

Bose-Einstein Condensate

Super Solids



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1 Bose Einstein Condensate

1.1 Questions

- $\int \frac{1}{r^n} d^3r$ divergent just for $n \leq 3$? **Yes.** Apart from an angular part, the integral goes as $\int dr \frac{1}{r^{n-2}}$. If you integrate till R (which you will tend to infinity), then you can easily see that for $n = 3$ it diverges as $\ln(R)$. For any other n it goes as R^{3-n} , and hence it converges for $n > 3$, and diverges for $n < 3$. **Thanks.**
- What is an s-wave? **s-wave means angular momentum $l = 0$.** **Thanks.**
- if $l < \frac{n-3}{2}$, and like k^{n-2} otherwise (Landau and Lifshitz, 1977). For a van der Waals-like potential ($n = 6$), only $l = 0$ (s-wave) matters at low energies. What's with $l = 0$? Lifshitz is a typo? **$l = 0$ is special. For $l = 0$ the centrifugal barrier $\frac{\hbar^2 l(l+1)}{2mr^2}$ vanishes.** Intuitively you can see that for short-range potentials, like $1/r^6$ you need to be close to $r = 0$, such that the particles see each other. If the kinetic energy is too low, then the kinetic energy is not enough to overcome the centrifugal barrier. As a result, only $l = 0$, i.e. the s-wave, contributes. **Thanks.**
- In the notes: eq. 1.8 to 1.9 the commutator $[\psi, \psi^\dagger]$ was used, but from 1.9 to 1.10 Bogoliubov approximation was used, why not directly on 1.8? **It could have been done directly in 1.8.** **Thanks.**
- What is the derivation of 1.11? It's a FFT, but what are the exact steps? **First $\hat{\psi}(\vec{r}) = \sum_{\vec{p}} \hat{a}_{\vec{p}} \frac{e^{i\vec{p} \cdot \vec{r}/\hbar}}{\sqrt{V}}$, where V is just a quantization volume, just to have the proper units. We will also Fourier-Transform the potential $V(\vec{r} - \vec{r}') = \sum \tilde{V}(\vec{q}) e^{i\vec{q} \cdot (\vec{r} - \vec{r}')}.$ Then, you see that**

$$\int d^3r \int d^3r' V(\vec{r} - \vec{r}') \hat{\psi}^\dagger(\vec{r}) \hat{\psi}^\dagger(\vec{r}') \hat{\psi}(\vec{r}) \hat{\psi}(\vec{r}')$$

becomes of the form of Eq. (1.11), after using that $\frac{1}{V} \int d^3r e^{i\vec{k} \cdot \vec{r}} = \delta(\vec{k})$. **Thanks.**

$$= \int d^3r \int d^3r' a_{p_3}^\dagger a_{p_4}^\dagger \delta(p_2 - p_4 - q) \delta(p_1 - p_3 + q) a_{p_2} a_{p_1} = a_{p_1+q}^\dagger a_{p_2-q}^\dagger a_{p_2} a_{p_1}$$

- What are the python packages to use operators like $\hat{\psi}$? There is some sympy implementation, but probably there is a better one? **I'm unsure what do you mean here.**
We surely will use calculation with more complicated operators than diagonal matrices (complicated Hamilton operator). Which python packages you know for that? I know that sympy has implementations for operators for symbolic calculations.

- how to get from 1.11 to 1.12? Is it a commutator expansion? Why q is disappearing? This is based on the so-called Bogoliubov approximation. Since the system condenses in $\vec{p} = 0$, you approximate \hat{a}_0 and \hat{a}_0^\dagger by $\sqrt{N_0}$, with N_0 the number of condensed atoms. This is because $N_0 \gg 1$. Actually $N_0 \simeq N$, and hence the number of non-condensed atoms is much smaller than N_0 . Then you will do the following. You will consider up to second order, i.e. up to terms with at most two operators $a_{\vec{p}}$ with $\vec{p} \neq 0$ (you will see that there are no terms with just one operator with $p \neq 0$). Let's have a look to the interaction term in Eq. (1.11) (I forget here hats and vectors in order to ease the notation) $a_{p_1+q}^\dagger a_{p_2-q}^\dagger a_{p_2} a_{p_1}$. Let's see which combinations you have that have at most 2 operators with a non zero momentum: $(p_1 + q, p_2 - q, p_2, p_1) = (0, 0, 0, 0), (0, 0, x, x), (x, x, 0, 0), (0, x, x, 0), (x, 0, 0, x), (0, x, 0, x), (x, 0, x, 0)$, where x means that it is not zero. The first term gives you the energy at order zero: $E_0 = \frac{U(0)}{2V} N_0^2$. Note that $N = N_0 + \sum_{p \neq 0} a_p^\dagger a_p$, and hence up to second order $N^2 = N_0^2 + 2N_0 \sum_{p \neq 0} a_p^\dagger a_p$. As a result: $E_0 = \frac{U(0)}{2V} N^2 - U(0)n_0 \sum_{p \neq 0} a_p^\dagger a_p$. The 3rd and 4th terms give $U(0)n_0 \sum_{p \neq 0} a_p^\dagger a_p$ which cancels exactly with the term in E_0 . The other terms will give you what you find in Eq. (1.12). Alright, but why are there just p and $-p$ in Eq. (1.12), but p_1, p_2, q Eq. (1.11)?
- In 1.17 ω_p part has a factor 2, but ω_z not, despite beeing symmetric in ψ . Why? This is because of the two directions on the plane. Of course. Thanks.
- What is variable a ? Why should $a > 0$ as repulsive short-range interactions stabilize the BEC (p.10)? a is the s -wave scattering length. It characterizes the short-range part of the interaction. As mentioned above the contact interaction is just described by what happens at s -wave. At low energies, this means that the short-range interactions are given by a single parameter, which is the scattering length. A positive scattering length means repulsive interactions, i.e. the interaction energy increases when the density increases, i.e. when the distances decreases (i.e. the particles repel each other). The opposite is true for $a < 0$, for which the particle attract each other. The dipole-dipole interaction is partially attractive and partially repulsive. Attraction is dangerous, because the system tends to collapse. Adding repulsive contact interaction may compensate the dipolar attraction, hence preventing collapse. Thanks. May compensate, means this is still under investigation?
- "When the atomic density grows due to the attractive interaction, three-body losses predominantly occur in the high-density region." What does three-body losses mean? Three-body losses means that three particles meet at close distances (which becomes more and more probable for larger and larger densities). When this occurs, two of the particles may fall into a bound state, giving the excess energy to the third one. As a result all three particle are loss from the system, and hence the term three-body loss. Thanks.

- “As the collapse occurs mainly in the x-y direction due to anisotropy of the DDI (in the absence of inelastic losses, the condensate would indeed become an infinitely thin cigar-shaped cloud along z), and therefore the condensate explodes essentially radially, producing the anisotropic shape of the cloud.” Why is the collapse not along z axis? Note that the dipoles attract each other when placed head with tail, i.e. in this case when they are placed aligned on top of each other along the z-direction. Hence collapse produces a thin “cigar” along z. However, before this “cigar” becomes infinitely thin, three-body losses kick out. When this occurs, the kinetic energy on the xy plane, which is huge (due to the strong compression, recall Heisenberg uncertainty), is released and the BEC explodes. A note here: the key point of all the droplet business in recent works (starting in 2016) is that the collapse in a dipolar condensate may be actually stopped due to the stabilizing effect of quantum fluctuations, but this wasn’t known when I wrote the notes! Thanks.
- How are the regions stable, metastable, unstable derived in Figure 1.5, here Figure 2? It comes from the two solutions of Eq. (1.25). Thanks.
- Typo in “we obtain a 1D equation similar to the a GP equation”, just the or a
- “ground-state wave-function is independent of the in-plane coordinates ” Why? This is because there is no trap on the plane. Then, the wavefunction is flat over xy. This is so if the interactions are repulsive, which is assumed. This is Ok as long as the roton (the energy minimum in Fig. 1.6) remains stable (i.e. remains such that the energy² ≥ 0 (i.e. the energy is not imaginary).
- 1.26 to 1.27, where does the U_{dd} go? This is because you take that the density is equal to $|\psi(\vec{r}, t)|^2 = (1/S)|\psi_0(z, t)|^2$, with S some quantization surface, to get the units right. The Fourier Transform is then $n(\vec{k}, t) = (2\pi)^2 n(k_z, t) \delta(k_x) \delta(k_y)$. When you integrate, using convolution theorem:

$$\int d^3r U(r-r') |\psi(r')|^2 = \int d^3k / (2\pi)^3 (4\pi d^2/3) (3 \cos^2 \theta_k - 1) n(k, t) e^{ikr} = \int dk_z / 2\pi (4\pi d^2/3) * 2n(k_z, t) e^{ik_z r_z}$$

