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Tongcang Li

Fundamental Tests of Physics with Optically Trapped Microspheres



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Tongcang Li

Fundamental Tests of Physics with Optically Trapped Microspheres

Doctoral Thesis accepted by the University of Texas at Austin, USA



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Dedicated to my parents, Daogan Li and Zhangman Huang, my wife, Ruyi Li, and my sister, Xiaoping Li, for their endless love and support

Supervisor's Foreword

I am very pleased to introduce this Ph.D. Thesis by Dr. Tongcang Li. Tongcang was a graduate student under my supervision, and completed his Ph.D. in May 2011. He was the senior student on an experiment to study the motion of micronsized beads of glass held in optical tweezers. This system has provided spectacular experimental results, and I credit Tongcang with the vision and the ability to pull it off. We were able to resolve, for the first time, the instantaneous velocity of a Brownian particle in air. In 1907, Albert Einstein analyzed this problem and showed that such a measurement could be used to test the energy equipartition theorem, one of the basic tenets of statistical mechanics. However, Einstein concluded in his paper that the time-scale was too short to be measured in practice! This prediction held for over 100 years, but Tongcang Li was able to meet the challenge of Albert Einstein, and to measure the instantaneous velocity of a Brownian particle. We used the velocity data to verify directly the energy equipartition theorem for a Brownian particle. The work has attracted great interest and press, partly due to the historical challenge by Einstein, and also because it addresses fundamental questions on statistical mechanics of small systems.

Following that work, Tongcang led an experiment to cool the center-of-mass motion of a bead in vacuum close to the absolute zero of temperature. This looks like an ideal system to test quantum mechanics at the boundary between the microscopic and macroscopic, and answer fundamental questions about the collapse of the wave function due to the environment.

In parallel, Tongcang started our efforts to observe the instantaneous velocity of a bead in water, and we expect to be able to complete this work in the near future. I believe that this experiment will serve as a new testing ground for statistical mechanics on short time scales. It will also be a new probe into the nature of complex fluids, such as water, a topic of great interest to chemists and biologists.

In recognition of these accomplishments, Tongcang was awarded a Biedenharn Graduate Fellowship in the physics department. He also won the Jean Bennett Memorial Student Travel Grant from the Optical Society of America Foundation (the only one out of 80 applicants), and the Chinese Government Award for Outstanding Self-financed Students Abroad. Tongcang was recognized by the UT Physics Department outstanding Ph.D. award.

In summary, this Ph.D. Thesis by Tongcang Li is simply outstanding, and is a worthy addition to the Springer Thesis Series.

Austin, August 2012

Prof. Dr. Mark Raizen

Acknowledgments

First, I would like to thank my supervisor, Dr. Mark Raizen, for providing me the opportunity to work in an outstanding group. Mark is a great person to work with and is full of ideas. I am impressed by his vision and confidence. Mark often proposes ambitious ideas that may appear suspicious to most people at first. During my time in his group, I had the privilege to witness these ideas ultimately realized in ground-breaking experiments. I could not have asked for a better advisor to learn how to do creative research.

I joined the lab in November 2004 to work on the sodium Bose–Einstein condensate (BEC) experiment with Kevin Henderson and Hrishikesh Kelkar. Kevin was very helpful and excellent in teaching. I learned a lot of experimental skills from him, including driving a car. Hrishi was extremely patient and hard working. He aligned a lot of difficult parts of the optics, some of which took several weeks of continuous work. David Medellin joined this experiment in 2006. David is fantastic with mathematics and physical concepts, which is very helpful to our work. He is always trying his best to make his work perfect. I also enjoyed the dinners with Hrishi and David when we were working together. Kirsten Viering joined this experiment as a Würzburg exchange student and later returned back after working on a Rubidium experiment. She is a very good experimentalist and works very efficiently.

I started the new experiment on optical trapping and cooling of microspheres with the help of David Medellin. A particular thanks goes to Simon Kheifets. He joined this experiment just after we were able to trap microspheres in vacuum and has participated in almost all aspects of this experiment since then. He worked very hard and helped me to make this experiment fruitful. I also enjoyed the discussions with him on our new ideas almost every day. I fully expect that he will push the experiment to an unprecedented level. I would also like to thank Ivo Popov, Ioannis K. Charidakos, and Isaac Chavez for their contributions to this experiment.

I would like to thank other students in the lab who have provided much help to me. They are Braulio Gutierrez-Medina, Todd Meyrath, Chih-Sung Chuu, Gabriel Price, Travis Bannerman, Ed Narevicius, Adam Libson, Rob Clark, Tom Mazur, Melissa Jenkins, and Jianyong Mo. I would also like to thank Charlotte Sanders, Francisco Camargo, Daniel Ellsworth, Julia Majors, Christoph Schaeff, Christian Parthey, Max F. Riedel, Artem M. Dudarev, Chuanwei Zhang, and Shoupu Wan for making the environment of our lab friendly and enjoyable.

I would like to thank Jack Clifford, Allan Schroeder and all other members of the machine shop, and Ed Baez and Larry Sandefur in the cryo lab for their excellent work. I would like to thank Olga Vera, Elena Simmons, Marybeth Casias, and other members of CNLD for their administrative help.

I also appreciate the friendships of Shaoping Lu, Hui Chen, Ning Kong, Huai Huang, Jiaxin Han, Tianyi Yang, Zhangqi Yin, Ming Lei, Kin Chung Fong, and more.

Finally, I would definitely like to thank my wife, Ruyi Li. She is very supportive and provides me a sweet home. I also want to thank my parents and my sister for their endless love and encouragement.

28 April 2011, Austin, TX, USA

Tongcang Li

Contents

1	Introduction			
	1.1 Macroscopic Quantum Mechanics	1		
	1.2 Instantaneous Velocity of Brownian Motion	3		
	1.3 Contents of this Dissertation	5		
	References	5		
2	Physical Principle of Optical Tweezers	9		
	2.1 Ray Optics Approximation	10		
	2.2 Rayleigh Approximation	11		
	2.3 Generalized Lorentz-Mie Theory	15		
	References	19		
3	Optical Trapping of Glass Microspheres in Air and Vacuum	21		
	3.1 Launching Microspheres.	22		
	3.2 Trapping Microspheres	27		
	3.2.1 Optical Levitation Trap	28		
	3.2.2 Counter-Propagating Dual-Beam Optical Trap	30		
	3.3 Vacuum System	35		
	References	37		
4	Measuring the Instantaneous Velocity			
	of a Brownian Particle in Air	39		
	4.1 Theories of Brownian Motion	39		
	4.1.1 A Free Particle	39		
	4.1.2 A Trapped Microsphere	41		
	4.2 A Fast Detection System	46		
	4.3 Measured Power Spectra	47		
	4 Measurement of the Instantaneous Velocity			
	of a Brownian Particle in Air	50		
	References			

5	Towards Measurement of the Instantaneous Velocity				
	of a	Brownian Particle	e in Water	59	
	5.1	Motivation		59	
	5.2	Hydrodynamic Th	neories of Brownian Motion	59	
		5.2.1 A Free Par	rticle in Water	59	
		5.2.2 An Optical	Ily Trapped Microsphere in Water	61	
	5.3	Requirements for	Measuring the Instantaneous Velocity	66	
	5.4	A Simple Optical	Tweezer in Water	70	
	5.5	Interferometer-Enl	hanced Optical Tweezers	75	
	Refe	rences		79	
6	Mill	ikelvin Cooling of	an Optically Trapped Microsphere		
	in V	acuum		81	
	6.1	Background		81	
	6.2	Principle of Feedb	back Cooling	82	
		6.2.1 Feedback	Cooling	83	
		6.2.2 Feedback	Amplification	84	
		6.2.3 Heating D	ue to Light Scattering	84	
		6.2.4 Damping I	Due to the Residual Gas in Vacuum	86	
	6.3	A 3D Split Detect	tion System	88	
	6.4	1D Optical Feedba	ack Cooling	90	
	6.5	Electrostatic Force	es	95	
	6.6	Millikelvin Coolin	ng with 3D Optical Feedback	99	
		6.6.1 Experimen	ntal Setup	99	
		6.6.2 Results of	3D Optical Feedback Cooling	102	
	6.7	Loss of Microsphe	eres in Vacuum	105	
	Refe	rences		108	
7	Том	ards Quantum Gr	round-State Cooling	111	
	7.1	Principle of Cavity	y Cooling	111	
	7.2	3D Ground-State	Cooling with a Single Cavity	113	
	7.3	Heating Effects of	f Laser Noise	116	
	7.4	Applications of Co	ooled Microspheres in Vacuum	117	
		7.4.1 Measuring	Weak Forces	117	
		7.4.2 Measuring	the Impact of Single Molecules	118	
		7.4.3 Searching	for Gravity-Induced		
		Quantum-S	State Reduction	120	
	Refe	rences		121	
Ap	opend	ix: Physical Prope	erties of Some Common Materials	123	
Cı	ırricu	lam Vitae		125	

Chapter 1 Introduction

1.1 Macroscopic Quantum Mechanics

It is well known that the dynamics of microscopic particles such as photons, electrons and atoms follow quantum mechanics, while the dynamics of macroscopic objects such as ping-pong balls follow classical mechanics. Why is there such a transition?

A simple explanation for the absence of quantum behavior of macroscopic objects is that the de Broglie wavelength λ of an object is inversely proportional to the momentum p of the object, i.e. $\lambda = h/p$, where h is the Planck constant. Macroscopic objects usually have large momenta, thus their de Broglie wavelengths and quantum behaviors are too small to be observed.

Since p = Mv, where *M* is the mass of an object and *v* is the velocity of the object, we can increase the de Broglie wavelength of an object by reducing its velocity. Thus macroscopic objects should behave quantum mechanically if we can reduce their kinetic energies to low enough values. An important characteristic of quantum mechanics is the possibility of superposition of two spatially distinct states. Creation of quantum superpositions of macroscopic objects ("Schrödinger cats") will provide opportunities to study untested regimes of quantum mechanics.

One additional motivation to study the macroscopic quantum mechanics is that the apparent conflict between general relativity and quantum mechanics remains one of the unresolved mysteries of the physical world [1, 2]. According to recent theories [2–4], this conflict results in gravity-induced quantum state reduction of "Schrödinger cats". It is argued that the perturbing effect of the mass distribution on the space-time structure leads to an essential uncertainty in the energy of a superposed state. So a macroscopic quantum superposition of two different mass distributions is intrinsically unstable.

The finite lifetime of a quantum superposition due to the gravity-induced state reduction is on the order of $\tau = \hbar/\Delta E$, where \hbar is the Planck constant/ 2π . ΔE is the gravitational self-energy of the difference between the mass distributions of the two superposed states [2, 5, 6]:

1 Introduction

$$\Delta E = \xi G \int \int \frac{[\rho(\mathbf{x}) - \rho'(\mathbf{x})] \cdot [\rho(\mathbf{y}) - \rho'(\mathbf{y})]}{|\mathbf{x} - \mathbf{y}|} d\mathbf{x} d\mathbf{y}, \qquad (1.1)$$

where G is Newton's gravitational constant, ρ and ρ' are the mass densities of two members of the superposed state. ξ is a dimensionless number that represents the strength of the quantum state reduction process, which is expected to be on the order of 1. The lifetime τ of a quantum superposition of two separated states of a 1-nm-radius particle is on the order of 10^{13} s, which is too long to be measured in an experiment. It is on the order of 10^{-17} s for a 1 mm particle, which is too short to be measured. For a 1 μ m particle, the lifetime of a spatially separated quantum superposition is on the order of 10^{-2} s, which seems realistic to be measured in an experiment.

In order to observe quantum behaviors in a mechanical oscillator, we need to cool the oscillator significantly. The energy of the quantum states of a mechanical oscillator is quantized in units of $\hbar\Omega_M$, where Ω_M is the angular frequency of the oscillator. The mean thermal occupation number (phonon number) is $\langle n \rangle = k_B T / \hbar\Omega_M$, where k_B is the Boltzmann constant, and *T* is the temperature of the oscillator. At room temperature, we have $\langle n \rangle \gg 1$ for almost all mechanical oscillators. We need $\langle n \rangle < 1$ for the oscillator to have a high probability to be in the quantum ground state and to have its dynamics be dominated by quantum mechanics.

In recent years, great progress has been made in cooling mechanical oscillators towards their quantum mechanical ground states [7–9]. The thermal occupation number of a 2.7 kg pendulum mode around 130 Hz was reduced from about 5×10^{10} at room temperature to about 234 at an effective temperature of $1.4 \,\mu\text{K}$ by feedback cooling [10]. Combining conventional cryogenic cooling and laser cavity cooling, the thermal occupation number of a mechanical deformation mode of a deformed microsphere around 100 MHz was reduced to about 40 [11], the thermal occupation number of the fundamental mechanical mode of a micromirror (the end mirror of a Fabry-Pérot cavity) at 945 kHz was reduced to about 30 [12]. Remarkably, quantum ground-state cooling has been achieved for a dilatational resonant mode of a membrane at 6.2 GHz by conventional cryogenic cooling [13], and for the fundamental mechanical mode of a superconducting aluminium membrane at 10.56 MHz by microwave cavity cooling [14].

These results are important steps towards the creation of Schrödinger cats in the laboratory, and the study of their destruction by decoherence. A direct test of the gravity-induced state reduction scenario may therefore be within reach. However, a recent analysis shows that for these mechanical oscillators, quantum superpositions are destroyed by environmental decoherence long before gravitational state reduction takes effect [6]. An alternate candidate may be a Bose-Einstein condensate (BEC) of ultracold atoms in vacuum. It is well isolated from the thermal environment. However, its mass density is too small. Even worse, the atoms in a Bose-Einstein condensate are only weakly interacting. They do not move together. Thus only mass fluctuation, not total mass, contributes to the ΔE . Theoretical calculation shows that a Bose-Einstein condensate formed from 10^{29} atoms is required to study the

gravity-induced state reduction [6]. This number is too large to be achieved in the foreseeable future.

Chapters 3 and 6 of this dissertation describe our experimental efforts on optical trapping and cooling of glass microspheres in vacuum [15]. An optically trapped microsphere in vacuum provides an ideal candidate for studying the interface between quantum mechanics and general relativity, and for studying objective collapse models of the wavefunction [16]. It is nearly perfectly isolated from the thermal environment, and allows quantum ground state cooling from room temperature [17–19]. The mass density of a glass microsphere is many orders larger than that of a BEC, and atoms contained in the microsphere move together as a whole. We have been able to trap glass microspheres in vacuum with high oscillation frequencies, and cool their center-of-mass motion from room temperature to a minimum temperature of about 1.5 mK.

1.2 Instantaneous Velocity of Brownian Motion

Brownian motion is the apparently random movement of particles suspended in a fluid (liquid or gas). It was discovered by Robert Brown (1773–1858) in 1827 when he used a simple microscope (a double convex lens with a focal length of about 0.8 mm) to study the action of particles contained in the grains of pollens [20]. The size of those particles was about 5 μ m. He "observed many of them very evidently in motion". He also observed the same motion with powders of many other materials, such as wood and nickel, suspended in water.

Persistence and randomness are two key characteristics of Brownian motion. The trajectories of a Brownian particle are classic examples of fractals. They are commonly thought to be continuous everywhere but not differentiable anywhere. Since its trajectory is not differentiable, the velocity of a Brownian particle is undefined.

At short time scales, however, the dynamics of a Brownian particle is expected to be dominated by its inertia and its trajectory cannot be self-similar. This is termed "ballistic Brownian motion" to be distinguished from the common "diffusive Brownian motion". Figure 1.1 shows a 2D trajectory of a Brownian particle. The black curve is assumed to be a true trajectory of the particle. Red dots are measured positions. In Fig. 1.1a, the sampling rate is too small to measure the velocity of the Brownian particle. The measured trajectory (red curve) is completely different from the true trajectory, and appears chaotic. It is impossible to obtain the velocity of the particle from the measured trajectory in Fig. 1.1a. In Fig. 1.1b, the sampling rate is much larger. Now the measured trajectory is very close to the true trajectory of the particle. If the measured displacement of the particle is $\Delta \vec{x}(t)$ during time Δt , then the velocity of the particle is approximately $\vec{v} = \Delta \vec{x}(t)/\Delta t$.

In 1900, F. M. Exner made the first quantitative study of Brownian motion by measuring the velocity of the Brownian motion of particles suspended in water [21, 22]. He found that the measured velocity decreased with increasing particle size and increased with increasing water temperature. However, his measured velocities of Brownian particles were almost 1000-fold smaller than those predicted by the energy

1 Introduction



Fig. 1.1 A 2D trajectory of a Brownian particle. The *black curve* is assumed to be a true trajectory of the particle. *Red dots* are measured positions, and *red curves* are measured trajectories. The sampling rate of \mathbf{b} is 10 times of that of \mathbf{a}

equipartition theorem [21]. The reason of this discrepancy was not understood until A. Einstein developed a kinetic theory about Brownian motion in 1905 [23].

In 1907, Einstein published a paper entitled "Theoretical observations on the Brownian motion" in which he considered the instantaneous velocity of a Brownian particle [24, 25]. Einstein showed that by measuring this quantity, one could prove that "the kinetic energy of the motion of the centre of gravity of a particle is independent of the size and nature of the particle and independent of the nature of its environment". This is one of the basic tenets of statistical mechanics, known as the equipartition theorem. However, Einstein concluded that due to the very rapid randomization of the motion, the instantaneous velocity of a Brownian particle would be impossible to measure in practice [24, 25]:

"We must conclude that the velocity and direction of motion of the particle will be already very greatly altered in the extraordinary short time θ ,¹ and, indeed, in a totally irregular manner. It is therefore impossible—at least for ultramicroscopic particles—to ascertain $\sqrt{v^2}$ by observation."

The resolution required to measure the instantaneous velocity is striking. For a 1 μ m diameter silica (SiO₂) sphere in water at room temperature, the momentum relaxation time is about 0.1 μ s and the root mean square (rms) velocity $v_{rms} = \sqrt{k_B T/M}$ is about 2 mm/s in one dimension. To measure the instantaneous velocity with 10% uncertainty, one would require 2 pm spatial resolution in 10 ns.

Recent experiments on Brownian motion in liquid [26–29] and gaseous environments [30–32] with fast detectors have observed nondiffusive motion of a Brownian particle. However, no previous experiment has been able to observe the instantaneous velocity of a Brownian particle and verify the energy equipartition theorem directly.

Chapter 4 of this dissertation describes our efforts to measure the instantaneous velocity of a Brownian particle suspended in air [33]. Due to the lower viscosity of gas as compared to liquid, the momentum relaxation time τ_p of a particle in air is much

¹ In Einstein's paper, θ was defined as the time in which the velocity falls to a tenth of its original value. $\theta = 2.30 \tau_p$, where τ_p is the momentum relaxation time of the particle. Einstein mistakenly obtained $\theta = 330$ ns for a 50-nm-diameter platinum nanosphere in water. The correct value should be $\theta = 6.8$ ns for his example.

larger. This lowers the technical demand for both temporal and spatial resolution. The main difficulty of performing high precision measurements of a Brownian particle in air, however, is that the particle will fall under the influence of gravity. We overcome this problem by using optical tweezers to simultaneously trap and monitor a silica bead in air and vacuum, allowing long duration ultra-high-resolution measurements of its motion.

We have successfully measured the instantaneous velocity of a Brownian particle in air. We used the velocity data to directly verify the Maxwell-Boltzmann velocity distribution, and the equipartition theorem for a Brownian particle. The ability to measure instantaneous velocity enables new fundamental tests of statistical mechanics of Brownian particles [34–37]. It is also a necessary step towards cooling of a particle to the motional quantum ground state in vacuum.

We are currently building a new detection system to measure the instantaneous velocity of a Brownian particle in water (Chap. 5). In contrast to air, the water has significant inertia and hydrodynamic memory effects [26, 29], which may cause deviations from the energy equipartition theorem.

1.3 Contents of this Dissertation

Chapter 2 introduces the basic principle of optical tweezers, and the differences between trapping microspheres in air and in water. Chapter 3 provides the details of launching glass microspheres to air, and trapping microspheres in air and vacuum with a counter-propagating dual-beam optical tweezer. We also describe the vacuum system and the first generation of our detection system. Chapter 4 covers the theory of Brownian motion in air at short time scales, a home-built detection system with ultrahigh resolution, and the results of our measurement of the instantaneous velocity of a Brownian particle in air. Chapter 5 discusses the motivation and challenges of measuring the instantaneous velocity of a Brownian particle in water. It also presents some of our on-going efforts on developing a new detection system to meet the challenges. Chapter 6 presents different approaches to cool the center-of-mass motion of an optically trapped microsphere in vacuum, and the results of 3D optical feedback cooling. We also discuss the trapping lifetime of optically trapped microspheres in vacuum. Finally, Chap. 7 discusses a scheme to perform 3D ground-state cooling of a optically trapped nanosphere with a single cavity, and the potential applications of cooled microspheres in vacuum.

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Chapter 2 Physical Principle of Optical Tweezers

The radiation pressure of light was first deduced theoretically by James C. Maxwell in 1873 based on his electromagnetic theory [1, 2], and measured experimentally by Lebedev [3], and Nichols and Hull in 1901 [4]. The radiation pressure force exerted on a totally reflecting mirror by an incident beam of light perpendicular to the mirror is $F_{mirror} = 2P/c$, where P is the power of the light and c is the speed of light in vacuum [5]. The factor of 2 in the formula is due to reflection. The force is about 7 nN for 1 W of light, which is tiny and had almost no application before the invention of the laser. In contrast to classical light sources, a laser beam can be strongly focused onto a small particle with a diameter on the order of 1 μ m. Due to the small mass of the particle, the radiation force of a 1 W laser can be 10⁵ times larger than the gravitational force on the particle, and can therefore have huge effects on the motion of the particle.

In 1970, Arthur Ashkin published a seminal paper [6] demonstrating that one could use focused laser beams to accelerate and trap micrometer-sized transparent particles. Optical levitation of oil droplets and glass microspheres in air [7] and vacuum was demonstrated several years later [8]. The laser radiation pressure was soon used to cool and trap atoms [9–12], leading to dramatic breakthroughs in atomic, molecular and optical physics, including a new generation of atomic clocks, and realization of Bose–Einstein condensation and degenerate Fermi gas. In 1986, Ashkin et al. [13] observed stable trapping of dielectric particles with the gradient force of a strongly focused laser beam. This technique was soon used to trap and manipulate viruses and bacteria [14, 15], and became a standard tool in biophysics [5].

In this chapter, we will first explain the principle of optical trapping of microspheres with ray optics, which is valid when the size of the microspheres is much larger than the wavelength of the trapping laser. This will be followed by theoretical calculations of the optical forces on a particle with the Rayleigh approximation, and numerical results of Lorentz-Mie theory. The differences between trapping microspheres in air and in water will be discussed.

2.1 Ray Optics Approximation

When the size of a microsphere is much larger than the wavelength of the trapping laser (usually $R > 10\lambda_0$, where R is the radius of the microsphere and λ_0 is the wavelength of the laser in vacuum), the optical forces on the microsphere can be calculated by ray optics [16].

A qualitative view of optical trapping of microspheres in the ray optics regime is shown in Fig. 2.1 [13, 16]. If we neglect surface reflection from the microsphere, then the microsphere will be trapped at the focus of the laser beam as shown in Fig. 2.1b. If the microsphere moves to the left of the focus (Fig. 2.1a), it will deflect the laser beam to the left and thus increase the momentum of photons to the left. The counter force from the deflected photons will push the microsphere to the right, i.e. back to the focus of the laser beam (Fig. 2.1c), it will focus the laser more strongly and thus increase the momentum of photons. The counter force from the deflected photons along the propagation direction. The counter force from the deflected photons along the propagation direction. The counter force from the deflected photons will push the microsphere back to the focus of the laser beam. The same thing will happen if the microsphere moves away from the focus in other directions. Thus a focused laser beam forms a stable optical trap in 3D.

The above discussion neglected surface reflection from the microsphere. In reality, we have to consider the effect of this surface reflection. The photons reflected back by the surface of a microsphere will push the microsphere forward. If this force is larger than the restoring force due to refraction (Fig. 2.1c), the microsphere will be pushed away from the focus, and thus cannot be trapped. The surface reflection depends on the relative refractive index of the microsphere and the medium $m = n_p/n_{md}$, where n_p is the refractive index of the microsphere and thus greater difficulty in trapping the microsphere with an optical tweezer [17]. *m* is about 1.10 for a silica microsphere in air ($n_{air} = 1.46$) in water ($n_{water} = 1.33$), and is about 1.46 for a silica microsphere in air ($n_{air} = 1.00$) (see Table A.1 for more information). Thus it is more difficult to trap microspheres in air than in water.

To increase the restoring force, the laser beam should be strongly focused by a high numerical aperture (NA) objective lens. The typical NA of objective lenses used for creating optical tweezers is about 1.2 and 0.95 in water [13, 17] and air [18], respectively.



Fig. 2.1 Qualitative view of optical trapping of dielectric spheres. **a** Displays the force on the particle when the particle is displaced laterally from the focus. **b** Shows that there is no net force on the particle when the particle is trapped at the focus. **c** Displays the force on the particle when the particle sphere at the focus.

2.2 Rayleigh Approximation

If the size of a nanosphere (microsphere) is much smaller than the wavelength of the trapping laser (usually $R < \lambda_0/10$), the nanosphere can be approximated as a dipole. In this regime, the optical force on the nanosphere can be calculated analytically with the Rayleigh scattering theory [13, 19]. Here we will calculate the optical forces in the Rayleigh regime following the formulas of Ref. [19].

We consider a nanosphere with radius R and a refractive index n_p being illuminated by a laser beam propagating along the z axis in the positive direction, as shown in Fig. 2.2. The power of the laser beam is P. The refractive index of the medium in which the nanosphere is suspended is n_{md} . The laser beam is a linearly polarized Gaussian beam (*TEM*₀₀) with beam waist radius ω_0 at the focus. The polarization direction of the electric field of the laser is parallel to the x axis. The center of the laser beam is located at the origin, and the center of the nanosphere is at $\vec{r} = (x, y, z)$.

The wavefront of a Gaussian beam is flat at the focus, and its waist $(1/e^2 \text{ radius})$ spreads in accordance with [20]:

$$\omega(z) = \omega_0 \left[1 + \left(\frac{\lambda_{md} z}{\pi \omega_0^2} \right)^2 \right]^{1/2}, \qquad (2.1)$$

where $\lambda_{md} = \lambda_0 / n_{md}$ is the wavelength of the laser in the medium.

The Rayleigh range (z_R) , defined as the distance over which the beam radius spreads by a factor of $\sqrt{2}$, is given by

$$z_R = \frac{\pi \omega_0^2}{\lambda_{md}}.$$
 (2.2)

The intensity distribution of the Gaussian beam is



Fig. 2.2 Schematic of a nanosphere near the focus of a laser beam

2 Physical Principle of Optical Tweezers

$$I(x, y) = I_0 e^{-2(x^2 + y^2)/\omega^2} = \frac{2P}{\pi\omega^2} e^{-2(x^2 + y^2)/\omega^2},$$
(2.3)

where $\omega = \omega(z)$ and *P* is the power of the laser beam.

The numerical aperture $(1/e^2)$ points in k-space) of a Gaussian beam is

$$NA = \frac{\lambda_0}{\pi\omega_0} = n_{md} \frac{\lambda_{md}}{\pi\omega_0}.$$
(2.4)

The optical force of the focused laser beam on the nanosphere can be separated into two components: the scattering force $\vec{F}_{scat}(\vec{r})$ which is proportional to the intensity of the laser, and the gradient force $\vec{F}_{grad}(\vec{r})$ which is proportional to the gradient of the intensity of the laser. The scattering force is a nonconservative force and the gradient force is a conservative force. The gradient force forms a trapping potential for the nanosphere, and the scattering force tends to push the nanosphere out of the trap. In order to form a stable trap, the gradient force should be lager than the scattering force.

