We present our experimental setup to produce ultracold strongly correlated fermionic superfluids made of a two-component spin-mixture of $^6\text{Li}$ atoms. Employing standard cooling techniques, we achieve quantum degeneracy in a single-beam optical dipole trap. Our setup is capable of generating spin-balanced samples at temperatures as low as $T/T_F \approx 0.1$ containing up to $5 \times 10^4$ atomic pairs. We can access different superfluid regimes by tuning the interparticle interactions close to a broad magnetic Feshbach resonance. In particular, we are able to explore the crossover from the molecular Bose-Einstein condensate (BEC) to the Bardeen-Cooper-Schrieffer (BCS) superfluid regimes.

**Keywords:** Quantum degenerate gases; fermionic superfluidity; Bose-Einstein condensation; laser cooling and trapping.

**PACS:** 67.85.-d; 67.85.Lm; 67.25.D-  
DOI: https://doi.org/10.31349/RevMexFis.66.388

1. **Introduction**

Quantum gases are macroscopic quantum many-body systems that represent a unique scenario to study quantum phenomena such as superfluidity and macroscopic quantum excitations [1]. Moreover, ultracold atoms have emerged as ideal quantum simulators of many-body phenomena, becoming effective testbeds of quantum Hamiltonians. Indeed, the combination of ultracold atoms and optical potentials has opened up a new way of studying condensed matter problems with unprecedented clarity [2]. This is possible thanks to the high level of controllability that quantum gases offer. The dimensionality and geometry of the system can be precisely modified by tailoring trapping potentials with laser light and magnetic fields. The thermodynamic properties of the gas, such as density, temperature, and volume can be easily manipulated. Full control of interparticle interactions is possible via magnetic Feshbach resonances [3]. Even the quantum statistics of the system can be changed by choosing fermionic or bosonic atoms. These are very powerful tools that distinguish ultracold atomic gases from ordinary condensed matter systems.

At ultralow temperatures, diluted gases composed of alkali-metal atoms interact predominantly through the $s$-wave scattering channel, since at such low energies, higher order collision channels are highly suppressed. In the case of ultracold bosons, Bose-Einstein condensation is possible and superfluidity emerges as long as the $s$-wave scattering length $a_s$ has a non-vanishing value [1].

The case of ultracold identical fermions is strikingly different. In this case, $s$-wave scattering is also suppressed due to Pauli blocking, making these systems nearly non-interacting and, in consequence, they do not exhibit superfluid behavior even at zero temperature. The quantum degenerate state corresponds to an ideal Fermi gas, also known as Fermi sea [4].

However, it is possible to introduce interactions into the system by creating a two-component spin mixture since Pauli blocking occurs only between identical fermions, while atoms in different spin states still interact via $s$-wave scattering. The absolute value of $a_s$ determines the interaction strength and its sign defines whether the interaction is effectively repulsive ($a_s > 0$) or attractive ($a_s < 0$). In fermionic systems, the interaction strength is usually described by introducing the dimensionless interaction parameter $(k_F a_s)^{-1}$, where $k_F$ is the Fermi wave vector [5]. Additionally, the existence of magnetic Feshbach resonances allows controlling the value of the scattering length practically at will by applying a constant magnetic field into the system. Therefore, varying the value of this external field makes the creation of different interaction regimes possible, from weakly to strongly interacting systems, from a repulsive to an attractive gas [3].

A very important consequence of having such control on interatomic interactions is the possibility of creating different types of bound states among the atoms. For repulsive interactions, $(k_F a_s)^{-1} > 0$, a molecular bound state exists in the interaction potential. In this case, it is possible...
to associate diatomic molecules composed by two identical atoms which, consequently, will exhibit bosonic statistics making possible the emergence of Bose-Einstein condensation of tightly bound molecules [6–8]. On the other hand, for attractive interactions, \((k_F a_s)^{-1} < 0\), the corresponding bound state occurs in momentum space due to many-body correlations at the Fermi surface, giving rise to loosely bound Cooper-like pairs whose behavior is well described by the BCS theory [9–11]. In this way, the Feshbach resonance allows to continuously transit from the BEC to the BCS regimes through the so-called BEC-BCS crossover [12–14]. The intermediate regime in which the scattering length diverges, \((k_F a_s)^{-1} = 0\), known as unitary limit, is particularly intriguing because the system is strongly interacting and strongly correlated giving rise to interesting physics. Indeed, being a universal regime, physics across the BEC-BCS crossover is interesting due to its relationship with other important phenomena such as high-\(T_c\) superconductivity [15] and other strongly correlated superfluids such as neutron stars or quark-gluon plasma [16–18]. Figure 1 shows the Feshbach resonance for the case of the two lowest hyperfine Zeeman levels of \(^6\)Li, corresponding to the two Zeeman components of the absolute ground state \(2^2S_1/2 F = 1/2, m_F = -1/2\) and \(m_F = +1/2\), which we denote as \(|1\rangle\) and \(|2\rangle\) respectively. Figure 1 also specifies the different superfluid states for the different interaction regimes. This Feshbach resonance is particularly broad, having a width of the order of 300 G, enabling a very fine and precise control of the scattering length.

In this paper we describe our newly built setup to produce ultracold atomic gases composed by fermionic \(^6\)Li atoms in a balanced spin-mixture of the states \(|1\rangle\) and \(|2\rangle\). We employ a standard Zeeman slower to decelerate an atomic beam coming from an effusive oven. The decelerated atoms are trapped and further cooled down in a magneto-optical trap in which subsequently sub-Doppler cooling is used [19]. These laser cooled atoms are transferred into a single-beam optical dipole trap [19,20]. Finally, quantum degeneracy is achieved by runaway evaporation. We produce ultracold samples in the different superfluid regimes across the BEC-BCS crossover containing about \(5 \times 10^4\) pairs at an approximate temperature of \(T/T_F = 0.1\).

The article is divided as follows. In Sec. 2 we provide details of our experimental setup, this includes the ultra-high vacuum system, the laser system, the magnetic field generation system, and the creation of conservative potentials. Section 3 is devoted to the procedures used to cool down the gas to quantum degeneracy: laser cooling and evaporative cooling techniques, as well as details on the production of a superfluid sample in different interacting regimes. Finally, in Sec. 4, we present our conclusions and future perspectives.

## 2. Experimental Setup

### 2.1. The ultra-high vacuum system

Our ultra-high vacuum (UHV) system is divided in three main sections, namely (i) the effusive oven; (ii) the differential pumping stage, and (iii) the Zeeman slower system and the main chamber where the sample is produced and the experiments performed. Each of these sections is pumped by a 200 l/s pumping system composed by a combination of an ion pump and a non-evaporable getter (model NEXTor® D 200-5 from SAES getters Inc.). Figure 2 shows a scheme of our UHV, including a detailed cut of our main chamber.

