Quantum Kinetic Theory of Condensate Growth—Comparison of Experiment and Theory

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In a major extension of our previous model 1 of condensate growth, we take account of the evolution of the occupations of lower trap levels, and of the full Bose-Einstein formula for the occupations of higher trap levels. We find good agreement with experiment, especially at higher temperatures. We also confirm the picture of the “kinetic” region of evolution, introduced by Kagan et al., for the time up to the initiation of the condensate. The behavior after initiation essentially follows our original growth equation, but with a substantially increased rate coefficient $W^*$.

Although the first Bose condensed atomic vapor was produced in a magnetic trap only in 1995 2,3,4, the kinetics of condensate formation has long been a subject of theoretical study 5,6. There is now intense theoretical work on Bose-Einstein condensation, which is excellently summarised in 5. Most theoretical studies of condensate growth either have not treated trapping, or have considered only traps which are so broad that the behavior of the vapor is not essentially different from the untrapped situation. Furthermore, they have given only qualitative estimates of condensate growth. Our previous paper 1 introduced a simplified formula for the growth of a Bose-Einstein condensate, in which growth resulted from stimulated collisions of atoms in a thermal reservoir, where one of the atoms enters the lowest trap eigenstate, whose occupation thus grows to form a condensate. We thus included the trap eigenfunctions as an essential part of our description, and gave the first quantitative formula for the growth of a condensate. The growth rate was of the order of magnitude of that estimated from experiments current at that time.

This direct stimulated effect must be very important once a significant amount of condensate has formed, but in the initial stages there will also be a significant number of transitions to other low lying trap levels whose populations will then also grow. As well as this, there will be interactions between the condensate, the atoms in these low lying levels, and the atomic vapor from which the condensate forms. This paper will extend the description of the condensate growth to include these factors, and will compare the results with experimental data on condensate growth obtained at MIT 3, with which good agreement is found.

As in our previous work, we divide the states in the potential into the condensate band, $R_C$, which consists of the energy levels significantly affected by the presence of a condensate in the ground state, and the non-condensate band, $R_{NC}$, which contains all the remaining energy levels above the condensate band. The division between the two bands is taken to be at the value, $e_{max}$. The situation is illustrated in Fig. 1. The picture we shall use assumes that $R_{NC}$ consists of a large “bath” of atomic vapor, in thermal equilibrium, whose distribution function is given by a time-independent equilibrium Bose-Einstein distribution \( \exp[(E - \mu)/k_BT] - 1 \) with positive chemical potential $\mu$. The value of $e_{max}$ will be assumed to be small enough for the majority of atoms to have energies higher than $e_{max}$, so that this part of the bath can be treated as being undepleted by the process of condensate growth.

![Fig. 1: a) Representation of the change in the energy spectrum due to the growth of the condensate. b) Occupations of the levels considered - leftmost is the condensate level, followed by several discrete energy bands, with the constantly occupied $R_{NC}$ at higher energies.](image)

In the following we will use the notation of $m$ as the mass of an atom with s-wave scattering length $a$, and $n_0$ for the number of condensate atoms.

The levels in $R_C$ must have time dependent energies due to the effects of the interaction with the growing condensate. The energy of the ground state is \( \mu(n_0) \), the chemical potential, for which we use a modified Thomas-Fermi approach \( \mu(n_0) = \alpha(n_0 + \nu)^{2/5} \) where $\alpha = (15\omega_x\omega_y\omega_z m^{2/5}h^2/\sqrt{2})^{2/5}$ and $\nu$ is a constant chosen so that $\mu(0) = b(\omega_x + \omega_y + \omega_z)/2$. As $\mu(n_0)$ rises with an increase in $n_0$, the energies of higher energy
levels must also rise. The exact nature of this rise does not affect the results greatly. An approximate treatment arises by leaving the levels with energies above $2\mu(n_0)$ unchanged (including all levels in $R_{NC}$), and compressing the spacing of the levels under $2\mu(n_0)$, so that the energies are given by

$$e_m = e_m^0 + \theta(2\mu(n_0) - e_m^0)(2\mu(n_0) - e_m^0 - \mu(0))/2$$

where $e_m^0$ are the non-interacting harmonic potential energy levels, and $\theta(x)$ is the step function. Note that $e_m^0 = \mu(0)$, and thus $e_0 = \mu(n_0)$. The levels used in this model are represented graphically in Fig. 1. To simplify the equations we also group the levels in narrow bands of mean energy $e_k$, $g_k$ levels per group, and $n_k$ particles per group. This corresponds to the ergodic assumption used in $[1,12]$.