The scattering force of the laser on a nanosphere is [19]:

$$\vec{F}_{scat}(\vec{r}) = \hat{z}(\frac{n_{md}}{c})C_{scat}I(\vec{r}) = \hat{z}\frac{128\pi^5 R^6}{3c\lambda_0^4} \left(\frac{m^2 - 1}{m^2 + 2}\right)^2 n_{md}^5 I(\vec{r}), \qquad (2.5)$$

where *c* is the speed of light in vacuum and C_{scat} is the scattering cross section. Because of the larger relative refractive index *m*, the scattering force on a nanosphere in air is about 4.2 times greater than the scattering force on the same nanosphere in water with the same laser intensity. The number of scattered photons per second is

$$N_{scat} = \frac{\lambda_{md}}{h} |\vec{F}_{scat}|, \qquad (2.6)$$

where h is the Planck constant.

The gradient force on the nanopshere is [19]:

$$\vec{F}_{grad}(\vec{r}) = [\vec{p}(\vec{r},t) \cdot \nabla] \vec{E}(\vec{r},t) = \frac{2\pi n_{md} R^3}{c} \left(\frac{m^2 - 1}{m^2 + 2}\right) \nabla I(\vec{r}), \qquad (2.7)$$

where $\vec{p}(\vec{r}, t)$ is the induced dipole of the nanosphere due to the instantaneous electric field $(\vec{E}(\vec{r}, t))$ of the laser. The gradient force forms a trapping potential:

$$V(\vec{r}) = -\frac{2\pi n_{md} R^3}{c} \left(\frac{m^2 - 1}{m^2 + 2}\right) I(\vec{r}).$$
(2.8)

The total force on the nanosphere is $\vec{F}(\vec{r}) = \vec{F}_{scat}(\vec{r}) + \vec{F}_{grad}(\vec{r})$. The minimum force along the *z* axis $F_z^{min} = \min(F_z(\vec{r}))$ must be negative in order to form a stable trap. Otherwise the force of the laser will always push the nanosphere forward and

there will be no trap. Because the scattering force is proportional to R^6 while the gradient force is proportional to R^3 , the scattering force decreases much faster than the gradient force when the size of the nanosphere decreases. Thus it is easier to achieve a negative F_{τ}^{min} for a small nanosphere than a large particle.

In order to trap a nanosphere stably, the well depth should be at least 10 times larger than the average kinetic energy of the nanosphere. This is due to the fact that the kinetic energy of a nanosphere follows the Maxwell–Boltzmann distribution at thermal equilibrium. The nanosphere has a significant probability for its instantaneous kinetic energy to be much larger than its average kinetic energy. According to the energy equipartition theorem, the average kinetic energy of a nanosphere is $k_BT/2$ in each direction, where k_B is the Boltzmann constant and T is the temperature of the medium. While the average kinetic energy is independent of the size of the nanosphere, the well depth of the trapping potential decreases as the size of the particle decreases. Thus it is difficult to trap a nanosphere if its size is too small.

Fig. 2.3 shows the calculated potentials and forces on a silica nanosphere in air with a focused laser beam. Since the refractive index of air is very close to the refractive index of vacuum, the potential and force on a nanosphere in air is practically the same as that in vacuum. For the calculations yielding Fig. 2.3, the wavelength of the laser is 1064 nm, the power of the laser is 200 mW, and the waist of the laser at the focus



Fig. 2.3 Optical potentials and forces on a nanosphere in air when the waist of the trapping laser is $1.5 \,\mu$ m. The power of the trapping laser is 200 mW and the diameter of the nanosphere is 50 nm



Fig. 2.4 Optical potentials and forces on a nanosphere in air when the waist of the trapping laser is 0.5μ m. The power of the trapping laser is 200 mW and the diameter of the nanosphere is 50 nm

is 1.5 μ m, corresponding to NA = 0.22. The Rayleigh range of the laser is 6.6 μ m. The diameter (D = 2R) of the nanosphere is 50 nm. The calculated well depth of the trap is 367 K. The laser will therefore only be able to trap a 50 nm nanosphere at a temperature much lower than room temperature. The potential is approximately harmonic near the bottom of the trap. The oscillation frequency is about 42 kHz in the radial direction and 6.7 kHz in the axial direction for a 50 nm nanosphere trapped near the bottom of the potential. The scattering force is zero in the radial direction (Fig. 2.3c) and is positive along the axial direction (Fig. 2.3d). The gradient force is negative at positive coordinates, and positive at negative coordinates. Thus it will always pull back the nanosphere to the center of the trap.

Fig. 2.4 shows the calculated potentials and forces on a silica nanosphere in a laser beam with a much smaller waist. The waist of the laser beam is $0.5 \,\mu$ m, which corresponds to NA = 0.68. Other conditions are the same as in Fig. 2.3. Because of the smaller waist, the well depth becomes large enough (3310 K) to trap a 50 nm nanosphere at room temperature. The trapping frequency is about 376 kHz in the radial direction and 180 kHz in the axial direction. The scattering force is negligible compared to the gradient force. Comparing Figs. 2.4 and 2.3, it is clear that a laser



Fig. 2.5 Optical forces on a microsphere in water along axial (**a**) and radial (**b**) directions of a laser beam focused by objective lenses with different numerical apertures (*NA*'s). The power of the laser is 100 mW, and the diameter of the microsphere is 3 μ m

beam focused by an objective lens with a larger NA is much better for trapping nanospheres.

2.3 Generalized Lorentz-Mie Theory

In most experiments with optical tweezers, the sizes of the dielectric particles are comparable with the wavelength of the trapping laser ($R \sim \lambda_0$). In this case, neither ray optics nor the Rayleigh approximation is appropriate. Instead the electromagnetic theory of light has to be used. For optical trapping of homogeneous and isotropic microspheres, one can use the generalized Lorenz-Mie theory. The mathematical calculation of the generalized Lorenz-Mie theory is quite complex. Here we will only introduce this method briefly, and use the computational toolbox developed by Nieminen et al. [21] to obtain some numerical results of the optical force on a microsphere.



Fig. 2.6 Optical forces on microspheres in air along axial (a) and radial (b) directions of a laser beam focused by an objective lens with NA = 0.95. The power of the laser is 100 mW. The diameter of the microsphere is 2.4 μ m for the *dotted lines*, 3.0 μ m for the *dashed lines*, and 3.6 μ m for the *solid lines*

The optical force on a microsphere comes from the momentum of photons (electromagnetic field) from a laser. It can be obtained by calculating the change of the momentum of the electromagnetic field scattered by the microsphere. A natural choice of coordinate system for calculating the light scattering by a micropshere is spherical coordinates (r, θ, ϕ) centered on the trapped microsphere. The incoming and outgoing fields can be expanded in terms of incoming and outgoing vector spherical wavefunctions [21]:

$$\mathbf{E}_{in} = \sum_{i=1}^{\infty} \sum_{j=-i}^{i} a_{ij} \mathbf{M}_{ij}^{(2)}(k\mathbf{r}) + b_{ij} \mathbf{N}_{ij}^{(2)}(k\mathbf{r}), \qquad (2.9)$$

$$\mathbf{E}_{out} = \sum_{i=1}^{\infty} \sum_{j=-i}^{i} p_{ij} \mathbf{M}_{ij}^{(1)}(k\mathbf{r}) + q_{ij} \mathbf{N}_{ij}^{(1)}(k\mathbf{r}), \qquad (2.10)$$



Fig. 2.7 Optical forces on microspheres in air as a function of the diameter of the microspheres and the NA of laser beams

where $\mathbf{M}_{ij}^{(1)}$ and $\mathbf{N}_{ij}^{(1)}$ are outward-propagating TE and TM multipole fields, and $\mathbf{M}_{ij}^{(2)}$ and $\mathbf{N}_{ij}^{(2)}$ are the corresponding inward-propagating multipole fields.

The optical force on the microsphere along the axial direction is [21]:

$$F_{z} = \frac{2n_{md}P}{cS} \sum_{i=1}^{\infty} \sum_{j=-i}^{i} \frac{j}{i(i+1)} \operatorname{Re}(a_{ij}^{*}b_{ij} - p_{ij}^{*}q_{ij}) - \frac{1}{i+1} \left[\frac{i(i+2)(i-j+1)(i+j+1)}{(2i+1)(2i+3)} \right]^{1/2} \times \operatorname{Re}(a_{ij}a_{i+1,j}^{*} + b_{ij}b_{i+1,j}^{*} - p_{ij}p_{i+1,j}^{*} - q_{ij}q_{i+1,j}^{*}), \quad (2.11)$$



Fig. 2.8 Minimum optical forces on a microsphere in air along the axial direction as a function of the diameter of the microspheres. The power of the laser is 100 mW

where

$$S = \sum_{i=1}^{\infty} \sum_{j=-i}^{i} (|a_{ij}|^2 + |b_{ij}|^2).$$
(2.12)

Figure 2.5 shows the calculated optical forces on a microsphere in water from a laser beam focused by objective lenses with three different NA's. The wavelength of the laser is 1064 nm, the power of the laser is 100 mW, and the diameter of the microsphere is 3 μ m. The optical forces along the radial direction are similar for all three NA's (NA = 0.85, 1.0, 1.25) as shown in Fig. 2.5b. On the other hand, the optical forces along the axial direction are very different for different NA's. This is because the scattering force is only along the axial direction. The microspheres will be trapped at positions where the total optical force changes its sign. The scattering force affects the trapping position (Fig. 2.5a).

Figure 2.6 shows the calculated optical forces on microspheres in air exerted by a laser beam focused by an objective lens with NA = 0.95. The maximum value of the NA for an objective lens in air is 1.0, while it is 1.33 for an objective lens in water. Because of a larger relative refractive index in air than in water, the scattering force on a microsphere in air is much larger than that in water. This makes Fig. 2.6 appear very different from Fig. 2.5. The optical forces along the axial direction are asymmetric, because the scattering forces are in the forward direction.

Figure 2.7 shows more calculation results of optical forces on microspheres in air along the axial direction. For a $D = 3.0 \,\mu\text{m}$ microsphere (Fig. 2.7a), the minimum axial force (F_z^{min}) is positive when NA = 0.85 or NA = 0.9, and is only slightly negative when NA = 0.95. Thus a laser beam focused by an objective lens with NA less than 0.95 can not trap a 3 μ m silica microsphere. The situation becomes

better for smaller microspheres. The minimum axial force is negative for a 0.5 μ m microsphere in a laser beam focused by objective lenses with all three different NA's (Fig. 2.7d).

The minimum axial forces on a microsphere in air as a function of the diameter of the microsphere are shown in Fig. 2.8. The minimum forces oscillate as the diameter of the microsphere changes. This is because of the interference between the scattered light and un-scattered light. The oscillation period is about half the wavelength of the laser inside of the microsphere, which is $\lambda/(2n_p) = 364$ nm. A microsphere can not be trapped if the minimum force is positive. For NA = 0.85, only microspheres with certain diameters can be trapped. This serves as a selection process and can be used for sorting microspheres. The size distribution of the trapped microspheres will be different from the size distribution of the microspheres before trapping. For example, if the microspheres prior trapping have a large diameter distribution ranging from 0.7 to 1.7 μ m, the diameter of microspheres trapped by a NA = 0.85 laser will always be about 1.4 μ m.

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Chapter 3 Optical Trapping of Glass Microspheres in Air and Vacuum

Optical levitation of dielectric particles in air by an upward-propagating laser beam was first demonstrated by Ashkin and Dziedzic in 1971 [1]. A few years later, optical levitation of microspheres in vacuum at pressures down to 10^{-6} torr was achieved [2]. An optical levitation trap is formed by the balance between the scattering force from an upward laser and the gravitational force on a particle. The trapping frequency of an optical levitation trap is usually very small (about 20 Hz) [2]. Optical trapping of a dielectric particle in air with the gradient force of a single laser beam was first demonstrated in 1997 [3], 11 years after the invention of the single beam gradient force optical trap in water [4]. There was no report on optical trapping of a microsphere in vacuum by a single beam gradient force trap (without the help of the gravitational force) before this thesis.

Optical trapping of a microsphere in air with the optical gradient force is much more difficult than optical trapping of a microsphere in water [5]. First, the scattering force on a microsphere from a laser beam is much larger in air than that in water, because of the larger relative refractive index of a microsphere in air (m = 1.46) than that in water (m = 1.1). There will be no stable optical trap if the scattering force is larger than the gradient force, as discussed in detail in Chap. 2. Second, microspheres in air will fall rapidly due to the influence of gravity. On the other hand, microspheres in water can be suspended for a long time. In air, we have to launch a lot of microspheres to air first and capture one of them by an optical trap passively. In water however, we can move a microsphere to the focus of a laser beam (by moving the water chamber with a translation stage) and trap it actively.

In this chapter we will first describe the method used to launch microspheres into air; we will then show our results of trapping microspheres in air and vacuum. Finally, we will describe the structure of our vacuum system.

3.1 Launching Microspheres

The silica (SiO₂) microspheres (bought from Bangs Laboratories, Inc.) are initially stuck on a glass surface and with each each other by the van der Waals' (attractive) force. The minimum required force to pull a microsphere off from a surface is called the "pull-off force". The pull-off force between a glass microsphere and a flat glass surface is predicted by the model of Derjaguin, Muller, and Toporov (DMT) to be [6, 7]:

$$F_{sphere-flat} = 4\pi R\gamma, \qquad (3.1)$$

where *R* is the radius of the microsphere and γ the effective solid surface energy. The pull-off force between two identical glass microspheres is

$$F_{sphere-sphere} = 2\pi R\gamma, \qquad (3.2)$$

which is half of the pull-off force between a microsphere and a flat surface.

The pull-off force between two 1 μ m-diameter silica microspheres is measured to be about 88 nN [7], and the pull-off force between a 1 μ m-diameter silica microsphere and a flat silica surface is about 176 nN. The gravitational force on a 1 μ m-diameter silica microsphere is only about 10 fN, and the maximum force from a typical optical tweezer is about 0.1 nN. Thus the gravitational force and a typical optical force are too small to pull off a microsphere from a glass surface. In our experiment, we launch microspheres to air by ultrasonic vibration. The required acceleration to break the van der Waals' bond between a microsphere and a flat surface is:

$$a = \frac{F_{sphere-flat}}{M} = \frac{4\pi R\gamma}{\frac{4}{3}\pi R^3 \rho_{bead}} \propto \frac{1}{R^2},$$
(3.3)

where *M* is the mass of the microsphere and ρ_{bead} the density of the microsphere. The smaller the microsphere, the larger acceleration is required to break the van der Waals bond. For a 1 µm-diameter silica microsphere, the required acceleration is about 1.8 × 10⁸ m/s²!

Because the pull-off force between two microspheres is only half of the pull-off force between a microsphere and a flat surface, the microspheres will be separated from each other before being pulled off from a glass surface by ultrasonic vibration. Thus, most of particles launched out by ultrasonic vibration are single microspheres, even if the microspheres are not mono-dispersed initially. This is a very good property for us, because we want to trap only a single microsphere at a time.

We launch single microspheres into air by a home-built ultrasonic transducer (Fig. 3.1a, also see Fig. 3.7 for a schematic of the ultrasonic generator). The transducer is a sandwich structure with a piezoelectric ring and a glass slide clamped together by two copper plates. The main component of the transducer is the piezoelectric ceramic ring with an outer diameter of 38 mm, an inner diameter of 13 mm and a thickness of 6.35 mm. It is made of lead zirconate / lead titanate (PZT) ceramic



Fig. 3.1 **a** A home-built ultrasonic transducer for launching glass microspheres into air. It has a sandwich structure consisting of a piezoelectric ring and a glass slide between a pair of copper plates. The microspheres are held on the surface of a coverslip by the van der Waals force. **b** Glass microspheres distributed on the surface of a glass coverslip after ultrasonic vibration. The diameter of the microspheres is $3.0 \,\mu$ m. The picture was taken under a $40 \times$ objective lens



(APC International Ltd.). The two flat surfaces of the piezoelectric ring are coated by silver layers that serve as two electrodes. One flat surface is in direct contact with a copper plate that is grounded. The other flat surface is isolated from the other copper plate by the glass slide whose thickness is about 1 mm. A vacuum compatible kapton coated copper wire is soldered to this flat surface for applying a high voltage to the piezoelectric ring.

A No. 2 coverslip (thickness: 0.19-0.25 mm) is attached to one end of the glass slide by an ultrahigh-vacuum compatible epoxy (Epoxy Technology, Inc. Model: EPO-TEK H77). The two-component epoxy becomes a hard ceramic after mixing and curing at 120 °C for 2 h. After curing, it can be used continuously at temperatures as high as 250 °C without bond failure. The epoxy ceramic is as hard as the glass slide so it will not reduce the mechanical quality of the transducer. The purpose of using a coverslip attached to the glass slide is to increase the vibration amplitude. The microspheres are distributed on the coverslip at the side that is facing the piezo. We found that the two sides of the coverslip behave very differently for launching microspheres. We have tested several different types of ultrasonic transducers and find that this transducer is the best one among those we have tested.

A much simpler, but less efficient ultrasonic transducer for launching glass microspheres into air is shown in Fig. 3.2. A glass slide and a glass coverslip



Fig. 3.3 Glass microspheres distributed on the surface of a glass coverslip after ultrasonic vibration. The mean diameter of the microspheres is 3.0 μ m. The *left photo* was taken under a 4× objective lens, and the *right photo* was taken under a 10x objective lens, of the region labeled 'A'

are attached to the two sides of a piezoelectric ring with an epoxy ceramic. The microspheres are applied on top of the coverslip. This ultrasonic transducer needs 2–4 times more ultrasonic power to launch microspheres than the transducer shown in Fig. 3.1a. However, this is not a problem for launching microspheres that are larger than 3 μ m. This transducer is compatible with a commercial optical microscopes. The whole transducer can be mounted on a microscope stage like a glass slide. The piezoelectric ring has a hole at the center for optical access. We studied the launching process with this ultrasonic transducer before switching to more complex transducers. Photos in Figs. 3.1b and 3.3 were taken with this transducer under a commercial microscope.

We apply dry silica microspheres directly on the coverslip of an ultrasonic transducer. If the microspheres are supplied in aqueous solution, we pipet a few drops of the aqueous solution onto a glass slide. The glass slide is heated up to about 70 °C for several minutes to get rid of water. We then scratch some microspheres from the dried glass slide and apply them to the coverslip of an ultrasonic transducer. The microspheres should be dry when they are applied to the coverslip. We find that microspheres deposited on a coverslip directly from a solution (water or acetone) cannot be launched [8, 9].

The required ultrasonic power for launching particles depends strongly on the size of the particles. We can first remove large clusters of microspheres by applying low ultrasonic power for a few seconds. When the ultrasonic power is large enough, the bonds between different microspheres are broken and the microspheres become mono-dispersed. During ultrasonic vibration, a standing wave is formed on the coverslip. Some regions of the coverslip will have enough vibration to launch microspheres while other regions of the coverslip do not have enough vibration to break the van der Waals' bonds.

Figure 3.3 shows glass microspheres distributed on the surface of a coverslip after ultrasonic vibration. The mean diameter of the microspheres is 3.0 μ m. In the left photo of the figure, we can observe two distinct regions. In region 'A', microspheres



are mono-dispersed because the ultrasonic vibration is large enough to break the van der Waals' bonds and redistribute the microspheres. In region 'B', microspheres are not mono-dispersed because the vibration in this regime is too small.

Magnified photos of microspheres in region 'A' are shown in the right photo of Figs. 3.1b and 3.3. They show that most microspheres are isolated from each other after ultrasonic vibration, thus most of the launched particles are single microspheres. The launched spheres fall under gravity, and eventually a sphere approaches the laser focus and is trapped. A coverslip fully covered with a monolayer of microspheres can be used to launch and trap single microspheres several hundred times before there are too few beads left. Since a microsphere can be trapped for many hours, we only need to reapply microspheres every few months.

Figure 3.4 is the measured impedance of a free piezoelectric ceramic ring driven by a sine wave. We have also measured the impedance of piezoelectric rings mounted in the ultrasonic generators (Figs. 3.1, 3.2). The measured impedances are almost the same as the one shown in Fig. 3.4. As shown in Fig. 3.4, the impedance has many resonance dips corresponding to different modes of ultrasonic vibrations [10]. The circumference vibration mode (expansion of the ring circumference) has a resonant frequency of 44 kHz and impedance of about 17.9 Ω at resonance. The mechanical quality factor (Q) of this mode is 362. The thickness vibration mode has a resonant frequency of about 340 kHz and impedance of about 12 Ω at resonance. The Q of this mode is 402. We use this mode for launching microspheres to air.

In order to achieve high acceleration of the coverslip without breaking the ultrasonic transducer, we drive the transducer with a high power square wave at 340 kHz for a short duration (about 10 ms). A square wave causes much larger acceleration than a sine wave (the real shape of our driving wave is very complex because the piezoelectric ring has a large capacitance and inductance). The ultrasonic power required for launching 3.0 μ m diameter microspheres is about 4 W, and it is about 130 W for 1.5 μ m diameter microspheres (Table 3.1). These are nominal values. The
Diameter (µm)	Mass (kg)	Ultrasonic power (W)	
4.7	1.1×10^{-13}	1.3	
3.0	2.8×10^{-14}	4.2	
1.9	7.2×10^{-15}	80	
1.5	3.5×10^{-15}	133	
1.0	1.0×10^{-15}	Above 500 the piezo ring starts to brea	k

 Table 3.1
 The masses of silica microspheres with different diameters and the required ultrasonic powers for launching them

The driving frequency is about 340 kHz. The ultrasonic powers shown here are nominal values. The real required powers also depend on the air pressure and number of microspheres remaining on the coverslip



required power also depends on air pressure and density of microspheres remaining on the slide. The damage threshold of the piezoelectric ceramic ring and the coverslip is several hundred watts. Thus we are only able to launch microspheres with diameter larger than 1 μ m in the present experiment.

We drive the ultrasonic transducer with a home-built high-power pulsed generator (Fig. 3.5). Since the resonant impedance of the piezoelectric ring is only about 12 Ω , we need to drive it with a large current to achieve high ultrasonic power. The main components of the pulsed generator are two 8 mF capacitors (Digi-key, Model: 338-1236-ND) to store energy, and a power MOSFET to generate pulses. The limit voltage of the capacitors is 450 V. The capacitors are connected to an external power supply which charges them. We change the output voltage of that external power supply to regulate the charge on the capacitors.

The power MOSFET is IRFPS40N50L (Vishay Siliconix). The limit voltage and current of the MOSFET are 500 V and 46 A, respectively. Its "on" gate-source voltage is 10 V, and its "on" resistance is 0.087 Ω . This power MOSFET has a very large input capacitance (8110 pF), so we cannot drive it directly by a signal generator. We use a high-current high-speed MOSFET driver (Microchip: TC4422) to drive it through the "Control" terminal in Fig. 3.5. The TC4422 has a peak output current of 9 A and can drive the power MOSFET at the required speed (340 kHz). The TC4422 itself is driven by a signal generator (Agilent 33250A) that can generate arbitrary waveforms from DC to 80 MHz. The whole system is controlled by a digital output



Fig. 3.6 A home-built high power pulsed generator. The two *blue cylinders* are capacitors (8 mF, 450 V) for storing energy to generate the high power electronic pulses. The diameter of each capacitor is 7.6 cm, and the height is 22 cm

from a computer. Since the Q of the piezoelectric ring is about 402, it is enough to drive the system with pulses at 340 kHz for 10 min (about 3,400 total pulses) to achieve maximum acceleration.

A photo of our high-power pulse generator is shown in Fig. 3.6.

3.2 Trapping Microspheres

Once microspheres are launched into air, they will fall under the influence of gravity. We can capture a microsphere with an optical trap when the microsphere passes near the optical trap. Because of air damping, a microsphere that goes into an optical trap will lose its kinetic energy and becomes trapped. After a microsphere is trapped, we can reduce the air pressure and keep the microsphere trapped in vacuum.

Optical trapping of dielectric particles in air with a single beam gradient force trap was first demonstrated in 1997 [3], requiring an objective lens with NA = 0.95. So far it is still not feasible to trap a microsphere in vacuum with a single beam gradient force trap, due to the lack of vacuum-compatible objective lenses with NA of about 0.95.

The simplest way to trap a microsphere in air and vacuum is using an optical levitation trap, which utilizes the gravitational force to assist trapping [1]. However, the optical levitation trap is not very stable and its trapping frequency is very small (on the order of 10 Hz). Thus it is not suitable for quantum ground-state cooling. We eventually decided to use a counter-propagating dual-beam gradient force trap to trap microspheres in air and vacuum. In the following sections, we will describe our experimental apparatus and results of trapping microspheres with an optical levitation trap and a dual-beam trap.



Fig. 3.7 A schematic of an optical levitation trap and an ultrasonic transducer for launching microspheres. The trap is formed by the balance between the scattering force from an upward laser beam and the gravitational force on the microsphere

3.2.1 Optical Levitation Trap

The first optical trap that we used to trap microspheres in air was an optical levitation trap. We used a diode pumped solid-state CW laser (Coherent Inc. model: Verdi V10) for trapping. Its wavelength is 532 nm, and its maximum power is 10 W. As shown in Fig. 3.7, a 532 nm laser beam with size of about 1 cm is focused by an achromatic lens whose focal length is 35 mm. The focal point of the laser beam is inside a glass cell. The width of the glass cell is about 1.0 cm and the height is about 5 cm. The thickness of the walls of the glass cell is about 1 mm. The silica microspheres are initially stuck on a coverslip above the glass cell. The gap between the coverslip and the glass cell is very small (~0.5 mm). Thus the coverslip and the glass cell form an almost sealed chamber to protect the optical trap from air flows.

We use the ultrasonic transducer to launch the microspheres to air. After launching, the microspheres fall down due to the gravitational force. The air will exert a frictional force on the microspheres, $F_{air} = 6\pi\eta_{air}R \cdot v$, where η_{air} is the viscosity of air (see Table A.1 for the values) and v is the velocity of the microsphere. The terminal speed of a microsphere is

$$v = \frac{Mg}{6\pi\eta_{air}R} \tag{3.4}$$

where *M* is the mass of the microsphere, and *g* the gravitational acceleration on the Earth. The settling speed is about 0.22 mm/s for a 1.9 μ m diameter silica bead, 0.57 mm/s for a 3.0 μ m diameter silica bead, and 1.4 mm/s for a 4.7 μ m diameter silica bead. So it takes about 3 min for a 1.9 μ m microsphere to reach the optical trap after launching. Once a microsphere is near the focus of the laser beam, it will be captured by the laser beam. In the vertical direction, the trap is formed by the balance



Fig. 3.8 A 4.7 μ m diameter silica microsphere levitated in air inside a glass cell by an upward laser beam. The *bright dot* near the center of the photo is the trapped microsphere. It appears much larger than the real size of the microsphere because of the overexposure of the camera

between the scattering force from the upward laser beam and the gravitational force on the microsphere; in the horizontal direction, the trap is formed by the gradient force from the laser beam.

A photo of a 4.7 μ m diameter microsphere levitated by a laser beam in air is displayed in Fig. 3.8. We can change the laser power from 2 W to 0.4 W while keeping the microsphere trapped. As expected, the vertical position of a trapped microsphere changes when we change the laser power. When we increase the laser power, the trapped microsphere moves up. Surprisingly, we found that we could also change the vertical position of a trapped microsphere by driving the ultrasonic transducer. By driving with different ultrasonic frequencies, we could drag the microsphere either up or down. This suggests that the pressure force from the ultrasonic wave may be enough to levitate the microsphere. We have not studied this phenomenon in detail.