The effusive oven consists of a cylindrical recipient which contains 5 gr of purified \(^6\)Li. The oven is heated to a temperature of 450°C, at this temperature the vapor pressure of lithium is about \(1 \times 10^{-4}\) Torr [21]. The oven is connected to the rest of the UHV system through a 4 mm diameter nozzle. The vapor produced in the oven passes through this nozzle generating an atomic beam that propagates through the rest of the system. We estimate that the atomic flux of the beam effusing out from the nozzle is of the order of \(6 \times 10^{15}\) atoms/s [22].

The pressure right after the nozzle reaches a value well above \(10^{-9}\) Torr, which is too high for producing quantum degenerate samples. To keep a sustained pressure difference between the oven and the region in which experiments are performed, we have placed a differential pumping stage which consists of two aligned tubes separated by 25 mm from each other: the first one, facing the oven, has a 4.6 mm inner diameter and a second one, facing the Zeeman slower, is 7.7 mm inner diameter. This scheme is designed to keep a pressure difference across the differential pumping stage up to five orders of magnitude. In this way, the pressure in the main chamber is of the order of \(10^{-11}\) Torr.

The main chamber is connected to the differential pumping stage by a 56 cm long and 16.5 mm inner diameter tube. Around this tube there is a conical solenoid which is used to create a spatially inhomogeneous magnetic field which is required to implement a Zeeman slower system (more details are provided in Sec. 2.3.1.).

Finally, the main chamber is a stainless steel custom-made octagon chamber from Kimball Physics Inc. This

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**Figure 1.** Feshbach resonance for the two lowest hyperfine Zeeman levels of \(^6\)Li. Different superfluid regimes are possible depending on the value of the scattering length \(a_s\).
The main chamber contains eight CF40 viewports on its sides; two CF100 vertical viewports, and ten CF16 viewports connected to the chamber by arms extruded from it at an angle of 13° from the horizontal plane. The Zeeman slower tube is connected to the main chamber by one of these arms while the Zeeman slower laser beam enters through another one diametrically opposed with respect to the center of the chamber. All viewports have anti-reflection coating for all the wavelengths used in our experiment (532 nm, 671 nm and 1064 nm).

We have placed on both CF100 flanges reentrant viewports of high optical quality whose inner face is very close to the atoms, at a distance of only half an inch. This opens the possibility of building a large numerical aperture optical system to produce high resolution images of the sample.

2.2. Laser system

2.2.1. Optical cooling scheme

To implement the different laser cooling techniques in our experiment we use both the D2 and D1 optical transitions of 6Li [23]. We use two separate diode lasers, one for each line. The emission frequency of these lasers is locked-in into an atomic reference using a spectroscopy cell containing 5 gr of purified 6Li heated at 320°C where we implement standard saturated absorption spectroscopy (SAS) [24]. We use the same cell to lock-in both lasers.

In the experiment, the D2 line is used first to implement the magneto-optical trap (MOT) and later, in the optical molasses cooling stage, while the D1 line is subsequently used to apply a sub-Doppler cooling stage. The natural linewidth of both lines is $\Gamma = 2\pi \times 5.87$ MHz [25]. The main optical frequencies employed in our experiment are shown in Fig. 3.

To produce the D2 frequencies we have an extended-cavity diode laser (cat-eye configuration, model CEL002 from MOGLabs) which pumps an optical tapered amplifier (model MOA002 also from MOGLabs). We divide the amplified beam into two beams and independently shift the frequency of each of them using two different acousto-optic modulators (AOM). Next, each one of these beams pumps another tapered amplifier and in this way we generate two high power beams (power of $\sim$500 mW each) at a wavelength of approximately 670.9 nm with a frequency difference between them of 228.2 MHz, which corresponds to the hyperfine splitting of the ground state $^2S_1/2$ of 6Li. One of these beams, the one with lower frequency, corresponds to the cooling frequency which is red-detuned from the $^2S_1/2(F = 3/2) \rightarrow ^2P_{3/2}$ transition by 8.5 $\Gamma$ (about 50 MHz). The second beam is used as repumper frequency and is red-detuned from the $^2S_1/2(F = 1/2) \rightarrow ^2P_{3/2}$ transition by 8.5 $\Gamma$. Note that we do not specify the hyperfine level of the excited state $^2P_{3/2}$ in either the cooling nor repumper frequencies. This is because the energy levels of these hyperfine states are too close together, their separation is less than $\Gamma$, and therefore we can not resolve them in our spectroscopy cell. We superimpose both beams using a 50:50 non-polarizing beam splitter which, hence, produces two beams with the same power, each one carrying both cooling and repumper frequencies. One of these beams is used to generate the light for the MOT. To do so, we subsequently divide it into three equally powered beams and couple each one into a polarization maintaining optical fiber which brings the light directly to the experimental region. The second beam coming from this 50:50 beam splitter is additionally red-shifted by 76 $\Gamma$ using an additional AOM. In this way we produce the Zeeman slower beam (which also arrives into the experiment by a polarization maintaining optical fiber). This large frequency shift is chosen to match the Doppler and Zeeman shifted optical fiber which brings the light directly to the experimental region. The second beam coming from this 50:50 beam splitter is additionally red-shifted by 76 $\Gamma$ using an additional AOM. In this way we produce the Zeeman slower beam (which also arrives into the experiment by a polarization maintaining optical fiber). This large frequency shift is chosen to match the Doppler and Zeeman shifted optical fiber which brings the light directly to the experimental region.

On the other hand, to implement the D1 sub-Doppler cooling stage, we require the two frequencies that are shown in the right side of Fig. 3. We have a cooling frequency blue-detuned from the transition $^2S_{1/2}(F = 3/2) \rightarrow ^2P_{1/2}(F' = 3/2)$ by 5 $\Gamma$ (about 30 MHz), and a repumper frequency blue-detuned from the transition $^2S_{1/2}(F = 1/2) \rightarrow ^2P_{1/2}(F' = 3/2)$ also by 5 $\Gamma$. Note that we never use both D2 and D1 lines at the same time and that the cooling and repumper frequencies of the D1 line are, evidently,
EXPERIMENTAL SETUP FOR THE PRODUCTION OF ULTRACOLD STRONGLY CORRELATED FERMIONIC SUPERFLUIDS OF $^6$Li

As can be seen, we essentially set all the required frequencies using three AOMs in double-pass configuration [26]. These AOMs are also used to dynamically change the frequency of these beams and implement the $D_2$ optical molasses and $D_1$ sub-Doppler cooling stages, as explained in Sec. 3.1.

2.2.2. Generation of probing light

The most important diagnostic tool in cold atoms experiments is imaging the samples using laser light. In our case the preferred technique is absorption imaging due to its simplicity and reliability [27, 28].

