reachable score of a single particle. One should also note that controls with duration $T = 10.2ms$ does not perform as well

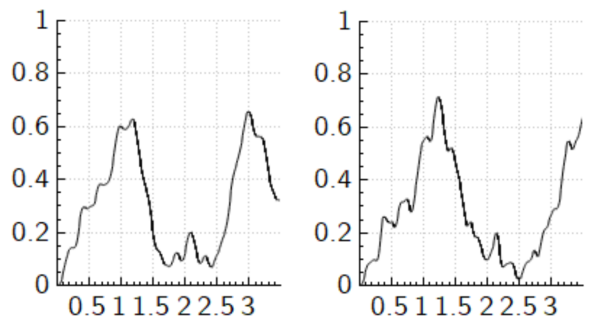


FIG. 9: Fidelity as a function of time for the control function in Eq. (8) for two different values of β . Left is $\beta = 7$, right is $\beta = 12$. As can be seen, the overall trend is the same for the 2 systems and so it becomes a matter of 'timing' the final time to reach a higher fidelity if one refrains from using GRAPE to optimize controls.

as the other 2 shown durations. This is suspected to be caused by the long control duration allowing for states to be excited beyond the targeted first excited state, which the smaller control durations are not as susceptible to. Another explanation could be that the increased control duration makes the optimization landscape more complex and so more sensitive to the initial control that is being optimized on.

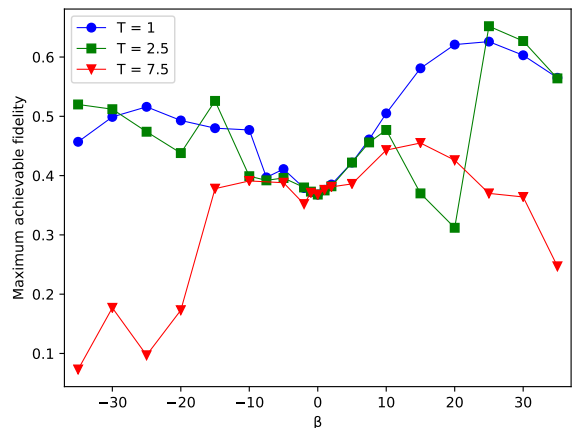


FIG. 10: The result of varying β and T for a BEC in the harmonic potential. As it can be seen, there exists 'resonant' combinations where it is possible to achieve above 50% higher fidelity compared to a single particle (at $\beta = 0$), but many also combinations that are on par with or worse than a single particle.

It was attempted to estimate the Quantum Speed Limit for this control problem and its dependence on the interaction strength of the BEC, β . The methods for determining seeds are discussed in section V. The seeds chosen were then optimized for 100 iterations where it was then simulated to see if it could surpass a fidelity

of 0.99. The resulting estimates of the QSL for different values of β can be shown on Fig. 11. Notice how it seems that the speed limit is lower for attractive BECs and increasingly higher for increasingly more repulsive BECs. It was not possible to estimate the QSL for values of β below -10 or above 25 , since no control function could be optimized to 0.99 within 100 iterations.

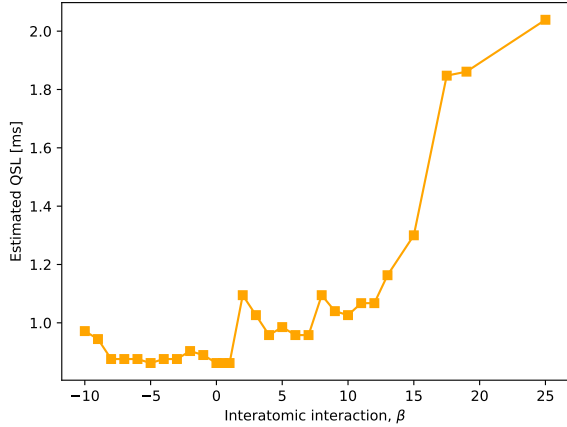


FIG. 11: The estimated Quantum Speed Limit for a BEC in a quartic potential as a function of β .

It is clear after having explored the behavior of BECs in different numerical setups that the behavior the BECs are not symmetric with respect to β . This is not unexpected, since the sign of β constitutes whether the atom-atom interaction happening inside the BEC is repulsive ($\beta > 0$) or attractive ($\beta < 0$) and so it would not be unjustified to say that the sign of β describe entirely different species of bosons. Since attractive BECs have proven difficult to optimize on for $\beta < -5$, it was explored what the lower limit of β would be with respect to reaching an optimizable fidelity of at least 0.99. The results of this exploration is seen on figure 12, where it can be seen that for values of $\beta < -10$ it is no longer possible to reach $F \geq 0.99$. While this lower bound could be seen as a numerical limitation of Composer as discussed in Sec. IV A, it is also relevant to consider that there exists a lower bound for how attractive a BEC can be before the atoms collapse in on themselves in a process known as a Bose-Nova[7].