The optical levitation trap is pretty simple. However, it has several drawbacks. First, the laser beam must be upward. This is a problem because thousands of microspheres must be launched into air for each microsphere which is trapped. Those microspheres will fall to the bottom of the glass cell. They will be in the path of the laser beam and make the optical trap unstable. We can only launch and trap microspheres a few times before the bottom of the glass cell becomes too dirty. Second, the optical levitation trap tends to trap several microspheres at one time [11]. It is very difficult to trap only one microsphere at a time, or control the number of the trapped microspheres. Third, the trapping frequency of the optical levitation trap is too small. It is usually on the order of 10 Hz [2]. This is very close to the mechanical vibration frequencies of experimental instruments, and can cause heating of the motion of a trapped microsphere. It is important that the trapping frequency be much higher than the frequencies of seismic vibration in order to achieve significant cooling.

3.2.2 Counter-Propagating Dual-Beam Optical Trap

The ideal optical trap for our experiment will be a single beam gradient force trap. However, it requires an objective lens with a NA of about 0.95, which is currently not available for use in vacuum. A high NA objective lens usually has a very short working distance (usually about 200 μ m), which is not good for us. We need the working distance to be long enough to allow optical accesses for feedback cooling and cavity cooling.

On the other hand, a low NA lens usually has a long working distance but can not focus a laser beam strongly enough to create a gradient force trap. The scattering force from a weakly focused laser will be larger than the gradient force. One way to overcome this problem is using two counter-propagating laser beams (Fig. 3.9). The scattering forces from the two beams cancel, and the gradient forces forms a stable 3D trap for the microsphere.

We built a simple dual-beam optical trap (Fig. 3.9) to test its suitability for using in vacuum. As shown in Fig. 3.9, we use two achromatic lenes to focus two laser beams that are propagating in the horizontal direction. The focal lengths of the two lenses are 35 and 30 mm, respectively. Because the launched microspheres fall down to the bottom of the glass cell and do not affect the light path, this dual-beam trap is much more stable than the optical levitation trap. We can trap microspheres with diameters of 1.9, 3.0, and 4.7 μ m in air repeatedly. We have also filled the glass cell with water and trapped microspheres in water with this dual-beam trap. This is remarkable because the walls of the glass cell and the water distort the laser beam significantly. Dual-beam optical tweezers are also suitable for trapping particles with high-refractive index in water [12].

The major disadvantage of a counter-propagating dual-beam trap is that the two laser beams must be aligned correctly. Ideally, the two beams should be exactly counter-propagating and focused at the same point. If the two beams are misaligned from each other, the resulting trap will not be a simple harmonic trap. The scattering force from the laser beams is not conservative, and can do work on the microsphere under certain conditions [13, 14]. When the two beams are misaligned (Fig. 3.10),



Fig. 3.9 A counter-propagating dual-beam optical trap in air



Fig. 3.10 A possible motion of a microsphere trapped in a misaligned dual-beam optical tweezer

the scattering force from the left laser beam will push the microsphere to the right, and the scattering force from the right beam will push the microsphere to the left. This results in a cycled motion. In air (or water), the resulted complex motion can be stable due to the air (or water) damping. We have observed this phenomenon in both air and water. In vacuum, however, the mechanical energy of the microsphere will keep increasing until the microsphere escapes from the trap eventually. This problem can be minimized by aligning the two beams correctly.

As the dual-beam trap in air was much more stable than the optical levitation trap in air, we decided to use the dual-beam trap for trapping microspheres in vacuum. Fig. 3.11 shows a simplified schematic of the first generation of our



Fig. 3.11 A simplified schematic of the first generation of our counter-propagating dual-beam optical trap for trapping microspheres in vacuum, and a detection system to monitor the position of a trapped microsphere. The two 1,064 nm laser beams are orthogonally polarized and have different frequencies to avoid interference

counter-propagating dual-beam optical trap in vacuum. The two 1,064 nm laser beams for trapping are from a fiber laser whose maximum output power is 10 W. The two laser beams are orthogonally polarized to avoid interference. Their powers are controlled by two 80 MHz acousto-optic modulators (AOM's). The power of beam No. 1 can be changed from 3 mW to 2 W, and the power of beam No. 2 can be changed from 3 mW to 3 W. One beam is the +1 order of the output beams from the AOM, and the other beam is the -1 order of the output beams from the AOM. Thus the frequencies of the two laser beams differ by 160 MHz. The two laser beams are focused by two identical aspheric lenses whose focal lengths are 3.1 mm. The two lenses are inside a vacuum chamber.

We collect the scattered light from a trapped microsphere with a lens from the side to monitor the microsphere by a camera. We also use a weak 532 nm laser beam (about 2 mW) and a quadrant detector to monitor the motion of the microsphere with high resolution. The quadrant detector splits the 532 nm beam into four parts. The difference between the left (top) two parts and the right (bottom) two parts provides the position information of the microsphere in the horizontal (vertical) direction, and the summation of the four parts provides the position information of the microsphere along the axial direction [15]. Thus the quadrant detector can monitor the position of a trapped microsphere in 3D.

The setup inside our vacuum chamber is shown in Fig. 3.12. The ultrasonic transducer (Fig. 3.1a) is mounted vertically. The coverslip with microspheres is above the optical trap by about 5 cm. A pair of aspheric lenses (Newport, model: KGA330-B) with focal length of 3.1 mm are fixed on stainless steel structures with vacuum compatible epoxy. The stainless steel structures are mounted on an aluminum holder by stainless steel screws. The separation between the front surfaces of the two lenses is about 3.5 mm.



Fig. 3.12 Top view of the setup inside the vacuum chamber. A pair of *aspheric lenses* with focal length of 3.1 mm, and a *piezoelectric ultrasonic transducer* are inside the vacuum chamber

3.2 Trapping Microspheres



We align the laser beams in three steps. First, we send a parallel 1,064 nm laser beam through the two lenses. One of the two lenses is fixed. We change the position of the other lens to make sure that the output beam after the two lenses is still parallel. Second, we align the directions and divergences of all three laser beams (two 1,064 nm beams and one 532 nm beam) to make sure that all beams are on top of each other. Third, we align all laser beams with a pinhole to make sure that the foci of the three beams are at the same point (Fig. 3.13).

The setup for aligning a laser beam and measuring its waist with a pinhole is shown in Fig. 3.13. This setup is good for aligning the trapping beam No. 1 and the detection beam. We move the photodiode and the lens in front of it to the other side of the vacuum chamber for aligning the trapping beam No. 2. The diameter of the pinhole aperture is 1.0 \pm 0.5 μ m. The pinhole is mounted on a 3-axis translation stage. The translation stage is controlled by 3 manual differential actuators (Thorlabs: DM10) with graduation of $0.5 \,\mu$ m. In the vertical direction, we also have a piezoelectric actuator to fine tune the pinhole. We first align the pinhole to the focus of the trapping beam No. 1 (by aligning the pinhole to maximize the output power of the beam from the pinhole). The pinhole from National Aperture is easier to align than the high energy pinhole from Newport, because the pinhole from National Aperture is located at the center of a small (about 200 μ m) countersink, which can be seen by the naked eye and acts as a guide for alignment. Once the pinhole is aligned to the focus of the trapping beam No. 1, we fine tune the directions and divergences of the detection beam and the trapping beam No. 2 to maximize their output powers from the pinhole. With this method, we can align the focuses of all three laser beams to the same point with an accuracy of about $0.5 \,\mu\text{m}$.

After alignment, we can scan the pinhole to measure the profiles of the laser beams. Fig. 3.14 shows an example of measured profiles of the beam No. 1 along the horizontal and vertical directions. The measured beam profiles agree with Gaussian functions very well. The measured waists of the beam are 2.01 μ m in the horizontal direction and 2.12 μ m in the vertical direction. The real waists of the beam should be smaller than these values because of the finite size of the pinhole aperture.

A 4.7 μ m diameter microsphere trapped inside a vacuum chamber by a counterpropagating dual-beam optical tweezer is displayed in Fig. 3.15. The trapping laser is infrared. A weak green laser (about 2 mW) is used for illumination. The bright spot near the center of the photos is the trapped microsphere. It usually take us about 10 min to trap one microsphere. Once a microsphere is trapped, the system is very



Fig. 3.14 An example of measured beam profiles of the beam No. 1 in the *horizontal* and *vertical* directions. The *red curves* are Gaussian functions



Fig. 3.15 A 4.7 μ m diameter microsphere trapped inside a vacuum chamber by a counterpropagating dual-beam optical tweezer. The wavelength of the trapping beams is 1064 nm. A *weak green* (532 nm) laser is used for illumination



Fig. 3.16 *Left:* A single optically trapped 3.0 μ m diameter bead; *Right:* multiple (2 or maybe 3) optically trapped 3.0 μ m beads. The air pressure is 752 torr. The powers of the two counterpropagating beams are 119 and 100 mW

stable. We have tested it by trapping a 4.7 μ m bead in air continuously for 46 h, during which time the power of both laser beams was varied between 5 mW and 2.0 W to test the stability of the trap. The trap becomes less stable in vacuum. The lowest pressure at which we have trapped a bead without requiring extra stabilization is about 0.1 Pa.

We find that the waists of the two counter-propagating laser beams determine how many microspheres will be trapped by the dual-beam optical tweezer at one time. When the two beams have the same beam waist, we usually trap more than one microsphere (usually two microspheres) at a time. This is because the two counterpropagating laser beams form a double-well potential when their foci are not exactly on top of each other. When the waist of one of the laser beams is larger than the other (for example, 3.0 μ m for one beam, and 2.0 μ m for the other beam), we usually trap only one microsphere at a time. The left photo in Fig. 3.16 shows a single 3.0 μ m diameter bead trapped by the dual-beam optical tweezer, and the right photo in Fig. 3.16 shows multiple (2 or maybe 3) 3.0 μ m beads trapped at the same time.

3.3 Vacuum System

The science chamber (the vacuum chamber in which we trap microspheres) and most of optic components are mounted on a 3×3 foot breadboard. The breadboard is supported by elastomers (Newport, Model: NewDamp) on an air-floated optical table to minimize the mechanical vibration. The peak-to-peak vibration amplitude of the lab floor is about 30 μ m due to seismic vibrations. The peak-to-peak relative vibration amplitude of optical elements on an optical table is about 0.1 μ m (in 10 s) when the table is sitting on rigid legs directly. This is reduced to about 10 nm when the optical table is supported by pneumatic isolators (air-floated). Thus it is essential to use pneumatic isolators to reduce vibrations.



Fig. 3.17 A schematic of the vacuum system

Figure 3.17 shows a schematic of our vacuum system. The science chamber on the breadboard is connected to a small ion pump on the optical table by a flexible stainless steel bellows to isolate mechanical vibrations. The ion pump is connected to roughing pumps on the lab floor by a bellows. We initially use a sorption pump for rough pumping to avoid mechanical vibrations. The sorption pump reduces air pressure by absorbing molecules within a porous material (molecular sieve) which is cooled by liquid nitrogen. There are no mechanical movable parts in the sorption pump. Thus it has almost no mechanical vibration. The lowest pressure that we can obtain with a single sorption pump is about 3 mtorr (1 mtorr = 0.133 Pa). Using two sorption pumps to pump the vacuum chamber sequentially, we can reduce the pressure from 760 torr to about 10^{-5} torr in 1 h.

The sorption pump needs to be heated up to about 200 °C to drive off absorbed water before using and requires liquid nitrogen to cool it during use, thus it is not convenient to use the sorption pump. We later switched to using a mechanical roughing pump in place of the sorption pump. There are 4 bellows separated by 3 rigid structures between the mechanical pump and the science chamber. They isolates the mechanical vibration sufficiently. There is no observed effect on the motion of a trapped microsphere when we turn on and off the mechanical roughing pump. The mechanical roughing pump can reduce the air pressure from 760 torr (1 atm) to 1 mtorr in a few minutes.

The ion pump can reduce the air pressure from 1 mtorr to about 10^{-6} torr in 1 h without bakeout. At first, the ion pump tended to kick out the trapped microsphere immediately after the pump was turned on. This was caused by momentary electrical arcing (accompanied by a sudden pressure burst) in the ion pump [16]. We later moved the pump further away, and this problem disappeared after we used the ion pump for a few months. With bakeout, one should be able to achieve pressures much

lower than 10^{-6} torr with the ion pump. The ion pump is not ideal for using with pressures above 10^{-6} torr. Since we are able to isolate the mechanical vibration, a turbomolecular pump will be a better choice in future.

We have a cold cathode gauge and a combination gauge near the science chamber to measure the air pressure inside the science chamber. The cold cathode gauge (Kurt J. Lesker Company, model: KJLC 943) can measure the pressure between 10^{-10} and 10^{-2} torr. The combination gauge (Kurt J. Lesker Company, model: KJLC 910) can measure the pressure between 10^{-5} and 1,500 torr. The combination gauge has a Piezo sensor and a Pirani sensor. The Pirani sensor is gas-type sensitive as it measures pressure based on the thermal conductivity of the gas. The Piezo sensor measures the absolute pressure independent of the gas type. The gauge reads the Piezo sensor when the pressure is above 15 torr, reads the Pirani sensor when the pressure is below 5 torr, and reads both sensors when the pressure is in between. The accuracy of the measurement is 1 % of the reading when the pressure is in the range of 10^{-1} ,000 torr, and is about 10 % of the reading for lower pressures. We also have a thermocouple gauge (TC gauge) near the sorption pump to monitor the pressure in the rough pumping region.

There are several manual angle valves between the science chamber and the roughing pumps that can be use to isolate different parts of the system. The vacuum chamber is also connected to a pure nitrogen (or air) gas tube by a gas shut off valve (Kurt J. Lesker Company, model: Nupro BK, SS-4BK). With these valves and vacuum pumps, we can control the pressure in the science chamber from 10^{-6} to 1,500 torr easily. We can achieve much lower pressure with bakeout. The ability to control air pressure allows us to study the Brownian motion of a trapped microsphere at different pressures, and cool the motion of a trapped microsphere in vacuum.

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Chapter 4 Measuring the Instantaneous Velocity of a Brownian Particle in Air

Brownian motion of particles affects many branches of science. The Brownian motion was discovered by Brown in 1827 [1], and explained correctly by Einstein in 1905 [2]. The trajectories of a Brownian particle are commonly considered to be not differentiable. This is accompanied by the fact that the instantaneous velocity of a Brownian particle has not been measured successfully before. Here we will present our measurement of the instantaneous velocity of a Brownian particle in air.

In this chapter, we will first introduce some theories about the Brownian motion of free particles and trapped microspheres. This will be followed by a description of our home-built detection system that can study the Brownian motion of an optically trapped microsphere in air with an ultrahigh resolution. We will show some measured power spectra of the motion of trapped microspheres at different air pressures. Finally, we will show the results of our measurement of the instantaneous velocity of a Brownian particle in air. Our results provide direct verification of the Maxwell-Boltzmann velocity distribution, and the equipartition theorem for a Brownian particle.

4.1 Theories of Brownian Motion

4.1.1 A Free Particle

Einstein's theory of Brownian motion predicts that

$$\langle [\Delta x(t)]^2 \rangle \equiv \langle (x(t)) - (x(0))^2 \rangle = 2Dt, \qquad (4.1)$$

where $\langle [\Delta x(t)]^2 \rangle$ is the mean-square displacement (MSD) of a free Brownian particle in one dimension during time *t*, and *D* is the diffusion constant [2]. The diffusion constant can be calculated by $D = k_B T / \gamma$, where *T* is the temperature, and $\gamma = 6\pi\eta R$ is the Stokes friction coefficient for a sphere with radius *R*. Here η is the viscosity of the fluid. The mean velocity measured over an interval of time *t* is $\bar{v} \equiv \sqrt{\langle [\Delta x(t)]^2 \rangle}/t = \sqrt{2D}/\sqrt{t}$. This diverges as *t* approaches 0, and therefore does not represent the real velocity of the particle [3, 4].

The equation $\langle [\Delta x(t)]^2 \rangle = 2Dt$, however, is only valid when $t \gg \tau_p$, i.e., in the diffusive regime. Here $\tau_p = M/\gamma$ is the momentum relaxation time of a particle with mass M. At very short time scales $(t \ll \tau_p)$, the dynamics of a particle is dominated by its inertia, and the motion is ballistic. The dynamics of a Brownian particle with mass M over all time scales can be described by a Langevin equation [5–7]:

$$M\frac{d^2x}{dt^2} + \gamma \frac{dx}{dt} = F_{therm}(t), \qquad (4.2)$$

where

$$F_{therm}(t) = (2k_B T \gamma)^{1/2} \zeta(t) \tag{4.3}$$

is the Brownian stochastic force. $\zeta(t)$ is a normalized white-noise process. Hence for all t and t',

$$\langle \zeta(t) \rangle = 0$$
, and $\langle \zeta(t)\zeta(t') \rangle = \delta(t-t')$. (4.4)

Let v(t) = dx/dt, and divide the Eq. 4.2 by M, we obtain

$$\frac{dv(t)}{dt} = -\Gamma_0 v(t) + A(t), \qquad (4.5)$$

where $\Gamma_0 = \gamma/M = 1/\tau_p$ is the damping coefficient and $A(t) = F_{therm}(t)/M$ is the fluctuating acceleration. The velocity and position of the particle at time t = 0 are $v(0) = v_0$ and $x(0) = x_0$, respectively. Then its velocity at time t is [7]:

$$v(t) = v_0 e^{-\Gamma_0 t} + e^{-\Gamma_0 t} \int_0^t e^{\Gamma_0 s} A(s) ds.$$
(4.6)

Taking the mean over an ensemble of particles, which have started at t = 0 with the same velocity v_0 , and using Eq. 4.4 we get:

$$\langle v(t) \rangle_{v_0} = v_0 e^{-\Gamma_0 t},$$
(4.7)

$$\langle v(t)v(0)\rangle_{v_0} = v_0^2 e^{-\Gamma_0 t}.$$
 (4.8)

Taking a second average over v_0 and using the energy equipartition theorem $\langle M v_0^2/2 \rangle = k_B T/2$, we obtain the velocity autocorrelation function:

$$\langle v(t)v(0)\rangle = \frac{k_B T}{M} e^{-\Gamma_0 t}.$$
(4.9)

By integrating Eq. 4.6 again, we get the position of the particle [7]:

$$x(t) = x_0 + \frac{v_0}{\Gamma_0} (1 - e^{-\Gamma_0 t}) + \int_0^t e^{-\Gamma_0 s_1} ds_1 \int_0^{s_1} e^{\Gamma_0 s_2} A(s_2) ds_2.$$
(4.10)

Taking the average, we get:

$$\langle \Delta x(t) \rangle_{x_0} = \langle x(t) - x(0) \rangle_{x_0} = \frac{v_0}{\Gamma_0} (1 - e^{-\Gamma_0 t}).$$
 (4.11)

The MSD for a Brownian particle at thermal equilibrium with the air is [7]:

$$\langle [\Delta x(t)]^2 \rangle = \frac{2k_B T}{M\Gamma_0^2} (\Gamma_0 t - 1 + e^{-\Gamma_0 t}).$$
(4.12)

At long time scales, the MSD is the same as the prediction of Einstein's theory:

$$\langle [\Delta x(t)]^2 \rangle = 2Dt \quad \text{for} \quad t \gg \tau_p.$$
 (4.13)

At very short time scales, the MSD is

$$\langle [\Delta x(t)]^2 \rangle = \frac{k_B T}{M} t^2 \quad \text{for} \quad t \ll \tau_p.$$
(4.14)

Although the above equations are derived for an ensemble of identical particles, the ergodic theorem dictates that they are also valid for measurements of a single particle taken over a long time.

At very short time scales, the motion is ballistic and its instantaneous velocity can be measured as $v = \Delta x(t)/t$, when $t \ll \tau_p$ [7]. The ballistic Brownian motion is different from a simple ballistic motion. For a simple ballistic motion with velocity u, we have $\Delta x(t) = ut$ and $[\Delta x(t)]^2 = u^2 t^2$. The velocity u can be any value and usually has no relation with the temperature of the environment.

The 1D Maxwell-Boltzmann distribution of the velocity of a particle in thermal equilibrium is

$$f_v(v_i) = \sqrt{\frac{M}{2\pi k_B T}} \exp\left(-\frac{M v_i^2}{2k_B T}\right),\tag{4.15}$$

where v_i is the velocity of the particle along any direction *i*.

4.1.2 A Trapped Microsphere

For small displacements, the effect of optical tweezers on the microsphere's motion can be approximated by that of a harmonic potential. The equation of the Brownian motion of a microsphere in a harmonic trap is:

$$\frac{d^2x}{dt^2} + \Gamma_0 \frac{dx}{dt} + \Omega^2 x = \Lambda \zeta(t), \qquad (4.16)$$

where $\Omega = \sqrt{\kappa/m}$ is the natural angular frequency of the trapped microsphere when there is no damping, and $\Lambda = (2k_B T \Gamma_0/M)^{1/2}$. The cyclic frequency of the damped oscillator is $\omega_1 = \sqrt{\Omega^2 - \Gamma_0^2/4}$. The system is underdamped when ω_1 is real $(\Omega > \Gamma/2)$, critically damped when $\omega_1 = 0$, and overdamped when ω_1 is imaginary $(\Omega < \Gamma/2)$.

4.1.2.1 Displacement and Velocity

The MSD of a Brownian particle in an underdamped harmonic trap in air can be obtained by solving Eq. 4.16 [8]:

$$\langle [\Delta x(t)]^2 \rangle = \frac{2k_B T}{M\Omega^2} \left[1 - e^{-t/2\tau_p} \left(\cos \omega_1 t + \frac{\sin \omega_1 t}{2\omega_1 \tau_p} \right) \right].$$
(4.17)

The position autocorrelation function is related to the MSD by:

$$\langle [\Delta x(t)]^2 \rangle = 2\langle x^2 \rangle - 2\langle x(t)x(0) \rangle, \qquad (4.18)$$

where $\langle x^2 \rangle = k_B T / (M\Omega^2)$. The rms amplitude is $x_{rms} = \sqrt{k_B T / (M\Omega^2)}$. The normalized position autocorrelation function (PACF) of the particle is [8] (Fig. 4.1):

$$\frac{\langle x(t)x(0)\rangle}{\langle x^2\rangle} = e^{-t/2\tau_p} \left(\cos\omega_1 t + \frac{\sin\omega_1 t}{2\omega_1 \tau_p}\right).$$
(4.19)

The normalized velocity autocorrelation function (VACF) of the particle is [8]:

$$\frac{\langle v(t)v(0)\rangle}{\langle v^2\rangle} = e^{-t/2\tau_p} \left(\cos\omega_1 t - \frac{\sin\omega_1 t}{2\omega_1\tau_p}\right). \tag{4.20}$$

Both the position autocorrelation function and the velocity autocorrelation function oscillate for an underdamped system.

For an overdamped system, ω_1 is imaginary. We can rewrite the above formula for the normalized position and velocity autocorrelations as [9]:

$$\frac{\langle x(t)x(0)\rangle}{\langle x^2\rangle} = \frac{1}{2|\omega_1|\tau_+} e^{-t/\tau_-} - \frac{1}{2|\omega_1|\tau_-} e^{-t/\tau_+}, \qquad (4.21)$$



Fig. 4.1 The rms amplitude (x_{rms}) of a microsphere in a harmonic trap with trapping frequency $\Omega/(2\pi)$ at thermal equilibrium with the environment

$$\frac{\langle v(t)v(0)\rangle}{\langle v^2\rangle} = -\frac{1}{2|\omega_1|\tau_-}e^{-t/\tau_-} + \frac{1}{2|\omega_1|\tau_+}e^{-t/\tau_+},\tag{4.22}$$

where

$$\tau_{\pm} = \frac{2\tau_p}{1 \pm 2\tau_p |\omega_1|}.$$
(4.23)

Thus the correlations in an overdamped system decrease as a double-exponential with characteristic times τ_{\pm} . A numerical simulation of the dynamics of harmonic oscillator with different damping can be found in Ref. [9].

4.1.2.2 Power Spectrum Analysis

Similar to the optical spectrum of an atom, the power spectrum of the Brownian motion of a trapped microsphere contains a lot of information about the system. The power spectral density (PSD) of a variable is the squared modulus of its Fourier transform [8-10].

The Fourier transformations of x(t) and $\zeta(t)$ are:

$$\widetilde{x_k} = \int_{-T_{rec}/2}^{T_{rec}/2} e^{i\omega_k t} x(t) dt, \qquad (4.24)$$

$$\widetilde{\zeta}_k = \int_{-T_{rec}/2}^{T_{rec}/2} e^{i\omega_k t} \zeta(t) dt, \qquad (4.25)$$

where $\omega_k = 2\pi k/T_{rec}$, k is an integer, and T_{rec} is the time duration of the recorded positions of a trapped microsphere.

Fourier transforming Eq. 4.16 gives

$$-\omega_k^2 \widetilde{x}_k - i\omega_k \Gamma_0 \widetilde{x}_k + \Omega^2 \widetilde{x}_k = \Lambda \widetilde{\zeta}_k.$$
(4.26)

Thus

$$\widetilde{x_k} = \frac{\Lambda \widetilde{\zeta_k}}{\Omega^2 - \omega_k^2 - i\omega_k \Gamma_0}.$$
(4.27)

From Eq. 4.4 we have $\langle \tilde{x}_k \rangle = 0$ and $\langle \tilde{\zeta}_k \tilde{\zeta}_l \rangle = T_{rec} \delta_{kl}$. The PSD of an experimentally recorded x(t) is

$$S_{k}^{rec} \equiv |\tilde{x}_{k}|^{2} / T_{rec} = \frac{|\tilde{\zeta}_{k}|^{2}}{T_{rec}} \frac{2k_{B}T}{M\Omega^{2}} \frac{\Omega^{2}\Gamma_{0}}{(\Omega^{2} - \omega_{k}^{2})^{2} + \omega_{k}^{2}\Gamma_{0}^{2}}.$$
 (4.28)

The expected values of the PSD is

$$S(\omega) \equiv \langle S_k^{rec} \rangle = \frac{2k_B T}{M\Omega^2} \frac{\Omega^2 \Gamma_0}{(\Omega^2 - \omega^2)^2 + \omega^2 \Gamma_0^2}.$$
(4.29)

The real measured PSD of a recorded x(t) is S_k^{rec} . It has a $|\zeta_k|^2/T_{rec}$ term, which is a random number. Thus an experimental PSD will appear noisy. Averaging many measured S_k^{rec} will result in a spectrum close to the expected spectrum $S(\omega)$. Another way to reduce the noise in a measured spectrum is "blocking" [10]. Here, a "block" of consecutive data points $(\omega_{k_1}, S_{k_1}^{rec}) \dots (\omega_{k_2}, S_{k_2}^{rec})$ is replaced with a single new "data point" $(\overline{\omega_k}, \overline{S_k^{rec}})$ which are the block averages.

Let us define a new function to show the shape of the spectrum

$$f_S(\omega) \equiv \frac{\Omega^2 \Gamma_0}{(\Omega^2 - \omega^2)^2 + \omega^2 \Gamma_0^2}.$$
(4.30)

We have

$$\int_0^\infty f_S(\omega)d\omega = \frac{\pi}{2},\tag{4.31}$$

thus

$$\int_0^\infty S(\omega)d\omega = \pi \frac{k_B T}{M\Omega^2} = \pi < x^2 > .$$
(4.32)



Fig. 4.2 Calculated spectra $(f_S(\omega))$ for systems with different damping. The plot is displayed in linear scales



Fig. 4.3 Calculated spectra $(f_S(\omega))$ for systems with different damping. The plot is displayed in log-log scales

So the integral of the PSD is proportional to the square of the rms amplitude. Some calculated spectra $f_S(\omega)$ for systems with different damping are shown in Fig. 4.2 (in linear scales) and Fig. 4.3 (in log-log scales). As dramatically displayed in the linear scale plot (Fig. 4.2), the spectra have large values at low frequencies for an overdamped system, and have large values at the resonant frequency for an underdamped systems.