Absorption imaging consists in probing the sample using a collimated laser beam whose frequency is resonant to some atomic transition. To produce the image, we pulse this light on the atoms during a short time (of the order of 5 µs). The atoms will absorb some of the light, generating an absorption profile on the beam. Finally, the light is collected by a telescope that creates an image of such absorption profile on a CCD camera (model MANTA G-145 NIR from Allied Vision Technology GmbH). The density profile of the gas can be extracted from this image [27].

In our experiment we want to produce samples at different interaction regimes across the BEC-BCS crossover. This is done by applying an external magnetic field that changes the value of the scattering length by means of a Feshbach resonance. This magnetic field, in turn, will also cause a Zeeman splitting on the electronic levels of the atoms. Hence, probing

Figure 3. Level scheme (not to scale) for $^6$Li showing (left) the $D_2$ and (right) the $D_1$ hyperfine structures and the transitions used for the laser cooling processes. See text for details.

Figure 4. Simplified scheme of the laser cooling and imaging optical setup showing the main features of the system. Lenses and waveplates have been omitted for clarity. See text for details.
the atoms at different interaction regimes poses the necessity of generating different light frequencies to keep the imaging light resonant with the atoms.

To do so, we use the Zeeman slower beam which already has a considerable shift of $-76 \Gamma$. We divert a fraction of this beam using a polarizing beam splitter before it is coupled into the Zeeman slower optical fiber, as shown in Fig. 4. Next, this diverted beam passes through additional AOMs that will further shift the frequency to match it to the specific magnetic field in which we want to probe the atoms. This configuration of AOMs allows to tune the frequency of the probing light at different values within the range from 0 to $-220 \Gamma$ from the $D_2$ transitions. In this way, we are able to produce images at practically any magnetic field from 200 G to 1200 G and also at the vicinity of zero magnetic field. In this way, as can be seen in Fig. 1, we can image the sample in all the superfluid regimes across the BEC–BCS crossover, and also at weakly interacting regimes in which the gas is simply a Fermi degenerate gas but not a superfluid.

Finally, it is important to mention that the magnetic field used to access the BEC–BCS crossover is high enough to ensure that the hyperfine splitting of the atoms is well within the Paschen-Back regime, where the separation between the $|1\rangle$ and $|2\rangle$ states remains almost constant at approximately 76 MHz. For this reason, we can probe both spin states in any magnetic field through the Feshbach resonance.

2.3. Magnetic field generation system

We employ three different sets of coils to generate all the required magnetic fields to trap and manipulate the atoms. We describe each of them in the following sections.

2.3.1. Zeeman slower magnetic field

We use a Zeeman slower stage to decelerate the atomic beam coming out from the effusive oven. As mentioned in the previous section, the detuning of the Zeeman slower laser beam is $\delta_Z = -76 \Gamma$ for both cooling and repumper frequencies. The corresponding magnetic field along the direction of propagation of the atomic beam (direction $z$) is designed to decelerate atoms with velocities up to $v_0 = 960$ m/s at a constant deceleration $a \approx \hbar k / 2m$ [29] through the formula [19]:

$$B(z) = \frac{\hbar}{\mu_B} \left( \delta_Z + k \sqrt{v_0^2 - 2az} \right),$$  \hspace{1cm} (1)

where $\mu_B$ is the Bohr magneton and $k$ is the wavevector of the cooling frequency of the slower light. In this formula we only consider the cooling frequency since it is the one responsible for the deceleration of the atoms.

The mean velocity of the atoms coming out from the oven is $\bar{v} \approx 1540$ m/s, which is higher than the maximum velocity $v_0$ we can decelerate in our slower. This means that, in the best case, we can only slow down about 20% of the atoms. This is not a problem since the flux of atoms effusing from the oven is very large, about $6 \times 10^{15}$ atoms/s, so we can still efficiently load our magneto-optical trap.

The magnetic field of the slower is generated by a succession of eight size-decreasing coils connected in series and an extra ninth coil at the end of the slower in which the current circulates in opposite direction, inverting in this way the direction of the magnetic field. This is known as “spin-flip configuration” [30, 31]. All nine coils are wound directly onto the slower UHV tube using 1 mm diameter cooper wire. The coils are held together using a thermal conducting and electric insulating ceramic epoxy (Duralco™ 128). The total current passing through each coil is of the order of 2.0 A to generate a field which goes from a maximum around 600 G to a minimum of about $-250$ G.

Figure 5a) shows a scheme of the coil configuration of our Zeeman slower. Figure 5b) presents the generated magnetic
field. Finally, Figure 5c) exhibits the calculated velocity profile of the decelerated atoms through their propagation along the slower.

2.3.2. Magnetic quadrupole

To produce the magneto-optical trap we use a quadrupole magnetic field whose axial gradient at the center of the trap is \( \partial B_z(z)/\partial z|_{z=0} \approx 28 \text{ G/cm} \). This field is generated by two small coils of \( 6 \times 4 \) windings connected in anti-Helmholtz configuration. Each of these coils is mounted in a cylindrical water-cooled support to prevent them from heating. This support is made of Teflon\textsuperscript{TM}, which is a non-magnetic and insulating material, this prevents eddy currents from being induced in it when the quadrupole field is abruptly varied. Both supports are mounted inside the reentrant viewports of the main chamber, along the vertical direction. The coils are wound with strip-shaped copper wire of \( 4 \text{ mm} \times 1 \text{ mm} \) and held together with ceramic epoxy (Duralco\textsuperscript{TM} 128). Figure 2 shows the position of these coils in the main chamber.

2.3.3. Feshbach resonance magnetic field

As already mentioned, one of the important advantages of ultracold lithium gases is the possibility of controlling interatomic interactions with a high degree of precision by means of a Feshbach resonance [3]. \(^6\text{Li}\) presents several Feshbach resonances whose characteristics depend on the internal state of the interacting atomic pair. We will use the resonance between states \(| 1 \rangle \) and \(| 2 \rangle \), shown in Figure 1, centered at 832 G. So we need an extra set of coils that are able to produce an uniform magnetic field with any value from zero to 1000 G in order to have full control of all interaction regimes. To do so we have a pair of coils connected nearly at Helmholtz configuration. We deliberately move slightly away from the Helmholtz configuration so the magnetic field presents a small curvature, which will be useful to confine the atoms along the weak direction of our optical dipole trap; right at the Feshbach resonance, at 832 G, this curvature along the coils axis direction is \( B''_{z}(0) = 6.2 \text{ G/cm}^2 \), while the corresponding magnetic gradient is nearly zero (see Sec. 2.4 for more details).

The Feshbach coils are made by 4 mm square section copper wire. This wire is hollow, with an internal diameter of 2 mm, which allows cooling the coil by circulating cold water inside the wire. These coils were fabricated by the company Oswald Elektromotoren GmbH and each of them is embedded in a non-magnetic and insulating resin that avoids the induction of undesired eddy currents. We can circulate a current above 200 A without noticing any significant heating of the coils. This thermal stability together with a PID feedback loop makes possible to produce magnetic fields with a stability of one part in \( 10^4 \). We place these coils along the vertical direction, colinear to the quadrupole field coils. Figure 2 shows each of the employed set of coils and their position in the experimental setup.