- Typo If: “roton momentum. if this were so”
- Why should a modulation with a finite wavelength allow superfluids? I guess you mean supersolids. A density modulation with a finite wavelength (not a diverging length of the order of the size of the whole system, as it is the case for a phonon instability), means that you have a crystal-like modulation, and hence a supersolid. The problem is that in this theory, once the modulation starts to develop there is no way to stop it, and what one gets is actually local collapses.
- Typo repeatance: “the width of the width”

- What are the spin-F matrices? It is just the spin matrices for a spin $S = F$: $S_{x,y,z}$. Defining $S_{\pm} = S_x \pm iS_y$, then $S_+|m\rangle = \sqrt{F(F+1) - m(m+1)}|m+1\rangle$, $S_-|m\rangle = \sqrt{F(F+1) - m(m-1)}|m-1\rangle$, and $S_z|m\rangle = m|m\rangle$.
- Is the occurrence of these spin textures in Figure 4 special? Special in which sense? They are the result of spin excitations.
- “In 3D system there exists a regime where no collective modes are below the chemical potential, which corresponds to the particle emission threshold \Rightarrow excitation leads to self-evaporation of the droplet to zero temperature” Is this a good/useful or is it just counter-productive for experiments? Meaning switching between this regime and a normal one to get probably higher densities? It is interesting, because this would mean that the droplet would have zero entropy. One can then use it to e.g. cool other system. For example, you pass energy of a given system into the droplet, then the droplet immediately evaporates the extra energy, and this would cool. It may be useful in this sense, but it is not in my opinion the most relevant feature. In dipolar droplets (rather than those in binary non-dipolar mixtures) there is no regime without any excitation.
- In Figure 7 b, if the theoretical values would fit, then the dashed line would be where the darkest color is? If yes, why is it not? That’s right. It’s not there because the calculation is based on linear response theory, i.e. it is an approximation. the agreement is however pretty good, I would say.
-
- What are bright and dark solitons? Bright solitons are wavepackets that maintain their shape without spreading. A dark soliton is of the form $\psi(x) = \tanh(x/dx)$. Then you see that there is a phase slip of ϕ between left and right, and the density has a dip (down to zero density) at $x = 0$. A dark soliton does not move, but if the density does not fall to zero, then it may move, and you can visualize it (in terms of the density) as a constant density minus a bright soliton. It is not exact, but it gives you a rough idea.
- What is meant by crossover between bright solitons and droplets? It is important to understand the difference between droplets and solitons (I mean here bright solitons). In the case of solitons, the wavepackets keeps its shape due to the competition of kinetic energy (which pushes the wavepacket outwards) and attractive interaction energy (which pushes the wavepacket inwards). In 1D (and only in 1D for contact interacting systems), the kinetic energy (which scales with $\frac{1}{\text{length}^2}$) can eventually counter-rest the interaction energy (which scales as $\frac{-1}{\text{length}^D}$, with D the dimension). As a result you get as soliton.

A droplet is an altogether different issue. There you have the compensation of attractive mean-field interaction energy and the repulsive effect given by quantum fluctuations (LHY correction). One can visualize it as follows. Quantum fluctuations introduce a pressure outwards (this is what is called a Casimir pressure). This Casimir pressure compensates the tendency to collapse. As a result the droplet keeps its shape, i.e. it is self-bound even in 3D.

- “anisotropy of droplets leads to a frustration of the dipolar quantum droplet when compressed along the magnetic field direction” What does frustration mean in that context? Frustration is used in the following sense. In dipolar droplets, the droplet gets more elongated (along the dipole direction) when the number of particles increases. However, if one cannot get more elongated because there is a trap along the dipole direction, then the system will find more favorable to form two droplets instead of one. This leads eventually to the formation of a droplet array.
- “multi-droplet states appear upon crossing into the bistable region, due to the fragmentation of the system following a modulational instability \Rightarrow multi-droplet states are not the ground state” Why is this following (what is the argument here)? In the experiments they drive an initially stable condensate into instability. When the BEC becomes unstable, it starts developing ripples (modulational instability). These ripples lead eventually (although the nonlinear dynamics involved is pretty complicated) to the observed array of droplets. The droplets are hence formed dynamically. They are not the ground state. There may be however droplet arrays which are the ground-state of the system. There are experiments which have directly cooled into that ground state.
- What are in-situ density profiles? In-situ profiles means that they measure them while in the trap. The most common technique to image cold atoms is actually to release the trap, let the atoms fall and image them after some time of free fall (time-of-flight measurement, TOF). This is practically advantageous because when expanding the cloud becomes much larger and hence easier to image. The drawback is that what one measures in TOF is not the spatial density distribution but rather the momentum distribution. The reason is that expansion acts as a sort of Fourier Transform. You can intuitively understand that by thinking that the larger the initial particle velocity the farther it will move while falling. Hence the final spatial distribution after falling matches the initial velocity distribution.
- “interference of multiple quantum droplets, allowing the characterization of the nearest- and next-nearest neighbour coherence” Can this be seen in Figure 11, if yes how? If a side peak appears in momentum space, this means that there is coherence between the droplets. Otherwise you will see no side peak

but just the Fourier transform of each droplet, which will give you only a central peak. In the picture they plot the variance of the peak, which shows that it is reproducible from shot to shot, and hence that the droplets are coherent.

- When running `main.py` with a rectangle pulse, centered around the potential, the result is NOT the ground state, instead an excited mode. Why? I would say that the problem lies in the kinetic energy. First of all, did you check that the `fft` gives the correct ordering of the momentum distribution? As I told you the `fft` scrambles the ordering of momenta. Second, in the kinetic energy you have $(-0.5 * \text{self.k} ** 2)$. It should be $0.5 * \text{self.k} ** 2$. I'm not sure if these are the only problems, but these are in any case an issue.


```

75     self.x = np.linspace(-self.L, self.L, self.resolution)
76     k_over_0 = np.arange(0, resolution/2, 1)
77     k_under_0 = np.arange(-resolution/2, 0, 1)
78     self.k = np.concatenate((k_over_0, k_under_0), axis=0) * (np.pi / L)
79
80     if imag_time:
81         # Convention: $e^{\{-iH\}} = e^{\{UH\}}$
82         self.U = -1
83     else:
84         self.U = -1.0j
85
86     if psi_0 or V:
87         x_real = Symbol('x', real=True)
88
89     if V:
90         self.V = lambdify(x_real, V, "numpy")
91     else:
92         V = 1 / 2 * x ** 2
93         self.V = lambdify(x_real, V, "numpy")
94
95     self.psi = norm.pdf(self.x, loc=-0.1 * L, scale=1.0)
96
97     self.H_kin = np.exp(self.U * (-0.5 * self.k ** 2) * self.dt)
98
99     # Here we use half steps in real space, but will use it before and after H_kin with normal steps
100    self.H_pot = np.exp(self.U * (self.V(self.x) + self.g * np.abs(self.psi) ** 2) * (0.5 * self.dt))

```

Figure 1: The method from the first draft was implemented, but the resulting `split_time_real.mp4` looks wrong (imaginary time works). Probably the error lies in the equations, that were used. Note that Strang-Splitting was used as in 14: Per timestep H_{pot} was used before and after H_{kin} , but with just $0.5dt$. Also in that reference they used \exp^{-iH} , which seems right, instead of \exp^{iH} in the draft. Note also the definition of `self.k` being a concatenation of `[0:16]` and `[-16:0]` as `fft` expecting (this fixed `split_time_imag.mp4`)

DANIEL SCHEIERMANN, [split_operator.py](#) (2020)

1.2 Summary

- dipol-dipol interaction (DDI):

$$U(r) = \underbrace{g\delta(r)} + \underbrace{U_{dd}(r)} \quad (1)$$

$$\frac{4\pi\hbar^2 a(d)\delta(r)}{m} \quad \frac{C_{dd}}{4\pi} \frac{(e_1 \cdot e_2)r^2 - 3(e_1 \cdot r)(e_2 \cdot r)}{r^5}$$

- Use pseudo potential as dipol-dipol interaction is anisotropic and all partial wave (different l) mix

- coupling of different channels generates short-range contribution in the s-channel $s = 0 \Rightarrow$ by changing DDI strength a gets modified too \Rightarrow shape resonances \Rightarrow virtual state transform into a new ground state
- for fermions s-channel does not exist, so just long-range
- FFT of U_{dd} using spherical harmonics Y_{lm} gives:

$$\tilde{U}_{dd}(k) = \int d^3r U_{dd}(r) e^{-ik \cdot r} = \frac{C_{dd}}{3} (3 \cos^2(\theta_k) - 1) \quad (2)$$

