High throughput system for magnetic manipulation of cells, polymers, and biomaterials


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In the past decade, high throughput screening (HTS) has changed the way biochemical assays are performed, but manipulation and mechanical measurement of micro- and nanoscale systems have not benefited from this trend. Techniques using microbeads (particles ~0.1–10 μm) show promise for enabling high throughput mechanical measurements of microscopic systems. We demonstrate instrumentation to magnetically drive microbeads in a biocompatible, multiwell magnetic force system. It is based on commercial HTS standards and is scalable to 96 wells. Cells can be cultured in this magnetic high throughput system (MHTS). The MHTS can apply independently controlled forces to 16 specimen wells. Force calibrations demonstrate forces in excess of 1 nN, predicted force saturation as a function of pole material, and powerlaw dependence of $F \sim r^{-2.7 \pm 0.1}$. We employ this system to measure the stiffness of SR2+ Drosophila cells. MHTS technology is a key step toward a high throughput screening system for micro- and nanoscale biophysical experiments. © 2008 American Institute of Physics. [DOI: 10.1063/1.2976156]

I. INTRODUCTION

In the past decade, high throughput screening (HTS) has changed the way assays, such as binding, cellular processes, and motility, are performed. Enthusiasm has separately yet simultaneously developed around manipulation and mechanical measurement of micro- and nanoscale systems, including cells and single molecules. This is because the mechanical properties of biological fluids and tissues are responsible for the structural integrity, function, and response of the organism.

There has been little intersection between these two trends: low throughput techniques dominate mechanical study of biological systems. One high throughput nanomanipulation device, IBM’s “Millipede” multip tip atomic force microscope, is a notable exception, but its primary application has been in data storage.

Among the systems for nanomanipulation, techniques using microbeads (particles ~0.1–10 μm) have been widely applied. Microbeads are used to measure mechanical properties of cells. Drug delivery strategies are being developed using magnetic micro- and nanoparticles. Transfection efficiency has been improved using “magnetofection,” where vectors are attached to magnetic particles that are pushed against cells by magnetic fields.

Microbead experiments of various biomaterials—including cells, actin, and fibrin—are frequently interpreted using the language of microrheology. It has been noted that microrheological measurements show promise as a technique for high throughput screening. We note that among microbead techniques, the “active” or “driven” techniques, where an external force is applied to a bead, can measure a wider range of moduli and are generally more flexible than the passive techniques, where the bead motion is due to thermal diffusion. Driven microrheology instrumentation includes laser traps (“optical tweezers”) and paramagnetic attraction (“magnetic tweezers”). A laser trap offers high spatial resolution and can be precisely calibrated. A magnetic system can drive multiple beads simultaneously, produce higher forces at a given bead size, and will neither heat nor otherwise interact with most biological specimens.

In pursuit of HTS nanomanipulation, we have expanded on instrumentation to magnetically drive microbeads. Here we demonstrate a biocompatible, multiwell magnetic force system compatible with HTS standards. This magnetic high throughput system (MHTS) can apply forces of ~1 pN –1 nN. It is based on a standard microplate geometry and is designed to be scalable to 96 wells. We demonstrate that this technology can be used to measure mechanical properties of cells. MHTS technology is a key step toward a high throughput screening system for micro- and nanoscale biophysical experiments.
II. MHTS DESIGN AND FABRICATION

A. Magnetics miniaturization and temperature control

Applying force to magnetic beads requires a magnetic field gradient. For an initially unmagnetized, permeable magnetic bead, the force on the bead is

$$F = \nabla (m \cdot B) = \frac{\pi d^3}{4 \mu_0 \mu_r} (B^2),$$

where $m$ is the magnetic moment of the bead, $d$ is its diameter, $\mu_r$ is its relative permeability, and $B$ is the flux density of the ambient magnetic induction $B$, as discussed in a previous work on a three-dimensional force microscope (3DFM).19 This relation is valid until the bead becomes fully magnetized at a saturation magnetization $M_{sat}$, where

$$m = \frac{\pi d^3}{6} M_{sat} \mathbf{m},$$

where $\mathbf{m}$ is the unit vector along $B$. With the bead saturated, the force equation simplifies to

$$F = \frac{\pi d^3}{6} M_{sat} \nabla B.$$

We desire a large magnetic field gradient to achieve high forces. In our work on the 3DFM and in Ref. 20, we employ a magnetic circuit design to achieve sufficient field gradient. Current-carrying drive coils magnetize a drive core, which preferentially directs magnetic flux to a pole of magnetic foil. The foil tapers to a pointed tip that sits submerged in the substrate is a pole flat affixed to glass. The quadrants on the substrate show the footprint of a 2×2 section of the 384-well tray. The three-legged structure is the corresponding section of the magnetics block. A magnetic circuit begins in the drive coil (front well), where current creates magnetic flux directed by the drive core into the pole. Flux jumps the specimen gap (back well) to the flat, which is connected by two return path ears (left and right wells) to the back of the coil. By changing the etching pattern in the specimen gap (back well), the force field can be changed. A through hole above the specimen gap provides access for dosing of the specimen during an experiment and illumination for brightfield transmission microscopy.