Equations of motion then follow from Quantum Kinetic theory $[11]$ and full derivation will be given elsewhere, but the result is essentially equivalent to modifying the Quantum Boltzmann (QBE) equation as follows.

(i) We use the QBE in an approximated ergodic form, where $e_{\min} = \min(e_m, e_n, e_p, e_q)$

$$\frac{\partial f(e_n)}{\partial \tau} = \sum_{e_m, e_p, e_q} \delta(e_m + e_n - e_p - e_q)g(e_{\min}) \times \left[ f(e_p)f(e_q)(1 + f(e_m))(1 + f(e_n)) \right.$$

$$\left. - f(e_m)f(e_n)(1 + f(e_p))(1 + f(e_q)) \right],$$

where $n_k = g_k f(e_k)$ is the number of particles with energy $e_k$, and $\tau = (8\mu^2\omega^2/\pi\hbar) \times t$.

(ii) We use the modified energy levels as given above, but otherwise do not change the QBE.

(iii) We sum out over all levels in $R_{NC}$ which is assumed time independent.

(iv) We omit any scattering between atoms which both have energies less than $e_{\max}$, which is reasonable if the number of atoms in the bath is almost 100% of the total number of atoms.

in Fig. 2. The evolution for the population of the $m$th level in $R_C$ is then

$$\frac{\partial n_m}{\partial \tau} = \dot{n}_m = \dot{n}_m|_{\text{growth}} + \dot{n}_m|_{\text{scatt}}$$

where the individual terms are as follows.

Scattering: A collision between an atom initially in an energy level below $e_{\max}$ and a bath atom transfers the first atom to another energy level below $e_{\max}$. This is described by

$$\dot{n}_m|_{\text{scatt}} = e^{\mu/k_B T}\Gamma(T) \times$$

$$\left\{ \sum_{k < m} \frac{1}{g_m} \left[ n_k(g_m + n_m)e^{-\hbar\omega_{km}/k_B T} - n_m(g_k + n_k) \right] \right.$$ 

$$+ \sum_{k > m} \frac{1}{g_k} \left[ n_k(g_m + n_m) - n_m(g_k + n_k)e^{-\hbar\omega_{km}/k_B T} \right] \right\}. \quad (4)$$

where $\omega_{km} = e_k - e_m$. The value of $\Gamma(T) \equiv \sum_{e_m > e_{\max}} e^{-e_m/k_B T}$ depends on the energy spectrum. We use the value for an isotropic 3-dimensional harmonic oscillator with frequency $\omega = (\omega_0^2 \omega_2^2 \omega_3^2)^{1/3}$. Thus $e_n = (n + 3/2)\hbar\omega$, so that we find

$$\Gamma(T) = \frac{e^{-e_{max}/k_B T}}{1 - e^{-\hbar\omega/k_B T}}. \quad (5)$$

However this value is not critical; similar results are obtained for any $\Gamma(T)$ greater than about 10% of $[1]$.

Growth: A collision between a pair of atoms initially in the bath of atomic vapor results in one of the atoms having a final energy less than $e_{\max}$. This gives

$$\dot{n}_m|_{\text{growth}} = 2[(n_m + 1)W^+(n_0) - n_m W^-(n_0)]$$

where

$$W^+(n_0) = \frac{1}{2} \int_{e_{\max}}^{\infty} de_1 \int_{e_{\max}}^{\infty} de_2 \int_{e_{\max}}^{\infty} de_3 f(e_1)f(e_2)$$

$$\times (1 + f(e_3))\delta(e_1 + e_2 - e_3 - e_m), \quad (7)$$

$$W^-(n_0) = \frac{1}{2} \int_{e_{\max}}^{\infty} de_1 \int_{e_{\max}}^{\infty} de_2 \int_{e_{\max}}^{\infty} de_3 (1 + f(e_1))$$

$$\times (1 + f(e_2))f(e_3)\delta(e_1 + e_2 - e_3 - e_m). \quad (8)$$

Because we assume the non-condensate band is in thermal equilibrium, with temperature $T$ and chemical potential $\mu$, we have