- Use DDI in Gross-Pitaevski Equation, FFT, approximate to 2nd order, diagonalize with Bogoliubov transform
- As a result the square root can be imaginary, so the BEC gets dynamically unstable for long-wave length (phonon-instability):

$$\epsilon(p) = \sqrt{\frac{p^2}{2m} \left[\frac{p^2}{2m} + 2n_0 (g + U_{dd}(p)) \right]} \quad (3)$$

$$= pc_s \sqrt{1 + \epsilon_{dd} (3 \cos^2 \theta_p - 1)} \quad (4)$$

$$\underset{p \rightarrow 0}{=} pc_s \sqrt{1 - \epsilon_{dd}} \quad (5)$$

- For dipolar BEC the trap geometry is crucial (for non-dipolar not)
- “pancake traps” can stabilize the phonon-instability
- qualitative features for $a_{crit}(\lambda)$ by gaussian ansatz, for exact numerical solution non-local Gross-Pitaevskii Equation needed

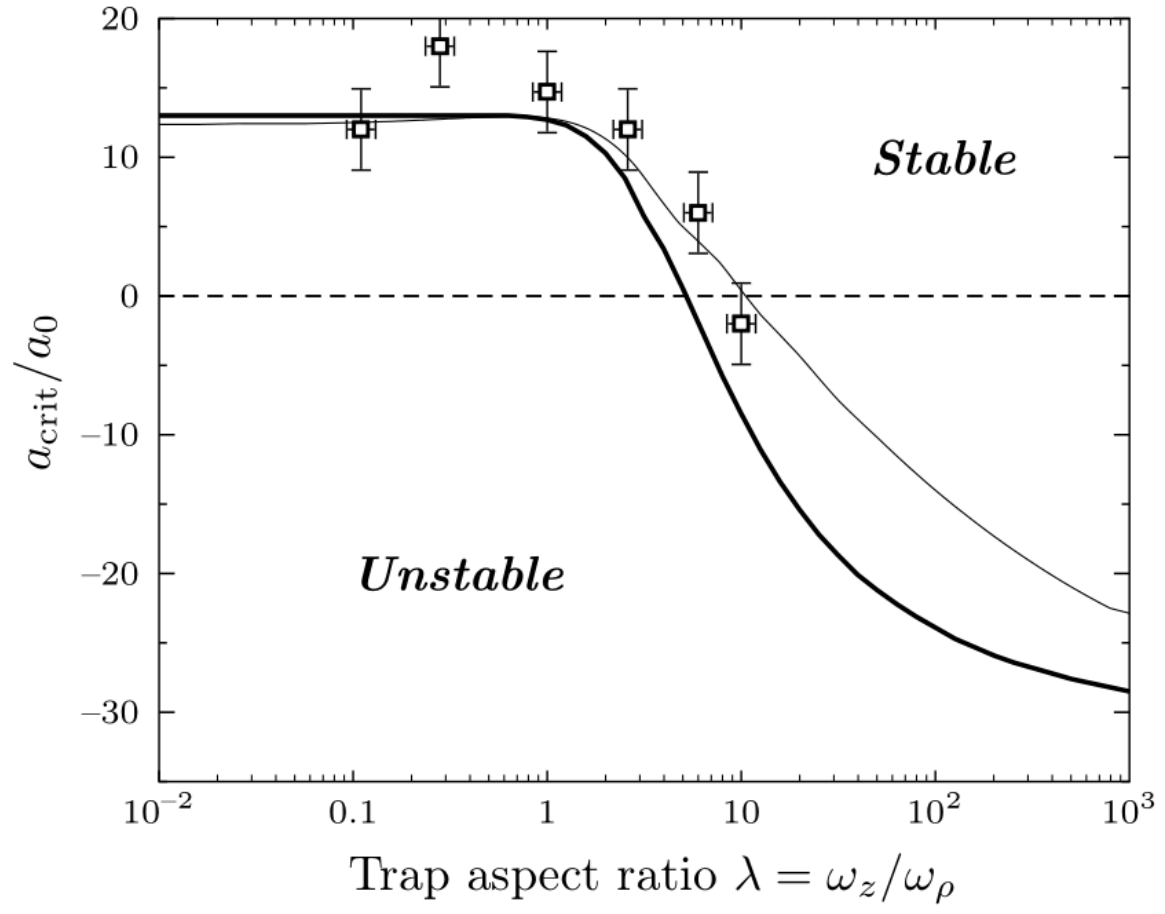


Figure 2: Logo
SANTOS, title (year)

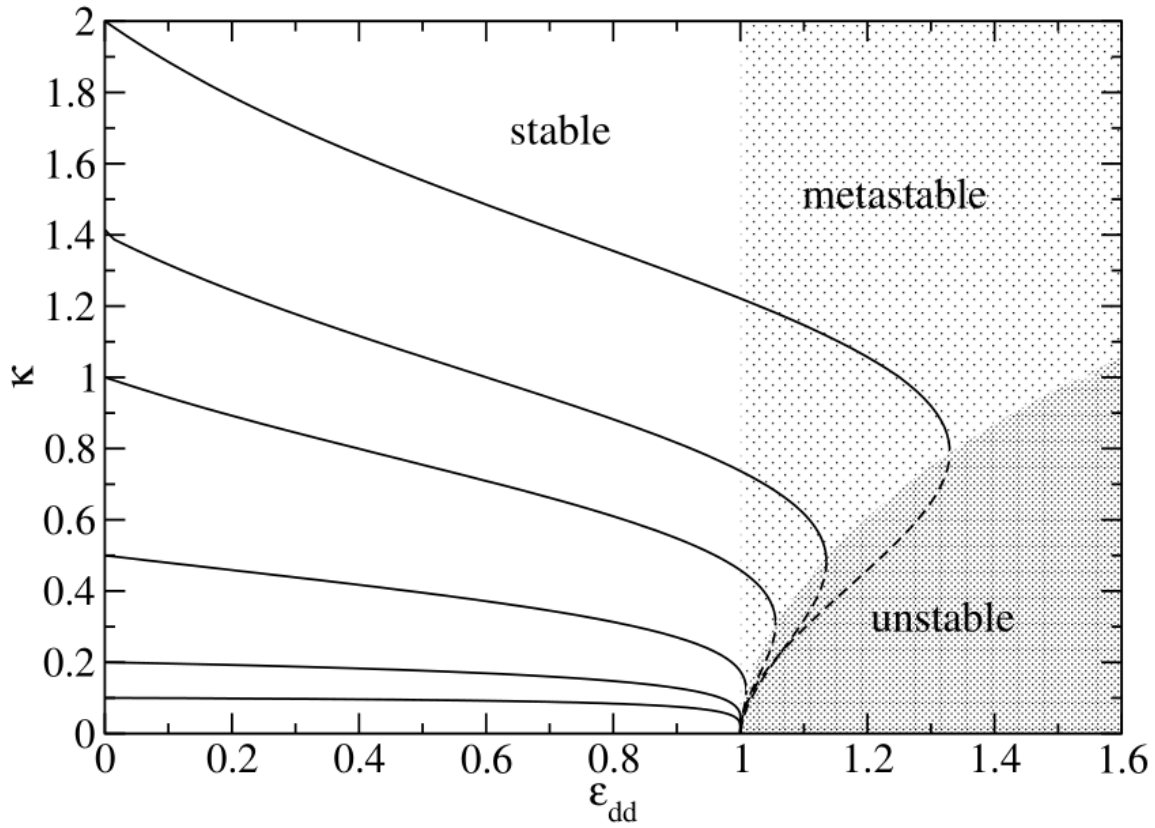


Figure 3: Logo
SANTOS, *title* (year)

- for sufficiently strong interactions, we may neglect quantum pressure, and consider the Thomas-Fermi (TF) regime
- TF solution for the trapped BEC has the same inverted parabola shape (as in non-dipolar case)
- BEC is prolat for $0 < \kappa < 1$ and $1 < \kappa$ oblat
- Bogoliubov-de Gennes Equation shows that the nonlocal character of the DDI causes a momentum dependent coupling constant, leading to a roton-like dispersion law, leading to dynamical instability, when the roton $\beta = \frac{g_d}{g}$ touches zero (experimentally not observed yet)
- by varying the density, the frequency of the confinement, and the short-range coupling, one can control the spectrum (roton minimum deeper/shallower)
- sequence of the non-local non-linearity 2D bright solitary waves may become stable under appropriate conditions (Pedri and Santos, 2005)