Collectively, the pole and flat are called a pole flat; this geometry can pull only toward the pole tip. Alternate pole geometries may be used for experiments that require different force fields. This thin-foil design accommodates a high numerical aperture, short working distance imaging objective, enabling high resolution imaging simultaneously with force application.

The present instrument, a MHTS, consists of two components: (1) a magnetics block, which generates magnetic flux for each well, and (2) a magnetic microplate, a custom-designed high throughput tray that incorporates a pole flat in each specimen well. The arrangement of these components is summarized in Fig. 1. The magnetics block contains a system of drive cores and drive coils; we report here on a 16-well prototype with a design that is scalable to 96 wells.

1. The magnetics block

Generating magnetic forces requires a complete magnetic system: drive coil, drive core, and magnetic return path. Figure 2 demonstrates how a complete magnetic system can fit within the area of one Society for Biomolecular Screening (SBS) microplate well. Each of the 16 magnetic systems includes a drive core and two ears that provide the magnetic return path. The magnetics block also has 16 through holes for brightfield illumination of the wells. To direct magnetic flux to the poles, the drive cores and ears extend past the bottom of the coils, so that when the magnetics block drops into the microplate, the drive cores and ears contact the pole flats at the bottom of the wells. The magnetics block is machined from a blank of ASTM A848 low-carbon magnetic iron. The drive coils are 58 turns of No. 26 AWG magnet wire, Part No. UNC-1 (Quality Coils, Inc., Bristol, CT).

Both biology and rheology are highly temperature sensitive, so we have built a temperature control system into the magnetics block. Fluid channels are drilled transversely to the optical paths, and aluminum manifolds are attached to the sides of the block as an interface to a heat exchanger and recirculation pump. In the limiting condition of all 16 wells at a maximum drive current, the temperature rise is 4 °C and the time constant is less than 1 min. The flow rate is within the laminar flow region of the channels, as calculated using a Moody chart.21

2. Fabrication of a biocompatible magnetic microplate

While we can place the magnetics above the specimen, the poles must be built into the HTS microplate. This constraint has three motivations. First, the force on beads increases with the pole tip’s proximity to the specimen. Second, when using a pole flat, forces are applied in the plane of the pole tip, which to record bead motion must be parallel to the imaging plane. (For experiments requiring forces parallel to the imaging axis, alternate pole geometries may be implemented.) Third, a magnetic microplate can be washed and reused. It is relatively inexpensive, and so can be produced in quantity to allow for simultaneous preparation of many different experiments.

We assemble the magnetic microplate by adhering a pole plate underneath a bottomless plastic 384-well superstructure (Thermo Fisher Scientific, Portsmouth, NH). The pole plate is an array of 16 pole flats fixed to a 4.33×2.93 in.2 No. 1–thickness coverglass. We summarize the process in Fig. 3.
We create the pole plate pattern using a commercial photolithographic etching process (Fotofabrication Corporation, Chicago, IL). The poles must be isolated from any surrounding magnetic material. Therefore, to maintain registration of these “floating” poles relative to the flats, we bond the 0.004 in. thick foil (Magnetic Shield Corporation, Bensenville, IL) to a sacrificial sheet of 0.005 in. thick polycarbonate backing (K-mac Plastics, Wyoming, MI). Here we report on an array of pole flats (see the inset of Fig. 3, step 2), but any shape of pole tip may be specified at this stage. The pole plate pattern is etched from the metal side of the foil on plastic.

To transfer the patterned foil to an optically clear substrate while maintaining the registration of the poles, we roll an UV-curable adhesive NOA 81 (Norland Optical Adhesives, Cranburg, NJ) onto the etched foil, cover it with No. 1–thickness coverglass, press flat with a glass plate and cure, then remove the plastic backing. This orientation places the sharp tip edge \( \sim 100 \, \mu m \) above the surface of the coverslip. A layer of parylene, vapor-deposited at this stage, inhibits pole degradation.