2.4. Conservative trapping potential

We produce the quantum degenerate fermionic system in a conservative trap generated by the combination of an optical potential and a magnetic curvature.

The optical potential consists in a far red-detuned single-beam optical dipole trap (ODT) created by focusing a gaussian infrared laser beam [20]. We use a single mode ytterbium-doped fiber laser from IPG Photonics Corp. (model YLR-200-LP) which delivers up to 200 W of continuum linearly polarized infrared light at \( \lambda = 1070 \text{ nm} \). The beam of this laser is coupled into an acousto-optic modulator and the first diffracted order is used to produce the optical trap. We use a quartz crystal AOM that withstands very high intensities, above 1 GW/cm\(^2\), from the company Gooch & Housego (model I-M080-2C10G-4-AM3). The diffraction efficiency of this AOM and, consequently, the power of the ODT, is manipulated by controlling the amplitude of the RF signal that drives the modulator using an external analog signal. To stabilize the power of this diffracted order, we employ a PID circuit driven by the signal of a photodiode (Thorlabs, model DET36A) that detects the small fraction of the light of this beam that is transmitted by a 99.9% reflection mirror.

Next, we collimate the beam at a diameter of approximately \( D = 5.5 \text{ mm} \) and finally use a f = 40 cm focal length lens to focus the light on the atoms. The beam waist at focus is \( w_0 = 2 \lambda f / \pi D \approx 50 \text{ µm} \), which corresponds to a Rayleigh length of \( z_R = \pi w_0^2 / \lambda = 7.34 \text{ mm} \).

The trap frequencies of this single beam ODT along the radial and axial directions are given, respectively, by [20]

\[
\omega_{rODT} = \sqrt{\frac{4U_0}{m w_0^2}} \quad \text{and} \quad \omega_{zODT} = \sqrt{\frac{2U_0}{m z_R^2}},
\]

where \( U_0 \) is the depth of the trapping potential and it is given by [20]

\[
U_0 = \left( \frac{3\pi e^2 \Gamma}{2 \omega_0^2 \Delta \omega} \right) I_0,
\]

where \( \omega_0 \) is the frequency of the lowest energy optical dipole transition of the \(^6\text{Li}\) atom, which corresponds to the \( D_1 \) transition, \( \Gamma \) is the natural linewidth of such transition and \( \Delta \omega = \omega_0 - \omega \) is the detuning between such transition and the ODT laser frequency \( \omega \). Finally, \( I_0 \) is the intensity of the ODT beam at focus, \( I_0 = 2 P / \pi w_0^2 \), where \( P \) is the power of the ODT laser.

This trap provides a tight confinement along the radial direction of the beam, however, along the axial (or propagation) direction it is very weak. For instance, at the end of the evaporative cooling where the power of the ODT laser is approximately \( P = 35 \text{ mW} \) (see Sec. 3.2.2), the radial and axial frequencies of the optical trap respectively are \( \omega_r = 2 \pi \times 163 \text{ Hz} \) and \( \omega_z = 2 \pi \times 0.94 \text{ Hz} \), which would provide an extremely elongated sample with an aspect ratio larger than 1:160.
For this reason, we add to the optical potential a magnetic curvature that provides a better confinement along the axial direction. As mentioned in Sec. 2.3.3, we produce such curvature by setting the Feshbach coils slightly off the Helmholtz configuration. In this way, we create a saddle-point magnetic potential of the form [28, 32]

\[ U_{mag}(r, z) = \frac{m}{2} \left( \omega_{mag}^2 z^2 - \omega_{mag}^2 r^2 \right), \tag{4} \]

where the trap frequencies are determined by the curvature of the field component along the corresponding direction, i.e., \( \omega_{mag}^2 = \mu B''(0)/m \) and \( \omega_{mag}^2 = \mu B''(0)/m \), being \( m \) the mass of \(^6\)Li atom and \( \mu \) the magnetic moment of the trapped state which, in general for the ground state of alkali atoms, is of the order of the Bohr magneton, \( \mu \sim \mu_B \).

Note from Eq. (4) that along the radial direction we have an “anti-curvature” which will tend to deconfine the atoms along that direction. The total frequencies of our hybrid trap will be given simply by

\[ \omega_r = \sqrt{\omega_{ODT}^2 - \omega_{mag}^2} \quad \text{and} \quad \omega_z = \sqrt{\omega_{ODT}^2 + \omega_{mag}^2}. \tag{5} \]

In our experiment, once the quantum sample is produced, we have that the radial optical confinement is much larger than the magnetic one \( (\omega_r \approx \omega_{ODT}) \), and, vice versa, along the axial direction the confinement is dominated by the magnetic component \( (\omega_z \approx \omega_{mag}) \).

The axial curvature generated for the fields used to access the BEC-BCS crossover is of the order of \( B''(0) = 6.2 \text{ G/cm}^2 \) which, superimposed to the ODT potential, translates into a total axial frequency of \( \omega_z \approx 2\pi \times 11 \text{ Hz} \). In this way we obtain a cigar-shaped quantum sample whose aspect ratio, of the order of 1:15, is appropriate for our goals.

3. Methods and results

In the following sections we provide details on the experimental procedures employed to produce the ultracold samples.

In a very general way, the production of the quantum sample can be divided into two main processes: (i) an initial laser cooling stage mediated by absorption and reemission of light, explained in Sec. 3.1, and (ii) transference into a conservative potential and cooling by forced evaporation, presented in Sec. 3.2

3.1. Implementation of laser cooling technique

In this first cooling process we are able to produce atomic samples at temperatures as low as 40 \( \mu \)K containing \( 4.5 \times 10^8 \) atoms with a density of the order of \( 4.5 \times 10^9 \) atoms/cm\(^3\), which correspond to a phase-space density of about \( 6.6 \times 10^{-6} \). We provide details on the laser cooling procedure in the following sections.

3.1.1. Zeeman slower and magneto-optical trapping

Zeeman slower operation

The quantum sample generation process starts by heating the lithium sample contained in the oven of our UHV system to 450°C. This generates a high temperature atomic beam that propagates through the UHV system towards the main chamber. The atoms of this beam undergo a first cooling process as they are decelerated by our Zeeman slower. Along the slower, a laser beam propagates in the opposite direction to the atomic beam. This laser carries two different frequencies, both of them red-detuned by 76 GHz (~446 MHz) from the cooling and repumper transitions of the \( D_2 \) line. Each of these frequencies has a power of 40 mW and carries positive circular polarization \( \sigma^+ \). In this way, we are able to decelerate all the atoms from velocities classes under 960 m/s to speeds of the order of 40 m/s, well below the 60 m/s capture velocity of the MOT, as shown in Fig. 5c).