- two instability regions for 2D solitons (against collapse and against unlimited expansion)
- $\tilde{g}_{cr}(\beta) \equiv \frac{g_{cr}}{2\pi l_z}$, so stable 2D anisotropic self-localised solitons exists just for $N < N_{cr}$
- non-dipolar BECs scatter elastically, the scattering of dipolar solitons is inelastic due to the lack integrability
- The solitons may transfer centre-of-mass energy into internal vibrational modes, resulting in intriguing scattering properties:
 - including soliton fusion (Fig. 1.8)
 - appearance of strong inelastic resonances
 - possibility of observing 2D- soliton spiraling as that already observed in photo-refractive materials
- Dipolar effects in spinor condensates
 - spinor BECs: we focus on an effect which resembles the Einstein-de Haas effect
 - Because of Zeeman sub-levels short-range interactions may occur in different s-wave scattering channels with different total angular momentum (for bosons even number) (spin-1 bosons we have just $F = 0$ and $F = 2$)
 - Each scattering channel has an associated s-wave scattering length a_F
 - short-range interactions necessarily preserve the spin projection S_z
 - DDI does not necessarily conserve the spin projection along the quantisation axis as DDI is anisotropic
 - for initially maximally stretched state ($m_F = -F$)
 - short-range interactions cannot induce any spinor dynamics (due to conservation of total magnetisation S_z)
 - DDI may induce a transfer to $m_F + 1$
 - for cylindrical symmetry around the quantisation axis, this violation of the spin projection is accompanied by a transfer of angular momentum to the centre of mass, resembling the well known Einstein-de Haas effect \Rightarrow initially spin-polarised dipolar condensate can generate dynamically vorticity
 - Einstein-de Haas effect is destroyed by weak magnetic fields (1 mG)

- the dominant Larmor precession, and invoking rotating-wave-approximation arguments, the physics must be constrained to manifolds of preserved magnetisation (2D optical lattices could help)
- Effect of DDI could be even observable under conserved S_z (alkali spinor condensates)
- spin-changing collisions: collisions that conserve S_z , but do not conserve the relative population of the different Zeeman components
- Spin-changing collisions are characterised by an energy scale proportional to the difference between scattering lengths at different channels
- this difference is very small, so can be significantly modified by the presence of other small energy scales (DDI) \Rightarrow helical spin textures

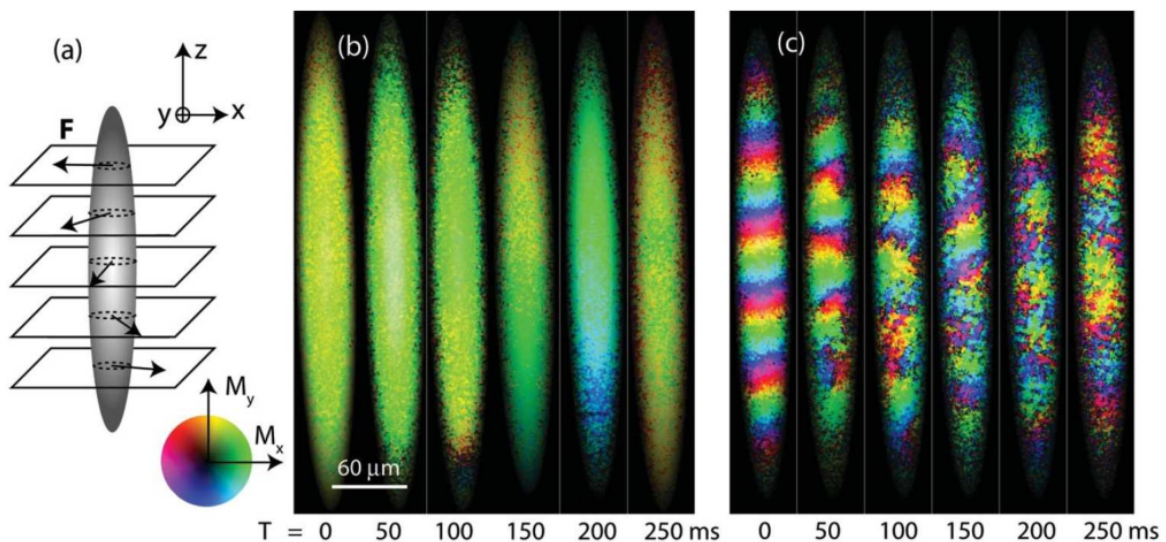


Figure 4: Is the occurrence of these textures special?

SANTOS, *title* (year)

2 Supersolids

- supersolid: features both the crystalline structure of a solid and the frictionless flow of a superfluid. In this state, every constituent atom is part of the solid and the superfluid simultaneously
- direct observation was limited to systems where the structure formation was mediated by external light fields

- beyond mean-field approximation leads to corrections to the ground state energy stemming from quantum fluctuations of the collective modes in a BEC (LHY-correction)
- In 2018 quantum droplets in a Bose-Bose mixture were observed
- Quantum droplets in Bose-Bose mixtures
 - mean- field energy depends on the difference of the two coupling constants $\delta(g) = |g_{rep}| - |g_{att}|$
 - LHY-correction depends on the individual coupling constants
 - For weakly attractive combination of interactions, a repulsive beyond mean-field correction can stabilize the BEC
 - after a peak density increasing the number of particles only leads to an increase in the size of the droplet
 - eGPE: kinetic energy, external trapping, and two-body interactions, LHY
 - beyond mean-field correction has only been calculated for a homogeneous system and can therefore only be included within a local-density approximation
 - QMC calculations in full many-body system verified the formation
 - intra-species scattering lengths a_{11} and a_{22} lead to different equilibrium densities $n_0^{(i)}$ for the two components of the mixture.
 - droplet forms an intrinsic imbalance in the atom numbers of the two components ($\frac{N_1}{N_2} = \sqrt{\frac{a_{22}}{a_{11}}}$)
 - larger density than in original BEC increases the rate of three-body loss \Rightarrow extra term in eGPE

$$i\hbar\partial_t\psi = \left[-\frac{\hbar^2\nabla^2}{2m} + V_{ext}(r) + \alpha n_0|\psi|^2 + \gamma n_0^{\frac{3}{2}}|\psi|^3 \right] \psi \quad (6)$$

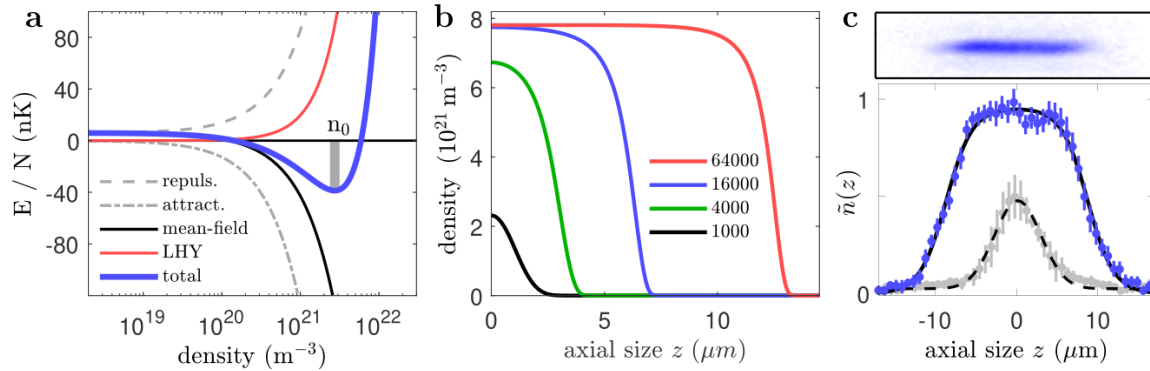


Figure 5: Peak density of the droplet saturates in z-direction
SANTOS, *title* (year)

- Dipolar Quantum droplets

$$i\hbar\partial_t\psi = \left[-\frac{\hbar^2\nabla^2}{2m} + V_{\text{ext}}(r) + g|\psi|^2 + \phi_{dd} + g_{qf}|\psi|^3 \right] \psi \quad (7)$$

- Liquid-like density saturation appears indicating very low compressibility
 - a droplet has $2.2 \cdot 10^4$ atoms
 - the dynamical state after 5ms of evolution is a Gaussian density distribution
 - most liquid-like aspect arises from the balance between attractive mean-field interaction and repulsive quantum fluctuations (self-bound nature)
 - BECs and degenerate Fermi gases gaseous (expand release from trap)
 - quantum droplets are self-bound ($N > N_{\text{crit}}$) caused by interplay between the binding mechanism and the kinetic energy cost of an inhomogeneous density distribution
 - kinetic energy leads to an increase E/N , which for small atom numbers can be strong enough to drive a liquid-to-gas transition
 - N_{crit} experimental sequence:
 1. prepare a quantum droplet with a high atom number
 2. external trapping potential switched off, and the unavoidable process of three-body decay leads to a rapid loss of atoms.
 3. crossing the phase boundary to the gaseous state, the droplet turns into a gas and rapidly expands
- ⇒ significant reduction in the density suppressing further losses

