

Step-by-step guide to the realization of advanced optical tweezers

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Since the pioneering work of Arthur Ashkin, optical tweezers (OT) have become an indispensable tool for contactless manipulation of micro- and nanoparticles. Nowadays OT are employed in a myriad of applications demonstrating their importance. While the basic principle of OT is the use of a strongly focused laser beam to trap and manipulate particles, more complex experimental setups are required to perform novel and challenging experiments. With this article, we provide a detailed step-by-step guide for the construction of advanced optical manipulation systems. First, we explain how to build a single-beam OT on a homemade microscope and how to calibrate it. Improving on this design, we realize a holographic OT, which can manipulate independently multiple particles and generate more sophisticated wavefronts such as Laguerre–Gaussian beams. Finally, we explain how to implement a speckle OT, which permits one to employ random speckle light fields for deterministic optical manipulation. © 2015 Optical Society of America

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1. INTRODUCTION

The study of the interaction between light and matter has been extremely prolific, leading to the invention of novel techniques that have had a profound impact in many different fields. In particular, the use of light to exert forces and torques on microscopic and nanoscopic particles has blossomed: since the pioneering work of Ashkin and co-workers [1,2], optical tweezers (OT) have become an indispensable tool in, e.g., biophysics, experimental statistical physics, and nanotechnology [3–6]. At the same time, starting from the relatively simple tightly focused laser beam employed by Ashkin *et al.* to confine a microscopic particle in three dimensions [2], experimental setups for optical trapping and manipulation have grown more complex, leading to OT capable of, e.g., trapping multiple particles [3], rotating particles by transferring spin [7,8] and orbital angular momentum [3,9], measuring forces in the femtonewton range [10,11], and sorting particles based on their physical properties [5,12,13].

Many OT setups are built on commercial microscopes, especially for experiments on biological samples such as DNA, molecular motors, and cells [14]. While commercial microscopes offer several advantages, e.g., high-quality imaging of the sample, easy integration with powerful imaging techniques such as phase contrast, fluorescence excitation, and differential interference contrast, they provide limited flexibility when dealing with more elaborate experiments. Thus, many research groups have started to develop homemade microscopes, which have the great advantages of being very versatile, much less expensive, and, if implemented properly, a superior mechanical stability.

In the literature, there are various articles that provide guidance on how to implement an OT setup. For example, Refs. [15,16] explain how to build and calibrate a single-beam OT setup for undergraduate laboratories, and Refs. [17,18] explain how to transform a commercial microscope into an OT. However, to the best of our knowledge, a detailed explanation

of how to build an homemade OT setup with advanced functionalities, such as multiple holographic optical traps and speckle OT, is still missing from the literature.

In this article, we provide a detailed step-by-step guide to the implementation of state-of-the-art research-grade OT. The only requirement to be able to follow the procedure we explain is a basic knowledge of experimental optics. We start by implementing a high-stability homemade optical microscope, and then we transform it into a series of OT with increasingly more advanced capabilities. In particular, we explain in detail the implementation of three OT setups with cutting-edge performance: (1) a high-stability single-beam OT capable of measuring forces to a few femtonewtons [11], (2) a holographic optical tweezers (HOT) [19,20] capable of creating multiple traps and advanced optical beams such as Laguerre–Gaussian beams, and (3) a speckle OT particularly suited to develop robust applications in highly scattering environments and in microfluidic devices [13,21]. For each setup, we also provide a time-lapse movie (see [Media 1](#), [Media 2](#), and [Media 3](#)) to make it easier to understand and follow each step of the implementation. The software employed for the data analysis and the generation of the holographic masks is freely available online [22].

In Section 2, we provide a brief overview of the setup. In Section 3, we provide a detailed step-by-step procedure for the realization of each setup: first, we build a homemade microscope capable of performing digital video microscopy (DVM) by analyzing the images acquired with a CCD camera; then, we show how to upgrade this microscope to a single-beam OT equipped with a position sensitive detection device based on a quadrant photodiode (QPD); finally, we describe how to build a HOT and a speckle OT. In Appendix A, we discuss how to calibrate the single-beam OT. In Appendix B, we explain the fundamentals of how spatial light modulators (SLM) work, and we provide some examples of the phase masks and algorithms most commonly employed with HOT.

2. OPTICAL TWEEZERS SETUP

A. Homemade Microscope

In the implementation of every OT experiment, it is crucial to start with a microscope that is as mechanically stable as possible. For this reason, our homemade microscope [Figs. 1(a) and 1(b)] is built on a stabilized optical table, which permits us to minimize environmental vibrations, and its structure is realized with AISI 316 stainless steel, which offers one of the lowest thermal expansion coefficients and a great rigidity. This structure consists of three levels. The first level, which lies on the optical table, is where the optical components necessary to focus the sample image on the camera (and to prepare the optical beam for the OT, Subsection 2.B) are hosted. In particular, there is a mirror arranged at 45° to reflect the illumination light toward the CCD camera (and the laser beam toward the objective lens for the OT, Subsection 2.B). Videos of the sample are recorded by a fast CCD camera (Mikrotron, MotionBLITZ EoSens Mini 1).

The second level is realized with a breadboard held on eight columns; here is where the objective mount as well as the stages to hold and manipulate the sample are hosted. The objective holder is mounted vertically and comprises a high-stability

translational stage (Physik Instrumente, M-105.10) with manual micrometer actuator (resolution 1 μm); if needed, this stage can be equipped easily with a piezo actuator for computer-controllable and more precise movements. The sample stage is realized by a combination of a two-axis (x and y) manual stage for coarse (micrometric) movement (Newport, M406 with HR-13 actuators) and a three-axis piezoelectric stage (Physik Instrumente, PI-517.3CL) with nanometer resolution.

Finally, the third level, which is built on the second-level breadboard using four columns, hosts the illumination lamp (and the position detector devices for the photonic force microscope (PFM), Subsection 2.C). To illuminate the sample, we opted for a cold-light white LED (Thorlabs, MCWHL5), which offers high light intensity without heating the sample. The white LED is collimated by an aspheric lens. The whole system is easily realized using modular optomechanics components (Thorlabs, cage system). As the condenser lens, we use an objective with magnification 10× or 20× mounted on a five-axis stage (Thorlabs, K5X1), which assures fine positioning and good stability of the lens.

B. Single-beam Optical Tweezers

The single-beam OT [Figs. 1(a) and 1(b)] is realized by coupling a laser beam to the homemade microscope described in Subsection 2.A. We use a solid state laser featuring a monolithic Nd:YAG crystal NPRO (Non-Planar Ring Oscillator) (Innolight, Mephisto 500, $\lambda = 1064$ nm, 500 mW max power), which exhibits excellent optical properties for optical manipulation experiments that require high stability (e.g., for the measurement of femtonewton forces), i.e., extremely low intensity noise and very good pointing stability. The output of this kind of laser is slightly diverging and features elliptical polarization. Therefore, we linearize its polarization with a zero-order quarter-wave plate. To change the power without altering the injection current of the pump diode laser, which could affect the laser beam emission quality and, therefore, the optical trap performance, we used the combination of a zero-order half-wave plate to rotate the linear polarization and a linear polarizer to finely tune the laser power. The laser beam is directed through the objective making use of a series of mirrors. A 5× telescope is added along the optical train to create a beam with an appropriate size to overfill the objective back aperture and to generate a strong optical trap, as typically the beam waist should be comparable to the objective back aperture to obtain the optimal trapping performance [5].

To focus the laser beam inside the sample cell and generate the optical trap, we use a water-immersion objective (Olympus, UPLSAPO60XW 60×/1.2) because water-immersion objectives present less spherical aberrations—and, thus, a better trapping performance—than oil-immersion objectives. Reference [23] provides a detailed comparison of the performance of various objectives.

To achieve the highest mechanical stability, the OT setup is realized keeping the optical train as short as possible. Nevertheless, the optical train can be modified according to the specific requirements of each experiment. For example, if a movable trap is needed, a beam-steering system can be added according to the procedure described in Ref. [24].