C. Clustering of solutions

In a recent article where this problem has also been studied, it was found that optimal solutions for increasing final times were dominated by cosine functions of increasing frequencies $k\pi t/T$. The figure from [8] is shown on Fig. ??.

To replicate this result in using Quantum Composer,

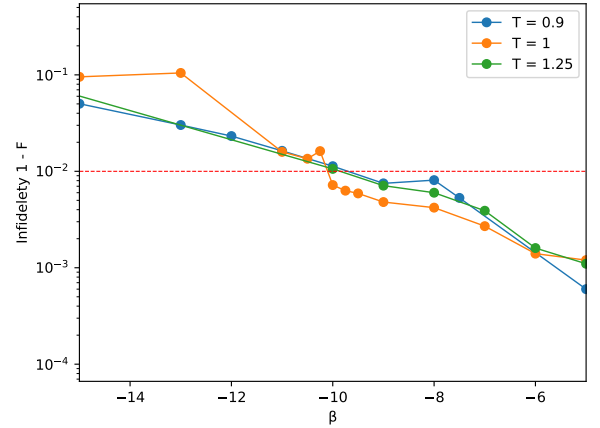


FIG. 12: Lowest reachable infidelities as a function of interatomic attraction β , shown for different timescales. It seems that $\beta = -10$ is the lower limit for reaching optimizable solutions $F \geq 0.99$. This lower limit is, however, suspected to be caused by numerical inaccuracies. The dashed red line shows is to help show where the systems cross $F \geq 0.99$.

the system conditions had to be rescaled to Composer, where the kinetic factor κ is locked at 0.5, while in the original system it was instead 0.36537. Thus, the different quantities of the system had to be reworked. This recalculation can be seen in App. A.

It was attempted to replicate this result by letting GRAPE optimize seeds on the form $u(t) = A \cos(k\pi t/T)$ for 750 iterations each. By varying k and A , the frequency and amplitude of the seeds were varied and their optimized solutions and reached fidelity noted. The results can be seen on Fig. 13. As can be seen on the figure, the results do not replicate those found in [8] where solutions dominated by different values of k outperform other k 's for some interval. Note in particular that the overall highest fidelity was reached for a seed with frequency $k = 2$ at $T \approx 0.93\text{ms}$, where $k = 5$ was expected to be the best performing frequency. However, note that both $k = 4$ and $k = 5$ reach solutions that outperform the other seeds for that particular timescale and that they do so in the expected order (4 before 5).

As is also shown on the figure, seeds with different frequencies do not have the same amplitude. The reason for this is that as the frequency of the seeds increased, it was found that having a too high amplitude caused the state to become numerically unstable and the simulation to not function correctly. This is also the reason that the $k = 4$ data spans a smaller time interval due to numerical errors in their solutions for $T > 0.8\text{ms}$.

As can be seen on figure 13, there are durations where the different control seeds reach similar levels of fidelity.

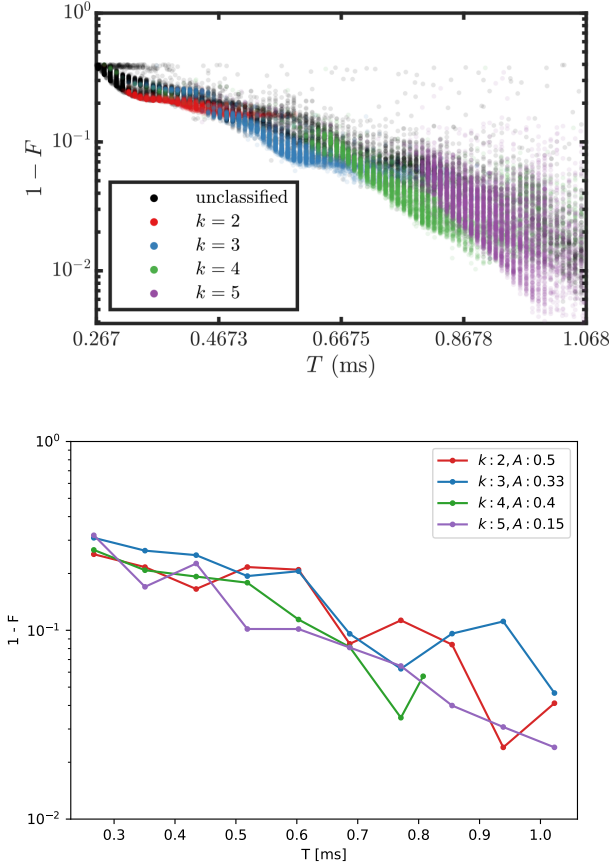


FIG. 13: **Upper:** This figure is taken from [8] and shows how optimized controls can be clustered by their projection onto $\cos(k\pi t/T)$. **Lower:** Infidelity of GRAPE optimized solutions with initial control $u(t) = A \cos(k\pi t/T)$. The first axis shows the control duration T .