We found that controlling independently the electric current of the spin-flip coil provides better results. Best results are obtained using a current of 2.0 A for the spin-flip coil and 2.9 A for all other coils, which optimize the number of loaded atoms into the MOT and minimize the corresponding loading time.

Loading of the magneto-optical trap: The decelerated atoms arrive into the main chamber where we capture them and further cool them in a magneto-optical trap (MOT) [19]. To implement the MOT we use three retroreflected mutually perpendicular laser beams with a diameter of \( D = 2.3 \text{ cm} \), as shown in Fig. 6.

The MOT beams carry two frequencies: a cooling frequency, red-detuned from the \( ^2S_{1/2}(F = 3/2) \rightarrow ^2P_{3/2} \) transition, and a repumper frequency, red-detuned from the \( ^2S_{1/2}(F = 1/2) \rightarrow ^2P_{3/2} \) transition. We use the standard \( \sigma^+ / \sigma^- \) polarization configuration. We determine the value

![Figure 6](image-url)
In order to further cool down the sample and increase its phase space density, the gas undergoes two different additional laser cooling processes. We first apply an optical molasses cooling process based on the $D_2$ laser line that allows approaching the Doppler limit temperature [19]. Next, we implement a gray-molasses technique, employing the $D_1$ line transitions to reach sub-Doppler temperatures [33–35]. We provide details in the two following sections.

**3.1.2. Doppler and sub-Doppler cooling**

In order to further cool down the sample and increase its phase space density, the gas undergoes two different additional laser cooling processes. We first apply an optical molasses cooling process based on the $D_2$ laser line that allows approaching the Doppler limit temperature [19]. Next, we implement a gray-molasses technique, employing the $D_1$ line transitions to reach sub-Doppler temperatures [33–35]. We provide details in the two following sections.

**D_2 optical molasses cooling:** The theoretical Doppler temperature limit for our sample is given by $T_D = \hbar \Gamma / 2 k_B = 140.9 \mu K$. To reach this limit it is necessary to lower the intensity of the MOT light to minimize light-scattering heating, so the MOT light intensity should be much lower than the saturation intensity $I_D = 2.54 \text{ mW/cm}^2$. Also, the cooling light must be detuned near to resonance, having an optimal value at $\delta_{\text{cool}} = -\Gamma / 2$. The process needs to be done in absence of any magnetic field.

After loading the MOT we abruptly switch off the quadrupole magnetic field (we also switch off the Zeeman slower magnetic field 400 ms before to guarantee the absence of any magnetic field in the sample region). Simultaneously, we decrease the intensity of the MOT beams and shift the value of cooling and repumper frequencies towards resonance. Figure 8a) shows the effect on $N$ and $T$ of the intensity reduction, while Figs. 8b) and (c) present the corresponding effect of the frequency shift of both MOT frequencies.

As we can see, an important temperature drop is observed when the intensity of the light decreases. Concerning the frequency shift, as long as we keep the detuning below $-2 \Gamma$, the number of atoms remains approximately constant while temperature decreases. We determine that the best values for intensity are $I_{\text{cool}} = 0.35 I_D$ for cooling light and $I_{\text{rep}} = 0.3 I_D$ for repumper, while the optimal frequency detuning is $\delta_{\text{cool}} = \delta_{\text{rep}} = -\Gamma$. We also found that the optimal duration of this molasses process is 850 $\mu$s; if shorter, the temperature does not reach the minimum possible value, and if longer we start losing atoms.

Under these conditions, we are able to cool down about $6 \times 10^8$ atoms to a temperature of about $500 \mu K$. The dashed black curve in Fig. 8b) shows the theoretical Doppler limit, compared to which our experimental points lie above for the entire range of the detuning of cooling light considered. In other elements, such as as rubidium or cesium, it is observed not only that the Doppler limit is reached but even sub-Doppler temperatures are attained due to the emergence of the Sisyphus sub-Doppler cooling mechanism [36]. For lithium this molasses scheme is not very efficient because the hyperfine levels of the state $2^3P_{3/2}$ cannot be well resolved, since their separation is smaller than $\Gamma$. This limits the efficiency of the cooling process and keeps the sample well above the Doppler limit. The increase of the phase space density is also not very good, and we improve only by a factor of 2, being of the order of PSD $= 1 \times 10^{-7}$. For this reason, we apply a second
In other words, we have a velocity selective process that protects the slowest atoms from light-assisted heating.

On the other hand, the $D_1$ light gets to the atoms through the same optical fibers used to produce the MOT (see Sec. 2.2), and hence they generate a 3D polarization gradient. This allows a Sisyphus-like cooling scheme between bright and dark states which decreases the momentum of the atoms. In this way, while the Sisyphus cooling mechanism decreases the momentum of the atoms of the gas, the VSCPT process accumulates the slowest atoms in a dark state. This significantly decreases the temperature of the sample.

In our experiment, we implement this cooling stage immediately after the $D_2$ molasses stage. We specifically use the $D_1$ transition frequencies $2^2S_{1/2}(F = 3/2) \rightarrow 2^2P_{1/2}(F' = 3/2)$, which we call “cooling” frequency, and $2^2S_{1/2}(F = 1/2) \rightarrow 2^2P_{1/2}(F' = 3/2)$, which we call “repumper”. This nomenclature is inherited from the standard molasses. Both frequencies are blue detuned, the cooling frequency by $\delta_1$ and the repumper light by $\delta_2$. Another important parameter is the difference between these detunings that we define as $\delta = \delta_1 - \delta_2$.

To characterize the gray-molasses we start by fixing $\delta_1 = +5.7\Gamma$ and keeping the repumper intensity low, at about $I_{rep} = 0.06I_{D1}^c$, and the cooling intensity at its maximum value of the order of $I_{cool} \approx I_{D1}^c$. The saturation intensity for the $D_1$ line is $I_{D1}^c = 7.59\text{ mW/cm}^2$. We then measure the number of atoms and the temperature of the sample as the detuning difference $\delta$ varies. The results are shown in Fig. 9.

We can see that the temperature follows a Fano-like profile, reaching a minimum at $\delta = 0$ (i.e. at $\delta_2 = \delta_1$), the so-called Raman condition, in which the temperature is as low as 40 $\mu$K. Although the number of atoms does not reach its maximum at the Raman condition but at $\delta \approx -0.25\Gamma$, we still have a very good efficiency of the process at $\delta = 0$, being able to cool about 75% of the atoms. These results are expected, as previously reported for the case of $^6$Li [35], and other atomic species such as $^{40}$K [33] and $^7$Li [34]. Notice that the plot of Fig. 9 has no data points in the interval $0.4 < \delta < 0.8$, as explained in Ref. 35, in this range the laser cooling technique that uses the $D_1$ line transitions, known as gray molasses, that allows true sub-Doppler cooling [34, 35].