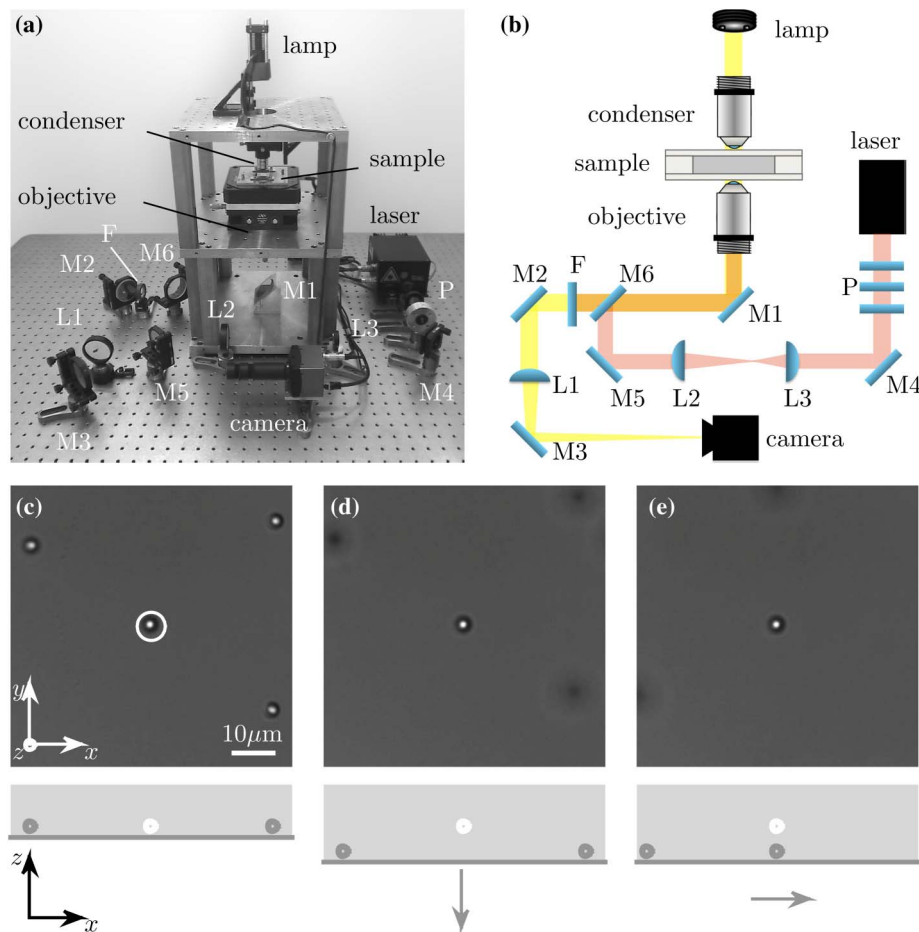


Fig. 1. Single-beam optical tweezers (OT). (a) Photograph and (b) schematic of the experimental setup. See [Media 1](#) for the complete building sequence. (c) A 1 μm diameter polystyrene bead (circled particle) is trapped by the OT and then the stage is moved (d) vertically and (e) laterally; notice how, as the stage is moved, the trapped particle remains in focus in the image plane, while the background particles get defocused. M1-M6, mirrors; P, polarization optics; L1-L3, lenses; F, short-pass filter.

C. Photonic Force Microscope

An optically trapped particle scatters the trapping light so that the light field in the forward direction is the superposition of the incoming and scattered light. This basic observation has been harnessed in the development of interferometric position detection techniques [25]. In our setup [Figs. 2(a) and 2(b)], a condenser (Olympus PlanC N, $10\times/0.25$) collects the interference pattern arising from the interference between the incoming and scattered fields, and a photodetector located on the condenser back focal plane (BFP) records the resulting signals [26]. As detector, we use a QPD based on an InGaAs junction (Hamamatsu, G6849) which, differently from silicon-based photodiodes [27], has a good high-frequency response in the infrared region of the spectrum. The output signals are acquired with an homemade analog circuit [28] and then analyzed it to obtain the three-dimensional particle position. These signals can then be used to calibrate the optical trap as explained in Appendix A.

D. Holographic Optical Tweezers

The experimental setup for HOT is slightly more complex than the one for a single-beam OT. HOT use a diffractive optical element (DOE) to split a single collimated laser beam into

several separate beams, each of which can be focused inside the sample to generate an OT [3]. These optical traps can be made dynamic and displaced in three dimensions by projecting a sequence of computer-generated holograms. Furthermore, non-Gaussian beam profiles can be straightforwardly encoded in the holographic mask.

In our implementation [Figs. 3(a) and 3(b)], the DOE is provided by a spatial light modulator (SLM) (Hamamatsu, LCOS-SLM, X10468-03). The SLM permits us to modulate the phase of the incoming beam wavefront by up to 2π . In practice, the phase mask that alters the beam phase profile is an 8 bit grayscale image generated and projected on the SLM using a computer (Appendix B). We placed the SLM after a $6\times$ telescope, which increases the beam size so that it completely fills the active area of the SLM. Then, we employ two lenses (with equal focal length, $f = 750$ mm) to image the SLM plane onto the BFP of the objective (and the necessary mirror to direct the optical train). In practice, the first lens is placed after the SLM at a distance equal to f , the second lens is placed at a distance equal to $2f$ from the first lens, and the BFP is at a distance f from the second lens. Thus, the total distance from the SLM to the BFP is $4f$ (for this reason this is often referred

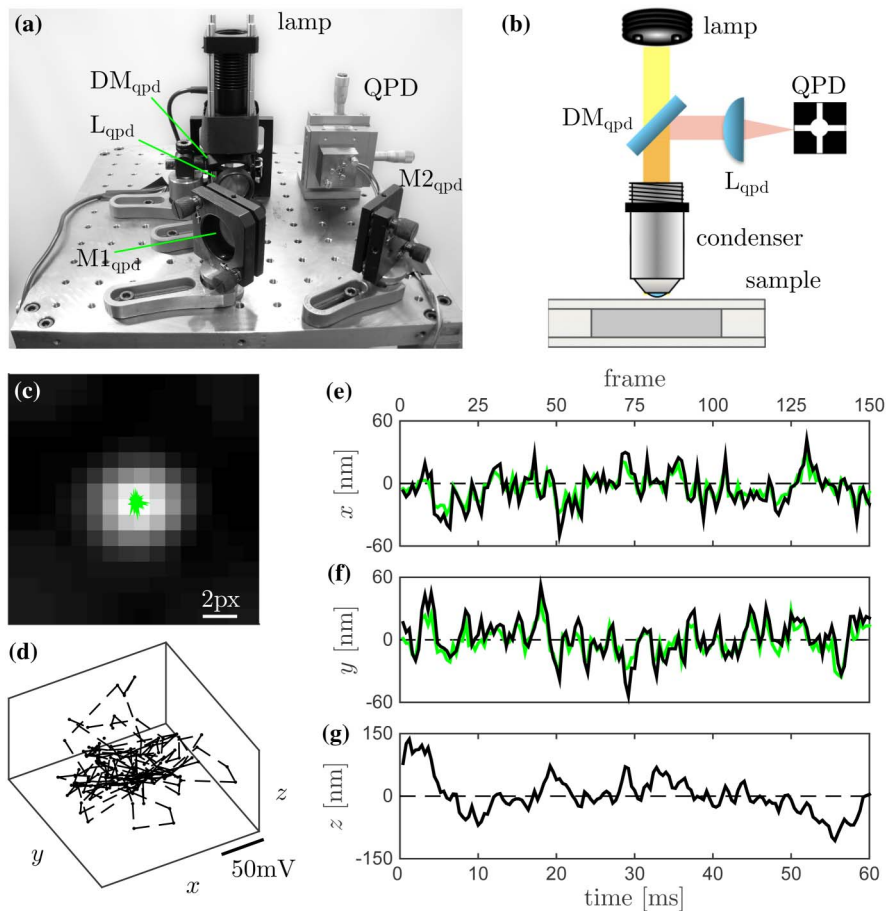


Fig. 2. PFM. (a) Photograph and (b) schematic of the detection apparatus that permits us to project the interference pattern at the BFP of the condenser lens onto a QPD. The position of the particle also can be monitored using (c) DVM or (d) the three interferometric signals received at the QPD. We obtain an excellent agreement between the positions measured with the two techniques as shown in panels (e) and (f), where the horizontal (x and y) coordinates of the particle using DVM (green/gray solid lines) overlap those obtained from interferometry (black solid lines). (g) The QPD also permits measurement of the vertical (z) coordinate of the particle. DM_{qpd}, dichroic mirror; L_{qpd}, lens; QPD, quadrant photodiode detector.

to as a $4f$ -configuration). We note that it is also possible to use lenses with different focal lengths to change the total path length and magnify/de-magnify the beam, for which we refer the readers to Ref. [24]. Therefore, the SLM (just like the BFP) is placed at a Fourier plane of the front focal plane (FFP) of the objective, and the field distribution in the FFP is (in first approximation) equal to the Fourier transform of the field distribution at the SLM [20].

We built our HOT on the same homemade microscope used for the single-beam OT. Apart from the addition of the SLM and the necessary optics in the optical train, the only modification concerns the illumination system: since the QPD is no longer needed, we prefer to increase the illumination intensity at the sample by moving the LED nearer to the condenser. High illumination intensity is needed to achieve high acquisition rates in digital video microscopy, as increasing the acquisition rate entails a decrease of the shutter time. For example, when we acquire images at 18000 frames per second (over a region of interest of 128×100 pixels), the maximum shutter time is only 55 μ s.