This could indicate that they find the same or a similar optimized solution. The optimized solutions from figure 13 with control duration $T \approx 0.685$ ms are shown in figure 14. As can be seen on this figure, these optimized controls do not look alike despite reaching fidelity. It is most likely due to how different their initial control seeds are and so, one could argue that these seeds are located away from each other in the fidelity landscape. What should be noted though is the connection between seed frequency in $k\pi$ and the number of 'shakes' on the corresponding optimized control function. We see that for $k = 2, 3, 4$ the number of shakes matches this frequency, which also strengthens the hypothesis for why these seeds don't converge to the same control.

While examining the optimized solutions it was found that seeds with the same frequency in $k\pi$ would sometimes converge to similar optimized controls. Figure 15 shows an example of this where the seed $A \cos(3\pi t/T)$ was optimized for different amplitudes A and different durations T , yet still reached a very similar optimized

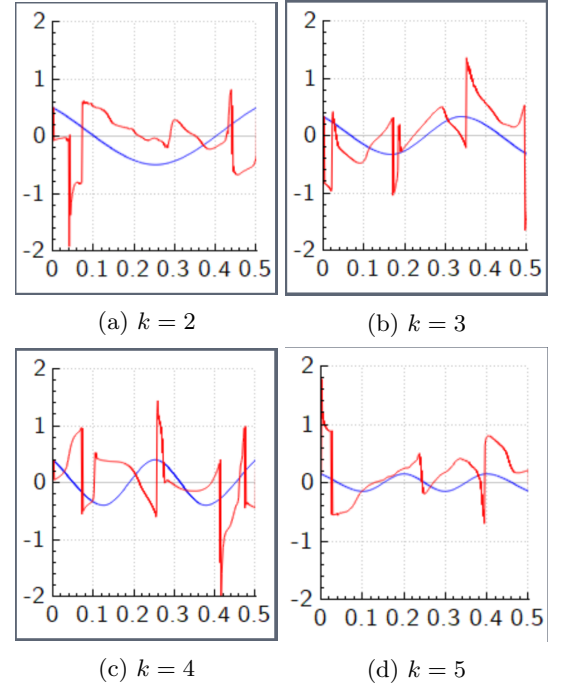


FIG. 14: Screenshots from Quantum Composer showing GRAPE optimized controls for the same seeds as in figure 13. First axis is time in scaling units, second axis is potential displacement from $x = 0$ in scaling units. Each figure shows the control seed (blue) and GRAPE optimized control (red) for that seed.

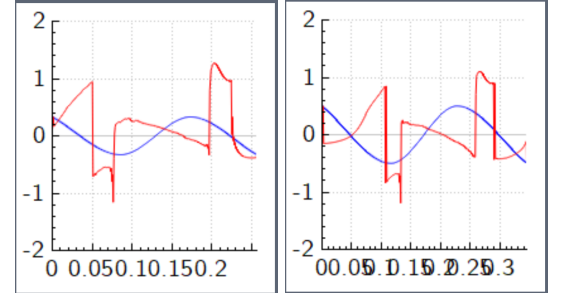


FIG. 15: Screenshots of optimized controls of a similar seed for 2 different durations. The axis are read in the same way as in figure 14. Left: $T \approx 0.35$ ms, $A = 0.33$. Right: $T \approx 0.47$ ms, $A = 0.50$.

solution. Notice also that both of these solutions have 2 shakes in the regime where it was found that optimal solutions were dominated by $\cos(2\pi t/T)$ frequencies [8].

D. Robustness of solutions

The robustness of optimized solutions wrt. interaction strength or potential scaling was studied in [4]. This is interesting since real experiments are often subject to the inherent uncertainties of their equipment e.g.

the frequency width of a laser. Since the interatomic interaction in a BEC is dependent on the number of atoms it consists of, it is not surprising that a specific factor β is difficult to replicate exactly in the lab. Likewise the trapping potential is created by standing waves of laser light and so it is also subject to some uncertainty in intensity and frequency.

The robustness of optimized solutions was replicated for the same system as the solution clustering. This replication was done by letting GRAPE optimize the same control seed, $u(t) = 0.15 \cos(5\pi t/T)$, for 300 iterations and for different durations $T = 0.5\tau$, $T = 0.875\tau$ or $T = 1.3\tau$, noting that these durations are respectively below, near and above the QSL of this system. These optimized solutions were then replayed in either a system with a modified interaction factor $\beta = a_I\beta$ or in a system with a scaled potential $\tilde{V}(x, u(t)) = a_V V(x, u(t))$. The results can be seen on figure 16. As can be seen on the figure, there seems to be an overall trend where the more optimized a solution is for the system, the more sensitive it is to small changes in either the potential or the interaction. Keeping in mind that better solutions are reached for longer durations, this could also be interpreted as how the 'fidelity landscape' becomes increasingly more complex the longer the control duration is. This could make sense since for small durations below the QSL, there could be many 'plateaus' near optimal fidelity. Small changes to the control could possibly change the shape/height of the plateau, but if the changes are small then the optimized solution should still be somewhere on this plateau. On the contrary, for long durations above the QSL, the landscape is more 'spiked' due to some specific solutions being able to reach very high fidelities and this spiked landscape would of course be more affected of changing parameters.