4.2 A Fast Detection System

In order to measure the instantaneous velocity of the Brownian motion of a trapped microsphere in air, we implemented an ultrahigh resolution detection system.

Our first generation of the dual-beam optical trap in vacuum (Fig. 3.11) used a 10 W fiber laser at 1064 nm for trapping, and a weak 532 nm laser for detection. We used a quadrant detector to monitor the position of a trapped microsphere in 3D. That system was not ideal for our experiment. The peak-to-peak intensity noise (in the range of 10 kHz–1 MHz) of the fiber laser is about 4 %, which is very big. The 532 nm laser has much less noise than the fiber laser. However, the quadrant detector has significant electronic noise and its bandwidth is only about 1 MHz.

In the second-generation experiment with the dual-beam optical trap in vacuum, we use an ultra-stable NPRO laser (Model: 126-1063-700, Lightwave Electronics (now JDSU)) to replace the fiber laser. Its rms intensity noise is <0.05 % over the range from 10 Hz to 2 MHz, and is shot noise limited above 10 MHz. It is a single frequency laser with a linewidth <5 kHz/ms and a coherence length longer than 1000 m. A detailed characterization of this type of laser can be found in Ref. [11]. These values are much better than those of the 532 nm laser. So we use this laser for both trapping and detection. This is achieved by using a polarizing beam splitter cube to reflect one of the trapping beams for detection (Fig. 4.4).

Our lab has previously developed a fast position-sensitive laser beam detector for studying the Brownian motion of particles in water at fast time scales [12]. The previous detector used a fiber-optic bundle that spatially splits the incident beam, and a fast balanced photodetector to measure the difference between the two halves of the beam. Here we simplify the detection system by using a mirror with a sharp edge (BBD05-E03, Thorlabs) to replace the fiber-optic bundle for splitting the beam. The sharp edge of a mirror is much smoother than the boundary between the two halves



Fig. 4.4 This simplified schematic shows our counter-propagating dual-beam optical tweezers, and a novel detection system that has 75 MHz bandwidth and low noise. The *s*-polarized beam is reflected by a polarizing beam splitter cube after it passes through a trapped bead inside a vacuum chamber. Then, for detection, it is split by a mirror with a sharp edge. The *p*-polarized beam passes through the cube

of a fiber-optic bundle. Thus a mirror with a sharp edge is not only much simpler, but also has less noise than a fiber bundle for splitting the laser beam [12].

We use a balanced detector (PDB120C, photodiode diameter 0.3 mm, Thorlabs) with a bandwidth of 75 MHz for detection. The detector is sensitive to light with wavelengths in the range of 800–1700 nm. It has a high transimpedance gain of 1.8×10^5 V/A. The detector measures the difference between the two halves of the beam, which is proportional to the particle excursion. The intensity noise of the laser is contained in both halves and is thus canceled in the measurement. This detection system enables us to monitor the real-time position of a trapped microsphere with Ångstrom spatial resolution and microsecond temporal resolution.

The major optical circuit of our second generation dual-beam trap and the detection system is shown in Fig. 4.5. The powers of the two trapping beams and the probe beam are controlled by AOM's. We use photodiodes to monitor the powers of the beams. With measured powers, we use analog proportional-integral-derivative (PID) circuits to stabilize the laser powers. This eliminates the drifts of the laser powers. The size of the beam No. 1 before entering the vacuum chamber is twice of the size of the beam No. 2. This ensures that the optical tweezer traps only one microsphere at a time. The green laser at 532 nm is helpful for aligning the laser beams and monitoring microspheres before they are trapped. Once a microsphere is trapped, we can see it on a VCR monitor connected to the CCD camera. The quadrant detector and the balanced detector give high resolution signals of the motion of a trapped microsphere. For studying the Brownian motion, we turned off the 532 nm laser, and use only the balanced detector to monitor the position of a trapped microsphere with ultrahigh precision.

4.3 Measured Power Spectra

Figure 4.6 shows a waveform and a spectrum of a 3.0 μ m diameter bead trapped by a dual-beam optical tweezer at 749 torr. The powers of the two trapping beams are 10.7 and 14.1 mW. The data were taken by a DAQ card (PCI 6133, National Instruments) controlled by a Labview program. The waveform and spectrum shown here were saved from the Labview program directly. The sampling rate was 2 MHz, and the time duration was 2 s. So the waveform has 4 million data points. As we can see, the waveform appears as a white noise and has very little information.

The power spectrum of the waveform, on the other, is much more instructive. The power spectrum in Fig. 4.6 is in log-log scales. Comparing it with the spectra in Fig. 4.3, we can find that its trapping frequency $(\Omega/(2\pi))$ is about 3 kHz, and the linewidth $(\Gamma_0/(2\pi))$ is close to the trapping frequency. At high frequencies near 1 MHz, the shape of the measured spectrum is very different from the shape of the expected spectrum as shown in Fig. 4.3. This is because at frequencies near 1 MHz, the real signal due to the Brownian motion of the trapped microsphere is smaller than the spectrum of noise (laser noise, electronic noises of the detector and the



Fig. 4.5 The optical circuit of the counter-propagating dual-beam optical trap and the detection system

DAQ card). So the displayed spectrum near 1 MHz is the spectrum of noise rather than the real signal.

Figure 4.7 shows a waveform and spectrum of a 3.0 μ m diameter bead trapped by a dual-beam optical tweezer at 20.6 torr. Other conditions are the same as those in Fig. 4.6. Because of less damping, a peak appears at the trapping frequency. The spectrum at high frequencies (around 100 kHz) is also smaller than those in Fig. 4.6, in agreement of the theoretical predications (Fig. 4.3).



Fig. 4.6 A waveform (*top*) and a spectrum (*bottom*) of a 3.0 μ m bead trapped at 749 torr. The powers of the two trapping beams are 10.7 and 14.1 mW

Sometimes we can trap more than one microsphere at the same time (Fig. 3.16). Figure 4.8 shows a waveform and spectrum of multiple (2 or 3) microspheres trapped at 752 torr. The powers of the two laser beams are 119 and 100 mW. The spectrum has new peaks at frequencies around 300 kHz. These peaks are not presented when there is only one microsphere. They are due to the relative motion between the trapped microspheres. The frequencies of relative motion are much larger than the frequency of the center-of-mass motion (about 1.5 kHz) in this example.

The spectra of the relative motion between multiple microspheres depend strongly on the details of the optical trap, such as the alignment of the laser beams and the size of the microspheres. Figure 4.9 shows the spectra of two 4.7 μ m beads trapped at 9.78 torr with different laser power. The frequency of the highest peak is about 70 kHz when the powers of the two trapping beams are 0.7 and 1.0 W, and is about 150 kHz when the powers of the two trapping beams are 2.0 and 2.5 W. The data in Fig. 4.9



Fig. 4.7 A waveform (*top*) and a spectrum (*bottom*) of a 3.0 μ m bead trapped at 20.6 torr. The powers of the two trapping beams are 10.7 and 14.1 mW

were taken in our first generation dual-beam trap with a probe beam (Fig. 3.11). The red curves are for the motion in the horizontal direction, and the blue curves are for the motion in the vertical direction. Under certain conditions, we can even observe such narrow peaks at 1 atm. It seams that the scattering forces from the trapping laser pump energy to the motion and amplify the motion.

4.4 Measurement of the Instantaneous Velocity of a Brownian Particle in Air

A simplified scheme of our setup for measuring the instantaneous velocity of Brownian particle in air is shown in Fig. 4.4, and a more detailed optical circuit is shown in Fig. 4.5. The trap is formed inside a vacuum chamber by two counter-



Fig. 4.8 A waveform and spectrum of multiple (2 or maybe 3) 3.0 μ m beads trapped at 752 torr. The powers of the two trapping beams are 119 and 100 mW

propagating laser beams focused to the same point by two identical aspheric lenses with focal length of 3.1 mm and numerical aperture of 0.68. The two 1064 nm laser beams are orthogonally polarized, and their frequencies differ by 160 MHz to avoid interference.

The two laser beams are aligned with the help of a pinhole aperture whose diameter is $1.0 \pm 0.5 \,\mu$ m. Because the pinhole has a finite thickness (13 μ m), it is difficult to align the foci of the two beams to the same point in the axial direction. We intentionally make the waist of one beam larger than the other to make this alignment less critical. The measured waists of the two beams are 2.2 and 3.0 μ m in the horizontal direction. The real waists should be smaller than these values due to the finite size of the pinhole. Also note that the waists of the two beams are measured at different axial positions separated by the thickness of the pinhole. Once a bead is



Fig. 4.9 Spectra of two 4.7 μ m beads trapped at 9.78 torr with different laser powers. The powers of the two trapping beams are 0.7 and 1.0W for the *top figure*, and 2.0 and 2.5W for the *bottom figure*

trapped, we keep the power of one beam constant, and tune the power of the other beam to maximize the trapping frequency.

When the bead deviates from the center of the trap, it deflects both trapping beams. We monitor the position of the bead by measuring the deflection of one of the beams, which is split by a mirror with a sharp edge. The difference between the two halves is measured by a fast balanced detector. This simple, yet novel, detection scheme has a bandwidth of 75 MHz and ultra-low noise [12, 13].

For the data included in this section, unless otherwise stated, the powers of the two laser beams are 10.7 and 14.1 mW, the diameter of the bead is 3μ m, the temperature of the system is 297 K, and the air pressure is 99.8 kPa (749 torr) or 2.75 kPa (20.6 torr). The trapping is very stable, and the heating due to laser absorption is negligible at these conditions. Examples of power spectra of a 3μ m microsphere trapped at these conditions have been shown in Figs. 4.6 and 4.7.



Fig. 4.10 One-dimensional trajectories of a 3 μ m diameter silica bead trapped in air at 99.8 kPa (a) and at 2.75 kPa (b). The instantaneous velocities of the bead corresponding to these trajectories are shown in (c) and (d)

The position signal of a trapped bead is recorded at a sampling rate of 2 MHz. Because of the limited spatial resolution, we are not able to obtain accurate instantaneous velocities of a bead at this rate. To reduce the noise, we average every 10 successive position measurements, and use these averages to calculate instantaneous velocities with time resolution of 5 μ s. Although this method reduces the temporal resolution by a factor of 10, it greatly increases the signal-to-noise ratio if both the trapping period ($2\pi/\omega_0$) and momentum relaxation time are much larger than 5 μ s. These conditions are satisfied here since the trapping period is about 320 μ s, $\tau_p = 48 \,\mu$ s at 99.8 kPa, and $\tau_p = 147 \,\mu$ s at 2.75 kPa.

Figure 4.10 shows typical samples of position and velocity traces of a trapped bead. The position traces of the bead at these two pressures appear very similar. On the other hand, the velocity traces are clearly different. The instantaneous velocity of the bead at 99.8 kPa changes more frequently than that at 2.75 kPa, because the momentum relaxation time is shorter at higher pressure.

Figure 4.11 shows the mean square displacements of a 3 μ m silica bead as a function of time. The measured MSD's fit excellently with Eq. 4.17 over three decades of time for both pressures. The calibration factor $\alpha = position/voltage$ of the detection system is the only fitting parameter of Eq. 4.17 for each pressure. *M* is calculated from the size and density of the microsphere. τ_p and Ω are obtained from the measured normalized velocity autocorrelation function (VACF). The two α 's obtained for these two pressures differ by 10.8%. This is because the vacuum



Fig. 4.11 a The mean square displacements of a 3 μ m silica bead trapped in air at 99.8 kPa (*red square*) and 2.75 kPa (*black circle*). They are calculated from 40 million position measurements for each pressure. The "noise" signal (*blue triangle*) is recorded when there is no particle in the optical trap. The *solid lines* are the theoretical predictions of Eq. 4.17. They fit with the measurements excellently. The prediction of Einstein's theory of free Brownian motion in the diffusive regime is shown in *dashed lines* for comparison. **b** MSD's at short time scales are shown in detail. The *dash-dot line* indicates ballistic Brownian motion of a free particle

chamber is distorted slightly when the pressure is decreased from 99.8 to 2.75 kPa. One may avoid this problem in future by coupling the trapping laser into the vacuum chamber with an optical fiber and a teflon feedthrough [14]. The measured MSD's are completely different from those predicted by Einstein's theory of Brownian motion in a diffusive regime. The slopes of measured MSD curves at short time scales are double of those of the MSD curves of diffusive Brownian motion in the log-log plot (Fig. 4.11a). This is because the MSD is proportional to t^2 for ballistic Brownian motion, and it is proportional to t for diffusive Brownian motion. Another important feature is that the MSD curves are independent of air pressure at short time scales, as is predicted by $\langle [\Delta x(t)]^2 \rangle = (k_B T/M) t^2$ for ballistic Brownian motion, whereas the MSD in the diffusive regime does depend on the air pressure. At long time scales, the MSD saturates at a constant value because of the optical trap. Figrue 4.11b displays more detail of the Brownian motion at short time scales. It clearly demonstrates that we have observed ballistic Brownian motion.



Fig. 4.12 The distribution of the measured instantaneous velocities of a $3 \,\mu\text{m}$ silica bead. The statistics at each pressure are calculated from 4 million instantaneous velocities. The *solid lines* are Maxwell-Boltzmann distributions. We obtained $v_{rms} = 0.422 \,\text{mm/s}$ at 99.8 kPa (*red square*) and $v_{rms} = 0.425 \,\text{mm/s}$ at 2.75 kPa (*black circle*) from the measurements. The rms value of the noise (*blue triangle*) is 0.021 mm/s, which means we have 1.0 Å spatial resolution in 5 μ s

The distributions of the measured instantaneous velocities are displayed in Fig. 4.12. They agree very well with the Maxwell-Boltzmann distribution. The measured rms velocities are $v_{rms} = 0.422 \text{ mm/s}$ at 99.8 kPa and $v_{rms} = 0.425 \text{ mm/s}$ at 2.75 kPa. These are very close to the prediction of the energy equipartition theorem, $v_{rms} = \sqrt{k_B T/M}$, which is 0.429 mm/s. As expected, the velocity distribution is independent of pressure. The rms value of the noise signal is 0.021 mm/s, which means we have 1.0Å spatial resolution in $5\,\mu s$. This measurement noise is about 4.8% of the rms velocity. Figure 4.12 represents direct verification of the Maxwell-Boltzmann distribution of velocities and the equipartition theorem of energy for Brownian motion. For a Brownian particle in liquid, the inertial effects of the liquid become important. The measured rms velocity of the particle will be $v_{rms} = \sqrt{k_B T / M^*}$ in the ballistic regime, where the effective mass M^* is the sum of the mass of the particle and half the mass of the displaced fluid [15]. In order to measure the true instantaneous velocity in liquid as predicted by the equipartition theorem, the temporal resolution must be much shorter than the time scale of acoustic damping, which is less than 1 ns for a 1 μ m particle in liquid [15].

Figure 4.13 shows the normalized VACF of the bead at two different pressures. They fit with Eq. 4.20 nicely. At 2.75 kPa, one can clearly see the oscillations due to the optical trap. Equation 4.20 is independent of the calibration factor α of the detection system. The only independent variable is time *t*, which we can measure with high precision. Thus the normalized VACF provides an accurate method to measure τ_p and ω_0 . Fitting the normalized VACF with Eq. 4.20, we obtained $\tau_p =$ $48.5 \pm 0.1 \ \mu$ s, $\omega_0 = 2\pi \cdot (3064 \pm 4)$ Hz at 99.8 kPa, and $\tau_p = 147.3 \pm 0.1 \ \mu$ s, $\omega_0 = 2\pi \cdot (3168 \pm 0.5)$ Hz at 2.75 kPa. The trapping frequency changed by 3 % due to the distortion of the vacuum chamber at different pressures. We can also calculate the diameter of the silica bead from the τ_p value at 99.8 kPa [16]. The obtained diameter for this microsphere is 2.79 μ m. This is within the uncertainty range given



by the supplier of 3.0 μ m silica beads. We use this value in the calculation of MSD and normalized VACF.

For a particle at a certain pressure and temperature, τ_p should be independent of the trapping frequency. We verified this by changing the total power of the two laser beams from 25 to 220 mW. The measured τ_p of a microsphere trapped at 19.6 and 749 torr as a function of the total laser power is shown in Fig. 4.14. Each data point has a few percent of uncertainty because we use a smaller data sets to calculate the τ_p than those used in the previous figures. Although the data points for each pressure are not perfectly on a line, it is clear that the τ_p 's are independent of the laser power within the experimental uncertainty. Fitting the data for each pressure with a straight line, we obtain $\tau_p = [151.3 + 0.00168 P/(1mW)] \,\mu s$ at 19.6 torr, and $\tau_p = [53.74 + 0.00275 P/(1mW)] \,\mu s$ at 749 torr for this microsphere, where P is the total power of the two trapping beams. Thus τ_p changed less than 1.3 % for both pressures when the total laser power is changed from 0 to 200 mW. This proves that the fitting method is very accurate, and the heating due to the laser beams (which would change the viscosity and affect τ_p) is negligible at these pressures.

In conclusion, we have observed the Brownian motion of a single particle in the ballistic regime, and measured its instantaneous velocity successfully for the first





time. The ability to measure the instantaneous velocity of a Brownian particle will be invaluable in studying nonequilibrium statistical mechanics [17, 18] and can be used to cool Brownian motion by applying a feedback force with a direction opposite to the velocity [19, 20].

In vacuum, our optically trapped particle promises to be an ideal system for investigating quantum effects in a mechanical system [13, 21], due to its nearperfect isolation from the thermal environment. Combining feedback cooling and cavity cooling, we expect to cool the Brownian motion of a bead starting from room temperature to the quantum regime, as predicted by recent theoretical calculations [22, 23]. We have directly verified the energy equipartition theorem of Brownian motion. However, we also expect to observe deviation from this theorem when the bead is cooled to the quantum regime. The kinetic energy of the bead will not approach zero even at 0 K because of its zero-point energy. The rotational energy of the bead should also become quantized.

An optically trapped microsphere can also be used to probe the properties of superfluid helium. Liquid helium has many special properties. Its density is only 0.129 g/cm^3 , and its refractive index is only 1.024, which is close to that of air (Table A.1). Thus optical trapping of microspheres in liquid helium is similar to optical trapping of microspheres in air. A microsphere in superfluid helium will do Brownian motion due to collisions between the microsphere and quasiparticles, such as phonons and rotons [24–26]. Thus we can probe the properties of superfluid helium by studying the Brownian motion of an optically trapped microsphere in it.

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Chapter 5 Towards Measurement of the Instantaneous Velocity of a Brownian Particle in Water

5.1 Motivation

In the previous chapter, we presented our measurement of the instantaneous velocity of a Brownian particle in air. A more interesting experiment would be to measure the instantaneous velocity of a Brownian particle in water (or other liquid). Brownian motion was first discovered in water, and it affects many aspects of the life of cells in water. More importantly, Brownian motion in water may reveal new physics that has not been explored before.

It is clear that the velocity distribution of a Brownian particle in air satisfies the Maxwell–Boltzmann distribution. Since the Maxwell–Boltzmann distribution is derived for an ideal gas (or a weakly interacting system), it is not clear whether it holds equally well in water. In water, a moving microsphere drags water along with it as it moves. If the moving microsphere is suddenly stopped, the water flow caused by the previous motion of the microsphere will drag the microsphere to keep it moving. Thus the presence of water adds a memory effect to the motion of the bead. The water will also add an effective mass to the microsphere, since accelerating the microsphere requires a force both on the microsphere and the water which it displaces as it moves.

5.2 Hydrodynamic Theories of Brownian Motion

5.2.1 A Free Particle in Water

The effective mass of the microsphere in water is the sum of the mass of the microsphere and half of the mass of the displaced water [1, 2]:

$$M^* = M_p + \frac{1}{2}M_f, (5.1)$$

59

where $M_p = (4/3)\pi R^3 \rho_p$ is the mass of the microsphere, $M_f = (4/3)\pi R^3 \rho_f$ is the mass of displaced water, ρ_p is the density of the microsphere, and ρ_f is the density of water. The energy equipartition theorem needs to be modified to:

$$\frac{1}{2}M^*\langle v^2\rangle = \frac{1}{2}k_BT \tag{5.2}$$

where v is the velocity of the microsphere in one dimension. Thus the rms velocity is $v_{rms} = \sqrt{k_B T/M^*}$. Because of the memory effect of water, the velocity autocorrelation function (VACF) of a free particle will not be $\langle v(t)v(0)\rangle = \frac{k_B T}{M}e^{-t/\tau_p}$ as in air, but [3–6]

$$\frac{\langle v(t)v(0)\rangle}{k_BT/M^*} = \frac{\alpha_+ e^{\alpha_+^2 t} \operatorname{erfc}(\alpha_+ \sqrt{t}) - \alpha_- e^{\alpha_-^2 t} \operatorname{erfc}(\alpha_- \sqrt{t})}{\alpha_+ - \alpha_-},$$
(5.3)

where

$$\alpha_{\pm} = \frac{3}{2} \cdot \frac{3 \pm (5 - 36\tau_p/\tau_f)^{1/2}}{\tau_f^{1/2}(1 + 9\tau_p/\tau_f)}.$$
(5.4)

 $\tau_p = M_p/(6\pi\eta R) = \frac{2}{9}R^2\rho_p/\eta$ is the momentum relaxation time of the particle due to its own inertia, $\tau_f = R^2\rho_f/\eta$ characterizes the effect of water. Here η is the viscosity of water and R is the radius of the microsphere.

At long time scales, Eq. 5.3 approaches

$$\frac{\langle v(t)v(0)\rangle}{k_B T/M^*} \propto \frac{1}{t^{3/2}} \text{ for } t \to \infty.$$
(5.5)

At short time scales, Eq. 5.3 approaches

$$\frac{\langle v(t)v(0)\rangle}{k_B T/M^*} = \exp\left(-b\sqrt{t/\tau_f}\right) \text{ for } t \to 0,$$
(5.6)

where

$$b = \frac{18}{\sqrt{\pi}(1 + 2\rho_p/\rho_f)}$$

For a silica microsphere in water, b = 2.03. The normalized VACF approaches 1 at short time scales as \sqrt{t} , rather than t. This is very different from the case in air.

The mean square displacement of a free microsphere in water is [7, 8]:

$$\langle [\Delta x(t)]^2 \rangle_{free} = 2Dt \left[1 - 2\sqrt{\frac{1}{\pi} \frac{\tau_f}{t}} + \frac{8}{9} \frac{\tau_f}{t} - \frac{\tau_p}{t} + \Xi \left(\frac{\tau_p}{\tau_f}, \frac{t}{\tau_f} \right) \right], \quad (5.7)$$

where $D = k_B T / (6\pi \eta R)$ is the diffusion coefficient. $\Xi(\frac{\tau_p}{\tau_f}, \frac{t}{\tau_f})$ is a correction term:

$$\Xi\left(\frac{\tau_p}{\tau_f}, \frac{t}{\tau_f}\right) = \frac{3}{t(5\tau_f - 36\tau_p)^{1/2}} \left(\frac{1}{\alpha_+^3} e^{\alpha_+^2 t} \operatorname{erfc}(\alpha_+\sqrt{t}) - \frac{1}{\alpha_-^3} e^{\alpha_-^2 t} \operatorname{erfc}(\alpha_-\sqrt{t})\right).$$
(5.8)

The behavior of the MSD at the long time limit is:

$$\langle [\Delta x(t)]^2 \rangle = 2Dt \quad \text{for} \quad t \gg \tau_p.$$
 (5.9)

Recently, the velocity autocorrelation function of a Brownian particle in water was measured successfully for $\frac{\langle v(t)v(0)\rangle}{\langle v^2 \rangle} < 0.35$ [6, 9]. However, the instantaneous velocity of a Brownian particle in water will be much more difficult to measure than the velocity autocorrelation function, and has not been measured to date. This is because the velocity autocorrelation function is a statistical average, which is insensitive to the high-frequency noise in the measurement. In our experiment, we hope that we can measure the instantaneous velocity of a Brownian particle in water for the first time, with particular interest to test the modified Maxwell–Boltzmann velocity distribution:

$$f_{v}(v_{i}) = \sqrt{\frac{M^{*}}{2\pi k_{B}T}} \exp\left(-\frac{M^{*}v_{i}^{2}}{2k_{B}T}\right).$$
(5.10)

5.2.2 An Optically Trapped Microsphere in Water

The optical trap provides a harmonic force $F_{trap} = -kx$ on the microsphere when the displacement of the microsphere is small. $k = M_p \Omega^2$ where Ω is the natural angular frequency of the trap. Clercx and Schram [5] gave analytical solutions for the MSD and VACF of a trapped Brownian particle in a liquid, and Berg-Sørensen and Flyvbjerg [10] gave a solution for the power spectrum density (PSD) of a trapped Brownian particle in a liquid. This section introduces their analytical solutions and provides some numerical results to visualize those solutions. These numerical results will serve as a guide for our experiment.

Because the velocity of the Brownian motion of a microsphere in liquid is much smaller than the speed of sound in the liquid, we can describe the fluid motion by the linearized incompressible time-dependent Navier–Stokes equation. The Langevin equation of the motion of a trapped microsphere in a liquid is [5]:

$$M^*\ddot{x}(t) = -kx(t) - 6\pi\eta R\dot{x}(t) -6R^2 \sqrt{\pi\rho_f \eta} \int_{-\infty}^t (t - t')^{-1/2} \ddot{x}(t') dt' + F_{therm}(t).$$
(5.11)
The first term after the equal sign of Eq. 5.11 is the harmonic force, the second term is the ordinary Stokes's friction, the third term is a memory term associated with the hydrodynamic retardation effects of the liquid, and the last term is the Brownian stochastic force.