**$D_1$ gray molasses sub-Doppler cooling:** Gray-molasses cooling is a two-photon process in $\Lambda$ configuration (see Fig. 3) which combines both, Sisyphus cooling [36] and Velocity Selective Coherent Population Trapping (VSCPT) [37] as cooling mechanisms. More details can be seen in references [35, 38]. In few words, the cooling process occurs in the following way. On the one hand, the $\Lambda$ scheme creates two coherent states, a so-called “bright state” that interacts with the light fields and a “dark state” which doesn’t. The transition probability from the dark to the bright state depends on the square of the momentum of the atoms, having as a consequence that the slowest atoms accumulate in the dark state.

**Figure 8.** Number of atoms $N$ (red dots) and temperature $T$ (black triangles) of the atoms of the MOT after the $D_2$ optical molasses as a function of a) the intensity of the cooling light and the detuning of b) the cooling light and c) the repumper light. The dashed black curve in b) corresponds to the theoretical Doppler limit for the temperature of our sample. In these plots, the error bars correspond to one standard deviation of ten independent measurements.

**Figure 9.** Number of atoms (red dots) and temperature (black triangles) of the sample as a function of the detuning between cooling and repumper light during gray molasses sub-Doppler cooling stage. The error bars correspond to one standard deviation of ten independent measurements.

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the energy of the dark state becomes larger than the energy of
the bright state and in consequence the VSCPT process sig-
nificantly heats the cloud. In this range, the temperature be-
tains for a duration time of 1 ms.

For the next stages, it is important to have all the atoms
loaded into the trap because at such high power the confinement provided
by the optical trap is much higher, thus excluding any loss of con-
finement. For these stages we ramp the power of the optical beam has reached
its maximum value, we ramp the Feshbach magnetic field to
its 40% value, and we ramp the ODT laser beam to 160 W in 7 ms. The beam
is focused right at the center of the atomic cloud, as shown in
Fig. 6. Once the power of the optical beam has reached
its maximum value, we ramp the Feshbach magnetic field to
832 G in 50 ms. This field corresponds to the unitary limit in
which the scattering length diverges, which is optimal for the
following evaporative cooling stage because the collision rate
is maximized and the thermalization process is optimized.

When the magnetic field is ramped up, the
F = 1/2 hyperfine state splits into the two states |1⟩ and |2⟩, where
|1⟩ has lowest energy for all magnetic fields. In the mag-
netic fields that we employ these states are well within the
Paschen-Back regime, so the energy difference between them
remains almost unchanged. Moreover, if the ramp of the
magnetic field is adiabatic, both states are nearly equally pop-
ulated, so we create a well balanced mixture.

The Feshbach field curvature provides an axial harmonic
confinement of about \( \omega_{mag} = 2\pi \times 11 \) Hz. This confine-
ment, of course, is negligible at the beginning of the ODT
loading since at such high power the confinement provided
by the optical trap is much higher, \( \omega_{ODT} \simeq 2\pi \times 10 \) kHz
and \( \omega_{ODT} \simeq 2\pi \times 87 \) Hz (see Eq. (5)), however, the mag-
netic confinement becomes more and more important as we
apply the evaporative cooling process in which the power of
the ODT laser beam is gradually decreased.

After the optical and magnetic fields have been ramped
up, we trap about \( 3 \times 10^6 \) atoms in the conservative poten-
tial, which means that our trapping efficiency is of the order
of 1%. We hold the atoms in this trap for 20 ms to let them
settle in the minimum of the potential. At this point we can

### Table I. Optimized parameters of the optical cooling stages.

| Cooling stage | Parameter | Optimal Value |
|---------------|-----------|---------------|
| MOT           | \( \partial B_z(z)/\partial z \mid_{z=0} \) | 28 G/cm |
|               | \( \delta_{cool} \) | -8.6 \( \Gamma \) |
|               | \( \delta_{rep} \) | -8.4 \( \Gamma \) |
|               | Loading time | 8.6 s |
| N             | 5 \times 10^9 atoms |
| T             | 7 mK |
| PSD           | 4.7 \times 10^{-8} |
| D_2 Molasses  | Duration | 850 \mu s |
|               | N         | 6 \times 10^8 atoms |
|               | T         | 500 \mu K |
|               | PSD       | 1 \times 10^{-7} |
| D_1 Gray molasses | Duration | 1 ms |
|               | N         | 4.5 \times 10^8 atoms |
|               | T         | 40 \mu K |
|               | PSD       | 6.6 \times 10^{-6} |

light, so we manage to concentrate nearly 95% of the atoms
in the \( F = 1/2 \) hyperfine level.

To summarize, after the whole laser cooling process, we
are able to produce a sample containing about \( 4.5 \times 10^8 \) atoms
in the hyperfine \( F = 1/2 \) state at a temperature of
40 \( \mu \)K. The phase space density increased considerably to
PSD \( \simeq 6.6 \times 10^{-6} \). This represents an excellent starting point
for the subsequent cooling stages.

Table I presents the list of all the parameters employed in
the laser cooling process.

### 3.2. Cooling the sample to quantum degeneracy

After the \( D_2 \) and \( D_1 \) cooling stages, the sample is ready to be
transferred into a conservative potential in which evaporative
cooling can be applied and quantum degeneracy is achieved.
In the following sections we explain how this process is done
in our setup.

#### 3.2.1. Transference into the conservative trap

As explained in Sec. 2.4, our trap is created as the com-
position of a single-beam optical dipole trap and a magnetic
curvature, which provide, respectively, radial and axial con-
finement.

During the \( D_1 \) cooling process we ramp the power
of the optical dipole trap (ODT) to 160 W in 7 ms. The beam
is focused right at the center of the atomic cloud, as shown in
Fig. 6. Once the power of the optical beam has reached
its maximum value, we ramp the Feshbach magnetic field to
832 G in 50 ms. This field corresponds to the unitary limit in
which the scattering length diverges, which is optimal for the
following evaporative cooling stage because the collision rate
is maximized and the thermalization process is optimized.

The Feshbach field curvature provides an axial harmonic
confinement of about \( \omega_{mag} = 2\pi \times 11 \) Hz. This confine-
ment, of course, is negligible at the beginning of the ODT
loading since at such high power the confinement provided
by the optical trap is much higher, \( \omega_{ODT} \simeq 2\pi \times 10 \) kHz
and \( \omega_{ODT} \simeq 2\pi \times 87 \) Hz (see Eq. (5)), however, the mag-
netic confinement becomes more and more important as we
apply the evaporative cooling process in which the power of
the ODT laser beam is gradually decreased.