V. STRATEGIES

Throughout working with this project, there has been two ways to determine the control function $u(t)$ in Composer: The control function is determined either by an analytical expression (using the Scalar expression node) or it is made by dragging the desired 'path' of displacement on a graph (using the Control node). These different nodes are shown in figure 17. One can use both of these approaches when working with quantum control and they both have their advantages and disadvantages.

Using an analytical expression as a control function, one gets the possibility of tweaking initial parameters to great accuracy such that the seed GRAPE gets already has a high fidelity. It is preferable to the graph approach in cases where a desired oscillation frequency is wanted e.g. the function $A \sin(E_{01}t)$. These kind of functions are difficult to mimic in the graph approach. When working

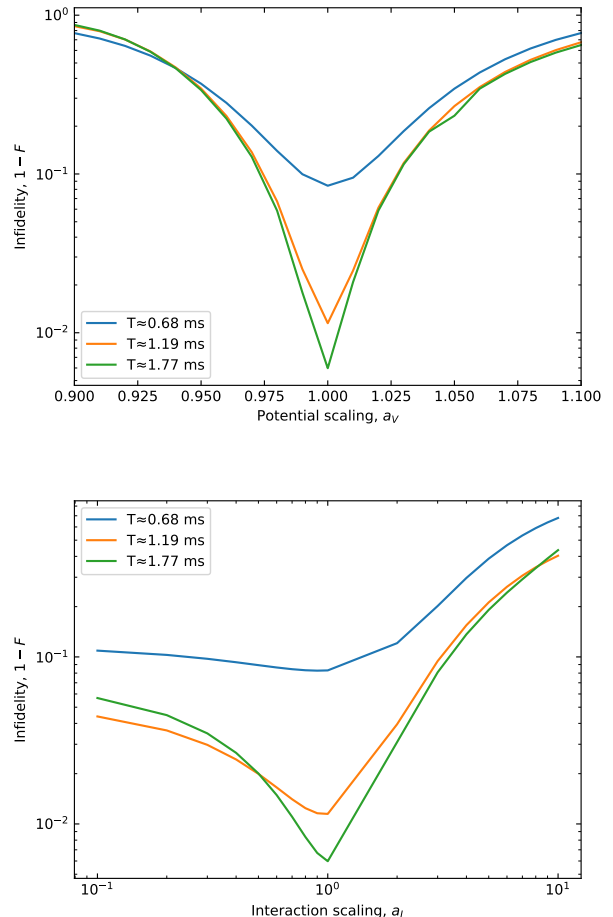


FIG. 16: Examining the robustness of an optimized solution wrt. scaling either the potential (top) or the atomic interaction (bottom).

with expressions independent of the final time T , adjusting this parameter translates into cutting the graph at different times. This is useful when exploring different functions and one finds that higher fidelity occurs at a different time than one currently works with. This method is shown in figure 18.

The Control graph node makes it possible to build up the control function as a series of steps. Here one can move one point on the graph and watch how the state evolves until that point, making it possible to optimize each step before going further. Since the graph node always spans the entire time interval $\{t_0, T\}$, this makes using the node useful when wanting to stretch or contract the entire function as shown on figure 19. As demonstrated with the BEC in a harmonic potential in figure 10, there exists a complex relationship between β and T and so being able to keep the shape of the control function the same while altering the total time makes the node a beneficial tool to use.

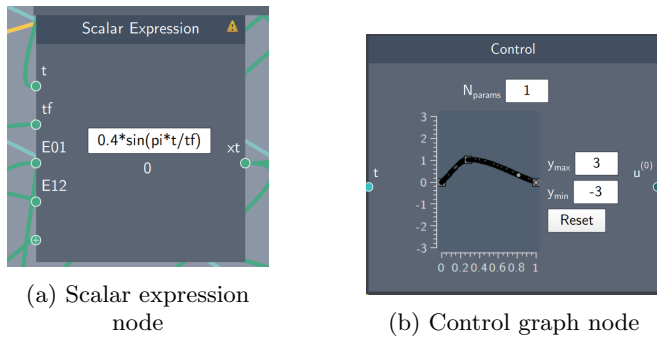


FIG. 17: Both nodes used for making a control function in Quantum Composer. The analytical function method (a) can take a number of input constants like final time $tf = T$, energy spacing between different states ($E01 = E_1 - E_0$) and variables such as time t . The graph method (b) always scales the first axis such that the first point on the graph is at $t = t_0$ and the final point is at $t = T$.