E. Speckle Optical Tweezers

The experimental setup for speckle OT is a simplified version of the setup for a single-beam OT. The main difference is that the condenser and illumination are substituted by a multimode optical fiber (core diameter 105 μ m, numerical aperture NA = 0.22, Thorlabs, M15L01) that delivers both the trapping laser light and the illumination light directly in close proximity to the sample upper surface. While the trapping beam must be spatially coherent and monochromatic to generate the speckle pattern through interference, the illumination should be generated with a spatially and/or temporally incoherent source to ensure a uniform illumination of the sample. The speckle light pattern is generated by directly coupling the laser beam into the multimode optical fiber [29]. The random appearance of the speckle light patterns [Fig. 5(d)] is the result of the interference of a large number of optical waves with random phases, corresponding to the various eigenmodes of the fiber. More generally, speckle patterns can be generated by different processes: scattering of a laser on a rough surface, multiple scattering in an

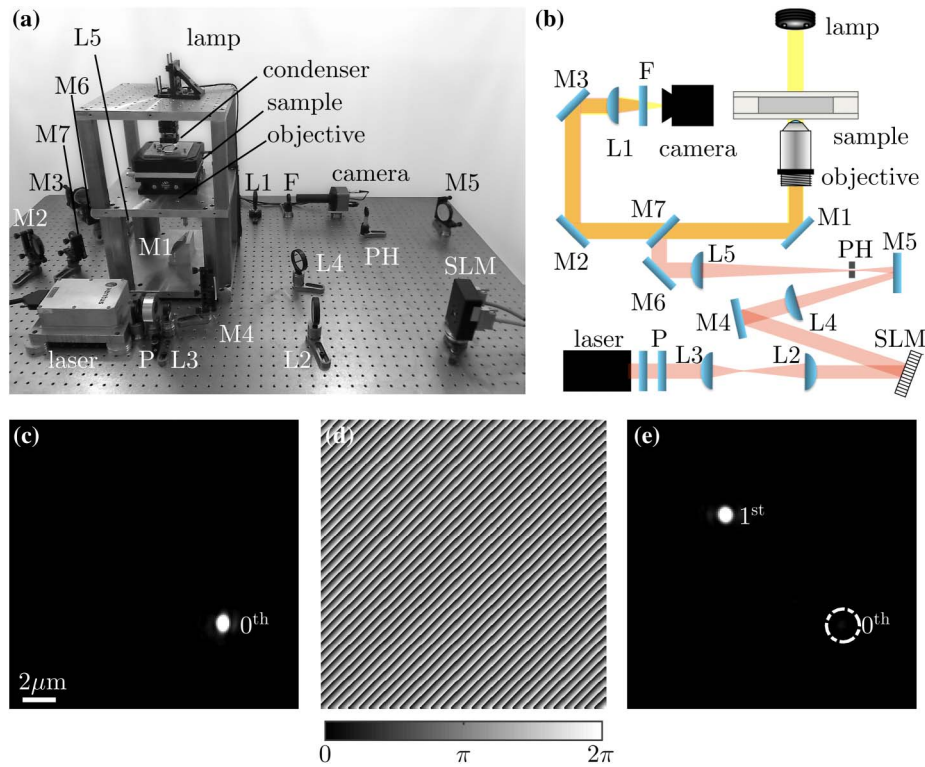


Fig. 3. Holographic optical tweezers (HOT). (a) Photograph and (b) schematic of the experimental setup. See [Media 2](#) for the complete building sequence. (c) In the absence of phase modulation on the SLM, the SLM works essentially as a mirror, reflecting almost the whole impinging light into a zero-order beam. When (d) a blazed grating phase profile is imposed on the SLM, (e) most of the beam light is deflected into a first-order beam, while there is only some residual light in the zero-order beam (circled). The amount of light in the zero-beam can be further reduced by the use of a pinhole placed at the beam waist between lenses L4 and L5. M1-M7, mirrors; L1-L5, lenses; P, polarization optics; SLM, spatial light modulator; PH, pin hole; F, short-pass filter.

optically complex medium or, as in this work, mode-mixing in a multimode fiber [29]. The choice of a multimode optical fiber provides some practical advantages over other methods, namely generation of homogeneous speckle fields over controllable areas, flexibility, and portability in the implementation of the device, as well as higher transmission efficiency.

In our setup [Figs. 5(a) and 5(b)], the fiber output is brought in close proximity of the upper wall of the sample by a micrometric three-axis mechanical stage. Optical scattering forces push the particles in the direction of light propagation toward the bottom surface of the microfluidic channel, so that they effectively confine the particles in a quasi two-dimensional space. The particles are then tracked by digital video microscopy.

3. STEP-BY-STEP PROCEDURE

A. Homemade Inverted Microscope

1. Build the first two levels of the microscope structure on a vibration insulated optical table. The first level can be either the optical table itself or, as in our case, a custom designed breadboard. The main advantage of using an independent breadboard as the first level is that the microscope is completely stand-alone, and can be easily moved if necessary. The optical components necessary to align the optical beam and to image the sample on the camera will be hosted on the optical table. Particular care should be taken in positioning the microscope: it

is preferable to mount it at the center of the optical table, since placing it on a side could decrease the damping performance of the table. The second level is composed of a breadboard held on eight columns; here is where the stages to hold and manipulate the sample will be hosted. Verify that these planes are stable and horizontal using a bubble-level. (The third level will be mounted later and will host the illumination white LED, Step 5, and the quadrant photodiode for particle tracking, Subsection 3.C.)

2. Fix a 45° gold mirror [M1, Figs. 1(a) and 1(b)] on the first level of the structure. This mirror will serve to deflect the white light from the microscope toward the camera (and to deflect the laser beam toward the back aperture of the objective, Subsection 3.B). The mirror center should be aligned with the central vertical axis of the structure (which will correspond to the axis of the objective). The height of the white light beam (and of the laser beam) above the optical table is determined by the height of the center of the 45° mirror. In our particular case, it is fixed at 7 cm.

3. Fix on the bottom side of the second level breadboard the vertical translation stage where the microscope objective will be mounted. The vertical stage has to be perfectly perpendicular to the breadboard. Use a right angle bracket to check the alignment (thanks to the length of the objective holder, angles as small as a few tenths of a degree can be easily detected). Since the vertical stage should be attached firmly to the second level, it is preferable to avoid kinematic mounts, even if of the best quality. Instead, to correct small misalignments, we suggest putting thin micrometric spacers between the mechanical parts and then

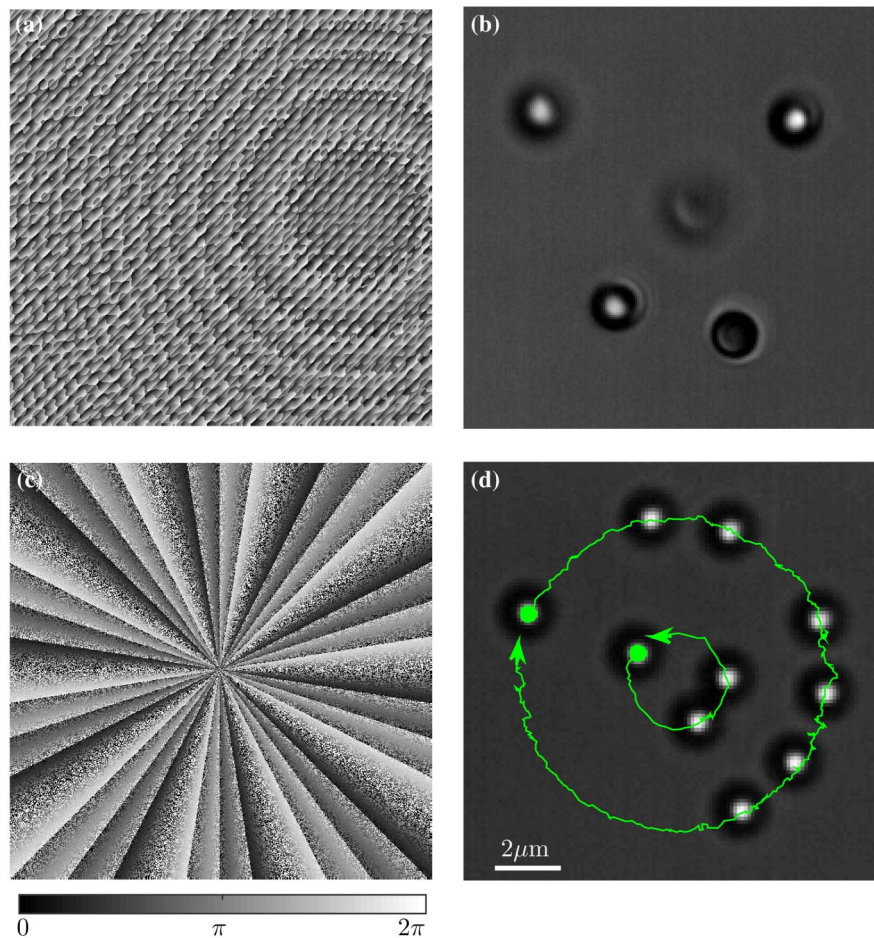


Fig. 4. HOT at work. (a) Phase mask obtained by a random superposition of diffraction gratings and Fresnel lenses, and (b) corresponding multiple optical traps placed at various lateral and vertical positions. (c) A phase mask to generate two Laguerre–Gaussian beams with a topological charge of $+10$ (inner circle) and -40 (outer circle), and (d) corresponding optically trapped particles; in this case, because of the strong scattering forces, the beads cannot be trapped in three dimensions but are pushed toward the top glass slide, where the particles rotate along two circles in opposite direction because of the transfer of orbital angular momentum (the solid lines represent the trajectory of two particles).

tightly fastening the screws. This technique will lead to a very precise vertical alignment of the objective with respect to the vertical axis of the microscope.

4. Mount the sample translation stage on the upper side of the second-level breadboard. In general, this translation stage can be either manual or automatized. Manual translation stages typically offer precision to a fraction of a micrometer over a range that can easily reach several millimeters. Automatized translation stages, which are often driven by mechanical or piezoelectric actuators, permit one to achieve subnanometer position precision over a range that typically does not exceed a few tens or hundreds of micrometers and have the advantage that they can be controlled in remote. In the realization of our setup, we combine a manual xy -translation (horizontal) stage for the rough positioning of the sample and an automatized piezoelectric xyz -translation stage for fine adjustment.

5. Proceed to build the third level of the structure, which is constituted of a breadboard held by four columns and will host the illumination and detection components.

6. Place a five-axis translation stage on the bottom side of the third level plane to mount the condenser. This stage permits one to align the condenser position along the lateral

(horizontal) directions and, therefore, to center the position of the collected illumination beam. The alignment along the vertical direction will be performed at Step 12 to optimize the illumination of the sample.