After the optical and magnetic fields have been ramped
up, we trap about \( 3 \times 10^6 \) atoms in the conservative poten-
tial, which means that our trapping efficiency is of the order
of 1%. We hold the atoms in this trap for 20 ms to let them
settle in the minimum of the potential. At this point we can

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implement the evaporative cooling process [39], which is the last step before reaching quantum degeneracy. Since the trap increases the density of the sample, we observe a considerable increase of the temperature of the sample to about 200 \( \mu \)K. Figure 10 shows an absorption image of the atoms from the sub-Doppler cooled sample transferred into the ODT beam.

### 3.2.2. Evaporative cooling

Evaporative cooling is performed by ramping down the ODT power while keeping the magnetic field at 832 G. To achieve runaway evaporation it is fundamental that the collision rate does not decrease as the atoms are evaporated, this means that the density of the cloud needs to increase as its temperature is reduced. To guarantee this condition, the evaporation process must be performed slow enough for thermalization to occur. At the same time, the evaporation has to be the main loss process, so it cannot be too slow for the background-vapor collisions with the sample to be important. A good quantity to evaluate the effectiveness of the evaporation process is the phase space density, \( \text{PSD} = n \lambda_{dB}^3 \propto n/T^{3/2} \), which must increase as the evaporation is applied [39].

The evaporative process is performed by concatenating three exponential ramps, as shown in the blue curves of Fig. 11. The first ramp goes from 160 W to 35 W in 300 ms having a characteristic time of \( \tau_1 = 125 \) ms (dotted curve in Fig. 11); the second ramp, from 35 W to 10 W in 1.0 s, with \( \tau_2 = 440 \) ms (dashed curve), and finally, a very slow ramp from 10 W to a variable value of the order of \( P_0 = 35 \) mW in 2.6 s, with \( \tau_3 = 2000 \) ms (solid curve). The total duration of the evaporation process is 3.8 s. These parameters are determined by maximizing the phase density of the system. The black data points in Fig. 11 shows how the measured PSD increases as the evaporation proceeds. Notice that PSD \( \geq 1 \) at the end of the last ramp, indicating the onset of quantum degeneracy.

![Figure 11](image-url)  
**Figure 11.** Blue curves: Plot of the evaporation ramps performed by decreasing the power of the optical dipole trap (not a measurement), see text for details. Black data points: Measurement of the phase space density of the system during evaporation. For these measurements, the uncertainty is of the order of 10\%, corresponding to one standard deviation of ten independent measurements, however, the error bars are not visible at the scale of the graph.

![Figure 12](image-url)  
**Figure 12.** Absorption images of the atomic samples (right pictures) and their corresponding integrated density profile (left graphs) as temperature is decreased. Upper panels: thermal gas above critical temperature \( T_C \). Middle panels: gas just below the critical temperature, notice the bimodal gaussian-parabolic distribution. Lower panels: molecular Bose-Einstein condensate well below the critical temperature, the parabolic distribution is dominant and the gaussian one is barely noticeable. The color gradient corresponds to the optical density of the gas. All pictures were taken after a time-of-flight of 15 ms. In the graphs, the dashed black line corresponds to a fitting of only the gaussian wings, while the orange solid line to the bimodal distribution.
At the end of the third evaporation ramp we adiabatically ramp the Feshbach field to the corresponding value in order to produce a sample in any desired interaction regime across the Feshbach resonance; this magnetic ramp lasts about 300 ms. The regimes that we are interested in exploring are within the interval of 670 to 900 G, which contains the BEC-BCS crossover.

By changing the value of the Feshbach field, we also modify the curvature of the magnetic field; however, it changes less than a 10% within the mentioned interval of interest, which means that we do not significantly modify the geometry of the trap as we change the scattering length. Of course, as can be seen from Eq. (5), the frequencies of the trap depend on the power of the ODT, which in turn, determines the temperature and degree of degeneracy of the sample.

After the evaporative cooling process we are able to produce quantum degenerate superfluid samples containing about $N = 5 \times 10^4$ atomic pairs at a temperature of the order of $T/T_F = 0.1$ (which corresponds for this value of $N$ to approximately 20 nK) and a phase space density well above the unity, of the order of $\text{PSD} \approx 10$, demonstrating the fully degenerate nature of our sample. The trap frequencies are $\omega_r = 2\pi \times 163 \text{ Hz}$ and $\omega_z = 2\pi \times 11 \text{ Hz}$, which means that our sample is cigar-shaped with an aspect ratio of the order of 1:15. The duty cycle of our experiment is shorter than 14 s.

### 3.2.3. Superfluids across the BEC-BCS crossover

As mentioned in the previous section, we select the interacting regime of the produced sample at the end of the last evaporation ramp by means of the Feshbach resonance that allows us to set the value of the scattering length $a_s$. As explained in Sec. 2.2.2, we are able to produce and probe samples at practically any magnetic field up to 1200 G. Specifically, as we explain below, we are able to produce ultracold superfluid samples within the interaction range of $-0.65 < (k_F a_s)^{-1} < 7.6$, which means that we can produce samples from the deep (weakly interacting) BEC regime to the strongly interacting BCS regime, passing, of course, through unitarity at $(k_F a_s)^{-1} = 0$. Clearly, we have access to most of the crossover region, $-1 < (k_F a_s)^{-1} < 1$, corresponding to the magnetic field interval 790 G to 900 G.

Evidently, the most important point here is to achieve, at every interacting regime, temperatures that are below the critical superfluid temperature, $T_C$. On the deep BEC side, $(k_F a_s)^{-1} > 1$, the critical temperature is approximately $T_C^{\text{BEC}} \simeq 0.52T_F$ and it is nearly independent of the scattering length [1, 40]. The minimum temperature attainable in our experiment, $T/T_F = 0.1$, remains well below $T_C^{\text{BEC}}$. In this case, the density profile of the cloud exhibits the very

![Figure 13](image-url)

**Figure 13.** Absorption images of quantum degenerate atomic samples (upper pictures) and their corresponding integrated density profile (lower graphs) as the scattering length is varied across the BEC-BCS crossover. Left panels: Bose-Einstein condensate of molecules at $(k_F a_s)^{-1} \approx 7.6$, the bimodal and gaussian fits are shown as a solid orange and black dashed lines, respectively. Middle panels: superfluid gas at unitarity at $(k_F a_s)^{-1} \approx 0.01$. Right panels: ultracold gas at the BCS side of the Feshbach resonance at $a_s$ at $(k_F a_s)^{-1} \approx -0.37$. The color gradient corresponds to the optical density of the gas. All pictures were taken after a time-of-flight of 20 ms.
characteristic bimodal distribution [1]. The condensed fraction presents a parabolic sharp density profile that arises from the Thomas-Fermi approximation, while the non-condensed thermal atoms follow a gaussian Maxwell-Boltzmann distribution, which we use to estimate the temperature of the cloud in time-of-flight (TOF) imaging [27]. These features can be seen in Fig. 12. The weakest interacting BEC that we can produce corresponds to a magnetic field of 670 G for which \( a_s = 1080 \alpha_0 \) and \( (k_F a_s)^{-1} = 7.6 \). For lower magnetic fields the lifetime of the molecular condensate is too short to perform any typical experiment (it is shorter than 100 ms, while in any other regime described here, it is of the order of 1.5 s).