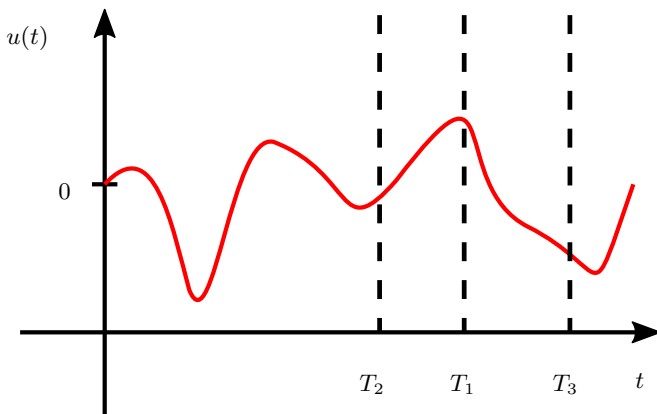


FIG. 18: This figure shows how using an analytical control function can be used to reach a higher fidelity seed by altering the final time T . In this example we use the control function shown in red and the final time T_1 . Evolving the system for a shorter(longer) time yields a higher fidelity though, and we can reach these fidelities by replacing the final time with T_2 (T_3). This method only works if the control function is independent of the final time.

When calculating the maximum reachable fidelity for some combination of β and T in the harmonic potential, the graph control node was used to create for GRAPE. The overall strategy using this tool involved moving points on the graph into certain shapes and looking at how the initial (i.e. no GRAPE) fidelity looked. I particularly liked the semi-circle shape of a $\sin(\pi t/T)$ function where the amplitude was varied. Another particular shape used in this work was generated by moving a single point around $0.1T$ or $0.9T$ to create a function which had a fast(slow) buildup to a maximum and then a slow(fast)

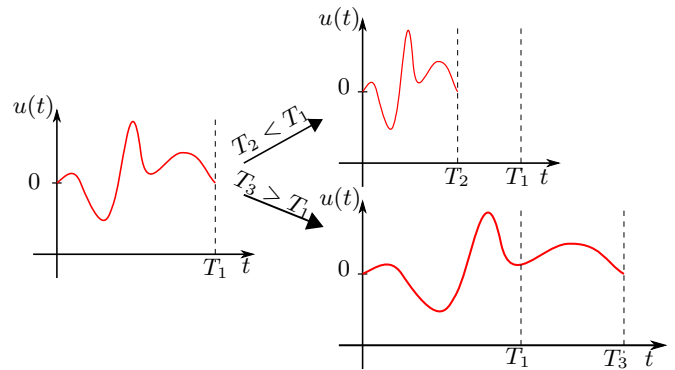


FIG. 19: A visualization of one method that can be used to optimize fidelity when working with the graph node. If one tries to change the final time T the resulting control function will be squeezed or stretched as a result. Altering the final time can be beneficial given the complex relationship between β and T , but also as a tool to regulate the speed at which the potential moves. Using this approach also has the practical advantage that we can draw the graph such that $u(T) = 0$ and so no matter the final time, the potential will always end up where it started. This is to be compared to the approach shown in figure 18 where this is not always the case and the state would possibly need to be transported back to its initial position.

return to the initial position. This wedge-like shape is shown on Fig. 17 (b).

Both of these strategies worked well and provided nice seeds for GRAPE to optimize on. Despite some cases where the fidelity changes significantly from small variations of β , the overall rule seems to be that something that works well for one value of β will have a similar fidelity for nearby values of β and so this was also used to find a good seed when β was larger than 10, since above this limit the system seemed a lot more sensitive to the initial control function with respect to reaching the highest possible fidelity for that particular combination of β and T .

Measuring the Quantum Speed Limit for the quartic potential involved a combination of 2 strategies: The self-dubbed "up-to-down" and "down-to-up" approaches.

UP-TO-DOWN:

1. Pick T_{up} to be a control duration estimated to lie above the QSL of the system
2. Find an optimizable solution capable of reaching F 0.99
 - If unable to do so, return to step 1. and choose a higher T_{up}
3. Gradually lower the duration in small steps, ensuring at each step that the control can still be optimized to F 0.99

- If the control is unable to reach F 0.99, one should try minor modifications to the control to see if this fixes it.

4. The lowest reachable T is then the estimate of the QSL for the system. These steps can be repeated for another control function.

The UP-TO-DOWN procedure was my mainly used method for estimating the QSL of the system shown on Fig. 11.

DOWN-TO-UP works in a similar way to UP-TO-DOWN, but in a reversed way.

1. Pick T_{down} such that it is estimated to lie below the QSL of the system.
2. Try a number of different control functions and choose the most promising (i.e. the seed capable of reaching highest fidelity)
3. T is raised in small steps, at each step optimized and modified to check if F 0.99 is reachable
4. The first time F 0.99 is reachable, the current T is the estimated QSL for the system.