7. Mount the objective to collect the image of the sample (and later to focus a laser beam for the OT, Subsection 3.B). At the beginning, it is convenient to use a low-magnification (e.g., $10\times$ or $20\times$) objective to simplify the alignment. Later, a high-numerical aperture water- or oil-immersion objective (in our case, a water-immersion objective, Olympus, UPLSAPO60XW) will be needed for optical trapping. Using objective lenses with the same parfocal length guarantees that switching between them will not modify the relative position of the image plane with respect to the objective stage (thus, it will not require any adjustment of the objective position).

8. Mount the condenser. This lens can be either a $10\times$ or $20\times$ infinity corrected objective or an aspheric condenser lens. The choice depends on the position detection system: the aspheric condenser lens is preferable for digital video microscopy because it permits higher and more uniform illumination intensity; the objective lens is preferable if the detection is based

on a QPD because it permits one to collect the scattered light over a larger angle (Subsection 2.C). The condenser position along the vertical axis will be optimized later (Step 12).

9. Place a sample on the sample stage. Start by using a calibration glass slide (best choice to set up the microscope), or a sample of microspheres stuck to a coverslip, or a glass coverslip to which some color has been applied on one side, e.g., by using an indelible marker.

10. Add an illumination source on the third level to illuminate and observe the sample. A white LED lamp is better than a halogen lamp because it will not heat the sample. The light from the LED can be collimated using an aspheric lens. The collimated light is then focused on the sample by the condenser. It is possible to see the resulting image by the naked eye by placing a white screen along the path of the light collected by the objective. The image of a calibration glass slide can be placed easily in the center of the circular spot of the light transmitted by the objective lens. With the illumination at the maximum intensity, project this image at the longest possible distance (>5 m). Adjust the focus position to have a clear image of the sample. This will set the objective in the correct position to image the sample to infinity, i.e., the objective is to have properly imaging objects that are in the image plane. Once this procedure is done, do not move the relative position between the objective and the sample until the CCD camera is placed in its final position (next step).

11. Add a CCD camera. When using infinity corrected objectives, it is necessary to use a tube lens (L1) to produce the image of the sample at a specific position. The CCD sensor should be placed at a distance from the tube lens equal to its focal length. Place the tube lens (L1) and then the camera. If needed, it is possible to use some mirrors (M2 and M3) to place the lens and the camera in more convenient positions. The alignment of the camera and the tube lens is crucial to have a clear image and to minimize distortions; in particular: (1) the white light spot must be in the center of the lens and the lens face must be perpendicular to the light path; (2) the heights of the tube lens and of the camera must be equal; (3) the mirrors along the image path should be adjusted so that the light beam makes 90° angles (the screw holes of the optical table may be useful as guides in the alignment procedure). The focal length of the tube lens determines the magnification of the microscope with a fixed objective: the longer the focal length, the higher the magnification. Nevertheless, increasing the focal length reduces the amount of light that reaches the camera and, thus, limits the acquisition speed. Therefore, a compromise between magnification and acquisition speed should be reached. To avoid damaging the camera sensor, perform this whole procedure with low lamp power, well below the camera saturation threshold.

12. Optimize the illumination brightness by adjusting the condenser axial position. More complex illumination configurations, e.g., Köhler illumination, are also possible if the coherence of the illumination as well its homogeneity need to be controlled.

13. It is useful to calibrate the CCD measuring the pixel-to-meter conversion factor. This can be done using a calibration glass slide or moving (by a known controllable displacement) a particle stuck to the bottom coverslip by means of a calibrated piezoelectric stage.

B. Single-Beam OT and Alignment of the Laser Beam

14. Fix the laser firmly on the optical table. In our case, we use a solid state laser with wavelength 1064 nm and maximum power 500 mW, even though typically we will only need 1 mW for trapping a single particle. Place the laser so that the beam height coincides to the height of the center of the 45° mirror [M1, Figs. 1(a) and 1(b)] on the first level of the microscope (Step 2). This is the height of the axis of the optical train (7 cm in our case). Importantly, keep in mind that correct safety procedures when working with lasers are of paramount importance; thus, we urge all users to undergo suitable safety training before starting to work with lasers and to observe all local safety regulations, especially in the use of appropriate eye safety equipment.

15. Since we use a diode-pumped solid state (DPSS) laser whose output beam has elliptical polarization, we place a quarter-wave plate to linearize the polarization just after the laser [the first element in P, Figs. 1(a) and 1(b)]. To maintain the power stability of the laser beam, we prefer not to alter the laser settings (and, in particular, its injection current) to change the laser output power. Therefore, we opt for an alternative way of controlling the laser power: we place along the beam path a half-wave plate and a beam polarizer [second and third elements in P, Figs. 1(a) and 1(b)], which permit us to tune the beam power by rotating the half-wave plate. The next step is to mount the optical train to direct the laser beam through the objective. To have a stable optical trap, keep the laser beam path as short as possible and consider using black plastic pipes to enclose the laser beam and to build a thermal and acoustic isolation enclosure for the whole experimental setup. For safety, and to avoid damaging the camera, perform this procedure with low laser power.

16. Use dielectric laser line mirrors to deflect the laser beam at a right angle [M4 and M5, Figs. 1(a) and 1(b)]. Check that the deflected beam travels at the same height of the incident beam. The last mirror of the optical train should be a dichroic mirror [M6, Figs. 1(a) and 1(b)], which permits one to split the light of the laser beam, which is routed toward the objective, and the light of the lamp, which is routed toward the camera. We used a short-pass dichroic mirror that reflects (reflection coefficient $>98\%$) in the region from 750 to 1100 nm and transmits shorter wavelengths.

17. Remove the objective and the condenser from their respective holders and, using only the last two mirrors [M5 and M6, Figs. 1(a) and 1(b)], align the laser beam so that it goes straight through the objective holder. This is a crucial point as the laser beam must be perfectly aligned along the vertical axis of the objective holder. To achieve this, you can use two alignment tools like pinholes (Thorlabs, VRC4D1) or fluorescing disks (Thorlabs, VRC2RMS). It is not critical to perfectly center the condenser at this point, because it will be optimized later while aligning the detection components.

18. Add a telescope to create a beam with an appropriate size to overfill the objective back aperture and to generate a strong optical trap. The beam waist can be measured by fitting a Gaussian profile to an image of the beam acquired by a digital camera (some neutral density filters might be necessary to prevent the camera from saturating). The typical beam waist (radius) of a DPSS laser is about 1 to 1.5 mm; therefore a telescope with magnification of about $5\times$ guarantees the proper

overflowing condition. The magnification can be easily obtained with lenses with focal lengths of 5 and 25 cm. To realize the telescope, first place the lens with the longer focal length [L2, Figs. 1(a) and 1(b)]. It should be positioned while controlling the position of the laser beam on the target placed on the objective lens mount: the position of the spot center must remain the same with and without the lens. Then place the lens with shorter focal length [L3, Figs. 1(a)–1(b)] between the first lens and mirror M4, verifying, at the same time, that the position of the laser beam on the target does not change, as done for the first lens, and the collimation of the laser beam. The latter can be done using a shear plate interferometer or checking that the size of the beam waist is constant on a long travel distance (>5 m).

19. Double-check the alignment of the laser beam along the vertical axis of the microscope using the alignment tools.

20. Add a short-pass filter (Thorlabs, FGS900) in the light path leading to the camera to remove any laser light that might saturate the camera [F, Figs. 1(a) and 1(b)]. It is preferable to mount the filter on a repositionable mount (Thorlabs, FM90) to easily block or unblock the laser light.

21. Mount again the objective, the condenser, and the sample cell. Allow the laser light to reach the CCD camera by removing the filter F [Figs. 1(a) and 1(b)].

22. Place a droplet of the immersion medium on the objective (water in our setup) and gently move the objective toward the sample cell until the focus is at the glass-air interface. This is best done by using a sample cell without any liquid to increase the amount of light reflected back to the camera, but it can also be done with a sample filled with a solution, even though the intensity of the back-scattered light will be significantly (about two orders of magnitude) lower. In practice, one has to slowly approach the objective to the sample, until the back-scattered pattern size is minimized. The resulting pattern (a series of concentric Airy rings, possibly with a cross-shaped dark area) should be as symmetric as possible. Furthermore, the pattern should remain symmetric with the same center as the relative distance between the focus and interface is changed.

23. Build a sample cell with a microscope slide and a coverslip (thickness #1, ca. 150 μm): (1) clean the slide and coverslip; (2) place two strips of Parafilm above the slide to work as spacers (Parafilm is a polymer resistant to many solvents and acids; it melts at about 80°C and it works like a glue when cooled); (3) place the coverslip above the Parafilm stripes; (4) heat with a hot air gun, or by placing the cell on a hot plate; (5) fill the cell with the sample solution and seal it with silicon grease, epoxy resin, or UV glue. The thickness of the cell is around 120 μm , but it is possible to have thicker cells overlapping two or three layers of Parafilm or thinner by stretching the Parafilm before placing the coverslip.

24. Place the sample cell filled with a mixture of water and microspheres on the stage (with the coverslip side below toward the objective) and adjust the position of the objective and sample holder such that the plane just above the glass-solution interface is on focus. The image recorded by the camera should show the sedimented particles, as in Fig. 1(c).