The mean-square displacement of a trapped microsphere in a liquid is [5, 8]

$$\langle [\Delta x(t)]^2 \rangle_{trap} = \frac{2k_B T}{k} + \frac{2k_B T}{M^*} \bigg[\frac{e^{z_1^2 t} \operatorname{erfc}(z_1 \sqrt{t})}{z_1(z_1 - z_2)(z_1 - z_3)(z_1 - z_4)} \\ + \frac{e^{z_2^2 t} \operatorname{erfc}(z_2 \sqrt{t})}{z_2(z_2 - z_1)(z_2 - z_3)(z_2 - z_4)} \\ + \frac{e^{z_3^2 t} \operatorname{erfc}(z_3 \sqrt{t})}{z_3(z_3 - z_1)(z_3 - z_2)(z_3 - z_4)} \\ + \frac{e^{z_4^2 t} \operatorname{erfc}(z_4 \sqrt{t})}{z_4(z_4 - z_1)(z_4 - z_2)(z_4 - z_3)} \bigg]$$
(5.12)

The coefficients z_1 , z_2 , z_3 , and z_4 are the four roots of the Eq. [8]

$$\left(\tau_p + \frac{1}{9}\tau_f\right)z^4 - \sqrt{\tau_f}z^3 + z^2 + \frac{1}{\tau_k} = 0,$$
(5.13)

where $\tau_k = 6\pi \eta R/k$. For $t \to \infty$, Eq. 5.12 approaches

$$\langle [\Delta x(\infty)]^2 \rangle_{trap} = \frac{2k_BT}{k}.$$

The normalized VACF of a trapped microsphere in a liquid is [5, 8]

$$A(t) = \frac{\langle v(t)v(0) \rangle}{k_B T/M^*} = \frac{z_1^3 e^{z_1^2 t} \operatorname{erfc}(z_1 \sqrt{t})}{(z_1 - z_2)(z_1 - z_3)(z_1 - z_4)} + \frac{z_2^3 e^{z_2^2 t} \operatorname{erfc}(z_2 \sqrt{t})}{(z_2 - z_1)(z_2 - z_3)(z_2 - z_4)} + \frac{z_3^3 e^{z_3^2 t} \operatorname{erfc}(z_3 \sqrt{t})}{(z_3 - z_1)(z_3 - z_2)(z_3 - z_4)} + \frac{z_4^3 e^{z_4^2 t} \operatorname{erfc}(z_4 \sqrt{t})}{(z_4 - z_1)(z_4 - z_2)(z_4 - z_3)}.$$
 (5.14)

The power spectral density is [8, 10]:

$$S(f) = \frac{D}{2\pi^2 f^2} \frac{1 + \sqrt{f/2\phi_f}}{(\phi_k/f - \sqrt{f/2\phi_f} - f/\phi_p - f/9\phi_f)^2 + (1 + \sqrt{f/2\phi_f})^2},$$
(5.15)

where f is the observation frequency, $\phi_k = 1/(2\pi\tau_k)$ is the corner frequency of the power spectrum due to the trap, and $\phi_{p,f} = 1/(2\pi\tau_{p,f})$. For $f \to 0$, Eq. 5.15 approaches

$$S(0) = \frac{2k_B T \gamma}{k^2},$$

where $\gamma = 6\pi\eta R$.

At $t \to 0$, Eq. 5.14 predicts $\langle v(0)v(0) \rangle = k_B T/M^*$, which is different from the energy equipartition theorem $\langle v(0)v(0) \rangle = k_B T/M_p$. This conflict is caused by the assumption in Eq. 5.11 that the liquid is incompressible. For $t < t_c$, we need to consider the liquid to be compressible. Here $t_c = R/c$ is the time required for a sound wave to travel a sphere radius, where *c* is the speed of sound in the liquid. The normalized velocity autocorrelation function at $t \sim t_c$ is [1]:

$$A(t) = \frac{\langle v(t)v(0) \rangle}{k_B T/M^*} = 1 + \frac{M_f}{2M_p} \left[\frac{1}{2} - \frac{iM^*}{(4M_p^2 - M_f^2)^{1/2}} \right] e^{-ix_1 t/t_c} + \frac{M_f}{2M_p} \left[\frac{1}{2} + \frac{iM^*}{(4M_p^2 - M_f^2)^{1/2}} \right] e^{-ix_2 t/t_c}, \quad (5.16)$$

where

$$x_1 = -i\frac{M^*}{M_p} + \left[1 - \frac{M_f^2}{4M_p^2}\right]^{1/2},$$
(5.17)

$$x_2 = -i\frac{M^*}{M_p} - \left[1 - \frac{M_f^2}{4M_p^2}\right]^{1/2}.$$
(5.18)

At very short time scales $t \ll t_c$, Eq. 5.16 approaches $A(0) = 1 + \frac{M_f}{2M_p}$. The short time limit $A(0) \neq 1$ because the normalization factor is $k_B T/M^*$, rather than $k_B T/M_p$.

Diameter	k	τ_p	τ_f	τ_k	τ_c
(µm)	(µN/m)	$\frac{2}{9}R^2\rho_p/\eta$ (µs)	$R^2 \rho_f / \eta$ (µs)	$6\pi\eta R/\kappa$ (µs)	R/c (ns)
1.0	100	0.11	0.25	94	0.34
3.0	33.3	1.0	2.2	851	1.01
4.7	21.3	2.45	5.51	2,083	1.58
10	10	11.1	25.0	9443	3.4

Table 5.1 Characteristic time scales of an optically trapped silica microsphere in water at 20° C

The second row shows the definitions of $\tau_{p,f,k,c}$. The second column (*k*) is the spring constant of the optical trap. It is assumed to be inversely proportional to the diameter of the microsphere when the laser power is constant



Fig. 5.1 Mean square displacement of an optically trapped silica microsphere in water at 20° C. Parameters are the same as those in Table 5.1



Fig. 5.2 Power spectra of an optically trapped silica microsphere in water at 20° C. Parameters are the same as those in Table 5.1

The MSD's of microspheres with different diameters in water are shown in Fig. 5.1, and the corresponding power spectra are shown in Fig. 5.2. Figure 5.3 displays the normalized velocity autocorrelation function (A(t)) of an optically trapped silica microsphere in water at 20° C. Figure 5.3a displays the VACF's with a linear-log scale to cover a large range of time scales, and Fig. 5.3b displays the VACF's in a log-linear scale to show the details at short time scales. The thin solid lines at short time scales ($t < 10^{-8}$ s) are calculated from Eq. 5.16, which includes the compressibility effects of water. The thick solid lines are calculated from Eq. 5.14, which treats the water as an incompressible fluid. The dashed lines are exponential



Fig. 5.3 Normalized velocity autocorrelation function of an optically trapped silica microsphere in water at 20° C. Parameters are the same as those in Table 5.1. **a** Covers large range of time scales, and **b** shows the details at short time scales. The thin solid lines ($t < 10^{-8}$ s) are calculated from Eq. 5.16, and the thick solid lines are calculated from Eq. 5.14. The dashed lines are exponential decays with $\tau_p = 1.0 \,\mu$ s, corresponding to a 3.0 μ m microsphere

decays with $\tau_p = 1.0 \,\mu$ s, corresponding to a 3.0 μ m microsphere. As clearly shown in Fig. 5.3, the VACF of a microsphere in water is very different from exponential decay because of the hydrodynamic memory effects of water.

The thin solid lines are expected to be correct for $t \sim t_c$, and the thick sold lines are expected to be correct for $t \gg t_c$. The intermediate regime $t_c < t < 100t_c$ is still poorly understood. It is suspicious that the thick solid curves approaches 1 so slowly at the short time limit. The normalized VACF a 4.7 µm diameter silica microsphere in water at 20 and 60° C is shown in linear–linear scales in Fig. 5.4. The VACF follows $\exp(-b\sqrt{t/\tau_f})$ at the short time limit, rather than $\exp(-t/\tau_p)$.



Fig. 5.4 Normalized velocity autocorrelation function of a 4.7 μ m diameter silica microsphere in water at 20 and 60° C. The *solid lines* are exact results of the hydrodynamic theory of Brownian motion in a liquid (Eq. 5.14). The *dashed lines* are $\exp(-b\sqrt{t/\tau_f})$, which is the short time limit of the hydrodynamic theory (Eq. 5.6). The *dashed dot line* is an exponential decay $\exp(-t/\tau_p)$

A recent experiment has measured the VACF of a Brownian particle in water at VACF < 0.35 [9]. A measurement of the VACF between 1 and 0.35 is required in order to better understand the hydrodynamic effects and compressibility effects of water on Brownian motion [11].

5.3 Requirements for Measuring the Instantaneous Velocity

In order to measure the instantaneous velocity of a Brownian particle in water, the temporal resolution of the detection system must be much shorter than the momentum relaxation time of the particle, and the spatial resolution of the detection system must be much smaller than the displacement of the particle during the measurement time.

Because of the inertia of the particle and the water, the effective momentum relaxation time is

$$\tau_p^* = \frac{M^*}{6\pi\eta R}.\tag{5.19}$$

In order to resolve the instantaneous velocity with 10% uncertainty, we need the temporal resolution to be at least $\Delta t = \tau_p^*/10$. The average displacement of the microsphere during Δt is $\tau_p^* v_{rms}/10$. Thus the required spatial resolution is $\Delta x = \tau_p^* v_{rms}/100$. The sensitivity of a detector is usually characterized by its noise spectral

Table 5.2 Minimum required detection resolution		20° C		(60° C		
for measuring the instantaneous velocity of a	Diameter (µm)	$\frac{\Delta t}{(\mu s)}$	Δx (pm)	$\frac{s_x}{(\text{fm/Hz}^{1/2})}$	$\frac{\Delta t}{(\mu s)}$	Δx (pm)	$\frac{s_x}{(\text{fm/Hz}^{1/2})}$
Brownian particle in water at	1.0	0.014	2.4	0.28	0.029	5.5	0.94
20 and 60° C	1.9	0.049	3.3	0.74	0.10	7.6	2.5
	3.0	0.12	4.2	1.5	0.26	9.6	4.9
	4.7	0.30	5.2	2.9	0.65	12	9.6
	10	1.4	7.6	8.9	2.9	17	30

density s_x . The spectral density of the detector should satisfy $s_x < \Delta x \sqrt{\Delta t}$ at the required frequency $1/\Delta t$ in order to measure the instantaneous velocity.

Table 5.2 shows the minimum required resolution for measuring the instantaneous velocity of a Brownian particle in water. To measure the instantaneous velocity of a 1 μ m diameter microsphere in water at 20° C, the noise spectral density of the detection system should be smaller than 0.28 fm/Hz^{1/2} around 71 MHz. The requirement is relaxed to 0.94 fm/Hz^{1/2} around 34 MHz if the temperature of the water is increased to 60° C. This is mainly because the viscosity of the water decreases when the temperature increases (Table A.1). Thus it is easier to measure the instantaneous velocity at 60° C than at 20° C. t_p^* is bigger for a larger particle, so the requirements for measuring the instantaneous velocity of a large microsphere is less demanding than that of a small microsphere. However, the sensitivity of the detection system varies for measuring the instantaneous velocity.

The resolution of particle tracking with an optical tweezer is fundamentally limited by the shot noise of the laser. Because the size of the microsphere is on the same order as the wavelength of the laser, an accurate calculation of the shot noise limited sensitivity is very complex. Here we only discuss a simple estimation. The shot noise limited noise spectral density is estimated to be [12]

$$s_x \approx 5 \times 10^{-16} G\left(\frac{R}{1\mu m}\right) \left(\frac{100 mW}{P}\right)^{1/2} \left(\frac{1\mu m}{\lambda}\right)^{1/2} m/\sqrt{Hz},$$
 (5.20)

where *G* is a geometrical factor that depends on R/λ , R/w, and the precise details of the Mie scattering by the sphere. λ is the wavelength of the laser, *w* is the waist of the laser beam, and *P* is the power of the laser.

If the wavelength of the laser is 1 μ m and the power is 100 mW, we need G < 1.1 to measure the instantaneous velocity of a 1 μ m diameter microsphere at 20° C. We need G < 2.5 to measure the velocity of a 4.7 μ m diameter microsphere at the same conditions, and if the temperature is 60° C the requirement is relaxed to G < 8.2. This means that our detection system must be very close to the shot noise limit in order to measure the instantaneous velocity of a Brownian particle in water.

The discussions in the previous paragraphs assume that the normalized VACF is $\exp(-t/\tau_p^*)$ at short time scales. However, the normalized VACF is most likely $\exp(-b\sqrt{t/\tau_f})$ at short time scales. This will require a temporal resolution of $\Delta t =$

 $\tau_f/(100 b^2)$ in order to measure the instantaneous velocity. Our optical tweezer will not be able to measure the instantaneous velocity of a Brownian particle in water if the normalized VACF follows $\exp(-b\sqrt{t/\tau_f})$ even at such short time scales. On the other hand, the normalized VACF is much easier to measure than the instantaneous velocity. We should be able to measure the normalized VACF between 0.35 and 1 even if we are not able to measure the instantaneous velocity.

In the presence of detection noise, the measured position of the microsphere can be expressed as

$$x_{msr}(t) = x_p(t) + x_n(t),$$
 (5.21)

where $x_p(t)$ is the real position of the microsphere, and $x_n(t)$ is the noise of the detection system. The mean square displacement (MSD) of the measured positions is [6]:

$$MSD_{msr}(t) = \langle [x_{msr}(t_0 + t) - x_{msr}(t_0)]^2 \rangle$$

= $\langle [x_p(t_0 + t) - x_p(t_0)]^2 \rangle + \langle [x_n(t_0 + t) - x_n(t_0)]^2 \rangle$
+ $2 \langle [x_p(t_0 + t) - x_p(t_0)] \cdot [x_n(t_0 + t) - x_n(t_0)] \rangle$
= $MSD_p(t) + MSD_n(t).$ (5.22)

The derivation assumes no correlation between the real position of the microsphere and the detection noise. In this case, the real MSD of the microsphere $(MSD_p(t))$ can be obtained by subtracting the noise $MSD_n(t)$ from the measured $MSD_{msr}(t)$, as is done in Ref. [9]. In reality, there may be a small correlation between the motion of the bead and the detection noise. Thus it is important to minimize the noise $MSD_n(t)$ in the measurement.

The measured velocity of the microsphere is

$$v_{msr}(t) = \frac{x_{msr}(t + \frac{\delta t}{2}) - x_{msr}(t - \frac{\delta t}{2})}{\delta t} = \frac{x_p(t + \frac{\delta t}{2}) - x_p(t - \frac{\delta t}{2})}{\delta t} + \frac{x_n(t + \frac{\delta t}{2}) - x_n(t - \frac{\delta t}{2})}{\delta t} = v_p(t) + v_n(t),$$
(5.23)

where $\Delta t \ll \tau_p$. Because the measured velocity contains a noise signal $v_n(t)$, the smallest possible δt is not the optimal value for measuring the velocity. The DAQ card creates noise when it converts an analog signal to a digital signal due to the finite number of bits. The minimum value of $x_n(t + \frac{\delta t}{2}) - x_n(t - \frac{\delta t}{2})$ is limited by the DAQ card, thus $v_n(t)$ may be larger than the real velocity of the microsphere $v_p(t)$ if δt is too small.

The measured velocity represents the real instantaneous velocity of the microsphere if $v_n(t)$ is negligible. This requires $\langle v_{msr}^2 \rangle \gg \langle v_n^2 \rangle$. One can check whether this condition is satisfied by comparing the signal when a microsphere is trapped in the optical tweezer and when there is not microsphere in the optical tweezer. The relation

between $\langle v_{msr}^2 \rangle$ and $\langle v_n^2 \rangle$ can also be obtained from the measured *MSD*:

$$\langle v_{msr}^2 \rangle = \left\langle \frac{\left[x_{msr}(t + \frac{\delta t}{2}) - x_{msr}(t - \frac{\delta t}{2}) \right]^2}{\delta t^2} \right\rangle$$

$$= \frac{MSD_{msr}(\delta t)}{\delta t^2}$$

$$= \frac{MSD_p(\delta t)}{\delta t^2} + \frac{MSD_n(\delta t)}{\delta t^2}$$

$$= \langle v_p^2 \rangle + \langle v_n^2 \rangle.$$

$$(5.24)$$

Thus $\langle v_{msr}^2 \rangle \gg \langle v_n^2 \rangle$ is equivalent to $MSD_{msr}(\delta t) \gg MSD_n(\delta t)$.

The measured velocity autocorrelation function is

$$\langle v_{msr}(t+t_0)v_{msr}(t_0)\rangle = \langle v_p(t+t_0)v_p(t_0)\rangle + \langle v_n(t+t_0)v_n(t_0)\rangle$$
(5.25)

Since the noise of the detection system has almost no correlation, the last term of this equation can be neglected. Thus

$$\langle v_{msr}(t+t_0)v_{msr}(t_0)\rangle \doteq \langle v_p(t+t_0)v_p(t_0)\rangle.$$
(5.26)

So the measurement of the velocity autocorrelation function is not sensitive to the noise of the detection system. On the other hand, the measurement of the instantaneous velocity is very sensitive to the noise of the detection system.

If the detection system samples the position of the microsphere every dt that is much shorter than the required temporal resolution Δt , we can reduce the noise in the measured velocity by using successively averaged positions to calculate the velocity. Let $\delta t = N dt$ ($N \ll \tau_p/dt$), then

$$x_{avr}(t) = \frac{1}{N} \sum_{j=1}^{N} x_{msr} \left(t + jdt - (N+1)\frac{dt}{2} \right).$$
(5.27)

The measured velocity becomes

$$v_{msr}(t) = \frac{x_{avr}(t + \frac{\delta t}{2}) - x_{avr}(t - \frac{\delta t}{2})}{\delta t}.$$
(5.28)

Then the velocity noise is

$$v_n(t) = \frac{1}{N^2 dt} \left[\sum_{j=1}^N x_n \left(t + j dt - (N+1) \frac{dt}{2} + \frac{\delta t}{2} \right) - \sum_{j=1}^N x_n \left(t + j dt - (N+1) \frac{dt}{2} - \frac{\delta t}{2} \right) \right].$$
 (5.29)

On average, the rms amplitude of $v_n(t)$ is $N\sqrt{N}$ times smaller than that of $[x_n(t + dt/2) - x_n(t - dt/2)]/dt$ if the position noise $x_n(t)$ is white noise.

5.4 A Simple Optical Tweezer in Water

Figure 5.5 shows a simplified schematic of an optical tweezer for trapping a microsphere in water and studying its Brownian motion with ultrahigh resolution. The trapping laser passes through a $\lambda/2$ waveplate and a polarizing beam splitter cube (PBS1), which control the power of the laser. The laser is then reflected by a dichroic mirror (DM2) and enters an objective lens (OB2) to form an optical tweezer inside a sample chamber at the focus of the objective lens. We use another objective lens (OB1) to collect the laser for detection. The sample chamber and the objective OB1 are both mounted on 3-axis translation stages. The motion of a trapped microsphere causes deflection of the trapping laser. Thus we can monitor the posi-



Fig. 5.5 Simplified schematic of an optical setup for trapping microspheres in water. PBS1 and PBS2 are polarizing beam splitter cubes, DM1 and DM2 are dichroic mirrors, OB1 and OB2 are $100 \times$ oil-immersion objective lenses, MX is a mirror with a sharp edge that splits the beam into two parts horizontally, and DAQ is a data acquisition card that is installed in a computer

tion of the microsphere by measuring the deflection of the laser beam. After passing through another $\lambda/2$ waveplate and beam splitter (PBS2) to reduce its power, the laser is split by a mirror with a sharp edge. The difference between the two halves is measured by a fast balanced detector [13, 14]. The output signal of the detector is proportional to the position of a trapped microsphere, and is collected by a fast data acquisition card (DAQ). We use a white LED for illumination to take bright field images by a camera for real-time imaging of the microspheres. The filter before the camera blocks the scattered light from the trapping laser.

The trapping laser is an ultrastable NPRO laser (Model: 126-1063-700, Lightwave Electronics (now JDSU)) with a wavelength of 1064 nm. The maximum power is about 800 mW. Its rms intensity noise is < 0.05 % over the range from 10 Hz to 2 MHz, and is shot noise limited above 10 MHz. PBS1 is a high-energy laser-line polarizing cube beamsplitter from CVI Melles Griot (PBSO-1064-50, CW damage threshold: 1 MW/cm²). Both OB1 and OB2 are 100x oil-immersion objective lenses with NA = 1.25. They are infinite-conjuncted objective lenses, and are designed for working with cover glasses with thickness of 0.17 mm. OB1 and OB2 are bought from the Microscope Store, LLC (www.microscope.com). Their working distance is about 0.1 mm. Thus the total distance between the front surfaces of the two objective lenses is about 0.54 mm. Two oil layers, two cover glasses, two epoxy layers and one water layer must be made to fit within this gap. We later use an objective lens with a longer working distance to make the alignment easier.

The sample chamber consists of three layers of coverslips bonded together by epoxy. The two outer layers are No. 0 coverslips (thickness: 0.08-0.13 mm), the center layer is a 1 cm² square of water layer surrounded by four small pieces of No. 1 coverslips (thickness: 0.13-0.16 mm). The coverslips are bought from Ted Pella, Inc. The microspheres are contained in the water layer. We use vacuum grease to seal the sample chamber. The sealed sample can be used for several hours. Otherwise the water will evaporate in about 10 min without sealing. The silica microspheres used for trapping are bought from Bangs Laboratories, Inc.

The mirror with a sharp edge (MX) is a D-Shaped mirror bought from Thorlabs, Inc. The detector is a balanced InGaAs photo detector. The two photodiodes are matched to cancel the common noise of the laser. The bandwidth of the detector is 100 MHz, enabling us to study the Brownian motion of a trapped microsphere at fast time scales. The DAQ card (Gage, Razor CompuScope 1622) has two channels with sampling rate of 200 MS/s and vertical resolution of 16 bits. This is one of the fastest high resolution digitizers currently available on the market. Our detection system can monitor the position of a trapped microshpere with sub-Ångstrom spatial resolution and sub-microsecond temporal resolution.

The maximum power of the LED is about 5 W. The camera is a USB color 3MP CMOS camera bought from Mightex Systems (Model: MCE-C030-U). It allows us to monitor the sample chamber in real time with spatial resolution of about 0.5 μ m.

Figure 5.6 shows successful trapping of a $4.7 \,\mu$ m diameter silica microsphere in water with an optical tweezer. Initially some microspheres undergo Brownian motion near the bottom of the sample chamber, while other microspheres are stuck to the surface of the coverslip due to the van der Waals force. We can use the optical



Fig. 5.6 A 4.7- μ m diameter silica microsphere (marked by an *arrow*) is trapped in water with an optical tweezer. The trapped microsphere does not move while other three microspheres are shifted when we move the glass slide perpendicular to the laser beam (**a** and **b**) or along the propagation direction of the laser beam (**b**, **c** and **d**). In **c** and **d**, the images of un-trapped microsphere are always sharp as it is trapped at the focus

tweezer to trap a microsphere that is not stuck on the surface. After trapping, the position of the trapped microsphere is fixed by the optical tweezer. We can move the glass slide in 3D while keeping the microsphere trapped, as shown in Fig. 5.6. An optical tweezer thus provides a tool to manipulate a microscopic object in 3D without physical contact.

Figure 5.7 shows the measured noise spectra of the digitizer and two balanced detectors (Thorlabs PDB120C, and PDB110C-AC). When no detector is connected to the DAQ card, the average noise spectral density is about -130 dB around 10 MHz. When a PDB120C detector is connected to the DAQ card, the average spectral density increases to about -107 dB around 10 MHz. When a PDB110C-AC detector is connected to the DAQ card, the average spectral density increases to about -107 dB around 10 MHz. When a PDB110C-AC detector is connected to the DAQ card, the average spectral density increases to about -115 dB around 10 MHz. Thus the PDB110C-AC detector has less noise than the



Fig. 5.7 Noise spectra of the DAQ card and two different detectors

PDB120C detector. So we use the PDB110C-AC detector for studying the Brownian motion in water.

Figure 5.8 shows a measured waveform and power spectrum of a $4.7 \,\mu$ m diameter microsphere trapped in water at room temperature. The waveform appears as a sine wave at 120 Hz. We found that it was due to mechanical vibration of the objective lens. The real signal of the Brownian motion of the microsphere is the high frequency



Fig. 5.8 A waveform and spectrum of a 4.7- μ m diameter microsphere trapped in water at room temperature

fluctuations in the "sine" curve. As a result, the power spectrum is very small at frequencies above 1MHz. The power spectrum at frequencies above 1 MHz basically comes from the noise of the laser, which is a problem for studying the Brownian motion of the microsphere.

Later we mounted a heavy copper block on the post of the top objective lens (OB1 in Fig. 5.5) to reduce the mechanical vibration. This improves the signal significantly. Figure 5.9 shows a measured waveform and power spectrum of a 4.7 μ m diameter microsphere trapped in water after we mounted a copper block on the post. It no longer appears like a sine wave. The high frequency fluctuation due to the Brownian motion of the microsphere dominates the signal. As a result, the power spectral density increased by about 7 dB at 1 MHz. The signal is about 95 dB at 2 MHz, and the noise of the detection system is about -103 dB. So the signal is only 8 dB above the detection noise at 2 MHz. This is not enough for measuring the instantaneous velocity of the Brownian motion. The signal needs to be increased by 10 dB or more



Fig. 5.9 A waveform and spectrum of a $4.7\,\mu$ m diameter microsphere trapped in water at room temperature. A copper block was mounted on the post of the objective lens to reduce mechanical vibration

in order to measure the instantaneous velocity of the 4.7- μ m diameter microsphere in water.

Figure 5.10 show preliminary results of the MSD of a 4.7 μ m bead in water. The measured $MSD_{msr}(t)$ is dominated by the noise when the time is shorter than 0.1 μ s. The $MSD_p(t) = MSD_{msr}(t) - MSD_n(t)$ agrees with the prediction of slope 2 at short time scales. However, this subtraction method is limited by the uncertainty on determining $MSD_n(t)$.

5.5 Interferometer-Enhanced Optical Tweezers

In order to measure the instantaneous velocity of a $4.7 \,\mu$ m microsphere in water, we need to improve the signal-to-noise ratio of the detection system by a factor of 10 or more at frequencies above 1 MHz.



Fig. 5.10 Preliminary results of the MSD of a $4.7 \,\mu$ m bead in water

We consider the detection of the position of a trapped microsphere along x axis that is perpendicular to the propagating direction of the laser. Suppose the position of the microsphere is x_p , and the electric field of the optical tweezer without the microsphere is $E_0(x)(1+\varepsilon_{int}+\varepsilon_{pt}(x))$, where ε_{int} represents the intensity fluctuation of the laser, and $\varepsilon_{pt}(x)$ represents the pointing fluctuation of the laser. The electric field with a trapped microsphere can be written as:

$$E_p(x) = E_0(x)(1 + \varepsilon_{int} + \varepsilon_{pt}(x))(1 + \xi(x_p))$$
(5.30)

where $\xi(x_p)$ represents the effect of the microsphere. If we assume $\varepsilon_{int} \ll 1, \varepsilon_{pt} \ll 1$ and $\xi(x_p) \ll 1$, the intensity of the laser beam after the microsphere is

$$I_p(x) \doteq I_0(x)(1 + 2\varepsilon_{int} + 2\varepsilon_{pt}(x) + 2\xi(x_p))$$
(5.31)

where $I_0(x) = E_0^2(x)$ is assumed to be a Gaussian function.

If we measure the position of the microsphere with a balanced detector (Fig. 5.5), we are measuring the difference of the power of the two halves of the laser beam:

$$\Delta P(x_p, x_{mir}) = \int_{x_{mir}}^{+\infty} I_p(x) dx - \int_{-\infty}^{x_{mir}} I_p(x) dx$$
(5.32)

where x_{mir} is the position of the split mirror. Let us choose a x_{mir} to satisfy

$$\int_{x_{mir}}^{+\infty} I_0(x)(1+2\varepsilon_{int})dx - \int_{-\infty}^{x_{mir}} I_0(x)(1+2\varepsilon_{int})dx = 0,$$
(5.33)

then the effect of the intensity fluctuation can be eliminated. The output signal $\Delta P(x_p, x_{mir})$ contains the position signal of the microsphere and the pointing fluctuation of the laser beam.

If we use a beam splitter to split a laser to two beams, the profiles of the two beams are the same. We can use one beam to trap the microsphere, and the other beam as a reference. The electric field of the reference beam is

$$E_{ref}(x) = E_0(x)(1 + \varepsilon_{int} + \varepsilon_{pt}(x))$$
(5.34)

If we add a phase delay to this reference beam, we can change its sign. Then we can combine the trapping beam and the reference beam together. The electric of the combined beam is

$$E_{cb}(x) = E_p(x) - E_{ref}(x) = E_0(x)(1 + \varepsilon_{int} + \varepsilon_{pt}(x))\xi(x_p)$$
(5.35)

Since $\varepsilon_{int} \ll 1$ and $\varepsilon_{pt} \ll 1$, we have

$$E_{cb}(x) \doteq E_0(x)\xi(x_p).$$
 (5.36)

The effects of the intensity fluctuation and the pointing fluctuation of the laser beam are almost eliminated from the combined beam. Thus the signal-to-noise ratio can be increased significantly with an interferometer enhanced optical tweezer. The difficulty of this method is that the reference laser must have the same pointing fluctuation as the trapping beam at high frequencies (> 1 MHz).