This procedure attempts to exploit that the optimization landscape for GRAPE to traverse is hopefully simpler due to the lower control duration that is then gradually raised in the hopes that the seed has found a potential fidelity peak in the landscape that becomes available for some slightly higher control duration.

While reproducing the clustering of optimized controls data seen in Fig. 13, it became apparent that using cosine functions instead of sine functions could be advantageous for small timescales. One can use that fact that cosine is displaced from 0 at $t = 0$ to ones advantage as it places the state on a steep slope of the potential. This generates motion of the state much faster than when initiating with a non-displaced control function. Similarly when the control is done, one can displace the potential such that the state feels rapid deceleration, thus reducing the velocity of the state much faster than if the control was constrained to end at $u(T) = 0$. This behavior can also be seen on the optimized controls shown on Fig. 14, where both end points experience rapid displacement away from $x = 0$. These advantages are similar to the "back-swing" solution for another quantum control problem described in [8].

VI. QENGINE

This section describes the work I've done using the QEngine, a C++ library that Composer is build on top of. This is the expert tool compared to Quantum Composer which fills out the role of amateur tool. Instead of the GUI that Composer offers it is library of C++ methods, which is inherently more flexible and

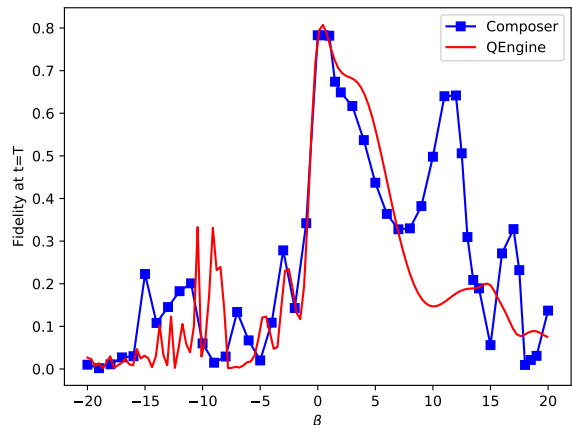


FIG. 20: Comparison of the results obtained using Composer and the QEngine to solve the same problem.

customizable than the graphical interface built onto it. The QEngine library comes with a number of example files that can be modified to set up a desired system and e.g. a control function with optimization with one of the three available optimizers. The `DataContainer` class can be assigned to save variables, control functions, etc. and can be saved in either `.mat` or `.json` format to later be read by either MATLAB or anything else, respectively. Similarly to the Composer case, the data gathered using the QEngine is further analyzed and plotted using PYTHON.

To start off an exploration into the advantages that QEngine offers compared to Quantum Composer, a previously examined problem is replicated using the QEngine. The replicated problem is the periodically shaken potential shown in Fig. 8 and the QEngine results are included in Fig. 20. While the 2 tools reach similar results between $\beta = -5, \dots, 5$ it cannot generally be said that the 2 tools get the same results when simulating this system.

In order to explain this difference in behavior, the energies of the first 3 energy eigenstates were calculated in the QEngine and compared with those calculated and plotted in Fig. 6. The difference between the Composer energy estimate and the QEngine estimate is plotted in Fig. 21. As can be seen there are small differences between the energies calculated by the 2 tools. This could explain why we observe the difference in fidelity shown on Fig. 20, since this indicates that the initial and final states are not exactly the same in both programs and thus we would expect them to evolve differently in time.

Despite this apparent difference in behavior between the 2 tools, the advantages of using QEngine for quantum control optimization and initial seed exploration is

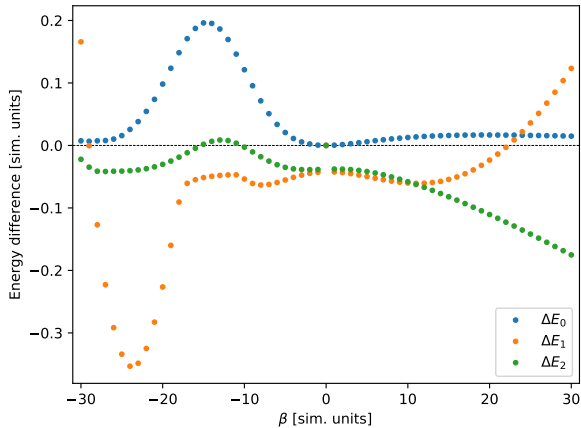


FIG. 21: The energy difference between Composer and QEngine for the same system as shown in Fig. 6. The striped black line marks $\Delta E = 0$.