25. Move the sample stage a few micrometers down (so that the laser focus effectively moves a few micrometers above the interface) and approach some of the particles. If a particle is free to move and not stuck to the glass, you will see the particle jump into the laser beam and get optically trapped. To get the image in focus, it might be again necessary to slightly adjust the

lens in front of the camera. It could happen that a bead is attracted by the laser beam, but instead of being trapped it will be pushed in the direction of the laser beam propagation. This means that the laser beam is not well aligned or is not overflowing the aperture sufficiently. Go back to steps 17–19 to check the alignment and overflowing. To double-check that the particle is actually optically trapped, try and move it around. First, move the sample stage farther down so that the optically trapped particle gets moved vertically above the glass-solution interface, as in Fig. 1(d): non-optically trapped particles on the sample cell bottom get out of focus, while the optically trapped particle remains in focus. Then, move the sample stage horizontally, as in Fig. 1(e): the image of the optically trapped particle remains on the same spot within the image captured by the camera, while the background particles sedimented on the coverslip appear to be displaced.

C. Photonic Force Microscope and Alignment of the Quadrant Photodetector

25. To use a QPD, the image plane of the condenser lens should be the same as the one of the objective, i.e., they should form a telescope. To achieve this, adjust the height of the condenser until the laser beam emerging from the condenser is well collimated.

26. Position a dichroic mirror on the third level below the illumination LED [DM_{qpd}, Figs. 2(a) and 2(b)].

27. Immediately after the dichroic mirror, place a 20 cm or 10 cm focal length lens [L_{qpd}, Figs. 2(a) and 2(b)] to project the image of the condenser BFP on the QPD.

28. Mount the QPD on a three-axis translation stage for precise alignment. Depending on the space available on the third level, use mirrors to deflect the laser light toward the QPD [two mirrors in our case, Figs. 2(a) and 2(b)]. The QPD sensitive area should be as perpendicular to the laser beam as possible.

29. Proceed to connect the output of the QPD to the computer or to a digital oscilloscope to observe the output signals.

30. Proceed to align the QPD. This involves first of all roughly centering the light collected by the condenser on the QPD and then proceeding to maximize the QPD sum signal while zeroing the differential QPD signals. This should be done without any trapped particle.

31. Pick a particle and observe the three signals. You should observe the thermal motion of the particles. In the xy -plane, the QPD output signals should have a small offset [Fig. 2(d)]; typically, the smaller the offset the better the alignment. The output signal in the axial z direction is the sum of the four quadrant and, therefore, has a large offset that depends on the laser power.

32. Acquire the position signals of a particle as a function of time [Figs. 2(e)–2(g)] and analyze them according to the procedures presented in Appendix A to calibrate the optical trap. It is also possible to compare the trajectories obtained with the QPD to the ones obtained through digital video microscopy [Figs. 2(e) and 2(f)]. The agreement between the two results is an indication of the quality and stability of the setup.

D. Holographic Optical Tweezers

33. To realize a HOT, we need to change the optical beam input train. In particular, we need to place the SLM in a plane conjugated to the back aperture of the objective using two lenses arranged in a $4f$ -configuration. To fit this optical train

within the optical table, it is necessary to carefully plan the laser beam path. In our case [Figs. 3(a) and 3(b)], the laser beam is first reflected by the SLM, then by a series of mirrors (M4, M5, M6, M7, and M1), and finally it reaches the objective. The total distance between the SLM and the objective back aperture is 3000 mm, so that we can realize a $4f$ -configuration using two lenses with $f = 750$ mm (L4 and L5, Step 41).

34. Change the illumination system to increase the light intensity. In particular, place the LED light close to the sample and use a couple of aspheric lenses to collimate the LED light.

35. Repeat Steps 14-15 to place the laser. Note that, to create multiple traps, more laser power is required; therefore we used a different DPSS laser with a maximum power of 3 W (Laser Quantum, Ventus 1064).

36. Expand the optical beam with a two-lens telescope [L2 and L3, Figs. 3(a) and 3(b)] as explained in Step 18, so that it fills the SLM active area. The laser spot should be perfectly centered on the SLM active area, lest the efficiency and the wavefront shape will be affected.

37. Rotate the SLM by a small (less than 10°) angle with respect to the incidence direction of the laser beam. Larger angles would result in a significant decrease of the SLM efficiency.

38. Display a blazed diffraction grating on the SLM. The SLM will split the incoming beam into several diffraction orders. Usually the strongest are the zero-order beam, i.e., the beam that is simply reflected by the active area, and the first-order diffracted beam, which is the one that will be used to generate the optical traps. The intensity of the first-order beam varies according to the efficiency of the SLM.

39. Use large optics (at least 2 inch diameter) in the optical train to collect as much as possible of the light modulated into the first-order beam.

40. Deflect the laser beam towards the objective back aperture. Align the laser beam so that it goes straight through the objective holder using mirrors M6 and M7 [Figs. 3(a) and 3(b)], as done in Step 17.

41. Add lenses L4 and L5 [$f = 750$ mm, Figs. 3(a) and 3(b)] to realize the $4f$ -configuration. Lens L5 must be placed at a distance from the back aperture of the objective lens exactly equal to f . While aligning L5, check that the position of the laser beam on the alignment tools does not change. Then, place L4 at a distance of 1500 mm from the first lens and of 750 mm from the SLM surface. Again, check that the position of the laser spot does not change after both lenses have been positioned.

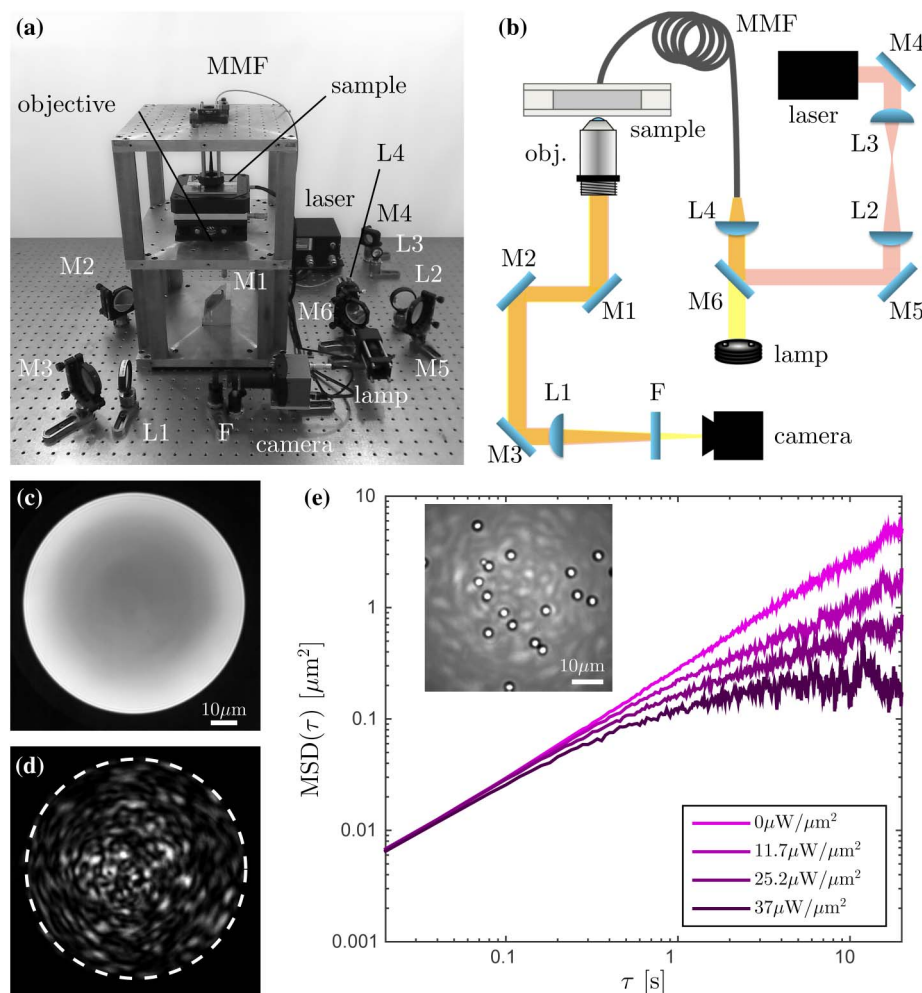


Fig. 5. Speckle optical tweezers. (a) Photograph and (b) schematic of the experimental setup. See Media 3 for the complete building sequence. (c) Illumination light and (d) speckle light field output from the optical fiber. (e) MSD of particles trapped at various laser powers; a transition from trapping at high laser powers to subdiffusion at low laser powers can be observed. M1-M6, mirrors; L1-L4, lenses; MMF, multi mode fiber; F, short-pass filter.

42. The middle point between L4 and L5 is conjugated to the objective FFP and, thus, it is possible to place a pinhole at this position to select only the first-order beam and cut out the zero-order beam and the higher diffraction orders.

43. Repeat Steps 20–24 to prepare the sample cell and observe the particles.

44. Observe how the optical trap can be moved in the lateral direction by using gratings with different orientations and pitches, and in the axial direction by using a Fresnel lens (Appendix B). Note also how, with a combination of gratings and Fresnel lenses, it is possible to displace a trap in three dimensions. Generate multiple optical traps and trap various particles, as in Figs. 4(a) and 4(b). Generate Laguerre–Gaussian beams and see the transfer of orbital angular momentum (OAM), as in Figs. 4(c) and 4(d).