Figure 5.11 shows a Mach-Zehnder interferometer enhanced optical tweezer and detection system. The two objective lenses change the electric field from E(x, y)



Fig. 5.11 A Mach-Zehnder interferometer enhanced optical tweezer and detection system



Fig. 5.12 A Wollaston interferometer enhanced optical tweezer and detection system. The W1 and W2 are two Wollaston prisms. Other components are the same as in Fig. 5.5

to E(-x, -y), so another pair of lenses is used to correct this. A mirror changes the electric field in one direction, i.e. from E(x, y) to E(-x, y). The trapping beam is reflected by three mirrors, and the reference beam is reflected by one mirror. So both beams are changed from E(x, y) to E(-x, y). We also need an EOM to change the phase of the reference beam. Because of low frequency mechanical vibration of mirrors, the phase of the reference beam must be actively locked to the trapping beam. A Sagnac interferometer enhanced optical tweezer may solve this problem [15], but its optics seem much more difficult to align.

We eventually decided to use a Wollaston interferometer enhanced optical tweezer to increase the signal-to-noise ratio. As shown in Fig. 5.12, its setup is very similar to a simple optical tweezer (Fig. 5.5). The Wollaston prism W2 (Karl Lambrecht Corp., Model: WQ-25-05-V 1064NM) separates the laser to two beams with different polarizations. The angle between the two beams is about 0.5° . These two beams form two optical tweezers that are separated by about $15 \,\mu$ m at the focus of the objective lens. One optical tweezer traps a microsphere, and the other optical tweezer serves as a reference. These two beams are combined together by another Wollaston prism [16]. We tune the angle of the $\lambda/2$ waveplate before the PBS2 so that the two beams interfere destructively at one port of the PBS2. The split detection system is at the dark port. Compared with the Mach-Zehnder interferometer, the Wollaston interferometer

is much easier to align. Both beams pass through the same optical path, so the reference and trapping beams contain the same pointing fluctuation. Thus the effect of the pointing fluctuation in detection can be eliminated.

We are currently working on this Wollaston interferometer enhanced optical tweezer. It should provide enough signal-to-noise ratio for measuring the normalized VACF of a Brownian particle in water between 0.35 and 1. It will also have many applications in biophysics.

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Chapter 6 Millikelvin Cooling of an Optically Trapped Microsphere in Vacuum

6.1 Background

Optical cooling and trapping of atoms [1-3] has led to dramatic breakthroughs in atomic, molecular and optical physics, including a new generation of atomic clocks, and realization of Bose–Einstein condensation and degenerate Fermi gas. Applying similar techniques to cool the mechanical motion of macroscopic objects towards the quantum ground state will benefit ultrahigh precision measurements and fundamental tests of macroscopic quantum physics [4-6]. A major obstacle to achieving groundstate cooling of most mechanical oscillators [7-13] is the thermal contact between oscillators and their environment. Recently, it was proposed that optical trapping of dielectric objects in vacuum would greatly reduce the thermal contact, and could even allow ground-state cooling from room temperature [14-22]. Besides providing ideal isolation from the environment, the optical trap can be switched off for timeof-flight measurements to perform full tomography of the mechanical state [21, 23]. Here we report optical trapping of SiO₂ microspheres in vacuum with high oscillation frequencies, and cooling of the center-of-mass motion to millikelvin temperatures with active feedback.

Feedback control has been used widely in industry and scientific experiments. The most commonly used feedback controller is a proportional-integral-derivative (PID) controller. A typical application of a PID controller is to stabilize the temperature of a system. In our lab, we routinely use PID controllers to stabilize the intensity of lasers.

A simple pendulum in air will oscillate if it is initially moved away from the equilibrium position. After oscillating for some time, the pendulum will decay to rest. In reality, however, the pendulum always vibrates with a small amplitude, due to external forces from seismic motion, and more fundamentally, due to the thermal Brownian stochastic force. Even at absolute zero temperature, the pendulum still oscillates because of the quantum zero-point energy. We can use feedback control to reduce the vibration amplitude. If the final vibration amplitude is smaller than the Brownian motion amplitude at thermal equilibrium, the feedback control is called

"feedback cooling" unless the reduction in amplitude corresponds to an increase in frequency. In order to do feedback cooling, the detection system must at least be able to resolve the Brownian motion of the system at thermal equilibrium.

According to the equipartition theorem, the root mean square (rms) amplitude of the Brownian motion of a trapped microsphere at thermal equilibrium is $x_{rms} = \sqrt{k_B T_0/(M\omega^2)}$, where T_0 is the environmental (air) temperature, M is the mass of the microsphere, and ω is the angular trapping frequency. The characteristic size of the quantum ground-state wavefunction is $x_{ground} = \sqrt{\hbar/(M\omega)}$, where \hbar is Planck's constant/ 2π .

Previous experiments have demonstrated optical levitation of a 20- μ m diameter sphere in vacuum with a trapping frequency of about 20 Hz [24], as well as feedback control of a trapped sphere which was used to increase the trapping frequency to several hundred hertz and stabilize its position to within a fraction of one micrometer [25]. However, the resolution of its detection system [25] was not sufficient to enable feedback cooling. For a 20- μ m diameter sphere trapped at 100 Hz, the rms amplitude is about 0.04 μ m at 300 K, and will be much smaller at lower temperature. The size of the quantum ground-state wavefunction is $x_{ground} = 0.14$ pm. Both values are far smaller than the resolution of the detection system of the previous experiment [25]. It is also important that the trapping frequency be much higher than the frequencies of seismic vibration in order to achieve significant cooling.

This chapter describes our efforts on feedback cooling of an optically trapped microsphere in vacuum. We use a dual-beam optical tweezer to trap a 3.0- μ m diameter sphere in vacuum with much higher oscillation frequencies (about 10kHz) to minimize the effects of instrumental vibration. We also demonstrate a detection system to monitor the motion of a trapped microsphere with a sensitivity of about 39 fm/ $\sqrt{\text{Hz}}$ over a wide frequency range. Using active feedback, we simultaneously cool the three center-of-mass (COM) vibration modes of a microsphere from room temperature to a minimum mode temperature of 1.5 mK, which corresponds to the reduction of the rms amplitude of the microsphere from 6.7 nm to 15 pm for that mode.

6.2 Principle of Feedback Cooling

An optically trapped microsphere in non-perfect vacuum will exhibit Brownian motion due to collisions between the microsphere and residual air molecules. When there is no feedback cooling, the equation of the Brownian motion of an optically trapped microsphere is:

$$\frac{d^2x_j}{dt^2} + \Gamma_0 \frac{dx_j}{dt} + \Omega_j^2 x = F_j^{th}, \tag{6.1}$$

where Γ_0 is the viscous damping factor due to air molecules, $\Omega_j/2\pi$ (j = 1, 2, 3) are the resonant frequencies of the optical trap along the three fundamental axes (x, y, and z axes), and $F_i^{th} = \zeta_j(t)\sqrt{2k_BT\Gamma_0/M}$ is the Brownian stochastic force.

6.2 Principle of Feedback Cooling

The damping term $\Gamma_0 \frac{dx}{dt}$ tends to stop any vibration, while the F_j^{th} term drives the motion. It is very interesting that Γ_0 is also contained in F_j^{th} . When the mechanical energy (sum of the kinetic energy and the potential energy) of the microsphere is larger than k_BT in one direction, the $\Gamma_0 \frac{dx}{dt}$ term will dominate and the mechanical energy of the microsphere will be reduced. On the other hand, the F_j^{th} term will dominate and increase the mechanical energy of the microsphere if its mechanical energy is smaller than k_BT . Thus the average mechanical energy of the microsphere will be k_BT in each direction at thermal equilibrium.

At thermal equilibrium, the power spectrum of COM motion of a trapped microsphere along each of the three fundamental mode axes is [7, 12]:

$$S_{j}(\omega) = \frac{2k_{B}T_{0}}{M} \frac{\Gamma_{0}}{(\Omega_{j}^{2} - \omega^{2})^{2} + \omega^{2}\Gamma_{0}^{2}},$$
(6.2)

where $\omega/2\pi$ is the observation frequency.

6.2.1 Feedback Cooling

To implement feedback cooling, we apply an external force on the trapped microsphere:

$$F_j^{cool} = -\Gamma_j^{cool} \frac{dx_j}{dt}.$$
(6.3)

The force is proportional to the velocity of the microsphere but with opposite direction. Thus it will slow down the motion of the microsphere. With feedback cooling, the equation of the Brownian motion of an optically trapped microsphere is:

$$\frac{d^2x_j}{dt^2} + (\Gamma_0 + \Gamma_j^{cool})\frac{dx_j}{dt} + \Omega_j^2 x = \zeta_j(t)\sqrt{\frac{2k_B T \Gamma_0}{M}}.$$
(6.4)

In contrast to the Γ_0 due to air molecules, Γ_j^{cool} is only contained in the damping term but not in the heating term. So "feedback cooling" is also called "cold damping".

With feedback cooling, the power spectrum of the COM motion of a trapped microsphere along each of the three fundamental mode axes is:

$$S_{j}^{cool}(\omega) = \frac{2k_{B}T_{0}}{M} \frac{\Gamma_{0}}{(\Omega_{j}^{2} - \omega^{2})^{2} + \omega^{2}(\Gamma_{0} + \Gamma_{j}^{cool})^{2}}.$$
 (6.5)

Let $\Gamma_j^{tot} = \Gamma_0 + \Gamma_j^{cool}$ be the total damping factor, and $T_j^{cool} = T_0 \Gamma_0 / \Gamma_j^{tot}$ be the effective temperature of the motion with feedback cooling, the power spectrum can be rewritten as:

6 Millikelvin Cooling of an Optically Trapped Microsphere in Vacuum

$$S_{j}^{cool}(\omega) = \frac{2k_{B}T_{j}^{cool}}{M} \frac{\Gamma_{j}^{tot}}{(\Omega_{j}^{2} - \omega^{2})^{2} + \omega^{2}(\Gamma_{j}^{tot})^{2}},$$
(6.6)

which has the same form as Eq. 6.2. The effective temperature along each axis may be different because Γ_j^{cool} can be different along different directions. The motion can be cooled significantly by applying a feedback damping $\Gamma_j^{cool} \gg \Gamma_0$. The lowest temperature will be limited by the noise in the detection system and feedback circuits, as well as coupling between different directions.

6.2.2 Feedback Amplification

Besides cooling, feedback control can also be used to amplify the motion. We can apply a force in the same direction as the velocity of the microsphere to amplify the motion:

$$F_j^{amp} = +\Gamma_j^{amp} \frac{dx_j}{dt}.$$
(6.7)

With feedback amplification, the equation of the Brownian motion of an optically trapped microsphere is:

$$\frac{d^2x_j}{dt^2} + (\Gamma_0 - \Gamma_j^{amp})\frac{dx_j}{dt} + \Omega_j^2 x = \zeta_j(t)\sqrt{\frac{2k_B T \Gamma_0}{M}}.$$
(6.8)

The power spectrum of COM motion of the microsphere is:

$$S_{j}^{amp}(\omega) = \frac{2k_{B}T_{0}}{M} \frac{\Gamma_{0}}{(\Omega_{j}^{2} - \omega^{2})^{2} + \omega^{2}(\Gamma_{0} - \Gamma_{j}^{amp})^{2}}.$$
 (6.9)

When $\Gamma_j^{amp} < \Gamma_0$, the system is stable. The effect of Γ_j^{amp} is to amplify the motion and decrease the linewidth of the vibration from Γ_0 to $\Gamma_0 - \Gamma_j^{amp}$. When $\Gamma_j^{amp} > \Gamma_0$, the system is not stable and the microsphere will be lost. We have observed both feedback cooling and feedback amplification in our experiment by inverting the velocity signal.

6.2.3 Heating Due to Light Scattering

For a microsphere trapped by an optical tweezer, there are also heating effects due to the light scattered by the microsphere. The heating effects of light scattering can be separated to two parts. The first part is because of the shot noise of the laser. This effect is very small and has been calculated in Refs. [14, 15].

6.2 Principle of Feedback Cooling

The second part is because the scattering force is not conservative. The scattering force can do net work on the microsphere when the microsphere moves over a closed loop under certain conditions [26]. If the two counter-propagating beams of our dual-beam optical trap is slightly misaligned along x axis (Fig. 3.10), the scattering force on the microsphere is along the axial direction (z axis), and is proportional to the displacement of the microsphere along x axis. We have $\vec{F}_{scat} \propto x\hat{z}$. For a microsphere moving in a loop in the x - z plane, the net work done by the scattering force over a loop is

$$W = \oint \vec{F}_{scat} dl \propto \sqrt{\langle x^2 \rangle} \sqrt{\langle z^2 \rangle}.$$
 (6.10)

The period of a harmonic oscillator is independent of the energy of the oscillator. The mechanical energy of the microsphere will increase due to the work done by the scattering force at a rate:

$$\frac{d}{dt}(E_x + E_z) \propto \sqrt{\langle x^2 \rangle} \sqrt{\langle z^2 \rangle} \propto \sqrt{E_x E_z}, \tag{6.11}$$

where E_x and E_z are mechanical energies of the microsphere. From this equation, it is clear that the scattering force couples the motion along different directions and can make the dynamics of the system very complex. Here we will try to obtain some qualitative properties of this heating effect. For simplicity, we assume that E_x is proportional to E_z , then $dE_x/dt \propto E_x$. So the scattering force will cause a heating that is proportional to the mechanical energy of the microsphere. This effect can be minimized by better alignment of the two counter-propagating laser beams.

A phenomenological description of the feedback cooling process with this heating effect can be written in an equation about the average mechanical energy of the microsphere along each axis:

$$\frac{dE_j}{dt} = -\Gamma_0 E_j - \Gamma_j^{cool} E_j + \Gamma_0(k_B T) + \alpha_j E_j.$$
(6.12)

The first term $(-\Gamma_0 E_j)$ describes the damping due to the air, the second term is due to feedback cooling, the third term describes the heating effect due to the air, and the last term describes the heating effect due to the scattering force. The final effective temperature of motion of the microsphere in each direction is

$$T_j^{cool} = E_j/k_B = \frac{\Gamma_0 T}{\Gamma_0 + \Gamma_j^{cool} - \alpha_j}.$$
(6.13)

The system will be stable when $\Gamma_0 + \Gamma_j^{cool} > \alpha_j$ and unstable when $\Gamma_0 + \Gamma_j^{cool} < \alpha_j$. This provides a method to estimate α_j experimentally. To estimate the α_j , we turn off the feedback cooling forces to let $\Gamma_j^{cool} = 0$. We first trap a microsphere at a high pressure. At high pressures, we have $\Gamma_0 > \alpha_j$, and the system is stable. We then reduce the air pressure to reduce Γ_0 . At a certain pressure, the system becomes unstable and microsphere is lost. Then we have $\alpha_j \approx \Gamma_0^{esc}$, where Γ_0^{esc} is the air damping factor at the moment when the microsphere is lost.

The heating rate due to the nonconservative force is proportional to the energy of the microsphere. As the energy of the microsphere is reduced by cooling, this heating effect becomes negligible and will not prevent ground state cooling. For cooling, we usually have $\Gamma_j^{cool} \gg \alpha_j$, thus the effect of α_j on the final temperature is very small (Eq. 6.13). The heating effect of the scattering force will be much smaller for a single-beam optical tweezer.

The above discussions are based on classical mechanics, which is sufficient for understanding our current experiment. Quantum mechanical description of feedback cooling will be necessary if the motion of the microsphere is cooled to near the quantum ground state [27–29].

6.2.4 Damping Due to the Residual Gas in Vacuum

The viscous damping factor due to air can be calculated by kinetic theory. Assuming the reflection of air molecules from the surface of a microsphere is diffusive, and the molecules thermalize with the surface during collisions, we have [30]

$$\Gamma_0 = \frac{6\pi\eta R}{M} \frac{0.619}{0.619 + Kn} (1 + c_K), \tag{6.14}$$

where η is the viscosity coefficient of the air, R is the radius of the microsphere, M is the mass of the microsphere, and Kn = l/R is the Knudsen number. Here l is the mean free path of the air molecules. $c_K = (0.31Kn)/(0.785 + 1.152Kn + Kn^2)$ is a small positive function of Kn [30]. At low pressures where $Kn \gg 1$, the viscous damping factor is proportional to the pressure. At high pressures where $Kn \ll 1$, the viscous damping factor is $\Gamma_0 = 6\pi\eta R/M$, which is the same as the prediction of Stoke's law.

Figure 6.1 shows the measured linewidth, $\Gamma_0/2\pi$, of the oscillation of a trapped 3µm microsphere at different pressures without feedback cooling. The powers of the two trapping beams are 120 and 100 mW, respectively. The linewidths are obtained by fitting the measured power spectra with Eq. 6.2. The measured linewidths agree very well with the prediction of kinetic theory (Eq. 6.14) from 10⁵ Pa down to 1 Pa. At pressures below 1 Pa, the measured linewidths are larger than the theoretical prediction. This linewidth broadening is due to power fluctuations of the trapping laser. The inset of Fig. 6.1 shows a power spectrum at 0.13 Pa. A more detailed power spectrum at 0.13 Pa (1 mtorr) is shown in Fig. 6.2. The trapping frequency $\omega_1/2\pi$ is 9756.4 ± 0.3 Hz, and the linewidth is 0.46 ± 0.06 Hz, giving a quality factor ($Q_j = \omega_j/\Gamma_0$) of 2.1 × 10⁴. This implies the power fluctuation of the trapping laser is smaller than 0.01 % during the measurement. An optically trapped microsphere provides a method to directly convert laser power to a frequency signal, which can be measured precisely. Stabilization of laser power to a trapped bead can find applications in laser



Fig. 6.1 Measured linewidths of the oscillation of an optically trapped 3- μ m diameter microsphere at different pressures. The *blue curve* is the prediction of a kinetic theory (Eq. 6.14). The *inset* is the measured power spectrum at 0.13 Pa. By fitting the spectrum with Eq. 6.2 (*red curve*), we obtain $\omega_1 = 2\pi \cdot (9756.4 \pm 0.3)$ Hz and $\Gamma_0 = 2\pi \cdot (0.46 \pm 0.06)$ Hz for this example. The same method is used to obtain linewidths for other pressures

physics, and can enable a more precise measurement of the Q for a second trapped bead in vacuum.

Before feedback cooling, we observe a sharp transition in the trap lifetime as a function of pressure. Above the transition pressure, the microsphere can be trapped stably for many hours. Below the transition pressure, the microsphere is lost within a few seconds. This is because of the heating effect due to the light scattering, which has been discussed in the previous section. The transition pressure depends critically



Fig. 6.2 The power spectrum of a trapped 3.0-µm diameter microsphere at 1 mtorr

on the alignment of the two counter-propagating trapping beams. We can reduce it to less than 0.1 Pa by aligning the two laser beams. Thus α_j in Eq. 6.12 can be smaller than $2\pi \times 0.5$ Hz. We also observed limit cycles (vortices) in the motion of a trapped microsphere when the trapping beams are intentionally misaligned. A 3D simulation of the system is required in order to fully understand these phenomena.

6.3 A 3D Split Detection System

In order to monitor the motion of a trapped microsphere with ultrahigh precision in all three dimensions, we built a 3D split detection system. As shown in Fig. 6.3, the X, Y, and Z detectors are fast balanced photo-detectors with bandwidth of 75 MHz. They have two matched photodiodes to cancel the common mode noise in the laser beams, allowing ultra-high precision measurements of the position of a trapped microsphere. When a trapped microsphere moves in the horizontal (vertical) direction, it deflects the trapping beam in the horizontal (vertical) direction. This changes the relative power between the two beams after the MX (MY) mirror, which is measured by the X (Y) detector. The motion of a trapped bead along the trap axis changes the divergence angle of the output beam, which changes the waists of incident beams



Fig. 6.3 Simplified schematic showing the detection system that can monitor the real-time position of a trapped microsphere with ultra-high precision in all three dimensions. One of the trapping beams (the other trapping beam is not shown) passes through a trapped microsphere inside a vacuum chamber and is reflected by a polarizing beam splitter cube (*PBS*). It is then split to three beams by two beam splitters (BS1 and BS2) for 3D detection. MX is a mirror with a sharp edge that splits the beam into two parts *horizontally*. MY is a mirror with a sharp edge that splits the beam into two parts *vertically*. BS3 is a beam splitter, ND is a neutral density filter, and LZ is a lens. The X, Y, and Z detectors are balanced detectors that have two matched photodiodes to cancel the common mode noise in the laser beams

on the Z detector. One photodiode of the Z detector is smaller than the waist of the incident beam. It measures only part of the power of the incident beam. Thus its output voltage depends on the waist of the incident beam, which is a function of the position of the microsphere in Z axis. The other photodiode of the Z detector is much larger than the waist of the incident beam. It measures the total power of the incident beam, and can serve as a reference signal [31]. A photo of our 3D detection system is shown in Fig. 6.4.

For small displacements of a trapped microsphere near the trap center, the voltage output (U_i) of each detector is proportional to the displacement (x_i^D) of the microsphere along the detection direction, i.e. $U_i = \beta_i x_i^D$, where i = 1, 2, 3 denotes X, Y, and Z detectors, and β_i is the calibration factor of the detector. We align the detection system carefully to make the detection direction of each detector be parallel to one of the trap's fundamental mode axes. In reality, however, there is always slight difference between the detection directions and the fundamental mode axes. Thus the voltage output from each detector is a combination of signals from each mode, i.e. $U_i = \beta_i (\alpha_{i1} x_1^M + \alpha_{i2} x_2^M + \alpha_{i3} x_3^M)$, where x_j^M (j = 1, 2, 3) is the displacement of the microsphere along the fundamental mode axis \hat{x}_j^M , and α_{ij} is the projection coefficient of \hat{x}_j^M to the detection direction \hat{x}_i^D . Usually only one term dominates as the detection directions are almost parallel to the mode axes.

The expected value of the power spectrum of the voltage output from each detector is [32]:

$$S_i^U(\Omega) \equiv \langle |\tilde{U}_i|^2 / T_{msr} \rangle = \beta_i^2 \langle |\alpha_{i1} \tilde{x}_1^M + \alpha_{i2} \tilde{x}_2^M + \alpha_{i3} \tilde{x}_3^M |^2 / T_{msr} \rangle, \qquad (6.15)$$

where T_{msr} is the measurement time, \tilde{U}_i and \tilde{x}_j^M are Fourier transforms of U_i and x_j^M , respectively. The expansion of $S_i^U(\Omega)$ has 9 terms, but the expected values of the



Fig. 6.4 A photo of our 3D detection system and a camera for imaging a trapped microsphere with the scattered light. *Red lines* are drawn on the photo to show the light paths of the detection system

cross correlation terms $\langle \tilde{x}_i^M \tilde{x}_j^M \rangle$ $(i \neq j)$ are 0 when there is no feedback, because the three components of the motion of a microsphere in a harmonic trap are uncorrelated. With active feedback, there can be small correlations between different directions if the feedback loops are coupled. In real experiments, the coupling is small and the trapping frequencies are different along different directions, thus we assume that the average values of the cross correlations are negligible. Then

$$S_i^U(\omega) = \beta_i^2 [\alpha_{i1}^2 S_1(\omega) + \alpha_{i2}^2 S_2(\omega) + \alpha_{i3}^2 S_3(\omega)],$$
(6.16)

where $S_j(\omega)$ is the power spectrum of COM motion along each fundamental mode axis. $S_j(\omega)$ is described by Eq.6.2 without feedback cooling, and by Eq. 6.5 with feedback cooling.

The detection system can be calibrated by fitting the measured power spectra at room temperature with the expected power spectra $S_i^U(\omega)$ to obtain calibration factor $\beta_i^2 \alpha_{ij}^2$ for each mode that is distinguishable in the power spectra. We can also obtain β_i^2 directly by the energy equipartition theorem, which says $\langle M v_{i/2}^2 \rangle = k_B T_0/2$, where v_i is the instantaneous velocity of the microsphere projected onto any axis. Since $U_i = \beta_i x_i^D$, we have $\beta_i^2 = \frac{M}{k_B T_0} \langle \left(\frac{dU_i}{dt}\right)^2 \rangle$. With $\beta_i^2 \alpha_{ij}^2$ and β_i^2 , we can easily obtain α_{ij}^2 to check the alignment of our detection system. In the experiment, each detector is used to monitor only one mode, so only three calibration factors ($\beta_1^2 \alpha_{11}^2$, $\beta_2^2 \alpha_{22}^2$, and $\beta_3^2 \alpha_{33}^2$) are required for measuring the mode temperatures with feedback cooling.

The mass of the microsphere is required to obtain the calibration factors. The pure silica (SiO₂) microspheres used in this experiment are from Bangs Laboratories, Inc. Their mean diameter is $3.0 \,\mu$ m, corresponding to a mean mass of 2.8×10^{-14} kg for each microsphere. The standard deviation of the size given by the supplier is 14%. The exact diameter of the microsphere is not important for feedback cooling. The temperatures of the feedback-cooled motion are obtained by comparing the power spectra of the same microsphere with and without feedback cooling, a measurement which is independent of the exact size of the microsphere. The viscous damping factor (Γ_0) of a microsphere in air, however, depends on the size of the microsphere. Using the measured damping factor shown in Fig. 6.1, we obtain the diameter of a microsphere by kinetic theory (Eq. 6.14) to be 2.7 μ m, which is within the uncertainty range given by the supplier.

6.4 1D Optical Feedback Cooling

Figure 6.5 shows the first feedback cooling scheme that we used in our experiment. The position signal of a trapped microsphere is sent through a preamplifier that has a bandpass filter, and a derivative circuit (d/dt) to provide a signal proportional to the velocity of the microsphere. This velocity signal is used to control the frequency



Fig. 6.5 Diagram of a 1D feedback cooling scheme. The position of a trapped microsphere is monitored by a home-built detection system. The position signal is sent through a preamplifier which has a bandpass filter (typically 100 Hz to 300 kHz), and a derivative circuit (d/dt) to provide a signal proportional to velocity. This velocity signal is used to control the frequency of a radio frequency (*RF*) AOM driver which modulates the direction of the laser beam. The data is digitized and stored on a computer by a data acquisition card (*DAQ*)

of the output signal of a radio frequency (RF) deflector driver (IntraAction, Model: DE-802M26). The output of the driver is a single frequency signal in the range of 60–100 MHz. The driver has an analog input to modulate the frequency of the signal. An analog input signal from 0 to 1 V will change the frequency of the output signal by 40 MHz. The RF signal drives an acousto-optic modulator (AOM) which controls the *p*-polarized trapping beam. The frequency of the RF signal determines the direction of the laser beam, and the power of the RF signal determines the power of the trapping beams in the horizontal direction. The laser beam will exert a force on the microsphere as a function of its velocity in the horizontal direction, which cools the motion of the microsphere.

The preamplifier (SR560, Stanford Research Systems) has a very low input noise (about $4 \text{ nV}/\sqrt{\text{Hz}}$), and a variable gain from 1 to 5×10^4 . In this experiment, we only need the gain to be about 2 or 5 because the output signal from the detector is pretty big already. The preamplifier contains two first-order RC filters that can be set to -3 dB cutoff frequencies chosen from a 1-3-10 sequence, from 0.03 Hz to 1 MHz. We typically set the low cutoff frequency to be 100 Hz and the high cutoff frequency to be 300 kHz. The preamplifier also has a function to invert the signal, which is very useful. The derivative circuit is simply the 'derivative' branch of an analog PID controller. We first used a commercial PID controller (SIM 960, Stanford Research Systems). It has a lot of powerful functions, including digital control of the P, I, D, and Offset. However, it picks up some external electronic noise through its power



Fig. 6.6 A derivative circuit for calculating the real-time velocity of a microshpere from its position signal

cord. We eventually decided to use our home-built derivative circuit for feedback cooling.