demonstrated with the following setup. Firstly, the system is set up using the *exact* same parameters as in the QM2 *Shake Up* since it is possible to set the kinetic factor κ to any value (Remember that in Composer κ is locked to 0.5). Furthermore, the control time was varied between $0.25ms$ and $1.25ms$ in steps of $1/32ms$. With the control durations set, an initial seed

$$u_{init}(t) = \sin(2\pi t/T) \quad (9)$$

was set, before a set of 3 seeds were derived from u_{init} but with added random noise. The noise was set to be a random number in the interval $[-0.7, 0.7]$ and was added to each point of the initial control. Each altered seed was then optimized using all three optimizers until the step size was below 10^{-8} . The resulting infidelity of these optimized solutions are shown on Fig. 22, where the best results from Composer in Fig. 13 have been included for reference. From this one seed with added noise, it is possible to calculate many more datapoints and optimize seeds to reach better fidelities than what is possible in Composer, even when GRAPE does the optimization. This is most likely a result of having no iteration cap on the optimizer in the QEngine, but solely have a lower bound on the size of the step taking in each iteration. This outperformance of Composer does come at the cost of a considerable computation time of around 10 hours to run the code on a laptop with a 1.51 GHz CPU.

VII. COMPARISON

As has been demonstrated, QEngine offers several advantages to Quantum Composer with respect to flexibility and computational power. However, this does not mean that Quantum Composer should be used for all things. Quantum Composer makes it very easy to set up

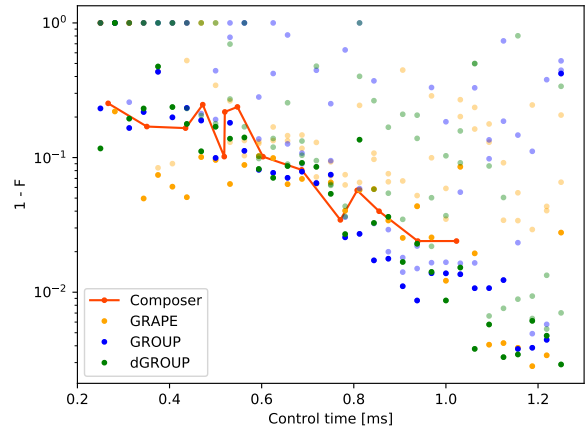


FIG. 22: Infidelities of solutions optimized using each of the 3 optimizers available in the QEngine. All seeds are generated from the same control function, u_{init} , where random noise is then added before optimization.

a desired system, run a simulation on that system and get the system plotted as the system evolves. This makes it preferable to use Composer when one wants to visually and (more or less) playfully explore the behavior of a system as the different parameters are tuned up or down.

Here, I will attempt to list some of the pros and cons of both tools.

QEngine

- + Flexible and customizable
- + 3 Optimization algorithms
- + Superior data gathering capabilities
- Data must be plotted separately
- Data gathering can be *very* time consuming
- Steep learning curve
- Difficult installation process

Quantum Composer

- + Many visualization options
- + Intuitive to explore a given system
- + Graph node to drag out a control function
- + Simulating and optimizing is usually fast
- Limited flexibility and customization
- Much data must be manually gathered
- Some data cannot be exported from the program

Based on the considerations above it is concluded that Quantum Composer is preferable to use as an exploration tool. It is fast to set up, relatively easy to use and most importantly it is very good at visualizing the simulation through its different kinds of plotting nodes. An example of this exploration could be a physicist trying to simulate an experiment to get a feeling for what kind of settings the experimental aperture should be capable of. With the current state of Quantum Composer, I find it advantageous to use the QEngine for any kind of simulation where more than around 10 datapoints are desired. While the QEngine has a vastly steeper learning curve, it is also that much more designed for generating large amounts of data and efficient calculations. It should also be mentioned that it is my personal experience that having worked or used Composer before working with the QEngine might have eased the learning curve of working with the QEngine. This is because many of the nodes in Composer correspond to a single class in the QEngine library and so one can use the intuition of how the nodes are connected in Composer to find out how to set up a system in the QEngine. The example files provided with the library are also of great help to get started.

It could also be advantageous to combine the playful, visual and fast elements that Composer has, with the more systematic, flexible and superior data gathering of the QEngine. One way to possibly exploit the best of both these tools could be to use Quantum Composer for globally exploring a given system through the intuitive and playful graph control node. Any promising candidates for good seeds could be exported from Composer in the .csv format to be read by a QEngine script, where noise and other more local optimization approaches could be utilized.

These considerations also give rise to a number of suggestions for desirable features for Quantum Composer in order to make it more attractive to use as an expert tool. Most of these points are a result of a brainstorming/discussion session with Shaeema and Tiantian.

Desired Composer features

- Conversion of real world data to dimensionless input scalars
- Exportable control functions to potentially be used in experiments
- Save data even when for-loops have run
- Import control functions from an experiment
- Enhance data gathering capabilities all around
- Select a time range for control durations
- Feed analytical control functions to GRAPE optimizer

- Add noise to the control function
- A more detailed description of how to use the GPE spectrum node
- Undo and redo buttons
- An ability to automate gathering data

VIII. DISCUSSION

As explained in section V, the Quantum Composer software in this workshop allows for 2 different ways to construct control functions. While both methods have been used to explore the behavior of BECs subject to a controlled potential, the files only allow optimization of control functions made using the graph method.