E. Speckle Optical Tweezers

45. Repeat Steps 14–15 to place the laser.

46. Mount a telescope to increase the size of the laser beam waist [L2 and L3, Figs. 5(a) and 5(b)].

47. It is necessary to overlap the illumination light on the laser beam to have them together at the fiber output. To do this, use mirrors [M5 and M6, Figs. 5(a) and 5(b)] to steer the laser beam toward the entrance port of the fiber. One mirror (in our case, M6) must be a dichroic mirror to allow the transmission of the white light of the illumination LED.

48. Align the laser beam into the fiber input using a lens [L4, Figs. 5(a) and 5(b)]. To maximize the coupling and to generate a fully developed speckle pattern, the focused beam should match the numerical aperture of the fiber. Since in our case the fiber has a numerical aperture of 0.20, we used a 20 mm focal length lens with a diameter of 12.7 mm and the beam waist was set to 4 mm. We suggest building a cage system to firmly hold the focusing lens and the fiber end. The fiber end, or equivalently the focusing lens, can be mounted on a translation stage (Thorlabs, SMZ1) to easily align the beam. Using a power meter, maximize the power at the output end of the fiber.

49. Place the illumination LED behind the dichroic mirror [M6, Figs. 5(a) and 5(b)] and observe the white light at the end of the fiber with the laser light off. Optimize the transmission of

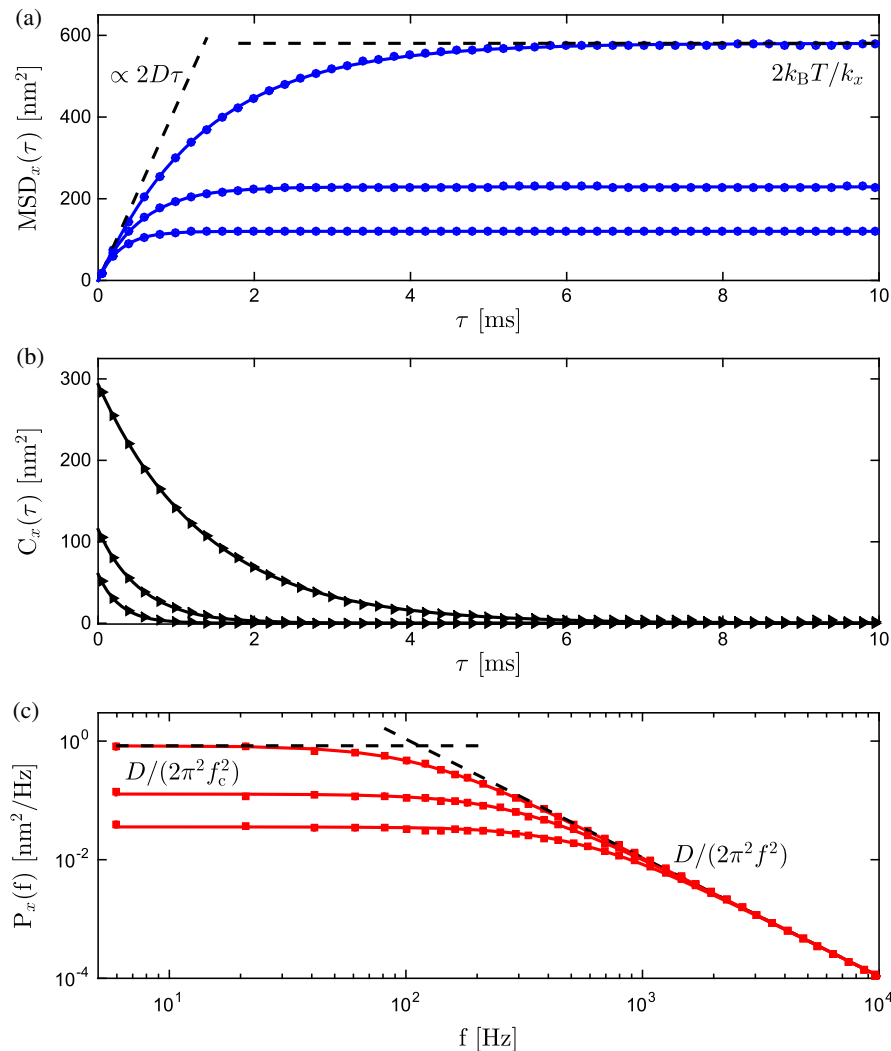


Fig. 6. Calibration of an optical trap using the (a) MSD, (b) ACF, and (c) PSD methods. The symbols correspond to the MSD, ACF, and PSD experimentally obtained from a 10 s trajectory of a 1 μm diameter polystyrene particle trapped by a single-beam optical tweezers. The solid lines correspond to the best fit to the theoretical MSD [Eq. (A1)], ACF [Eq. (A2)], and PSD [Eq. (A3)].

the illumination LED using the CCD camera. This can be easily done by moving the whole LED mount behind the dichroic mirror.

50. Using an IR card, you should now be able to observe a diverging speckle pattern from the output of the fiber, as shown in Fig. 5(d).

51. Build a cage system to host the output fiber adapter. It should be mounted vertically upside down just above the sample cell. Mount it on a precision three-axis translation stage to align the fiber output face along the optical axis of the observation objective [Figs. 5(a) and 5(b)] and to control the size of the average speckle grain by precisely translating the fiber vertically.

52. Place this system on the third level of the microscope [Fig. 5(a)].

53. To be able to observe the whole area corresponding to the fiber core, use an imaging objective with a relatively low magnification (we used a 40 \times objective).

54. The sample cell should be realized in a slightly different way to the one followed in Step 23: since the output fiber end must be very close to the sample (<300 μm), the 1 mm glass slide needs to be replaced by a much thinner coverslip. Place the sample cell on the piezoelectric translation stage and move the objective until you observe the particles on the bottom of the sample cell.

55. Change the height of the output fiber end until you can see its clear image [Fig. 5(c)]. Then move it back by a very small amount. Using the CCD camera, you should observe a blurred bright disk.

56. Repeat Step 20 to filter out the laser light. Note that, in this case, the laser light collected by the objective is sent toward the CCD. It could be necessary to use multiple filters to eliminate all the laser light.

57. Set the laser output power at a moderate level. Remove the short-pass filter from in front of the CCD camera. The speckle pattern will be now be visible inside the bright disk imaged by the camera, as shown in Fig. 5(d).

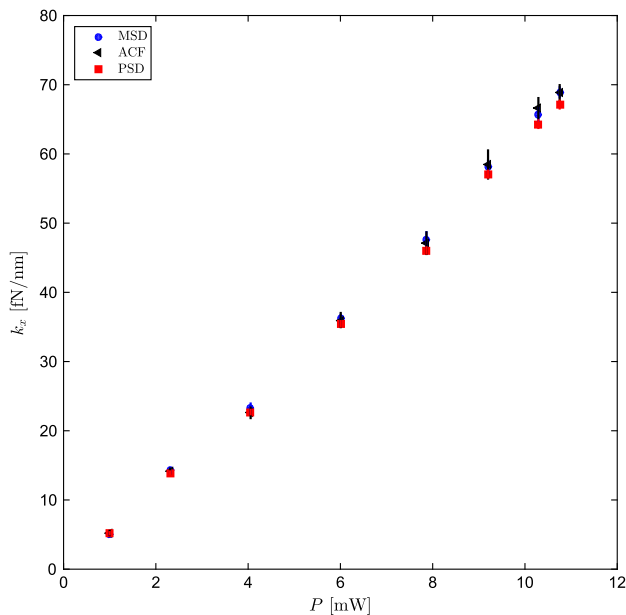


Fig. 7. Linearity of the trap stiffness k_x as a function of the laser power at the trap P . The calibrations obtained with MSD, ACF, and PSD show an excellent agreement.

58. At this point you should observe that some of the free diffusing particles on the bottom of the sample cell [inset in Fig. 5(e)] are attracted toward the high intensity maxima of the speckle pattern. Increasing the laser power will increase the number of hot spots with enough power to actually trap particles.

59. Measure the trajectories of the particles using digital video microscopy.

60. Observe how, by increasing the laser power, the particles are more and more confined in the speckle grains. Use, e.g., the mean square displacement analysis (see Appendix B) to quantify how much the particles are trapped, as shown in Fig. 5(e).

61. Moving the piezoelectric stage, observe how the speckle pattern captures particles as they approach the high intensity hot spots.

62. Prepare a binary or ternary mixture of particles of different size and/or different material. Observe how the speckle optical tweezers is able to selectively capture different particles as the piezoelectric stage is moved.

4. CONCLUSIONS

In this article we presented a detailed step-by-step guide for the construction of advanced optical manipulation techniques based on the construction of a homemade inverted optical microscope. Readers with some basic experience in experimental optics should be able to follow the explained procedure and build a single-beam OT, a HOT and, finally, a speckle OT. Readers will also be able to easily upgrade the designs we propose and realize in this article to fit their experimental needs.

APPENDIX A: CALIBRATION OF THE SINGLE-BEAM OPTICAL TRAP

Once the trajectory of a Brownian particle has been measured either by digital video microscopy or by interferometry, it is possible to use it to study quantitatively the optical potential. In calibrating an optical trap, the main objective is to determine the trap stiffness k_x along each direction, in this case, e.g., along x (one of the horizontal directions). A common secondary objective is to determine the conversion factor β_x from measurement units (e.g., pixels or volts) to physical units of length (meters). The knowledge of these two parameters allows to measure an external force applied to the trapped particle by measuring only the bead displacement from its equilibrium position. Several techniques can be employed for this task [4,30]. The easiest and fastest is based on the equipartition theorem which, for a harmonic potential, predicts an energy equal to $\frac{1}{2}k_B T$ for each degree of freedom. The trajectory along one coordinate has an energy of $\frac{1}{2}k_x \langle x^2 \rangle$; thus, from a direct comparison, it is possible to determine the trap stiffness. A more detailed determination uses the Boltzmann statistics, where the optical potential well is obtained from the probability distribution associated to a particle's trajectory. However, the most common and reliable methods employ the mean square displacement (MSD), the autocorrelation function (ACF), and the power spectral distribution (PSD). We will briefly discuss them in this appendix.