A detailed description of our home-built PID controllers can be found in Appendix C.1 of the Ph.D dissertation of Todd Meyrath [33]. The 'derivative' part of the circuit is shown in Fig. 6.6. The main component of the circuit is the low-noise operational amplifier, the capacitor C1 and the tunable resistor R2. The operational amplifier is OPA227 from Texas Instruments. It has a low noise level of $3 \text{ nV}/\sqrt{\text{Hz}}$ and a wide bandwidth of 8 MHz. The differentiation time of the derivative circuit is $C1 \times R2$. It is important that the OPA227 is unity-gain stable. We have tried to use an OPA228 amplifier which is faster than the OPA227, but the circuit is not as stable for low gain. The resistor R1 is much smaller than R2, and the capacitor C2 is much smaller than C1. R1 limits the differential gain, and C2 gives high frequency roll-off. R1 and C2 are necessary because the derivative circuit has a very large gain at high frequencies. For a pure derivative circuit, if the input signal is $V_{in} = \sin(\omega t)$, then the output will be $V_{out} = \omega \cos(\omega t)$, which will be infinite if ω is infinite. Thus a derivative circuit amplifies high frequency noise. So we use R1 to limit the gain, and C2 to serve as a low-pass filter to reduce the gain of high frequency noise. R3 is used to isolate the derivative circuit from other parts of the circuit so that they do not interfere with each other. Typical values of the resistors and capacitors are $R1 = 50 \Omega$, C1 = 2.2 nF, $R2 = 0-50 \text{ k}\Omega$, C2 = 0.1 nF, and $R3 = 1 \text{ k}\Omega$.

Figure 6.7 shows some results of the 1D feedback cooling. In Fig. 6.7a, the rms velocity of the microsphere is reduced from 0.43 mm/s to 0.090 mm/s by 1D feedback cooling. The temperature of the Brownian motion is reduced from room temperature (297 K) to 13 K. Figure 6.7b shows the power spectra of the microsphere with (blue curve) and without (red curve) feedback cooling at 208 mtorr. With 1D feedback cooling, the peak near 10 kHz is reduced by 2 orders, while the peaks near 800 Hz and 3 kHz do not change very much. Thus the 1D feedback cooling only cools the motion along one direction efficiently.

In the red curve of Fig. 6.7b, the peak at 10 kHz corresponds to the mode along the X axis, the tiny peak at 9 kHz corresponds to the mode along the Y axis. These



Fig. 6.7 a The normalized velocity distributions of a trapped 3.0- μ m diameter microsphere without feedback cooling at 26 torr (*red curve*), and with feedback cooling at 4 mtorr (*blue curve*). b The power spectra of the microsphere with (*blue curve*) and without (*red curve*) feedback cooling at 208 mtorr

two modes have almost the same frequency because the laser beam is only slightly elliptical. The peak at 3 kHz corresponds to the Z mode. We initially thought the peak at 800 Hz corresponds to the Z mode. After an extensive study, we believe that the 800 Hz peak is because of the frequency difference between the X and Y mode. This became clear when we do 3D feedback cooling. The Z cooling beam reduced the 3 kHz mode efficiently, but did not reduce the 800 Hz peak efficiently. The frequency (800 Hz) equals the frequency difference between the X and Y mode. We are also able to make the 800 Hz peak disappear by better alignment.



Fig. 6.8 a Modulation of the direction of beam No. 2 (the *p*-polarized beam) with an AOM in the *horizontal direction*. **b** Modulation of the direction of beam No. 2 with an AOM in the *vertical direction*

We later installed another AOM to control the direction of beam No. 2 (the p-polarized beam) in the vertical direction. Thus we can do 3D feedback cooling by modulating the intensity of beam No. 2, and the directions of beam No. 2 in horizon-tal and vertical directions. The beam No. 1 (the *s*-polarized beam) is not modulated because it is used for detection. We have tried such 3D feedback cooling method, but we were only able to cool the motion from 297 K to about 10 K.

We finally found out the problem after struggling for several months. In order to find out the problem, we use a quadrant detector to monitor the motion of a laser beam when we modulate its direction by an AOM. Figure 6.8 shows the motion of the center of beam No. 2 when we modulate its directions with an AOM (IntraAction, model: ATM-801A2) in the horizontal direction (a) and an AOM in the vertical direction (b). We use a sine signal with peak-to-peak voltage of 2 mV at 10 kHz to drive the analog input of the RF driver. This modulates the frequency of the RF signal (centered at about 80 MHz) by 80 kHz. Thus the frequency of the RF signal is $80 \text{ MHz} + 40 \text{ kHz} \cdot \sin(2\pi 10^4 t)$, where t is time. Ideally, the data points should be a straight horizontal line in Fig. 6.8a and a straight vertical line in Fig. 6.8b. However, the experimental points are not in a straight line, especially in Fig. 6.8b. This means that when we use an AOM to control the laser along the vertical direction, the laser beam also moves along the horizontal direction in a very complex way. Thus when we cool the motion vertically, we also heat the motion horizontally. This limits the final temperature of the feedback cooling with this method.

Figure 6.9 shows the motion of the center of beam No. 1 (the *s*-polarized beam) when we modulate its direction with an AOM (Isomet, model: 1205C-2-804) in the horizontal direction. We want it to be a straight line in the horizontal direction. However, the real motion of the laser beam is very complex.

Fig. 6.9 Modulation of the direction of beam No. 1 with an AOM in the horizontal direction



6.5 Electrostatic Forces

After we found the problem of modulating the direction of a laser beam with an AOM, we decided to try to perform feedback cooling using electrostatic forces. The natural charge of a microsphere is negligible. It is necessary to charge the microsphere first in order to apply sufficient electrostatic forces for feedback cooling.

Figure 6.10 shows our setup for air discharge and feedback cooling with an electrostatic force. The stainless steel mounts of the two aspheric lenses and the whole vacuum chamber are grounded. A thin stainless steel sheet is inserted half way between the two lenses. It is connected to a high-voltage amplifier that can deliver a



Fig. 6.10 Setup for air discharge and feedback cooling with an electrostatic force. The stainless steel holders of the two aspheric lenses and the whole vacuum chamber are grounded. The smallest separation between the two holders is about 4 mm. A thin stainless steel sheet is inserted at the center of the two lenses as an electrode. The edge of the steel sheet is about 1 mm away from where the microsphere is trapped. The steel sheet is connected to a high-voltage amplifier that can deliver $0-1 \, \text{kV}$ voltage output

0-1 kV voltage output. There will be a strong electric field between the steel sheet and the steel holders when a high voltage is applied to the steel sheet. At high voltage, the air breaks down and becomes conductive. This can damage the high-voltage amplifier. We connect a $1 \text{ M}\Omega$ resister in series with the steel sheet and the high-voltage amplifier to limit the peak current.

We first trapped a microsphere with the dual-beam optical trap at about 100 torr. We then reduced the pressure to about 0.5 torr. With the microsphere trapped, we applied an AC voltage in the form of $V(t) = V_0[\sin(2\pi f t) + 1]/2$ to the steel sheet. The frequency of the AC voltage was typically about 1 kHz. We increased the peak voltage slowly from a few volts to several hundred volts until air discharge occured. A spectrum of the motion of a trapped microsphere at 461 mtorr driven by a 1.5 kHz, $V_0 = 340$ V signal is shown in Fig. 6.11a. The peak at 1.5 kHz due to the AC signal is very small, because the natural charge of the microsphere is very small. When the peak voltage was increased to 680 V, air discharge occurred. The microsphere moved violently during the air discharge. Thus there are a lot of peaks in Fig. 6.11b. It is surprising to us that the optical tweezer is stable enough to trap a microsphere during air discharge. We reduced the peak voltage in a few seconds after air discharge occured to avoid the loss of the microsphere. The microsphere gained charge during air discharge. Figure 6.11c shows a spectrum of the motion of the microsphere driven by a 1.5 kHz, $V_0 = 340$ V signal after air discharge. The peak at 1.5 kHz after discharge is about 4 orders higher than the peak before discharge.

After air discharge, the microsphere maintains its charge, even at high pressures. Figure 6.12 shows a spectrum of the motion of a trapped microsphere at 205 torr after air discharge. The microsphere is driven by a 400 Hz, $V_0 = 510$ V AC voltage. The hight of the peak at 400 Hz is proportional to the square of the charge of the microsphere. By measuring the height of the peak, we can monitor the charge of the microsphere as a function of time. Figure 6.13 displays the charge of the microsphere as a function of time. Figure 6.13 displays the charge of the microsphere as a function of time over a period of 2 h. The charge fluctuates because there is a weak air discharge near the microsphere. So the microsphere gains and losses charges over time. The shape of the curve depends on the driving voltage and the air pressure. We have not been able to observe individual steps in the curve when the microsphere gains or loses one electron.

According to Coulomb's law, the force between two point charges q_1 and q_2 is:

$$F = k_e \frac{q_1 q_2}{r^2} \tag{6.17}$$

where $k_e = 8.99 \times 10^9 \,\text{N} \cdot \text{m}^2 \,\text{C}^{-2}$ is the Coulomb constant, and *r* is the distance between the two point charges. The force between two electrons separated by 1 mm and 1 μ m is 2.3 × 10⁻²² N and 2.3 × 10⁻¹⁶ N, respectively.

When N electrons e are distributed homogeneously on the surface of a microsphere with radius R, the energy required to add another electron is about:

$$E = k_e \frac{Ne^2}{R} \tag{6.18}$$



Fig. 6.11 Spectra of the motion of a trapped microsphere before (**a**), during (**b**) and after (**c**) air discharge. The pressure is 461 mtorr. The frequency of the AC voltage is 1.5 kHz, and the air discharge happens when the peak voltage is about 680 V. We reduce the peak voltage in several seconds after air discharge occurs to avoid the loss of the microsphere



Fig. 6.12 Spectrum of the motion of a trapped microsphere at 205 torr after air discharge. The micropshere is driven by a 400 Hz, $V_0 = 510$ V AC voltage. After air discharge, the microsphere maintains its charge even at high pressure

For a $R = 1.5 \,\mu\text{m}$ microsphere, the required energy is $k_B T/2$ at room temperature when N = 13. Thus the natural charge of a 3- μ m diameter microsphere is in the order of 13 e. We do not know the exact charge of the microsphere after air discharge, but estimate it to be on the order of 1000 e from the power spectrum (Fig. 6.11), which corresponds to E = 1 eV. The maximum electric field at the trap center is about 10 V/mm. Thus the maximum electrostatic force on the microsphere is in the



Fig. 6.13 Fluctuation of the charge of a 3- μ m-diameter microsphere trapped at 205 torr, driven by a 400 Hz, $V_0 = 680$ V signal. *a.u.* arbitrary unit
order of 1.6 pN. This is consistent with the observed shift of the trap center when we apply a DC voltage.

The air discharge is not very controllable. This makes the current setup (Fig. 6.10) not suitable for feedback cooling. We have tried to implement feedback cooling with electrostatic forces and were able to cool the motion from room temperature to about 10 K. The final temperature is limited by the fact that the charge of the microsphere fluctuates, and we do not know the real direction of the electrostatic force. These problems can be solved by a better design of the electrodes and a better charging method. The electrostatic forces. The microsphere can be charged by photoelectric charging with electrostatic forces. The microsphere can be charged by photoelectric charging with a ultraviolet lamp [34, 35], by using electrospray [36] or an electron gun. Combining an optical trap with an ion trap in the same location should be helpful in trapping and studying particles at ultrahigh vacuum.

6.6 Millikelvin Cooling with 3D Optical Feedback

Since significant efforts were required to improve feedback cooling with electrostatic forces, we decided to try feedback cooling with optical forces again. This time, we used AOM's to modulate the intensities of laser beams rather than the directions of laser beams to do 3D feedback cooling. This method turned out to work very well. It enables us to cool the center-of-mass motion of a trapped microsphere from room temperature to millikelvin temperatures in all three dimensions, with a minimum mode temperature of 1.5 mK [37].

6.6.1 Experimental Setup

A simplified scheme of our optical trap and cooling beams is shown in Fig. 6.14. The dual-beam optical trap is the same as described before. It is created inside a vacuum chamber by two counter-propagating laser beams focused to the same point by two identical aspheric lenses with a focal length of 3.1 mm and numerical aperture of 0.68. The wavelength of both trapping beams is 1064 nm. They are orthogonally polarized, and are shifted in frequency to avoid interference. The beams are slightly elliptical and approximately form a harmonic trap with three fundamental vibration modes along the horizontal, vertical and axial directions, denoted X, Y, and Z in Fig. 6.14. The motion of a trapped bead causes deflection of both trapping beams. We monitor the position of the bead by measuring the deflection of one of the trapping beams with ultrahigh spatial and temporal resolution in all three dimensions (Fig. 6.3).

Using the position signal, we can calculate the instantaneous velocity of the bead, and implement feedback cooling by applying a force with a direction opposing the velocity (Fig. 6.15). The feedback is generated by scattering forces from three orthogonal 532 nm laser beams along the axes as shown in Fig. 6.3. The average intensity



Fig. 6.14 Simplified schematic showing a glass microsphere trapped at the focus of a counterpropagating dual-beam optical tweezer, and three laser beams along the axes for cooling. The wavelengths of the trapping beams and the cooling beams are 1064 and 532 nm, respectively



Fig. 6.15 Diagram of the feedback mechanism for the *X axis*: The position of a trapped microsphere is monitored by a home-built detecting system. The position signal is sent through a bandpass filter (typically 100 Hz to 300 kHz) and a derivative circuit (d/dt) to provide a signal proportional to velocity. This velocity signal is used to control the output power of a radio frequency (*RF*) AOM driver which modulates the power of the X cooling beam. The data is digitized and stored on a computer by a data acquisition card (*DAQ*)



Fig. 6.16 A photo of the 3D optical feedback cooling system

of the cooling beams is about 1 % that of the trapping beams. The optical power of each cooling beam is controlled by an acousto-optic modulator (AOM). Each beam is modulated with a time-varying signal proportional to the instantaneous velocity of the bead, added to an offset. The proportional component generates the required cooling force, while the offset slightly shifts the trap center. A photo of the optics of our 3D optical feedback cooling system is displayed in Fig. 6.16. The green color in the photo is due to the scattered light from the cooling beams. The trapping beams are infrared and cannot be seen in this photo.

Figure 6.17 shows the power of the first order of a laser beam exiting the AOM as a function of the input voltage of the RF driver (IntraAction, model: ME-802) and the reading of the manual offset knob of the RF AOM driver. In general, the laser power is a nonlinear function of the input voltage. When the knob reading is at 5, the laser power depends on the input voltage linearly around 0. This is good for feedback cooling because we want the laser power to be proportional to the velocity of the microsphere. Thus we set the knob reading at 5, and use the velocity signal as the input voltage to control the laser power for feedback cooling.

The behavior of the system with three dimensional (3D) feedback cooling is straightforward to understand if we assume that there is no coupling between feedback forces and velocities in different directions. In this case, the feedback force in each direction adds an effective cold damping factor Γ_j^{fb} , and the total damping becomes $\Gamma_j^{tot} = \Gamma_0 + \Gamma_j^{fb}$. The power spectrum of the motion of a trapped microsphere with feedback cooling can be described by Eq. 6.5. The temperature of the motion with feedback cooling will be $T_j^{fb} = T_0 \Gamma_0 / \Gamma_j^{tot}$. Thus the motion can be cooled significantly by applying feedback damping $\Gamma_j^{fb} \gg \Gamma_0$. The lowest temperature will be limited by the noise in the detection system and feedback circuits, as well as coupling between different directions.



Fig. 6.17 Control of the laser power with an AOM. The AOM is driven by a RF driver which uses an analog input to control the RF power electronically and a knob to tune the RF power manually. The analog input accepts a voltage from 0 to 1 V, and the knob has a reading (carrier level) from 0 to 10. For the *black curve*, the analog voltage input is zero, and the manual knob is tuned from 0 to 10. The other curves are the power of the laser as a function of the analog input when the knob is set at different readings (*red* knob reading at 1; *green* knob reading at 2; *blue* knob reading at 5)

6.6.2 Results of 3D Optical Feedback Cooling

Figures 6.18, 6.19 and 6.20 show experimental results of feedback cooling. Before feedback is turned on, the resonant frequencies $(\omega_j/2\pi)$ are 8066 ± 5 Hz, 9095 ± 4 Hz, and 2072 ± 6 Hz for the fundamental modes at 637 Pa along the X, Y, and Z axes, respectively. At this pressure, the peaks in the power spectrum due to the three fundamental modes are distinguishable, and heating effects due to the laser are negligible. We can therefore use the measured power spectra at 637 Pa to calibrate the position detectors for the fundamental modes at room temperature. After we turn on feedback cooling, the temperature of the Y mode changes from 297 to 24 K at 637 Pa. The mode temperature is obtained by fitting the measured power spectrum with Eq. 6.6.

After switching on the feedback circuits, we reduce the air pressure while keeping the feedback gain almost constant, thus the heating rate due to collisions from air molecules decreases, while the cooling rate remains constant. As a result, the temperature of the motion drops. At 5.2 mPa, the mode temperatures are 150 ± 8 mK, 1.5 ± 0.2 mK, and 68 ± 5 mK for the x, y and z modes. The mean thermal occupation number $\langle n \rangle = k_B T_j^{fb} / \hbar \omega_j$ of the y mode is reduced from about 6.8×10^8 at 297 K to about 3400 at 1.5 mK. Figure 6.21 shows the temperature of the three fundamental modes as a function of pressure. At low pressure and when the feedback gain is constant, the mode temperature should be proportional to the pressure, which is shown



Fig. 6.18 Power spectra of a trapped 3- μ m diameter microsphere along the X axis as it is cooled. The *red curve* is the intrinsic spectrum at 637 Pa without feedback cooling, the *blue curve* is the spectrum at 637 Pa with feedback cooling, the *green curve* is the spectrum at 5.2 mPa with feedback cooling, and the *orange curve* is the noise signal when there is no particle in the optical trap. The *black curve* is the fit of a thermal model (see text for details). We obtain mode temperatures from these fits

as a straight line with slope 1 in the figure. The temperature of the y mode agrees with this prediction very well at pressures above 1 Pa.

At our lowest temperatures, the power spectra are still much larger than the noise level, and the minimum temperature is achieved at pressures above the minimum pressure we can obtain, thus the electronic noise (in detection and feedback circuits) and the pressure are not the limiting factor of the current experiment. The dominant



Frequency (kHz)



Fig. 6.20 Power spectra of a trapped $3-\mu$ m diameter microsphere along the *Z* axis as it is cooled. The meanings of the curves are the same as in Fig. 6.18

limiting factor is most likely residual coupling between the intensities and directions of the cooling beams. When we change the intensity of a cooling beam using an AOM, the direction and profile of the beam is also changed slightly. This causes heating of the motion of a microsphere perpendicular to the beam while cooling it parallel. This problem should be solved by replacing the AOM's with electro-optic modulators (EOM's). The final temperature limited by the present detection system will be about



Fig. 6.21 Temperatures of the three fundamental oscillation modes along X (*black squares*), Y (*blue circles*), and Z (*red triangles*) axes as a function of the air pressure. The *dashed line* is a *straight line* with slope 1 for comparison

0.1 mK. Currently, the laser beam is attenuated before entering the detectors because the laser power is larger than the damage threshold of the detectors. If we can utilize all of the signal contained in the laser beam for feedback cooling, the final temperature can be smaller than 0.01 mK, corresponding to a thermal occupation number in the order of 10 or less.

Our result is an important step toward quantum ground-state cooling of a trapped macroscopic object in vacuum by either cavity cooling [14, 15, 22] or feedback cooling with an improved detection and feedback scheme [27, 29]. Our three-dimensional cooling enables future work on quantum superposition and entanglement of the motion between different directions. For cavity cooling of a trapped object in vacuum, it is also important to use feedback cooling to pre-cool and stabilize the object, in order to have enough time to tune the cavity cooling laser to the correct frequency for efficient cooling.

6.7 Loss of Microspheres in Vacuum

With feedback cooling, we have been able to trap a microsphere for more than one hour at pressure below 10^{-4} torr at optimal conditions. Ashkin et al. [24] had observed similar lifetimes when they levitated a 20-µm diameter sphere in vacuum (The laser intensity of the levitation trap is much smaller than the laser intensity of our dual-beam trap, and the size of the microsphere is much larger than ours). Table 6.1 shows some examples of measured lifetimes of a trapped microsphere in vacuum under different conditions. These lifetimes should be long enough to perform cavity cooling [14, 15, 22] and many other interesting experiments. Ideally, however, we would like the lifetime of the optical trap to be infinite even in vacuum.

Several things can affect the lifetime of a trapped microsphere in vacuum. For example, the alignment of the trapping beams and the gains of the feedback circuits can significantly affect the lifetime. We usually turn on feedback circuits at about 20 torr. At this pressure, the damping due to air is still large enough to keep the optical trap stable even if the feedback circuits are not tuned correctly. If the amplitude of the bead's motion increases when the feedback is turned on, the sign (polarity) of the velocity signal is incorrect, and must be inverted for the feedback loop to cool the motion. After tuning the feedback circuits correctly at 20 torr, we reduce the pressure slowly. The motion of the trapped microsphere usually becomes unstable when the pressure is reduced below 50 mtorr. We need to fine tune the horizontal and vertical directions of beam No. 2 with the two AOM's to restabilize the motion of the microsphere. After this step, we can reduce the pressure to below 10^{-5} torr. Because we need to fine tune the laser beams and the feedback circuits for each individual microsphere, the process is not very reproducible. Thus the lifetimes can be very different microspheres (Table 6.1).

We also found evidence that the ion pump intermittently undergoes electrical arcing [38] which kicks out a trapped microsphere. We observed spikes in the motion of trapped microspheres when we turned on the ion pump (an old ion pump that had

No. Total power Total power Lifetime with Loss of trapping of cooling ion pump on ^a pressure				
No.	Total power of trapping beams (mW)	Total power of cooling beams ⁺ (mW)	Lifetime with ion pump on ^a (min)	Loss pressure (torr)
1	160	~60	7	3.8×10^{-5}
2	160	~ 60	21 ^b	2.4×10^{-4}
3	130	~ 60	5	$3.9 imes 10^{-5}$
4	130	~ 60	11 ^b	$2.6 imes 10^{-4}$
5	130	3.6	26	9.4×10^{-6}
6	130	6.5	22	7.7×10^{-6}
7	113	39	88	2.4×10^{-6}
8	82	19	69	2.5×10^{-6}

Table 6.1 Examples of the lifetimes of a trapped 3.0-µm diameter microsphere in vacuum under different conditions

⁺ The waists of the cooling beams are about 9μ m, so only parts of cooling beams pass through the trapped microsphere ^aThe ion pump was turned on at about 1 mtorr; and the pressure dropped below 1.0×10^{-4} torr within about 1 min after the ion pump was on ^bThe ion pump was not turned on; the lifetime was the trapping time at pressures below 1 mtorr

not been used for several years). Sometimes the microsphere was lost immediately after we turned on the ion pump. We moved the ion pump further away from the optical trap to alleviate this problem. This problem disappeared after the ion pump was used for a few months. However, the lifetime of the optical trap in vacuum was still only on the order of 10 min. Then we stopped using the ion pump for a while, and used two sorption pumps in sequence to achieve lowest pressures of about 10^{-5} torr. However, we still observed the loss of microspheres after trapping for about 15 min at low pressures. Then we considered the possibility that the sudden loss of trapped microspheres might be caused by floating dust in the air. A dust particle can cause fluctuations in the laser power and profile when it passes though the focus of a laser beam. So we used plastic sheets to seal the space between two lenses where a laser was focused in between. This method did not notably increase the lifetime of the trap in vacuum.

The final loss of a trapped microsphere is most likely caused by the heating due to light absorption. It seems that there is some dependance of the trap lifetime on the laser powers. The longest lifetime, 88 min, was observed when the total power of the trapping beams (1064 nm) was 113 mW and the total power of the cooling beams (532 nm) was 39 mW. Because the waists of cooling beams were much larger than the size of the microsphere, only a part of cooling beams passed through the microsphere. However, silica has much larger absorption at 532 than 1064 nm. So the heating effect of the cooling beams may be comparable to that of the trapping beams.

At pressures below 1 mtorr, the internal temperature of a trapped microsphere is mostly cooled by blackbody radiation [14, 24]. The wavelength of a photon with energy of $k_B T$ is $48 \,\mu m \,(\lambda = \frac{hc}{k_B T})$, where *h* is the Planck constant) at room temperature. This wavelength is much larger than the size of our microspheres. Thus we can treat the microsphere as a dipole in calculating the blackbody radiation when

the internal temperature of the microsphere is not much higher than room temperature. If we assume that the microsphere has a constant and temperature-independent permittivity $\epsilon(\omega) \approx \epsilon_{bb}$ across the blackbody radiation spectrum, the microsphere radiates blackbody energy at a rate [14]:

$$\frac{dE}{dt} \simeq -\frac{75}{\pi^2} \frac{V}{c^3 \hbar^4} \operatorname{Im} \frac{\epsilon_{bb} - 1}{\epsilon_{bb} + 2} (k_B T_{\text{int}})^5 \tag{6.19}$$

where V is the volume of the microsphere, and T_{int} is the internal temperature of the microsphere. Similarly, the microsphere absorbs blackbody radiation from the environment at a rate:

$$\frac{dE}{dt} \simeq \frac{75}{\pi^2} \frac{V}{c^3 \hbar^4} \operatorname{Im} \frac{\epsilon_{bb} - 1}{\epsilon_{bb} + 2} (k_B T_{\text{env}})^5$$
(6.20)

where T_{env} is the temperature of the environment.

In high vacuum, we can neglect the effect of background gas. The equilibrium internal temperature will be established when the sum of Eqs. 6.19 and 6.20 is equal to the heating rate due to light absorption of the laser beams. The absorbed energy of the microsphere from the laser beams is proportional to the power of lasers passing through the microsphere, and the optical absorption rate of the microsphere. Figure 6.22 shows the calculated internal temperature of a trapped 3.0- μ m diameter microsphere as a function of the environmental temperature, power of laser beams that pass through the microsphere, and the absorption rate of the microsphere.



Fig. 6.22 The equilibrium internal temperature of a trapped $3.0-\mu$ m diameter microsphere irradiated by 10 and 100 mW of 1064 nm laser power in ultrahigh vacuum as a function of the absorption coefficient of the microsphere

When the environment is at room temperature and the power of the laser passing through the microsphere is 100 mW, the internal temperature of the microsphere stays almost constant when the absorption coefficient is smaller than 10 dB/km (Fig. 6.22). However, the internal temperature increases significantly if the absorption coefficient is 1000 dB/km or higher. The microsphere will be lost when the internal temperature is above a certain threshold temperature. The silica microsphere is initially amorphous. It may undergo phase transition and become crystalline at high temperatures. Silica has many crystalline forms. The α -quartz converts to β -quartz at 846 K. At higher temperatures in vacuum, the silica microsphere will sublimate or melt and evaporate. The lifetime of the optical trap should be longer for a lower laser power.