As the scattering length increases, within the BEC-BCS crossover range, and specially right at unitarity, this well-defined bimodal distribution starts to wash out and becomes broader due to strong interactions [4, 41–43]. In this regime, it is not possible to discriminate between the superfluid fraction and the thermal fraction, and the density profile looks nearly Gaussian. However, we know that we are in the superfluid regime due to the following consideration. On the vicinity of the unitary limit the critical temperature is given by \( T_C \sim 0.167 T_F \) [44], which again, is above the temperature of our sample.

In contrast, on the BCS side of the crossover, the critical temperature is given by [4,45]:

\[
T_F^{BCS} \approx 0.28 T_F e^{-\pi/(2 k_F a_s)},
\]

so it exponentially decays as the quantity \( |k_F a_s|^{-1} \) increases. For instance, at \( (k_F a_s)^{-1} = -0.65 \), the critical temperature for the superfluid state is \( T_C^{BCS} / T_F \approx 0.1 \), which is comparable to the minimum achievable temperature of our setup. In consequence, we cannot access the deep (weakly interacting) BCS superfluid regime because the critical temperature is below the technical limit of our experiment. This means that in our setup, superfluid regimes are attainable within the range \(-0.65 \leq (k_F a_s)^{-1} \leq 7.6 \). Figure 13 shows a sequence of absorption images of a superfluid at \( T/T_F = 0.1 \) containing \( N = 5 \times 10^4 \) atomic pairs, as the scattering length changes from the BEC to the BCS regimes across the crossover.

Besides the considerations concerning the critical temperature that we have presented here, we have also performed an additional measurement that ensures that all the observed regimes present superfluidity. Right after releasing the atoms from the trap, we have performed a fast Feshbach magnetic field ramp from the strongly interacting regimes into the deep BEC side [46, 47]. As result of this ramp, the many-body wave function of the system is projected onto the far BEC side of the resonance. In all cases we observe the characteristic BEC bimodal distribution in the density profile, indicating that at unitarity and its vicinity we always have condensation of atomic pairs.

4. Conclusion and Future Perspectives

We have presented the experimental setup and methods we use to produce and study ultracold fermionic superfluid samples of \(^6\text{Li}\). We are able to generate samples containing \( 5 \times 10^4 \) atomic pairs at temperatures as low as \( T/T_F = 0.1 \) at any superfluid regime across the BEC-BCS crossover within a duty cycle shorter than 14 s. Our setup combines versatility and state-of-art techniques, which will allow us to study different aspects of quantum matter.

As a future perspective, we plan to study topological and hydrodynamic excitations such as quantized vortices (see for instance [48, 49] and references therein). We specifically want to understand how the dynamics of these systems depend on the interacting regime as well as on the temperature of the cloud. To carry out these experiments, we need to expand the capabilities of our imaging system. In particular, as a complementary technique to our current absorption imaging system, we will implement the non-destructive phase contrast imaging technique [27] that will allow us to perform several images of the same sample without perturbing it. This is very important to address the dynamics of the superfluid sample.

As a long term perspective, we plan to produce ultracold samples of \(^7\text{Li}\), a bosonic stable isotope of lithium. This is possible because we have also placed purified \(^7\text{Li}\) in our oven. The optical frequencies of the \( D_1 \) and \( D_2 \) lines of \(^7\text{Li}\) are very close to those of fermionic \(^6\text{Li}\) [23]. This means that with minor modifications on the optical cooling setup we should be able to produce, alternatively, bosonic ultracold samples of \(^7\text{Li}\). This is very interesting because this species also presents a broad Feshbach resonance, opening the possibility to study very weakly interacting bosonic systems, a regime that our current setup does not offer and which represents an excellent scenario to study the thermodynamic properties of the superfluid to normal gas transition.

Acknowledgments

Our experimental setup is the first apparatus able to produce quantum gases in Mexico, it is also the first one capable of producing fermionic ultracold systems in Latin America. We hope that our work helps to incentive the growth of this exciting and interesting research field in our country and in the Latin-American region.

The construction of our experiment has represented a major challenge. This has only been possible thanks to the support of numerous institutions, colleagues and different funding sources. We would like to use this space to make the corresponding acknowledgments.

We want to thank our colleagues from Instituto de Física, UNAM:

- To Manuel Torres Labansat, Víctor Romero-Rochín,
  Daniel Sahagún Sánchez, Rosario Paredes Gutiérrez,
Asaf Paris Mandoki and Carlos Villarreal Luján for their unconditional support and fruitful discussions.

- To Roberto Gleason Villagrán and Marco Antonio Veytia Vidana for their excellent work during the installation of the basic infrastructure of the laboratory.

- To the former members of our laboratory, Eduardo Ibarra García-Padilla, Jesús Ernesto Carro Martínez, Cristian Adán Mojica Casique, Iliana Isabel Cortés Pérez and Aurora Guadalupe Borges Sánchez, for their contribution during the early stages of construction of the experimental setup.

- To Giovanni Alonso Torres for his help in preparing Fig. 2.

We also thank our colleagues from INO-CNR and LENS:

- Alessia Burchianti and Chiara Fort for their technical advice in setting our automation and vacuum systems.

- Andreas Trenkwalder for his help in setting different analysis algorithms as well as his advice concerning our imaging setup.

- Marco de Pas for his advice in the construction of several electronic devices and instruments.

- In general, the entire Quantum Gases Group for their very useful advice.

We acknowledge as well Greg S. Engel (University of Chicago) for his advice in the installation of our air-conditioning system.

We are deeply grateful for the financial support provided by the following institutions:

- CIC-UNAM and CONACyT through the National Laboratories program, with grants number: LN232652, LN260704, LN271322, LN280181, LN293471 and LN299057.

- CONACyT through Ciencia Básica grants 255573 and 254942.

- Instituto de Física UNAM through grants PIIF-8 and PIIF-9.

- DGAPA-UNAM-PIAPIIT grant numbers IA101716, IN111516, IN107014 and IN103818.

- CONACyT through Redes Temáticas “Tecnología Cuántica”.

F.J.P.C. would like to thank the support from DGAPA-UNAM and from CLAF-SeCyTI for former postdoctoral fellowships. D.H.R., J.E.P.C., M.M.L., R.C.R., A.G.V. and S.A.M.R. acknowledge their scholarships from CONACyT, while D.H.R. also acknowledge a scholarship from DGAPA-UNAM-PIAPIIT.

We finally want to thank the company Seman Baker S.A. de C.V. for their generous donation of numerous machined pieces.

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