$$W^-(n_0) = \exp \left( \frac{(e_m - \mu)}{k_B T} \right) W^+(n_0) \quad (9)$$

The value of $W^+(n_0)$ (henceforth $W^+$) differs significantly from that used previously $[1]$ in that the integrals in $[7]$ are only performed over energy levels higher than $e_{\max}$.
(previously over all levels), and the Bose-Einstein distribution function is used over all the range of integration (previously approximated by the Maxwell-Boltzmann distribution). With these changes we find that

\[ W^+(n_0) = \frac{1}{2} \left( \frac{k_B T}{\hbar \omega} \right)^2 \left\{ \log(1 - z) \right\}^2 + z^2 \sum_{r=1}^{\infty} [z(n_0) \Phi(z, 1, r + 1)]^2, \]

where

\[ z = e^{\frac{\mu - \mu_{\text{max}}}{k_B T}} \quad z(n_0) = e^{\frac{\mu(n_0) - \mu_{\text{max}}}{k_B T}}. \]

The function \( \Phi \) is the Lerch transcendent [12], defined by \( \Phi(x, s, a) = \sum_{k=0}^{\infty} x^k (a + k)^s \). These changes result in a significant correction, increasing \( W^+ \) by about a factor greater than three (dependent on \( T, \mu \) and the trap parameters) compared to that used previously in [1], and producing correspondingly faster growth.

By making a further approximation, that \( W^+_m(n_0) \approx W^+(n_0) \), the calculations required are significantly simplified. We can do this because the \( W^+_m(n_0) \) are an average over all the levels contained in the \( m \)th group, and hence are expected to be of the same order of magnitude as \( W^+(n_0) \). As a validity check, it was found that the effect on the condensate growth rate was small when the \( W^+_m(n_0) \) were altered by a factor in the range 0.5 – 2. We now have, for the growth terms

\[ \dot{n}_m|_{\text{growth}} = 2W^+(n_0) \left\{ 1 - e^{-\frac{\mu - \mu_{\text{max}}}{k_B T}} \right\} n_m + g_m, \]

\[ \dot{n}_0|_{\text{growth}} = 2W^+(n_0) \left\{ 1 - e^{-\frac{\mu(n_0) - \mu_{\text{max}}}{k_B T}} \right\} n_0 + 1. \]

The overall evolution of the system can now be found from the numerical solutions to the set of equations [3]. The parameters used are \( \omega_x = \omega_y = 2\pi \times 82.3 \text{Hz} \) and \( \omega_z = 2\pi \times 18\text{Hz} \), as in [3], and \( \alpha = 2.75 \text{nm}, \epsilon_{\text{max}} \approx 2.2\mu(n_0(\text{eq})) \), where \( n_0(\text{eq}) \) is the final equilibrium value of \( n_0 \), which is defined by \( \mu(n_0(\text{eq})) = \mu \), the chemical potential of the vapor. The number of energy bands was set as large as possible, but it was required that there were at least four levels in the first group above the condensate, in order represent the fact that the levels are discrete.

The improvements to this model over that used in [1] speed up the condensate growth by up to an order of magnitude, depending on the exact parameters, as anticipated in [3]. The major cause of this is the correction to \( W^+ \) arising from the use of the correct Bose-Einstein distribution. The inclusion of the scattering terms does not change the overall rate of growth substantially (which is dominated by the bosonic stimulated growth), but does speed up the initial period of growth (dominated by spontaneous growth) and shortens the time before the stimulated term becomes significant. This gives a much sharper initial growth curve as compared to the smoother S-bend curves of [1]. In Fig. 3 we present an example of the results obtained for the growth of all the bands in \( R_C \), in which a number of features may be seen.

(i) The effect of the initial conditions used can be seen from the front corner. In this example the initial populations for all but the top ten bands were set at zero, whilst the top ten bands had populations determined by the same Bose-Einstein distribution function as for \( R_{\text{NC}} \). The figure shows that this initial condition is rapidly smoothed out by scattering processes. Different initial conditions merely generate a small change in the initial time defined below.

\[ \Delta t = \text{Time (ms)} \log_{10}(\epsilon_m) \]

Fig. 3: \( f(\epsilon_m) \) vs. energy \( \epsilon_m \) as a function of time. Note that the lines almost parallel to the time axis are not lines of constant energy, but rather lines of \( f(\epsilon_m) \) for constant level number whose energies change with condensate growth. The solid curve in the plane at the top of the axes represents the curve \( \log_{10}(\epsilon_m) \equiv \log_{10}(\mu(n_0)) \) as a function of time.