The highest force is in the plane of the pole edge, where the force ranges as high as \( \sim 1 \, nN \) on a 4.5 \( \mu m \) bead. This

![Diagram](image_url)
geometry is desirable for some applications, including rheometry of biomaterials. For other applications, including cell force experiments, it is desirable to have the bottom of the well level with the pole. To raise the cells to this height, we pot the foil in NOA 81, creating an optically clear surface within 10 μm of the plane of the poles. To create a surface conducive to cell adhesion, we vapor deposit 3-aminopropyltriethoxysilane (Sigma-Aldrich, St. Louis, MO) solution onto the layer of NOA 81.

In the final step, Double/Bubble epoxy (V. O. Baker Co., Mentor, OH) epoxy bonds the potted pole plate to the 384-well superstructure such that the wells will not leak. The result is a 384-well microplate where three quarters of the wells are used for the drive cores and ears, leaving 96 wells for specimens. To elucidate the arrangement of the pole plate, well superstructure, and magnetics block, the fully assembled MHTS is shown in Fig. 4.

3. Considerations for MHTS operation

The MHTS technology described above comprises 16 independently controlled magnetic force generation systems. This instrument may be operated in a variety of modes. For example, identical forces may be applied simultaneously to a range of specimens by running the coils in series. Alternatively, different force profiles (say, pulses of varying length) may be applied to each well, limited by the number of current supplies available. In the present work, we use a computer-controlled transconductance current source;20 manual switching runs this current in series through the desired coils.

The imaging subsystem in this work has a single field of view and a manual translation stage. Therefore, experiments that require a realtime record of bead motion must be performed sequentially. In addition, subsequent particle tracking and data analysis have been minimally automated. Nevertheless, experiments profit from the microplate geometry, which allows multiple specimens to be prepared simultaneously and more efficiently.

Using these nonautomated systems for a 15 s experiment, typical times are 2 min per well for data acquisition and 15 min per well for particle tracking and data analysis. Naturally, the MHTS technology presented here begs to be paired with existing HTS robotics and software. To this end, we are interested in implementing software to synchronize drive current commands with the position of a robotic microplate stage. The most urgent application for robotic data collection would be an automated force calibration routine. Additionally, control software to manage files associated with MHTS operation will greatly improve the utility of the MHTS. These files include force calibration, drive current, video logs, and subsequent tracking and analysis files.

III. FORCE CALIBRATION

The force field of an ideal pole can be modeled, but deviation of manufactured tips from the model makes the prediction inadequate for precise force calibration. To address this problem, we have developed variable force calibration (VFC) software to construct the force field around the pole tip as a function of both distances from the pole tip $r$ and the drive current $I$. We use VFC with the 3DFM to explore how forces are affected by the pole material, the drive current, and the bead position. We have also used VFC to calibrate the MHTS.

Forces are calculated from the velocity of beads pulled through a fluid of known viscosity. For Newtonian fluids with a low Reynolds number ($Re < 10^{-4}$), the force on a spherical object is equal to the Stokes drag multiplied by the particle’s velocity, $F = 6\pi \eta v r$, where $\eta$ is the sphere radius, $\eta$ is the dynamic viscosity of the fluid, and $v$ is the sphere’s velocity. Given the fluid viscosity and the bead radius, the bead’s velocity uniquely determines the force at a given position and drive current.

Calibration is typically performed using $2.5M$ sucrose ($\eta=0.12$ Pa s). Corn syrup (ACH Food Companies, Memphis, TN), a higher viscosity fluid ($\eta=3.4$ Pa s), can be used to calibrate higher force regimes.

A. Variable force calibration in the 3DFM

1. Magnet control and voltage sequencing

A graphical environment, written in MATLAB, allows the user to drive the magnetic system with a series of constant-current pulses. A single pulse sequence, shown in Fig. 5, can be repeated as many times as necessary to collect sufficient calibration data.

The magnetic foil exhibits remanent magnetization after the application of a magnetic field. We can measure this hysteresis as a velocity of the magnetic bead at zero drive current. The remanence magnetization following the application of high drive current can mask subsequent low drive currents. To degauss the poles, we apply a drive current

$$I(t) = I_{max} e^{-\frac{t}{\tau}} \sin 2 \pi f t,$$

where $I_{max}$ is the maximum applied current since the last degauss, $\tau$ is the decay time constant, and $f$ is the frequency of the sinusoid. In the MHTS, $f=1$ kHz and $\tau=0.012$ s. Figure 6 shows the efficacy of this degauss technique.