The MSD provides a measure of the displacement of a particle from its initial position. The MSD as a function of the

lag time τ is given by $\text{MSD}_x(\tau) = \langle (x(t+\tau) - x(t))^2 \rangle$. For free particles diffusing in a viscous medium, the MSD grows linearly with time, so that $\text{MSD}_x(\tau) = 2D\tau$, where D is the diffusion coefficient. A particle held in an optical tweezers is continuously pulled back by the restoring force; therefore, the MSD cannot grow indefinitely but reaches a plateau, so that

$$\text{MSD}_x(\tau) = \frac{2k_B T}{k} \left[1 - e^{-\frac{k\tau}{\gamma}} \right], \quad (\text{A1})$$

from which we can see that the particle diffuses inside the optical potential well at short times, i.e., until it starts feeling the trap restoring force. An OT can be calibrated by fitting the experimentally obtained MSD to Eq. (A1), as shown in Fig. 6(a). The plateau at long times depends on k , while the linear growth at very short times depends, at a first approximation [31,32], on D .

The ACF provides a measure of how fast a particle falls into the optical trap. The ACF on an optically trapped particle is

$$C_x(\tau) = \frac{k_B T}{k_x} e^{-\frac{k_x \tau}{\gamma}}, \quad (\text{A2})$$

where γ is the friction coefficient. The trap stiffness can be straightforwardly determined from a fit procedure of the ACF calculated from the experimental trajectories as shown in Fig. 6(b). As the trap stiffness increases the exponential decay of the ACF becomes faster. We note also that cross-correlation functions, i.e., correlations between the motion of a particle in two different directions, can be used to calibrate an optical trap as they can provide information about the presence of cross-talk [33] and the presence of torques [34].

The PSD is the Fourier transform of the ACF and, for an optically trapped particle, it is

$$P_x(f) = \frac{k_B T}{2\pi^2 \gamma} \frac{1}{f^2 + f_{c,x}^2}, \quad (\text{A3})$$

where $f_c = 2\pi\gamma k_x$ is the corner frequency, which represents the frequency beyond that the motion is purely diffusive. The calibration of the optical trap can again be achieved by fitting the experimentally obtained PSD to the theoretical one, as shown in Fig. 6(c). As the optical trap stiffness increases, f_c increases. One advantage of the PSD analysis is that it permits one to clearly identify various sources of noise, e.g., electrical interferences and mechanical drifts [33]. In particular, the flat plateau at low frequencies of the PSD in Fig. 6(c) is a clear sign of the high mechanical stability of the system.

The data reported in Fig. 7 show the linear dependence of the trap stiffness determined using the methods described above on the trapping laser power. The data are obtained using the single-beam OT with interferometric detection with the QPD. The agreement between the different techniques is excellent and demonstrates the quality of our setup.

APPENDIX B: GENERATION OF PHASE MASKS FOR HOT

The working principle of HOT is based on a DOE whose task is to shape the profile of an incoming optical beam. The easiest way to realize such an element is to use an SLM. An SLM is a transmissive or reflective device that is used to spatially modulate the amplitude and phase of an optical wavefront in two

dimensions. SLMs can be programmed to produce light beams with various optical wavefronts and, therefore, can be used in place of conventional optical elements, such as lenses, mirrors, and gratings. Furthermore, they can be used to create optical vortex and higher order Bessel beams, which feature an azimuthal phase variation. To accomplish these tasks, it is in principle necessary to display on the SLM a hologram obtained taking the inverse Fourier transform of the desired intensity distribution in the front focal plane. However, the simultaneous modulation of phase and amplitude are technically difficult and, furthermore, the amplitude modulation would remove power from the beam, reducing the modulation efficiency. Therefore, in practice, it is more convenient to employ a phase-only modulation hologram.

The SLM is typically placed in a plane conjugated to the objective BFP so that the light distribution generated on the FFP is essentially the Fourier transform of the phase mask imposed on the SLM (assuming an incoming beam with a flat wavefront). Therefore, imposing a phase mask corresponding to a blazed diffraction grating will change the angle of the beam emerging from the SLM and will displace the focus at the image plane; the angle of the grating determines the direction of the displacement, while the number of fringes determines the distance. To change the position of the focus along the axial direction, it is necessary to change the curvature of the wavefront, and this is accomplished applying the pattern of a Fresnel lens to the SLM. A combination of a diffraction grating and a Fresnel lens produces a shift of the trap in three dimensions; more precisely, the corresponding phase mask is given by

$$\phi^S(x, y) = \underbrace{\frac{2\pi}{\lambda_0 f} [xx_{to,1} + yy_{to,1}]}_{\text{Diffraction grating}} + \underbrace{\frac{\pi z_{to,1}}{\lambda_0 f^2} [x^2 + y^2]}_{\text{Fresnel lens}}, \quad (\text{B1})$$

where x and y are the Cartesian coordinates on the SLM, λ_0 is the vacuum wavelength, f is the focal distance of the lens, and $[x_{to,1}, y_{to,1}, z_{to,1}]$ is the desired position of the trap in the image plane. Since the generation of this holographic mask is very fast, this is a very effective way to generate a single optical trap that can be moved in three dimensions in real time.

To generate multiple (N) traps, one of the fastest, but also least efficient, algorithms is the random mask encoding algorithm [35]. For every pixel of the SLM, a phase shift is determined as if the hologram was made to generate only one of the N traps. This technique is very fast and permits one to achieve a good uniformity among traps. Nevertheless, the overall efficiency can be very low when N is large, because on average the number of pixels that interfere constructively to generate each trap decreases as $1/N$.

A straightforward algorithm, which achieves a better efficiency than the random mask encoding at a just slightly higher computational cost, is the superposition of gratings and lenses algorithm [36]. In this algorithm, the phase of each pixel is chosen equal to the argument of the complex sum of single-trap holograms. This algorithm has typically a good efficiency at the cost of poor uniformity. In fact, a typical problem that arises using this algorithm is that, when highly symmetrical trap geometries are sought, such as an array of traps, a consistent part of the energy is diverted to unwanted ghost traps.

To overcome these difficulties, it is necessary to use more sophisticated algorithms to compute the phase patterns. The most used are the Gerchberg–Saxton [37] and the adaptive-additive algorithms [38] that are able to realize a hologram that will lead to a very close approximation to any desired intensity distribution. The Gerchberg–Saxton algorithm is an iterative algorithm that permits one to find a phase distribution that turns a given input intensity distribution arriving at the hologram plane into a desired intensity distribution in the trapping plane by propagating the complex amplitude back and forth between these two planes and replacing at each step the intensity on the trapping plane with the target intensity and that on the SLM plane with the laser's actual intensity profile. It typically converges after a few tens of iterations, it is ideally suited to deal with continuous intensity distributions. The adaptive-additive algorithm starts with an arbitrary guess of the phase profile and an initial input wavefront; the Fourier transform of this wavefront is the starting estimate for the output electric field. The resulting error in the focal plane is reduced by mixing a proportion of the desired amplitude into the field in the focal plane; inverse-Fourier transforming the resulting focal field yields the corresponding field in the input plane. The amplitude in the input plane is replaced with the actual amplitude of the laser profile. Finally, the algorithm is iterated. The main advantage of this algorithm in comparison to the Gerchberg–Saxton algorithm is that it permits one to achieve a better uniformity over all the traps in the array.

An SLM can also be used to create non-Gaussian beams such as Hermite–Gaussian beams, Laguerre–Gaussian beams, non-diffracting beams, and continuous intensity distributions. In particular, the phase mask to generate Laguerre–Gaussian beams is given by

$$\phi^{\text{LG}}(\varphi, \rho) = i l \varphi, \quad (\text{B2})$$

where φ and ρ are the polar coordinates of a system of reference centered on the middle of the SLM and l is the order to the Laguerre–Gaussian beam.

APPENDIX C: MATERIALS

The advanced optical tweezers systems presented in this work are constructed using standard commercial products and custom-made pieces. In the following, we present a detailed list of all the parts used for each setup. All the optics holders were mounted on a vibration insulated optical table using custom-made pillar posts. The technical drawings of all the custom-made parts are freely available online [22].

To adjust the height, we used spacers of different thickness from 200 μm up to 5 mm. The optics mounts were fixed on the optical table using either custom-made clamps or clamping forks from Thorlabs (CF125 and CF175).

The particles used for experiments were silica, melamine, and polystyrene beads from Microparticles GmbH. The sample cells were made using a microscope slide and a coverslip or two coverslips from Knittel Glass.