- ⇒ settled at critical atom number N_{crit} of a self-bound droplet (actual value depends on the precise strengths of the two interactions involved)
4. by varying the effective scattering length using Feshbach resonances, the phase diagrams can be mapped out
 5. residual confinement along the direction of gravity leads to a significant reduction of N_{crit}

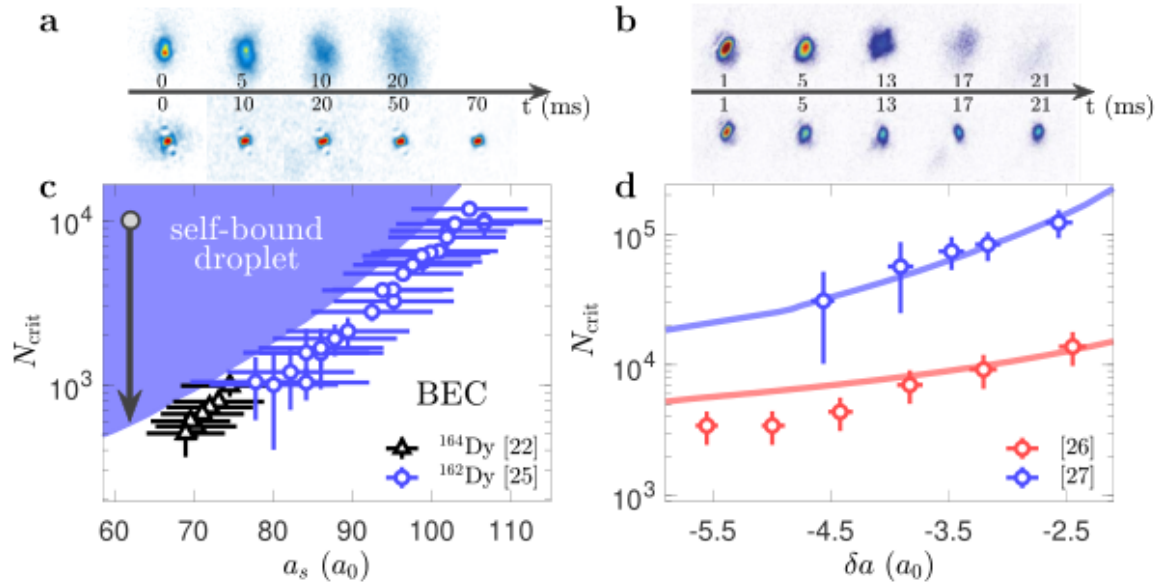


Figure 6: BEC and a self-bound quantum droplet in free space for a) the dipolar and (b) the Bose-Bose mixture.

In the droplet regime, imaging aberrations appear due to the high density and a spatial size of the droplets that is smaller than the experimental imaging resolution. Theoretical phase boundary between self-bound droplet and expanding BEC, together with the measured critical atom numbers for (c) the dipolar and (d) Bose-Bose droplets.

TILMAN PFAU ET AL., *New states of matter with fine-tuned interactions: quantum droplets and dipolar supersolids* (2020)

- For Bose-Bose mixtures excitations, where the components are in- or out-of-phase (to each other)
- collective modes would lead to exotic finite temperature behaviour of the binary droplets
- In 3D system there exists a regime where no collective modes are below the chemical potential, which corresponds to the particle emission threshold ⇒ excitation leads to self-evaporation of the droplet to zero temperature (just for Bose-Bose)

- out-of-phase modes are expected to be overdamped (too energetic to be bound)
 - No collective modes have been observed in Bose-Bose droplets (development of novel temperature probes needed)
 - quantum droplets are self-bound objects (time-of-flight expansion temperature measurements for cold atoms cannot be applied)
- In 1D quantum droplets the breathing mode always stays trapped (theoretically shown)
- dipolar case at least one collective mode is always expected to remain below the particle emission threshold
- In dipolar systems, measurements of collective excitations were observed
- an oscillation in the axial length of the droplet. Increase in the excitation energy upon approaching the transition (fits numerical simulations that include quantum fluctuations as the stabilization mechanism for the droplets)
- Scissors mode: breaking of the rotational symmetry causes an angular oscillation of the droplet around the polarizing magnetic field direction
- Scissor mode is an important marker of superfluidity (but moment of inertia of droplets does not differ significantly from the classical rigid due to the large anisotropy in their density distribution)
- scattering length comes from partial wave analysis, where one expands in the angular momentum components of the outgoing wave (similar to multipole expansion in Electrodynamics), it is related to the physically observable cross-section

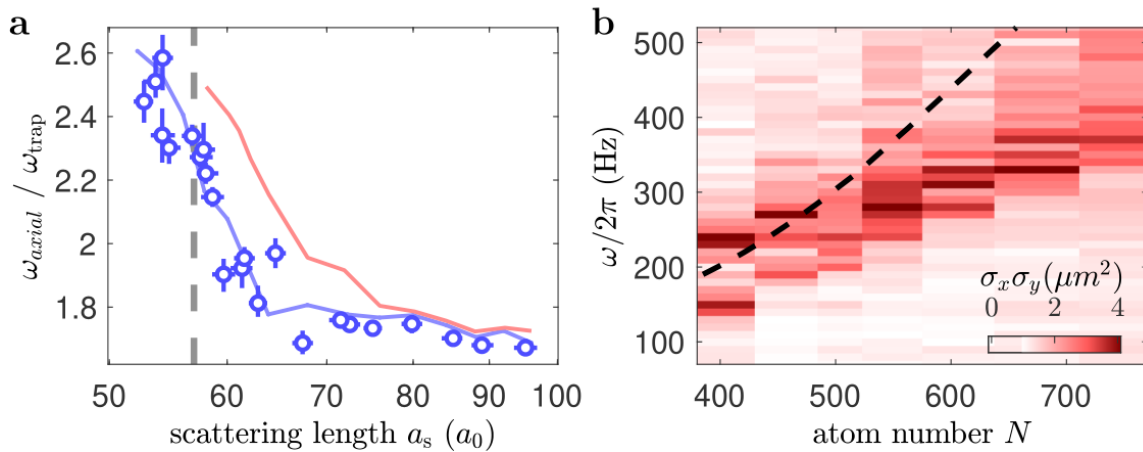


Figure 7: Measurements of the collective excitations of a dipolar quantum droplet.
a) With stabilizing contribution of quantum fluctuations (blue) and without (red).
b) Measurement of the scissors mode arising from the breaking of the rotational symmetry in dipolar quantum droplets. Experimentally, the orientation of the magnetic field is modulated at different frequencies and the response of the system is observed by looking at the droplet sizes $\sigma_{x,y}$. Theoretical scissors mode frequency expected from linear response theory (dashed).

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- solitons emerge as a single particle effect (dispersion in the waveguide stabilizes the system against collapse)
- quantum droplets are stabilized by many-body phenomenon (repulsive quantum fluctuations)
- solitons are only stable for effectively 1D gas (atom number $N < N_{crit_soliton}$)
- quantum droplets can exist as a purely self-bound state in 3D (atom number $N > N_{crit_droplet}$)
- Quantum droplets and solitons are distinct solutions of the eGPE

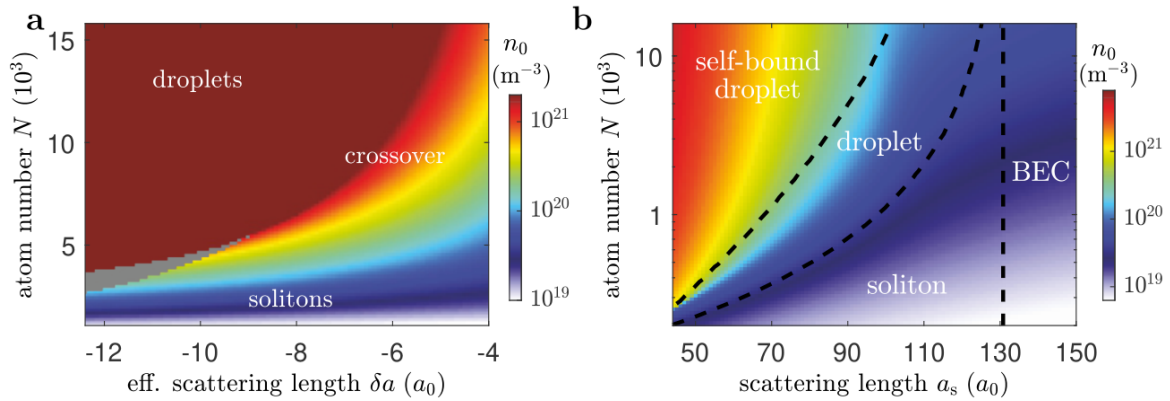


Figure 8: Simulated phase diagram of the ground-state peak density as a function of the atom number and the (effective) scattering length for
a) Bose-Bose mixture in a cigar-shaped confinement
b) dipolar quantum gas in a pancake-shaped confinement.
Solitons and droplets can coexist in a bistable region (gray) or can be smoothly connected by a crossover (dashed guide-line).