(ii) The \textit{initialization time} (here 60ms) is defined by the critical line \( C-C \). Up to this point the population of the condensate level is relatively small. The behavior up to the initialization time is similar to that found by Svidunov and Kagan et al. In particular the populations of the levels increase to approach a limiting dependence on energy of \( f(E) \propto E^{-1.61} \) on the critical line \( C-C \), which is in good agreement with their prediction of \( E^{-5/3} \) for the case of a harmonic trap.

(iii) After the initialization time the condensate grows enormously. However, the occupations of the other trap levels actually decrease quite rapidly to their equilibrium values, and then remain nearly constant while the condensate continues to grow by a factor of about 10 in this case. At the same time the energy spectrum of the trap levels in \( R_C \) changes according to [3]. This accounts for the small variation in occupations of these levels which is still evident in Fig. 3.

Fig. 4 compares with the experimental data of ref. [8], for two different temperatures. The MIT group fit-
ted their data to an uncorrected growth equation \( \dot{n}_0 = \gamma n_0 [1 - (n_0/n_0(\text{eq}))^{2/5}] \), and reported only values of the parameter \( \gamma \). To represent that the experimental curve is a fitted curve rather than the raw data, the curve has been plotted as a broad band. The MIT group used the initial population for their curves as free parameters in their fit. We have set the initial populations for the MIT curves so as to give the best agreement with the initiation of our growth.

We have given a description of condensate growth which covers the full range of behaviors, both before and after initiation of the condensate. It is quantitative and agrees quite well with experiment. The behavior before initiation is essentially as given by the solution of the quantum Boltzmann equation, and agrees with computations based on this equation \[10,7,15\]. But we are able to give a value for the initiation time which appears to be consistent with experiment. After the initiation of the condensate, the occupations of the non-condensate levels are clamped by their fast relaxation time to quasi-equilibrium values, which change with the rise in trap levels induced by the very much slower growth of the condensate to its final occupation.

We believe the future development of the theory for this problem will involve mainly refinements of our picture, such as including depletion of our fixed “bath” of vapor. But for quantitative comparison with experiment, it will be necessary to have more extensive data on spatial distributions at a range of temperatures.

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**Fig. 4:** Comparison of theoretical growth curves (black) with experimentally fitted curves (gray) from \[8\].

| Theory | Experiment |
|--------|------------|
| $T = 830 \text{nK}$, $n_0(\text{eq}) = 7.6 \times 10^6$; Experimental $T = 810 - 890 \text{nK}$, $n_0(\text{eq}) = 7.5 - 7.85 \times 10^6$; | $T = 830 \text{nK}$, $n_0(\text{eq}) = 2.3 \times 10^6$; Experimental $T = 580 - 610 \text{nK}$, $n_0(\text{eq}) = 2.1 - 2.5 \times 10^6$. |

In the $T = 830 \text{nK}$ case, Fig. 4a, the growth speed predicted agrees with that experimentally found. The $T = 590 \text{nK}$ case in Fig. 4b shows a theoretical growth rate some three times slower than is measured. The discrepancy between theory and experiment at lower temperatures is hard to evaluate using the data in the form presented in \[8\], which do not allow for direct comparison between the actual projected spatial distributions as given by phase-contrast microscopy, and theoretical spatial distributions. These are not difficult to calculate from our many-level growth curves—the methodology will be published elsewhere. The MIT method fits to a zero chemical potential vapor plus a non-zero chemical potential condensate - a reasonable estimate in the absence of any theoretical description of the spatial distribution of the vapor. But a detailed description might give quite different results for temperature, and for condensate and vapor numbers.

In summary, we have given a description of condensate growth which covers the full range of behaviors, both before and after initiation of the condensate. It is quantitative and agrees quite well with experiment. The behavior before initiation is essentially as given by the solution of the quantum Boltzmann equation, and agrees with computations based on this equation \[10,7,15\]. But we are able to give a value for the initiation time which appears to be consistent with experiment. After the initiation of the condensate, the occupations of the non-condensate levels are clamped by their fast relaxation time to quasi-equilibrium values, which change with the rise in trap levels induced by the very much slower growth of the condensate to its final occupation.