Single-Beam Optical Tweezers

1. Precision kinematic mirror mounts Thorlabs KS1/M—Fig. 1: M4; Fig. 2: M1_{qpd}, M2_{qpd}.

2. Precision kinematic mirror mounts Thorlabs KS2/M—Fig. 1: M2, M3, M5, M6.

3. Fixed lens mounts Thorlabs LMR1/M—Fig. 1: L3; Fig. 2: L_{qpd}.

4. Fixed lens mounts Thorlabs LMR2/M—Fig. 1: L1, L2.

5. Flip Mount Adapter Thorlabs FM90/M—Fig. 1: F.

6. Short-pass filter Thorlabs FES0700—Fig. 1: F.

7. N-BK7 AR coating: 650–1050 nm bi-convex lenses 1" diameter. $f = 25$ mm Thorlabs LB1761-B—Fig. 1: L3.

8. N-BK7 AR coating: 650–1050 nm bi-convex lenses $f = 25$ mm diameter. $f = 200$ mm Thorlabs LB1945-B—Fig. 2: L_{qpd}.

9. N-BK7 AR coating: 650–1050 nm bi-convex lenses 2" diameter. $f = 250$ mm Thorlabs LB1889-B—Fig. 1: L2.

10. N-BK7 uncoated bi-convex lenses 2" diameter. $f = 500$ mm Thorlabs LB1909—Fig. 1: L1.

11. N-BK7 uncoated bi-convex lenses 2" diameter. $f = 400$ mm Thorlabs LB1852—Fig. 1: L1 (optional instead of item n. 10).

12. Unmounted positive achromatic doublets AR coated: 400–700 nm 2" diameter. $f = 300$ mm Thorlabs AC508-300-A—Fig. 1: L1 (optional instead of item no. 10).

13. Unmounted positive achromatic doublets AR coated: 400–700 nm 2" diameter. $f = 200$ mm Thorlabs AC508-200-A—Fig. 1: L1 (optional instead of item no. 10).

14. Thorlabs 30 mm Cage System—Fig. 1: lamp.

15. CCD Mikrottron EoSens Mini1—Fig. 1: camera.

16. Diode pumped solid state laser Innolight—Mephisto 500 (500 mW maximum output) at 1064 nm—Fig. 1: laser.

17. Objective lens Olympus UPLSAPO60XW 60 \times /1.2 (Trapping Objective)—Fig. 1: objective.

18. Objective lens Olympus PlanC N 10 \times /0.25 (Condenser Lens)—Fig. 1: condenser.

19. White LED Thorlabs MCWHL5 with power supply LEDD1B—Fig. 1: lamp.

20. Zero-Order half wave plate at 1064 nm Thorlabs WPH10M-1064—Fig. 1: P.

21. Zero-Order quarter-wave plate at 1064 nm Thorlabs WPQ10M-1064—Fig. 1: P.

22. Glan-Thompson polarizer Thorlabs GTH10M—Fig. 1: P.

23. Broadband dielectric mirrors HR at 750–1100, 1" diameter. Thorlabs BB1-E03—Fig. 1: M4; Fig. 2: M1_{qpd}, M2_{qpd}.

24. Broadband dielectric mirrors HR at 750–1100, 2" diameter. Thorlabs BB2-E03—Fig. 1: M5.

25. Back side polished broadband dielectric mirrors HR at 750–1100, 2" diameter. Standa—Fig. 1: M6.

26. Gold protected mirrors 2" diameter. Thorlabs PF20-03-M01—Fig. 1: M1.

27. High-performance large aperture platform two-axis linear stage, M6 Newport M-406 with HR13 actuators—Fig. 1: sample stage.

28. Piezo stage Physik Instrumente PI-517.3CL—Fig. 1: sample stage.

Holographic Optical Tweezers

1. Precision kinematic mirror mounts Thorlabs KS2/M—Fig. 3: M2, M3, M4, M5, M6, M7.

2. Fixed lens mounts Thorlabs LMR1/M—Fig. 3: L3.

3. Fixed lens mounts Thorlabs LMR2/M—Fig. 3: L1, L2, L4, L5.

4. Flip mount adapter Thorlabs FM90/M—Fig. 3: F.
5. Short-pass filter Thorlabs FES0700—Fig. 1: F.
6. N-BK7 AR coating: 650–1050 nm bi-convex lenses 1" diameter. $f = 50$ mm Thorlabs LB1471-B—Fig. 3: L3.
7. N-BK7 AR coating: 650–1050 nm bi-convex lenses 2" diameter. $f = 300$ mm Thorlabs LB1917-B—Fig. 1: L2.
8. N-BK7 AR coating: 650–1050 nm bi-convex lenses 2" diameter. $f = 750$ mm Thorlabs LB1247-B—Fig. 3: L4, L5.
9. N-BK7 uncoated bi-convex lenses 2" diameter. $f = 500$ mm Thorlabs LB1909—Fig. 1: L1.
10. N-BK7 uncoated bi-convex lenses 2" diameter. $f = 400$ mm Thorlabs LB1852—Fig. 1: L1 (optional instead of item no. 9).
11. Unmounted positive achromatic doublets AR coated: 400–700 nm 2" diameter. $f = 300$ mm Thorlabs AC508-300-A—Fig. 1: L1 (optional instead of item no. 9).
12. Unmounted positive achromatic doublets AR coated: 400–700 nm 2" diameter. $f = 200$ mm Thorlabs AC508-200-A—Fig. 1: L1 (optional instead of item no. 9).
13. Aspheric condenser lens, uncoated Thorlabs ACL25416U—Fig. 3: condenser.
14. Iris diaphragms Thorlabs D25S—Fig. 3: PH.
15. Thorlabs 30 mm Cage System—Fig. 3: lamp.
16. CCD Mikrottron EoSens Mini1—Fig. 3: camera.
17. Diode pumped solid state laser Laser Quantum Ventus 1064 (3 W maximum output) at 1064 nm—Fig. 3: laser.
18. Objective lens Olympus UPLSAPO60XW 60 \times /1.2 (Trapping Objective)—Fig. 3: objective.
19. White LED Thorlabs MCWHL5 with power supply LEDD1B—Fig. 3: lamp.
20. Zero-Order half wave plate at 1064 nm Thorlabs WPH10M-1064—Fig. 3: P.
21. Glan-Thompson polarizer Thorlabs GTH10M—Fig. 3: P.
22. Broadband dielectric mirrors 2" diameter. Thorlabs BB2-E03—Fig. 3: M4, M5, M6.
23. Gold protected mirrors 2" diameter. Thorlabs PF20-03-M01—Fig. 3: M1, M2, M3.
24. Back side polished broadband dielectric mirrors 2" diameter. Standa—Fig. 3: M7.
25. High-performance large aperture platform two-axis linear stage, M6 Newport M-406 with HR13 actuators—Fig. 3: sample stage.
26. Piezo stage Physik Instrumente PI-517.3CL—Fig. 3: sample stage.
27. Spatial light modulator Hamamatsu LCOS-SLM (Optical Phase Modulator): X10468-03—Fig. 3: SLM.

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1. Precision kinematic mirror mounts Thorlabs KS1/M—Fig. 5: M4.
2. Precision kinematic mirror mounts Thorlabs KS2/M—Fig. 5: M2, M3, M5, M6.
3. Fixed lens mounts Thorlabs LMR1—Fig. 5: L3.
4. Fixed lens mounts Thorlabs LMR2—Fig. 5: L1, L2.
5. Flip mount adapter Thorlabs FM90/M—Fig. 5: F.
6. Short-pass filter Thorlabs FES0700—Fig. 5: F (2x).
7. N-BK7 AR coating: 650–1050 nm bi-convex lens 1" diameter. $f = 50$ mm Thorlabs LB1471-B—Fig. 5: L3.
8. N-BK7 AR coating: 650–1050 nm bi-convex lens 2" diameter. $f = 200$ mm Thorlabs LB1199-B—Fig. 5: L2.

9. N-BK7 uncoated bi-convex lens 2" diameter. $f = 500$ mm Thorlabs LB1909—Fig. 5: L1.
10. N-BK7 uncoated bi-convex lens 2" diameter. $f = 400$ mm Thorlabs LB1852—Fig. 5: L1 (optional instead of item no. 9).
11. Unmounted positive achromatic doublets AR coated: 400–700 nm 2" diameter. $f = 300$ mm Thorlabs AC508-300-A—Fig. 5: L1 (optional instead of item no. 9).
12. Unmounted positive achromatic doublets AR coated: 400–700 nm 2" diameter. $f = 200$ mm Thorlabs AC508-200-A—Fig. 5: L1 (optional instead of item no. 9).
13. Thorlabs 30 mm Cage System—Fig. 5: MMF holder, lamp.
14. CCD Mikrottron EoSens Mini1—Fig. 5: camera.
15. Diode pumped solid state Laser Innolight—Mephisto 500 (500 mW maximum output) at 1064 nm—Fig. 5: laser.
16. Objective lens Olympus UPlanFL 40 \times /0.75—Fig. 5: objective.
17. White LED Thorlabs MCWHL5 with power supply LEDD1B—Fig. 5: lamp.
18. Broadband dielectric mirrors HR at 750–1100, 1" diameter. Thorlabs BB1-E03—Fig. 5: M4.
19. Broadband dielectric mirrors 2" diameter. Thorlabs BB2-E03—Fig. 5: M5.
20. Gold protected mirrors 2" diameter. Thorlabs PF20-03-M01—Fig. 5: M1, M2, M3.
21. Back side polished broadband dielectric mirrors 2" diameter. Standa—Fig. 5: M6.
22. High-performance large aperture platform two-axis linear stage, M6 Newport M-406 with HR13 actuators—Fig. 5: sample stage.
23. Piezo stage Physik Instrumente PI-517.3CL—Fig. 5: sample stage.
24. Multimode fiber 105 μ m core NA = 0.22, Thorlabs M15L01—Fig. 5: MMF.

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