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- anisotropy of droplets leads to a frustration of the dipolar quantum droplet when compressed along the magnetic field direction and the emergence of arrays of dipolar quantum droplets
- multi-droplet states appear upon crossing into the bistable region, due to the fragmentation of the system following a modulational instability \Rightarrow multi-droplet states are not the ground state
- by confining a multi-droplet in a cigar-shaped trap it can be made the ground-state (confirmed experimentally)
- For large enough atom numbers, two-dimensional arrays of multiple quantum droplets eventually emerge as the ground state in cylindrically symmetric traps

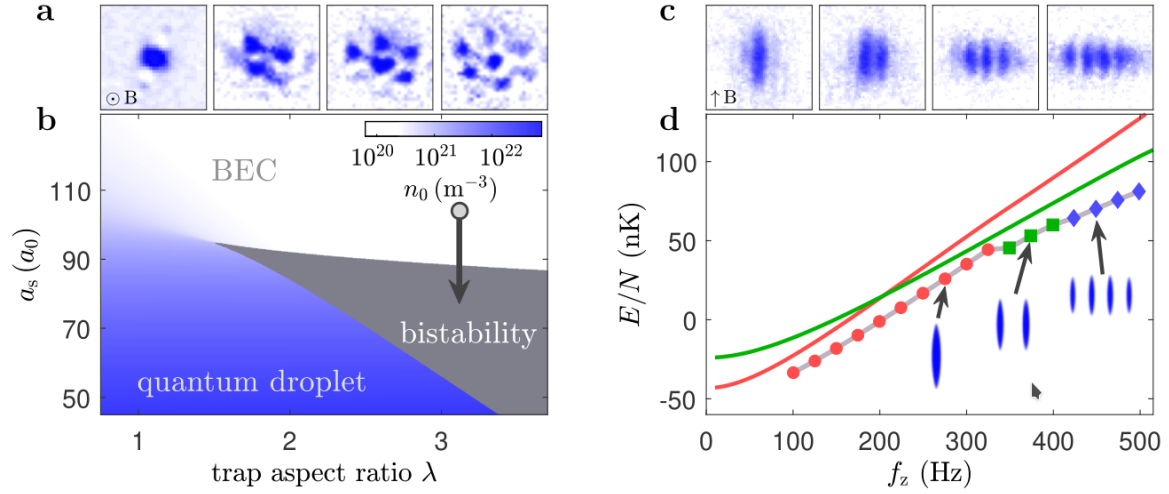


Figure 9: a) Example images of two-dimensional droplet arrays with increasing atom numbers in a cylindrically symmetric trap [19, 112]. (b) Phase diagram of an ensemble of 2.5×10^4 dysprosium atoms in a cylindrically symmetric trap with $\lambda = \frac{\omega_z}{\omega_r}$. b) In cigar-shaped traps, it is energetically favourable for the droplet number to increase with increasing confinement along the polarization direction. c) Examples for this behaviour upon changing the in-plane trap geometry. d) Total energy per particle E/N for a single droplet (red), two droplets (green) and four droplets (blue) in an external trap with $\omega_{\text{trap}} = 2\pi(70, 1000, f_z)$ Hz using a variational approach (lines) or numerical simulations of the eGPE (points).

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- collisions after loading a two-dimensional array into a one-dimensional waveguide with only weak confinement along one direction (Figure 10 a). The droplets initially repel each other due to the repulsive dipole-dipole interaction. However, the weak confinement along the waveguide axis forces the droplets to reverse their motion and collide with each other. Figure 10 b showing that the droplets bounce off each other twice. The oscillation of the droplets in the waveguide is strongly damped because of the relative motion between droplets and the residual background atoms, and inelastic collisions.
- separated BECs are typically realized with a double-well potential
- for collision the barrier needs to be removed
- Controlling the time at which the radial and vertical confinement is turned off, the velocity can be precisely controlled

- critical velocity v_c dividing the two outcomes of the collision (merging droplets and droplets bouncing off each other) depends on the mean atom number \tilde{N}_{coll} of the two droplets
 - $v_c(\tilde{N}_{coll})$ dependence is very different for small and large droplets
 - in a liquid-drop model the surface tension is the important energy scale for large droplets
 - small droplets there is no distinction between bulk and surface and as such the relevant energy scale is the binding energy
 - this can be seen as evidence of a crossover from a compressible quantum droplet at small atom numbers to a nearly incompressible droplet at large atom numbers

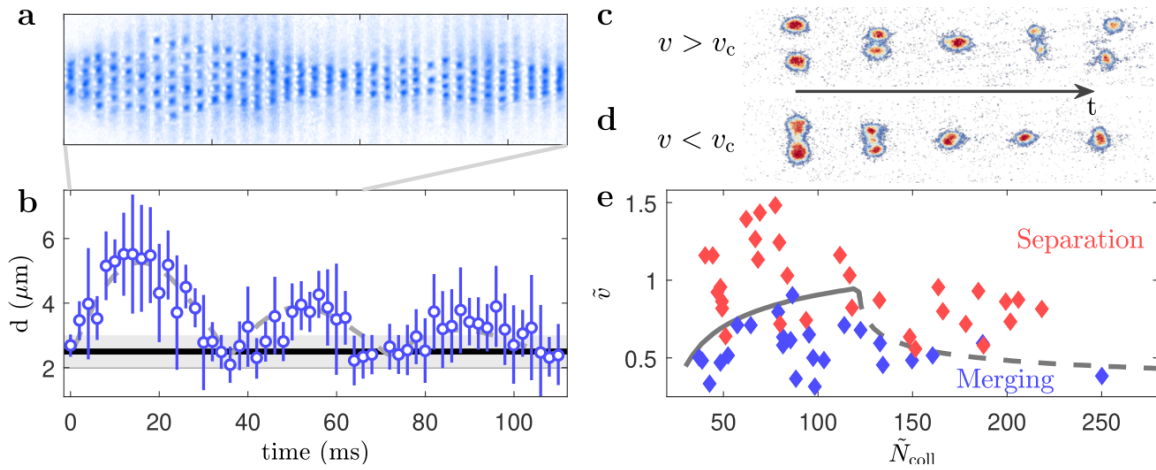


Figure 10: **Dipolar droplets:**

a) Single-shot images of colliding dipolar quantum droplets for different evolution times. b) Mean droplet distance d as a function of the evolution time together with a guide to the eye representing a damped bouncing motion at the trap frequency (dashed gray line).

Bose-Bose droplets:

(c), (d) Example images of collision measurements of independently created binary quantum droplets resulting in (c) a separation and (d) a merger of the droplets. (e) Outcomes of different droplet collision measurements as a function of the relative velocity \tilde{v} and the atom number \tilde{N}_{coll} at the time of the collision, with the blue (red) diamonds indicating a merger (separation) of the droplets.

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- self-organized droplet arrays break the continuous translational symmetry

- ⇒ supersolid state combines frictionless flow of a superfluid with the crystal-like periodic density modulation of a solid
- first experimental evidence for BECs coupled to external light fields, either exploiting cavity-mediated long-range interactions between the atoms or spin-orbit coupling
 - there two continuous symmetries are broken, the periodicity of the density modulation is set by the external light field, and therefore allowing no propagating phonon modes.
 - dipolar quantum gases can build droplet arrays in a self-organizing fashion and is based purely on the intrinsic interactions ⇒ propagating phonon modes allowed
 - to prove the coexistence of spatial order and superfluidity needs to be established
 - initial experiments states rapidly lost their global phase coherence
 - While each droplet itself is superfluid, the whole system is not
 - theoretically an infinitely extended system showed the coexistence of superfluidity and spatial order for a narrow range of the scattering length close to the phase transition
 - measurements showed indications of phase coherence in droplet arrays in a cigar-shaped trap geometry
 - experimental results undoubtedly proved the coexistence of spatial order and global phase coherence
 - in-situ density profiles in 3 regimes: isolated (incoherent arrays), phase-coherent droplet arrays, and a regular BEC
 - transition characterized by the strength of the density modulation Figure 11
 - interference of multiple quantum droplets, allowing the characterization of the nearest- and next-nearest neighbour coherence