Pure silica core optical fiber with loss of 0.148 dB/km at 1570 nm, 0.265 dB/km at 1310 nm, and 0.6 dB/km at 1064 nm have been reported [39]. Whispering-gallery modes in fused-silica microspheres with quality factor of $Q = 0.8 \times 10^{10}$ at 633 nm have been demonstrated experimentally [40]. This is close to the ultimate level determined by the fundamental material attenuation as measured in optical fibers (0.7 dB/km at 633 nm). The loss increases significantly if the silica contains OH [41]. Commercial monodisperse silica microspheres are produced by the chemical reaction of tetraalkoxysilanes (TEOS, Si(OC₂H₅)₄) in alcoholic solutions of water and ammonia [42, 43]. Thus they are expected to contain OH, C, and N which will increase their absorption coefficient.

In the future, we should directly measure the internal temperature of an optically trapped microsphere using Raman spectroscopy [44]. This should give us a better insight of the final loss mechanism of a trapped microsphere in vacuum. We may also be able to purify a trapped microsphere in situ at high pressures by heating it with a CO_2 laser [45]. Heating a trapped microsphere by a CO_2 laser can melt the microsphere and reduce the size of the microsphere by evaporation. This also provides a novel method to produce and trap a nanosphere in air and vacuum.

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Chapter 7 Towards Quantum Ground-State Cooling

Quantum ground-state cooling of an optically trapped microsphere in vacuum can be achieved by feedback cooling with an optimal detection and feedback scheme [1, 2]. However, it will be difficult to achieve this since the real detection system and feedback circuits have electronic noise. Another method to achieve quantum-ground state cooling is cavity cooling [3–7]. Cavity cooling has been implemented in many different micromechanical systems [8–17], and has been used to successfully cool the mechanical vibration of a 100 nm thick aluminum membrane with a diameter of 15 μ m to the quantum ground state recently [18]. Recent theoretical calculations show that cavity cooling can be used to cool the mechanical motion of an optically trapped dielectric particle in vacuum to the quantum ground-state from room temperature [19–26].

7.1 Principle of Cavity Cooling

As shown in Fig. 7.1a, we consider a microsphere trapped in a cavity by a dual-beam optical tweezer. We use a dual-beam trap here because a single-beam trap requires a NA ≈ 0.95 objective lens. The working distance of a common high NA objective lens is too short (usually about 0.2 mm) to integrate the optical cavity and the optical tweezer together. Thus we use two low NA lenses to create a dual-beam trap. In the system with which we have carried out feedback cooling, the distance between the two front surfaces of the two lenses is about 3.5 mm. This should be enough for the cavity cooling laser to pass though.

Let the vibration frequency of the microsphere along the *z* axis be ω_M , the frequency of the cooling laser be ω_L , the resonant frequency of the cavity be ω_C , the intrinsic cavity linewidth be κ , and the rate of a photon scattered by the microsphere be γ_{sc} . For simplicity, we assume the linewidth of the cooling laser be much smaller than the cavity linewidth.



Fig. 7.1 a Scheme of 1D cavity cooling. A microsphere is trapped inside an optical cavity with a dual-beam trap. **b** Principle of 1D cavity cooling. The frequency of the cooling laser (ω_L) is slightly smaller than the resonant frequency of the optical cavity (ω_C) . The mechanical vibration of the trapped microsphere at frequency ω_M induces two side bands of the laser at frequencies of $\omega_L - \omega_M$ and $\omega_L + \omega_M$.

When $\gamma_{sc} \ll \kappa \ll \omega_M$, the system is in the resolved-sideband limit. The mechanical vibration of the trapped microsphere will induce two side bands of the laser at frequencies of $\omega_L - \omega_M$ and $\omega_L + \omega_M$ (Fig. 7.1b). If the frequency of the cooling laser is red-detuned from than the resonant frequency of the cavity by ω_M , the blue sideband $\omega_L + \omega_M$ will be on resonance with the cavity and the red sideband $\omega_L - \omega_M$ will be further away from the resonance of the cavity. Photons in the blue sideband leak out of the cavity because they are on resonance, while photons in the red sideband are trapped in the cavity. Thus on average the photons which come out of the cavity have larger frequency than the photons which come into the cavity. These photons carry away energy that is from the kinetic energy of the microsphere. Thus the photons cool the vibration of the trapped microsphere.

Let us consider a microsphere at position z moving with momentum p along the z axis inside of a driven cavity. The microsphere causes the resonant frequency of a cavity with intrinsic resonant frequency ω_0^C to shift by an amount [19]:

$$\delta\omega_C = -\frac{1}{2} \frac{\int d^3 \mathbf{r} \delta P(\mathbf{r}) \cdot \mathbf{E}(\mathbf{r})}{\int d^3 \mathbf{r} \epsilon_0 \mathbf{E}^2(\mathbf{r})} \cdot \omega_0^C, \tag{7.1}$$

where $\mathbf{E}(\mathbf{r})$ is the bare cavity mode profile, and $\delta P(\mathbf{r})$ is the variation in permittivity introduced by the microsphere. Thus the resonant frequency of the cavity with microsphere is $\omega_C = \omega_0^C + \delta \omega_C$. If the diameter of the microsphere is much smaller than the wavelength of the laser, we can use Rayleigh approximation. In Rayleigh approximation, we have $P(\mathbf{r}') \approx \alpha_{\text{ind}} E(\mathbf{r}) \delta(\mathbf{r} - \mathbf{r}')$, where \mathbf{r} is the center-of-mass position of the microsphere, $\alpha_{\text{ind}} = 3\epsilon_0 V(\frac{\epsilon-1}{\epsilon+2})$ is its polarizability, V is its volume, and ϵ is the electric permittivity.

7.1 Principle of Cavity Cooling

The classical motion of the microsphere along the *z* axis can be described by the following coupled equations for the cavity field *E*, the particle momentum and its position [27]:

$$\dot{E} = [-\kappa - \gamma_{sc}(z) + i\Delta_C - i\delta\omega_C(z)]E - \alpha, \qquad (7.2)$$

$$\dot{p} = -|E|^2 \frac{d}{dz} \delta \omega_C(z) - M \omega_M^2 z, \qquad (7.3)$$

$$\dot{z} = p/M. \tag{7.4}$$

Here $\Delta_C = \omega_L - \omega_0^C$ is the detuning of the laser relative to the resonant frequency of the empty cavity, and α describes the external pump laser. Because the cavity mode is a standing wave, the scattering rate $\gamma_{sc}(z)$ and the frequency shift $\delta\omega_C(z)$ have the shape of $\cos^2(2\pi z/\lambda)$. This classical model predicts the microsphere will stop as $t \to \infty$.

Quantum mechanical calculations show that the microsphere will not stop completely. In the limit $\omega_M \gg \kappa$ and taking the cooling rate $\Gamma^{cool} \approx \kappa$ (can be achieved by control the power and detuning of the cooling laser), the steady-state phonon number of the vibration is [19]:

$$\langle n_f \rangle \approx \frac{\kappa^2}{16\omega_M^2} + \frac{\gamma_{sc}}{\kappa}.$$
 (7.5)

Thus quantum ground-state cooling ($\langle n_f \rangle < 1$) can be achieved if $\omega_M \gg \kappa$ and $\kappa \gg \gamma_{sc}$. The value of γ_{sc} depends on the size of the microsphere, the waist of the cavity mode and the length of the cavity. Calculations using Rayleigh approximation show that $\langle n_f \rangle < 1$ can be achieved at reasonable conditions for microspheres with diameter smaller than 400 nm when the wavelength of the cooling laser is 1064 nm [25]. However, the Rayleigh approximation is only valid when the size of the microsphere is far smaller than the wavelength of the laser. It overestimates the scattering rate by about 3 orders of magnitude when the diameter of the microsphere is the same as the wavelength of the laser [28]. Thus we believe that ground-state cooling with a cavity should be possible for microspheres with diameter of 1 μ m or slightly larger.

7.2 3D Ground-State Cooling with a Single Cavity

A microsphere will scatter the cooling laser to all three dimensions and cause 3D heating. The heating effects of laser noise are also 3D. If only one-dimensional motion is cooled efficiently, the others will be heated continuously and the microsphere will eventually be kicked out of the trap. In order to achieve ground state cooling of an optically trapped nanosphere, we must use a 3D cooling scheme. We can add two more cavities for cooling the other two dimensions, but the system will become too



Fig. 7.2 a A nanosphere is trapped by a dual-beam optical tweezer inside of a cavity. The cavity is driven by three lasers in TEM00, TEM01 and TEM10 modes to cool the 3D motion of the trapped nanosphere. **b** Three cooling modes TEM00, TEM01, and TEM10, and their radial distribution. The *black dot* represents the position of the trapped nanosphere. Figure is courtesy of Z. Q. Yin.

complex to be realized experimentally. A better method to cool and measure the 3D motion of a nanosphere is to use the TEM00, TEM01, and TEM10 modes of a single cavity [26]. The TEM01 and TEM10 beams can be generated from a TEM00 beam by two phase plates [29]. Each one of these three modes can be coupled to the motion of a trapped microsphere along one orthogonal axis. Thus they can be used to cool and detect the 3D motion of a microsphere.

Trapping single atoms in a high-finesse cavity driven by three lasers at TEM00, TEM01, and TEM10 modes simultaneously has been demonstrated in an experiment already [30]. A recent experiment also used a tilted cavity TEM10 mode to measure the trajectories of single neutral atoms deterministically [31].

For simplicity, we consider a microsphere with size much smaller than the laser wavelength. In this case, the microsphere is in fact a nanosphere. The nanosphere is trapped in a cavity by an optical tweezer, as shown in Fig. 7.2. The trapping frequencies are ω_1 , ω_2 , and ω_3 along the *z*, *x*, and *y* axes, respectively. The cavity is driven by three lasers in TEM00, TEM01 and TEM10 modes to cool the 3D motion of the trapped nanosphere. The TEM01 laser is *s*-polarized, and the TEM00 and TEM10 lasers are *p*-polarized. The TEM01 and TEM10 lasers have almost the same frequency, and the TEM00 laser has a frequency different from those of the TEM01 and TEM10 lasers. Thus the TEM01 laser can be separated from the other two lasers by a polarizing beam splitter, and the TEM00 and TEM10 lasers can be separated by a grating. The frequency differences between the TEM00 and TEM01 lasers can be very large, and the TEM01 and TEM10 modes are orthogonal in polarizations. Therefore the interference between the three cavity modes can be neglected.

The TEM00 mode laser is used to cool the motion of the nanosphere along the *z* axis, the TEM01 mode laser is for cooling the motion along the *y* axis, and the TEM10 mode laser is for cooling the motion along the *x* axis. The resonant frequencies of the cavity modes a_{c1} , a_{c2} , a_{c3} are ω_{c1} , ω_{c2} , ω_{c3} , respectively. The detunings between the lasers and the cavity modes are $\Delta_{cj} = \omega_c^j - \omega_L^j$ (j = 1, 2, 3). Let us use a_j to characterize the phonon mode along q_j direction with $q_1 = z$, $q_2 = x$, $q_3 = y$, Ω_j to be the driving strength of the lasers and U_j to characterize the coupling between the cavity mode a_{cj} and the nanosphere. In the limit that $\epsilon \gg 1$, we have [26]

$$U_{1} = -\frac{3V}{2V_{c1}} \exp\left(-\frac{2x^{2}+2y^{2}}{w^{2}}\right) \cos^{2}(k_{1}z+\varphi_{1})\omega_{c1},$$

$$U_{2} = -\frac{3V}{2V_{c2}} \frac{x^{2}}{w^{2}} \exp\left(-\frac{2x^{2}+2y^{2}}{w^{2}}\right) \cos^{2}(k_{2}z+\varphi_{2})\omega_{c2},$$

$$U_{3} = -\frac{3V}{2V_{c3}} \frac{y^{2}}{w^{2}} \exp\left(-\frac{2x^{2}+2y^{2}}{w^{2}}\right) \cos^{2}(k_{3}z+\varphi_{3})\omega_{c3},$$

where $V_{c1} = (\pi/4)Lw^2$ and $V_{c2} = V_{c3} = (\pi/16)Lw^2$.

Let us assume the trapping lasers to be much stronger than cooling lasers, and neglect the effects of cooling lasers on trapping. If we carefully choose the location of the trap, such as $z_0 = 0$, $x_0 = y_0 = 0.25w$, $\varphi_1 = \pi/4$, and $\varphi_2 = \varphi_3 = 0$, the gradients of the three light fields lie approximately along the three axes. The effective Hamiltonian is [26]

$$H_{eff} = \sum_{j=1}^{3} \left[\hbar \omega_j a_j^{\dagger} a_j - \hbar \Delta_j a_{cj}^{\dagger} a_{cj} + \frac{\hbar \Omega_j}{2} (a_{cj} + a_{cj}^{\dagger}) + \hbar g_j a_{cj}^{\dagger} a_{cj} (a_j + a_j^{\dagger}) \right],$$
(7.6)

where $g_j = q_j^{zero} \partial U(x, y, z) / \partial j|_{x=x_0, y=y_0, z=z_0}$ characterizes the coupling strength between the cavity mode and the oscillation of the nanosphere, and $q_j^{zero} = \sqrt{\hbar/2M\omega_j}$ is the zero-point fluctuation for the phonon mode a_j . In general, g_1 can be one to two orders larger than g_2 and g_3 . This effective Hamiltonian (7.6) is valid when the vibration amplitude of a trapped nanosphere is much smaller than the wavelength of the laser.

In the limit of resolved sideband cooling where $\omega_j \gg \kappa_j$, and when the driving strength is small, the final phonon number is [26, 32]

$$n_{mj} = -\frac{(\omega_j + \Delta'_{cj})^2 + (\kappa_j/2)^2}{4\omega_j \Delta'_{ci}},\tag{7.7}$$

where Δ'_{cj} is the effective detuning between the driving laser and the cavity mode a_{cj} . In the special case of $\Delta'_{cj} = -\omega_j$, the final phonon number is $n_{mj} = (\kappa_j/4\omega_j)^2 \ll$ 1. Thus the motion of the microsphere can be cooled to ground state in all three dimensions in the resolved sideband regime. The cooling rate is $\Gamma_j = g_j^2 |\alpha_j|^2 / [\kappa_j(1 + \frac{\kappa_j^2}{2})]$

 $\frac{\kappa_j^2}{16\omega_j^2}$)], where α_j is the amplitude of the cavity mode a_{cj} .

7.3 Heating Effects of Laser Noise

In a real experiment, the noise of lasers may cause significant heating of the motion of an optically trapped microsphere and may prevent ground-state cooling. The heating effect due to the photon shot noise of a laser has been found to be negligible [19]. However, the experiment is most likely to be limited by classical noise sources in the laser beams rather than the photon shot noise [26].

Let us first consider heating effects from the trapping laser [33]. In order to achieve ground-state cooling, the total heating rate from laser intensity fluctuation and laser pointing fluctuation should be much smaller than the cavity cooling rate. The relative intensity fluctuation of a laser is defined as $\epsilon(t) = (I(t) - I_0)/I_0$, where I_0 is the average laser intensity and I(t) is the laser intensity at time *t*. The heating rate due to intensity fluctuation can be obtained by using a first-order time-dependent perturbation theory [33]:

$$\Gamma_{\epsilon} \equiv \frac{\langle E \rangle}{E} = \frac{\pi}{2} \omega_j^2 S_{\epsilon}(2\omega_j) \tag{7.8}$$

where $S_{\epsilon}(\omega) = \frac{2}{\pi} \int_0^{\infty} d\tau \cos(\omega \tau) \langle \epsilon(t) \epsilon(t+\tau) \rangle$ is the one-sided power spectrum density of the relative intensity noise. For a trap frequency on the order of MHz, Γ_{ϵ} approaches the order of 10^{-1} Hz when the $S_{\epsilon}(\omega)$ is on the order of 10^{-14} Hz⁻¹.

The laser pointing fluctuation causes fluctuation of the position of the trap center. The heating rate due to pointing fluctuation is [33]:

$$\Gamma_j^p \equiv \frac{\langle \dot{E} \rangle}{\hbar \omega_j} = \frac{\pi}{2} M \omega_j^4 S_j(\omega_j) / (\hbar \omega_j), \qquad (7.9)$$

where $S_j(\omega)$ is the power spectrum density of position fluctuation of the trap center. Here the heating rate Γ_j^p denotes the increase of the phonon number per second. If we want the Γ_j^p to be on the order of 10^{-1} Hz, we must have $S_j(\omega_j)$ to be about 10^{-35} m²/Hz for $\omega_j \sim 1$ MHz. Experimentally, $S_j(\omega)$ has been able to be controlled to less than 10^{-34} m²/Hz for $\omega \sim kHz$ [34]. For frequencies much larger than the resonant frequencies of an instrument, $S_j(\omega_j)$ drops down quickly. Therefore, it should be possible to control the laser pointing fluctuation to be small enough to have $\Gamma_i^p \sim 0.1$ Hz.

For the cavity cooling laser, the dominate heating source is the phase noise of the laser [35–37]. The phase noise of the laser is related to the finite linewidth of the laser. The field of a laser with finite linewidth can be written as $\varepsilon(t) = \varepsilon e^{i\phi(t)}$, where the phase noise $\phi(t)$ is assumed to be Gaussian with zero mean value. For a Lorentzian noise spectrum with $S_{\dot{\phi}}(\omega) = 2\Gamma_L \gamma_c / (\gamma_c^2 + \omega^2)$, and correlation function $\langle \phi(s)\phi(s') \rangle = \Gamma_L \gamma_c \exp(-\gamma_c |s-s'|)$, where Γ_L is the linewidth of the laser and γ_c^{-1} is the correlation time of the phase noise, the minimum phonon number limited by the phase noise is [36]

$$n_{ph} = n_c \frac{\Gamma_L}{\kappa} \frac{\gamma_c^2}{\gamma_c^2 + \omega_i^2}.$$
(7.10)

If $\Gamma_L = 1$ kHz, $\gamma_c = 3$ kHz, $\omega_j = 10^6$ Hz, and $n_c = 10^7$, we have $n_{ph} \ll 1$. Thus the heating due to the phase noise of a cooling laser with linewidth on the order of 1 kHz will not prevent ground-state cooling.

7.4 Applications of Cooled Microspheres in Vacuum

7.4.1 Measuring Weak Forces

Optically trapped microspheres in water are sensitive to forces on the order of 10^{-12} N [38], and have found extensive applications in biophysical research. For example, optical tweezers have been used to measure the force exerted by single kinesin molecules and the binding force between two individual stands of DNA molecules [39, 40].

An optically trapped microsphere in vacuum provides an even more sensitive force detector. The microsphere (just as a cantilever of an AFM) is subject to thermal fluctuation forces from the environment. The minimum force that a microsphere (or a cantilever of an AFM) can detect is linked to the friction that it experiences from the environment. The minimum detectable force is [41]:

$$F_{min} = \sqrt{4M\Gamma_0 k_B T \Delta_b},\tag{7.11}$$

where Γ_0 is the viscous damping factor and Δ_b is the bandwidth of the measurement. From this equation, it is clear that a microsphere in vacuum is much more sensitive to a weak force than a microsphere in water because of the smaller damping Γ_0 . A microsphere is also more sensitive to a weak force than a normal AFM cantilever because of its smaller mass M. Feedback cooling or cavity cooling do not affect the force sensitivity [42], but they can localize the position of the microsphere and benefit force measurements.



Minimum detectable force by a microsphere in vacuum

Fig. 7.3 Minimum detectable force by a $3 \mu m$ diameter microsphere trapped in vacuum. The measurement bandwidth is 1 Hz

Figure 7.3 shows the calculated minimum detectable force by a $3 \mu m$ microsphere at different pressures with 1 Hz bandwidth. The force sensitivity increases when the pressure decreases. $F_{min} = 2.9 \times 10^{-15}$ N at 1000 torr, $F_{min} = 4.0 \times 10^{-18}$ N at 10^{-4} torr, and $F_{min} = 1.3 \times 10^{-21}$ N at 10^{-11} torr. The microsphere at 1000 torr is already much more sensitive than a common AFM cantilever, which can measure forces on the order of 1 nN.

This potential force sensitivity is remarkable. For comparison, the Newtonian gravitational force between two point-like objects is

$$F_g = \frac{G_N m_1 m_2}{r^2},$$
 (7.12)

where m_1 and m_2 are two masses separated by distance *r*. The gravitational force between two 1 g masses separated by 1 m is 6.67×10^{-17} N.

It has been proposed that an optically trapped microsphere can be used to search for non-Newtonian gravity forces via an enhanced sensitivity of 10^5-10^7 over current experiments at the 1 μ m length scale [43].

7.4.2 Measuring the Impact of Single Molecules

The 3D cavity cooling scheme (Fig. 7.2) can also be used to measure the 3D motion of the nanosphere. In the limit of resolved sideband cooling where $\omega_j \gg \kappa_j$, and when the driving strength is small $(g_j \alpha_j \ll \kappa_j)$, the output field from the cavity a_{cj}^{out} is related to the phonon mode of the motion of the nanosphere a_j by [26]



Fig. 7.4 Distribution of the average increase of the phonon number of the mechanical mode a_3 after single elastic collisions between the nanosphere and molecules. The mass of the molecules is assumed to be $m_a = 6.63 \times 10^{-26}$ kg (*blue curve*) or $m_b = 2.18 \times 10^{-25}$ kg (*red curve*). The mass of the nanosphere is $m_b = 1.03 \times 10^{-18}$ kg. The temperature of the gas is 300 K. Plot is courtesy of Z. Q. Yin

$$a_{cj}^{\text{out}} = -i \frac{2g_j \alpha_j}{\sqrt{\kappa_j}} a_j + a_{cj}^{\text{in}}.$$
(7.13)

Therefore the 3D motion of the nanosphere can be measured by detecting the output fields. In the resolved sideband limit, the output field is nearly vacuum, and will have a signal when there are collisions between the residual molecules in vacuum and the nanosphere.

If the collisions between the molecules and the nanosphere are elastic, and the nanosphere is in its ground state before the collisions, the average increase of the photon number for vibration mode a_i after single collisions is [26]

$$n_{j0} = \frac{2k_B T_{env} m_{mol}}{\hbar \omega_j M},\tag{7.14}$$

where T_{env} is the temperature of the residual molecules that is the same as the environmental temperature, m_{mol} is the mass the molecule, and M is the mass of the nanosphere. Since n_{j0} is proportional to m_{mol} , we can distinguish the molecules with different masses in the measurement. The effective temperature T_j of mode a_j after single collisions can be obtained from

$$\frac{1}{e^{\hbar\omega_j/k_B T_j} - 1} = n_{j0}.$$
(7.15)

The distribution of the average increase of the phonon number after single collisions is [26]

$$f(n_j) = \frac{2}{\sqrt{\pi}} \left(\frac{\hbar\omega_j}{k_B T_j}\right)^{3/2} \exp\left(-n_j \frac{\hbar\omega_j}{k_B T_j}\right).$$
(7.16)

Figure 7.4 shows the calculated distribution of mean phonon increase after single collisions with two different molecules. The two curves for the two different molecules are very different. Thus we can distinguish different molecules from the measurement. In the calculation, the radius of the nanosphere is r = 50 nm and the mass of the nanosphere is $M = 1.03 \times 10^{-18}$ kg ($\rho = 1.96$ g/cm³). The optical tweezer is formed by a laser with power of $P_t = 25$ mW and wavelength of $\lambda = 1500$ nm, focused by a lens with numerical aperture N = 0.9. The trap frequency is $(\omega_1, \omega_2, \omega_3)/2\pi \simeq (0.5, 0.5, 0.2)$ MHz [25]. The mass of the molecules is assumed to be $m_a = 6.63 \times 10^{-26}$ kg (blue curve) or $m_b = 2.18 \times 10^{-25}$ kg (red curve).

7.4.3 Searching for Gravity-Induced Quantum-State Reduction

A cooled microsphere in vacuum can also be used to study the gravity-induced quantum state reduction [44]. After cooling and creation of a superposition state in momentum, the optical trap can be switched off to let a microsphere undergo free-fall in vacuum [25]. The wavefuction will expand during free-fall and become a superposition state in space. The finite lifetime of a superposition due to gravity-induced state reduction is predicted to be on the order of $\hbar r/(GM^2)$ when the superposition is composed of states separated by a distance larger than the size of the microsphere [44, 45], where G is Newton's gravitational constant. The predicted lifetime is about 3 ms for a 3 μ m diameter microsphere, which is shorter than the environmental decoherence time in good vacuum and thus measurable.

Other sources of decoherence must be minimized in order to measure the gravityinduced decoherence. The laser beams are switched off during the free fall of the microsphere. So the decoherence due to laser scattering can be neglected [46]. A dominate environmental decoherence source is the residual air molecules in the vacuum chamber. The localization rate due to air molecules is $\Lambda_{air} = 8\sqrt{2\pi}m_a\bar{v}Pr^2/(3\sqrt{3}\hbar^2)$, where *P* is the air pressure, m_a is the mass of the air molecules and \bar{v} is their thermal velocity [46]. Λ_{air} can be reduced by reducing the air pressure and the environmental temperature.

There are many other applications of an optically trapped dielectric particle near the quantum ground state. Besides trapping glass microspheres, the dielectric particle can even be a virus or other living organism, in which case it would be possible to to produce Schrödinger cat states of living organisms [20].

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Appendix

Physical Properties of Some Common Materials

 Table A.1 Physical properties of some common materials. The values are taken from reference
 [1]

Material	Refractive index	Density	Dynamic
	(at 389 IIII)	(g/cm ²)	viscosity (µPa s)
Vacuum	1	0	0
Helium @ 0 °C, 1 atm	1.000036	0.000176	18.69
Air @ 0 °C, 1 atm	1.000292	0.00128	17.21
Air @ 27 °C, 1 atm	1.000265	0.00116	18.54
Liquid helium @ -269°C	1.0245 ^{<i>a</i>}	0.129	3.319
Liquid nitrogen @ -196 °C	1.199	0.8066	161.4
Water @ 20 °C	1.3337 ^b	0.9982	1002
Water @ 40 °C	1.3310	0.9922	652.7
Water @ 60 °C	1.3276	0.9832	466.0
Ethanol @ 20 °C	1.361	0.7893	1074 ^c
Acetone @ 20 °C	1.359	0.7845^{c}	306 ^c
Ethylene glycol @ 20 °C	1.432	1.114	16100 ^c
Ice (Ih) @ -7 °C	1.31^{d}	0.918 ^d	
Fused silica	1.458	2.21	
α-Quartz	$1.544(n_o), 1.553(n_e)$	2.65	
Silica microspheres ^e	1.43-1.46	2.0	
Polystyrene beads ^e	1.59	1.05	
Sapphire	$1.761(n_o), 1.769(n_e)$	3.97	
Diamond	2.418	3.51	

^aAt 546 nm

^bImaginary part of the refractive index of water and ice can be found at [2, 3]

^cAt 25 °C

^dValues from [4]

^eMaterials and values from Bangs Laboratories, Inc. [5]

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Curriculum Vitae

Tongcang Li received his Bachelor of Science degree in Physics from the University of Science and Technology of China in 2004, and his PhD degree in Physics from the University of Texas at Austin in 2011. He is currently a Postdoctoral Fellow at the University of California, Berkeley. He did some theoretical studies on spintronics and carbon nanotubes during 2002–2004. He was working on experiments in Bose–Einstein condensation of sodium atoms during 2005–2008. In Sep. 2008, he started a pioneering experiment to trap and cool glass microspheres in vacuum towards the quantum ground state of an optical tweezer, which is the content of this thesis. This work has led to several breakthroughs, including the measurement of the instantaneous velocity of a Brownian particle, a task that was said to be impossible by Albert Einstein in 1907. Recently, he proposed "space–time crystals" of trapped ions and a method to realize it experimentally by confining ions in a ring-shaped trapping potential with a static magnetic field. This work has generated huge interests in physics community and has a potential to open new directions in many-body physics.