While the graph method has its advantages as discussed above, the analytical function approach likewise has advantages that could be combined with GRAPE. Having multiple approaches to the optimization problem would most likely be beneficial both from a perspective of understanding and from the perspective of being able to fine tune parameters for generating good seeds.

From what has been observed partially during exploring the dynamics of BECs and partially while gathering data for Fig. 10 and the QSL in Fig. 11, it is safe to say that when working with high values of $|\beta|$, the behavior of the system becomes difficult to predict. As final time goes up this only becomes more apparent, which was the conclusion that was reached after optimizing BECs in the harmonic potential.

This raises a (probably realistic) concern that the data shown in those 2 figures have been very sensitive to the initial conditions (i.e. the seed) and that other values could have been reached given more time to try different control functions. This is of course one of the problems with optimizing complex problems in the first place: They can be incredibly sensitive to initial conditions and while finding a local minimum is not a problem, finding the global one is a (very) difficult one.

Throughout both the exploration and data collection parts of this workshop, the control functions that have been used have for the most part been simple harmonic functions like $A \sin(\pi t/T)$ or other simple approaches such as the function seen in Fig. 17 (b).

However, what has also become apparent while working with GRAPE is that it usually introduces (after a number of iterations) rapidly oscillating behavior on parts of the graph. It is understood that this is most likely due to the way GRAPE optimizes control functions, but it also shows that more complex control functions are not necessarily bad and can actually in some cases reach much higher fidelity than their 'simpler' counterparts. In any case the simpler control functions seem like a good way to explore the fidelity landscape (so-called

”global search”) while optimization algorithms such as GRAPE are then used to find the locally optimal control, usually more complex than the input seed.

As described in section IV A, some of the ways to explore this problem are blocked by numerical instabilities. What has not yet been tested is how changing the discretized space or discretized timescale (i.e. by adding more points and/or having a larger space) could potentially solve some of these problems. What has been found though, is that changing these parameters also introduce much higher variance to either ground or 1st excited state and so do not currently seem like viable strategy for solving this numerical problem, although it cannot be excluded either.

IX. CONCLUSION

It has been demonstrated that the behavior of BECs are sensitive to the parameters β and T . As it has been shown in Fig. 10, this changed behavior compared to a single particle can be beneficial and capable of reaching higher fidelity than is otherwise possible. On the other hand it is also apparent that β introduces an unpredictability into the system as well as numerical inaccuracy of the Hamiltonian eigenstates. It has also been shown that the β term in (1) increases the QSL up to a estimated value of 2 times that of a single particle for the case $\beta = 25$. It was also attempted to cluster the performance of $\cos(k\pi t/T)$ control seeds to the frequency k , although the data shown in Fig. 13 shown that the seed frequency and optimized performance are not connected in a straightforward manner. What seems more promis-

ing is analyzing the number shakes in the optimized solutions and clustering these. The robustness of some optimized solutions have been explored both regarding a scaled potential and a scaled atomic interaction, as is shown in Fig. 16.

Some strategies involving choosing a control function have also been explored. The differences, advantages and disadvantages of using either an analytical function node or a graph node to determine the control function have been discussed. This includes how varying T affects the control function, as has been illustrated on figures 18 and 19. Finally, 2 concrete strategies for determining the QSL in this problem have been explained, both using the graph node to create a control function.

Numerical boundaries of the software used have been demonstrated and their implications for exploring the challenge have been discussed.

Future aspects of this problem that could be interesting to research include:

- Increasing the resolution of Fig. 11, since it has been shown that $\beta = -10$ is the optimizable limit for this problem.
- Analyze the optimized controls from Fig. 13 to make a new (T, F) plot where each point is color-coded based on the number of shakes in its optimized solution. It is expected that this will replicate what was found in [8].
- Compare the robustness of optimized solutions with different number of 'shakes', but where both solutions reach similar fidelities.

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Appendix A: System parameters

chosen from the requirement $\hbar = m = 1 \Rightarrow \chi = 1\mu m, \tau \approx 1.36$ where χ and τ are the scales of length and time respectively, such that $x_{SI} = \chi x$ etc. The paper has on the other hand chosen scales such that $\chi = 1\mu m$ and $\tau = 1ms$ [5]. This results in a conversion factor of $\kappa = 1.36537$ compared to those found in [5] of the following quantities: $\beta_{old} = 1.8299 \rightarrow \beta = 2.5042 = \kappa\beta_{old}$, $p_i \rightarrow \kappa p_i$ where $V(x, u(t)) = \sum_{i=2,4,6} p_i(x - u(t))^i$ is the potential of the system and finally τ .