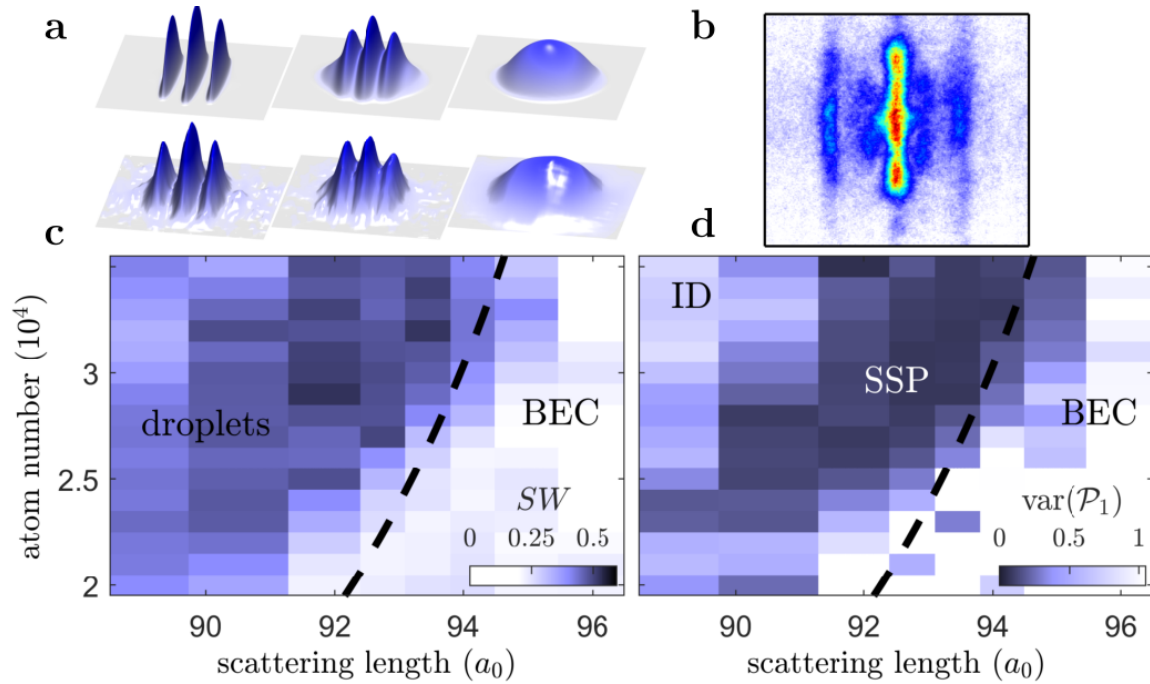


Figure 11: a) Comparison of the theoretically (top) and experimentally (bottom) observed density profiles of an array of isolated droplets (left side), an array of quantum droplets immersed in a condensate background (center) and a regular BEC (right side) [67]. (b) Example image of the multi-wave interference in [34], showing the principal and minor interference fringes at a different spacing. (c), (d) Experimental signatures of the phase diagram for (c) the in-situ density modulation and (d) the nearest-neighbour coherence. The strength of the density modulation in (c) is characterized by the spectral weight SW, which compares the contribution of Fourier amplitudes at finite momentum to the zero momentum contribution. (d) The Fourier transforms of the interference patterns reveal clear side peaks at the length scale corresponding to nearest- and next-nearest neighbours. Phase coherence leads to a well-reproducible interference pattern and thus a vanishing variance of the amplitude of these side peaks. The variance of the nearest-neighbour peak $\text{var}(\mathcal{P}_1)$ allows to differentiate between three regimes – isolated droplets (ID), phase-coherent droplets (supersolid phase, SSP) and a BEC. The black dashed line in (c) and (d) indicates the theoretical phase boundary obtained from numerical simulations of the eGPE. Adapted from [34, 67]. of a time-of-flight interference measurement is shown in Figure 7b.

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- to prove supersolidity (superfluid nature) elementary excitations needed

- The breaking of a continuous symmetry at the superfluid-supersolid phase transition fundamentally affects the spectrum of collective excitations
- ⇒ low-lying collective modes allow insights into the symmetry breaking and the supersolid nature
- energetically lowest mode is the dipole mode which is completely decoupled from interactions and always features an excitation energy corresponding to the trap frequency ω_x
- Upon decreasing the scattering length, a softening of the roton mode is observed
- roton mode is characterized by a minimum in the dispersion relation at a finite momentum and corresponds to a perturbative density modulation on top of the condensate density distribution
- In the considered trap geometry, the roton is comprised of two degenerate modes – a symmetric and an anti-symmetric roton mode. As these roton modes soften, avoided crossings are observed between pairs of modes with equal parity.
- softening of the roton triggers the phase transition to an array of quantum droplets (occurs in this finite system as a finite excitation energy of the roton modes)
- degeneracy of the even and odd roton modes is lifted with the emergence of a density modulation in the ground state
- At smaller scattering lengths the excitation energy of the symmetric mode rapidly increases, whereas the excitation energy of the anti-symmetric mode further decreases.
- symmetric mode features an oscillation between the array and condensate background (understood as Higgs amplitude excitation of the supersolid array)
- Close to the phase transition, the amplitude mode exists in an isolated state because of the energetic separation of the modes in the finite system
- The amplitude mode hybridizes with the other symmetric modes as its excitation energy increases.
- Close to the phase transition, collective modes with a larger excitation energy are affected by the rapidly increasing amplitude mode
- coupling of the higher symmetric modes with the amplitude mode leads to a bifurcation of the first quadrupole mode (after crossing the phase transition)

- higher-lying collective mode is scissors mode, decrease in the excitation energy upon crossing the phase transition (understood as a stiffening due to the emerging density modulation)

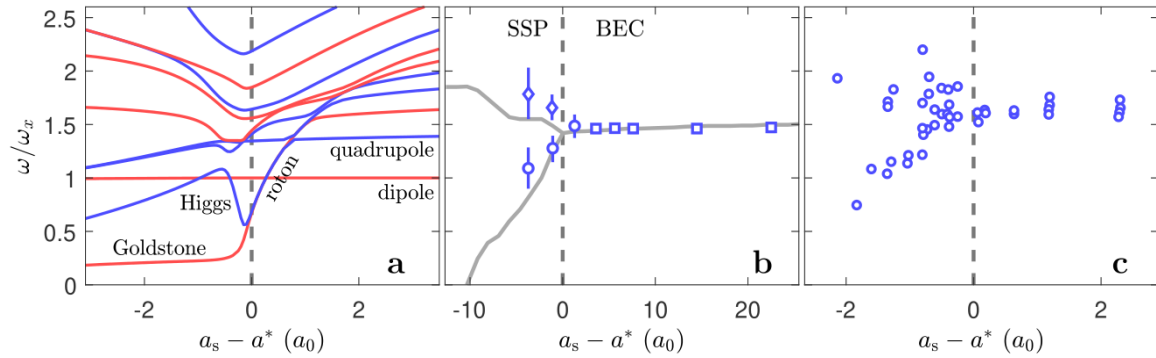


Figure 12: a) Calculated excitation frequencies ω_x of the lowest collective modes across the phase transition from a BEC to an array of quantum droplets in a cigar-shaped trap geometry with $\omega_{\text{trap}} = 2\pi[30, 90, 110]$ Hz. The blue (red) lines indicate an even (odd) parity of the density variation with respect to the weak trapping direction x . (b), (c) Signature of the phase transition in the excitation energy of the axial quadrupole mode measured with (b) ^{162}Dy and (c) ^{166}Er . The measurements were done using trap frequencies of $2\pi[19(2), 53(2), 81(2)]$ Hz and $2\pi[259(2), 30(1), 170(1)]$ Hz for the case of dysprosium and erbium, respectively.

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- most closely related to superfluidity is the low-energy Goldstone mode that emerges out of the anti-symmetric roton mode
- it features an out-of-phase oscillation between array and superfluid background, involving Josephson-like dynamics between droplets
- Goldstone theorem: connects existence of Goldstone mode to double symmetry breaking at the phase transition
- interplay between crystal motion and superfluid counterflow preserves the center of mass and leads to a linear correlation between the array displacement δx and the imbalance η between the side droplets
- finite lifetime of array prevents time-resolved measurement of the mode (excitation leaves a trace on the spatial density distributions)
- This trace can be statistically mapped out by repeating the experiment many times, leading to the observed correlation between the imbalance and the displacement

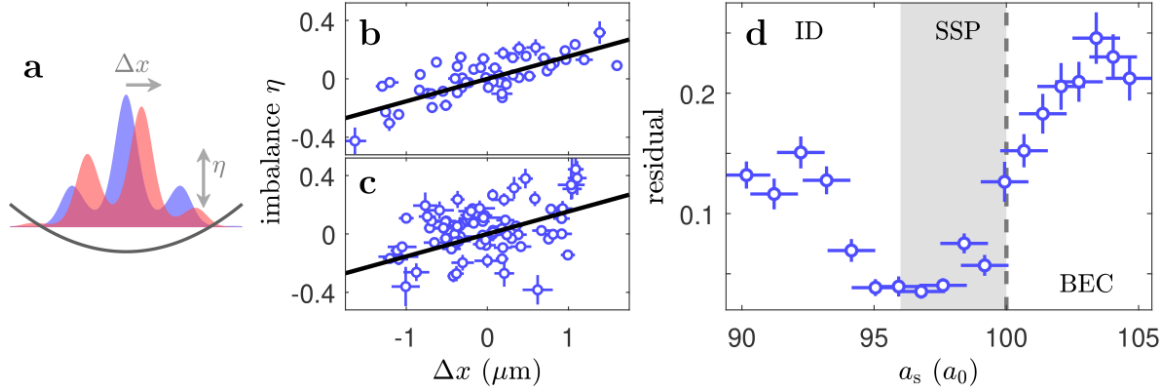


Figure 13: a) Schematic of the low-energy Goldstone mode, featuring an out-of-phase and center of mass preserving oscillation between the droplet array and the superfluid background. A displacement of the droplet array by δx leads to an imbalance η in the atom number of the side droplets. (b), (c) Measured imbalances η as a function of the measured displacements δx in (b) the supersolid region and (c) isolated droplet region. The black line shows the theoretically predicted correlation of the low-energy Goldstone mode. (d) The low residual of the experimental data with respect to the theoretical prediction close to the phase transition clearly proves the existence of the low-energy Goldstone mode. The gray area indicates the phase-coherent region.

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3 Outlook

- N_{crit} , roton softening in dipolar gases show discrepancies between theory and experiments
- mean-field of eGPE is limited to the perturbative regime at small gas parameters
- quantum depletion and quantum correlations start to play a role at the large quantum droplet densities
- beyond mean-field correction originating from quantum fluctuations is typically included in the description through a local density approximation
- validity may not always be given for:
 - long-range interaction
 - close to a phase transition

- dense systems
- large dipolar strengths ($\epsilon_{dd} > 1$)
- mean-field collapse leads to imaginary Bogoliubov modes, which are neglected (soft phonon mode for the Bose-Bose and roton mode for the dipolar system)
- Further improvement of theoretical description through:
 - diagrammatic Beliaev technique
 - behaviour in one-dimensional optical lattices
 - hypernetted-chain Euler-Lagrange method
 - Gaussian-state theory (includes squeezing effects)
 - bosonic pairing
 - inclusion of higher order corrections to the Bogoliubov speed of sound
 - quantum Monte-Carlo (intrinsically include particle correlations, quantum fluctuations, a finite system size, and a finite interaction range)
- these methods are limited to the usage of a simplified interaction potential because they cannot handle the bound molecular states in the complete interaction potential
- low-dimensional systems to circumvent these fundamental problems
- in 1D the energy functional of Bose-Bose mixtures does not suffer from the aforementioned imaginary Bogoliubov mode (quantum Monte-Carlo studies found small deviations from the predictions of the eGPE)
- measurements of N_{crit} and droplet density profile can be used as a sensitive benchmark for different theories in the near future
- in Bose-Bose self-evaporation to zero temperature has so far not been experimentally verified
- measurements of collective excitations are still lacking due to the finite lifetime in the experiments
- In dipolar droplets, the unavoidable presence of thermal fluctuations at finite temperatures has been proposed to increase N_{crit} of a self-bound droplet (large scattering lengths, large atom numbers)
- validity of the Born approximation for the dipolar scattering problem has been called into question, which would result in a temperature dependent systematic shift of N_{crit} to lower values

- Open questions:
 - How general is the emergence of a supersolid state with respect to variations in the experimental parameters?
 - How does finite temperature affect the supersolid phase transition?
 - Upon evaporating into the supersolid state, are both symmetries broken simultaneously or sequentially?
 - Do supersolid states exist in other systems, such as polar molecules ?
 - eGPE as a mean-field theory should also not be the correct description for arrays of isolated droplets with no wave function overlap \Rightarrow research needed for exact nature of supersolid-to-isolated-droplet-array transition

4 Simulations written in Python

4.1 First draft

- look into FFT artefacts
- GPE without Dipolar-Dipolar-Interactions

$$i\frac{\partial}{\partial t}\psi(x, t) = \left(\frac{-1}{2} \frac{\partial^2}{\partial x^2} + \frac{1}{2}x^2 + \tilde{g}|\psi(x, t)|^2 \right) \psi(x, t) \quad (8)$$

- dimensionless

$$\mu\phi_0(x) = \left(\frac{-1}{2} \frac{\partial^2}{\partial x^2} + \frac{1}{2}x^2 + \tilde{g}|\phi_0(x)|^2 \right) \phi_0(x) \quad (9)$$

- first use 1D later on 3D

4.2 Further ideas

- for split-operator more precise Trotter-Suzuki decomposition of time evolution operator $U(t) = \exp \frac{-iHt}{\hbar}$
- other method: Lanczos method (Krylov space)
- error $\propto E_{max} \Rightarrow$ spectrum centering to lower errors
- accuracy is not uniform for all eigenstates \Rightarrow Chebyshev method

- fft is implemented in scipy, sympy and numpy (research which one to use)

wavevector k . In the Hamiltonian shown above, we can split our system into position space components, $\hat{H}_r = [V(\mathbf{r}) + g|\Psi(\mathbf{r}, t)|^2] \Psi(\mathbf{r}, t)$, and momentum space components, $\hat{H}_k = [-\frac{\hbar^2}{2m} \nabla^2] \Psi(\mathbf{r}, t)$. I'll be honest, I didn't know what notation to use for \hat{H}_r because p is used to describe momentum. I settled on r for *real space*, but that is somewhat notationally ambiguous. In addition, k will indicate momentum space because it is a sum of all wavevectors, typically notated as k . Bad notation aside, let's continue.

If we assume a somewhat general solution to our quantum system:

$$\Psi(\mathbf{r}, t + dt) = \left[e^{-\frac{i\hat{H}dt}{\hbar}} \right] \Psi(\mathbf{r}, t) = \left[e^{-\frac{i(\hat{H}_r + \hat{H}_k)dt}{\hbar}} \right] \Psi(\mathbf{r}, t)$$

and assume we are simulating our system by a series of small timesteps (dt), we can perform similar splitting by using the Baker-Campbell-Housdorff formula:

$$\Psi(\mathbf{r}, t + dt) = \left[e^{-\frac{i\hat{H}_r dt}{\hbar}} e^{-\frac{i\hat{H}_k dt}{\hbar}} e^{-\frac{[i\hat{H}_r, i\hat{H}_k]dt^2}{2}} \right] \Psi(\mathbf{r}, t)$$

This accrues a small amount of error (dt^2) related to the commutation of the real and momentum-space components of the Hamiltonian. This is a relatively large error and that's not okay. In order to change the dt^2 error to dt^3 , we can split the system by performing a half-step in position space before doing a full-step in momentum space, through a process called *Strang Splitting* like so:

$$\Psi(\mathbf{r}, t + dt) = \left[e^{-\frac{i\hat{H}_r dt}{2\hbar}} e^{-\frac{i\hat{H}_k dt}{\hbar}} e^{-\frac{i\hat{H}_r dt}{2\hbar}} \right] \Psi(\mathbf{r}, t) + \mathcal{O}(dt^3)$$

We can then address each part of this solution in chunks, first in position space, then in momentum space, then in position space again by using [Fourier Transforms](#). Which looks something like this:

$$\Psi(\mathbf{r}, t + dt) = \left[\hat{U}_r \left(\frac{dt}{2} \right) \mathcal{F}^{-1} \left[\hat{U}_k(dt) \mathcal{F} \left[\hat{U}_r \left(\frac{dt}{2} \right) \Psi(\mathbf{r}, t) \right] \right] \right] + \mathcal{O}(dt^3)$$

where $\hat{U}_r = e^{-\frac{i\hat{H}_r dt}{\hbar}}$, $\hat{U}_k = e^{-\frac{i\hat{H}_k dt}{\hbar}}$, and \mathcal{F} and \mathcal{F}^{-1} indicate forward and inverse Fourier Transforms. Here's a flowchart

Figure 14: Logo

JAMES SCHLOSS, https://www.algorithm-archive.org/contents/split-operator_method/split-operator_method.html (year)

5 Goal

Solve:

$$i\hbar \frac{\partial}{\partial t} \psi(r, t) = \left(\frac{-\hbar^2}{2m} \nabla^2 + V(r) + g|\psi(x, t)|^2 \right) \psi(x, t) \quad (10)$$

6 Experiments

- If the initial wave function ψ_0 is chosen badly, the ground state is not found