2. Analysis pipeline

A datum of calibration is the bead’s velocity at each drive current in each pulse sequence. Consider a population of $n$ beads, which are each subjected to an average of $p$ pulse...
sequences. Each pulse sequence comprises $d$ pulses at a specified drive current. A calibration set then includes $npd$ data. Typical values are $n=20$, $p=10$, and $d=10$. We note that increasing $d$ improves resolution in measuring how force varies with drive current; increasing $n$ minimizes error due to bead-to-bead variation in force; shortening the dwell time of each pulse increases $p$ and also moves the beads a shorter distance during each pulse sequence, thereby improving spatial resolution in the force calibration.

The only beads included in the analysis are those within an angle of $\pm 30^\circ$ of the pole’s line of symmetry, as the force varies significantly outside of that cone. The force field’s radial symmetry is apparent in Fig. 7. We extend the bead trajectories to a point of common intersection within the area of the pole tip to determine the origin of this polar coordinate system. The coordinates of \([x(t),y(t)]\) are transformed into this system to find $r(t)$. A linear fit to $r(t)$ at each drive current $I$ reveals the velocity $v(r,I)$.

The force data $F$ may be displayed either as a function of $r$ or as a function of the drive current at a given distance, $F(I)$. The former representation is used in Fig. 6. The latter representation is used in Fig. 8 and provides insights into the pole plate material. At sufficient field strength, we expect the system’s magnetic materials to achieve some maximum magnetization. This manifests as a saturation in the force on the beads as a function of current. Figure 8 shows the saturation behavior of two pole plate materials. The foil types are Netic ($B_{sat}=2$ T) and CoNetic ($B_{sat}=0.8$ T). As expected, the material with higher flux density saturation shows a higher force at saturation.

In Ref. 19 we argued that a pole tip can be modeled as a...
magnetic monopole, such that $B$ drops quadratically with distance from the monopole. Using the force equation for a saturated bead, we note that

$$F \sim \nabla B \sim r^{-3}.$$  

Figure 9 demonstrates powerlaw behavior of $F \sim r^{-2.7}$. That the force model underpredicts the empirical powerlaw is encouraging. A real pole tip is not a magnetic monopole: it would more accurately be modeled as a distribution of magnetic charge. Such a distribution would have higher multipole moments than just the quadratic term, leading to a drop in the overall powerlaw.

B. MHTS calibration

We have used VFC to characterize the forces in the MHTS. Of greatest interest are the maximum force and crosstalk force. Figure 10 demonstrates that the maximum forces are $\sim 1$ nN, and with 15 wells on and one well off, the crosstalk is $3 \pm 1\%$.

Ideally, the pole tips would be uniform from well to well, ensuring identical force fields in all wells. However, the etching process that produces the pole plate pattern has not proven reliable in producing uniformly sized poles, and therefore the calibration varies from well to well by $\pm 40\%$.

IV. APPLICATION: MECHANICAL RESPONSE OF CELL MEMBRANE

There is a potential for a high throughput force system to have a dramatic impact on studies in cell mechanics and motility. For years, “cell rheology” has involved the study of biochemical and mechanical cellular responses under various mechanical (flow and prestress) and chemical (blockers and pH) conditions. 22,23 A recent trend in biological studies has been the development of libraries of small molecules, proteins, and RNAi for gene expression manipulation. These libraries offer enormous opportunities for understanding complex biochemical networks and for drug discovery, but impose daunting challenges to analytical techniques. A high throughput system can mitigate some of these challenges, including prep time, run time, and consistency in experimental procedure.

It has been a motivation for our development of the MHTS technology to explore gene-expression-inhibited and ligand-mediated cell mechanical responses of *Drosophila* (fruit fly) cell systems using a previously developed RNAi library. 22 Understanding these pathways will help reveal the genetic origins of diseases such as cancer, hypertension, and inflammatory disorders.

We have used the MHTS to study the mechanical response of cells under varying biochemistry. *Drosophila* derived cells (SR2+) were grown in SF 900 medium on potted, APTES treated, and Conconavalin A–washed magnetic microplates. That cell cultures grow in a magnetic microplate demonstrates MHTS biocompatibility; a representative well is shown in Fig. 12. The cultures were incubated at room temperature for 24 h before adding a suitable concentration of 2.8 μm volume-loaded superparamagnetic, COOH-functionalized beads (Invitrogen USA, Carlsbad, CA), which bind to the cell due to nonspecific electrostatic attraction. The cells were incubated for an additional hour, then washed to remove any unattached magnetic beads. Half of the wells with cells in them were treated with a protein tyrosine phos-
phatase (PTPase) inhibitor cocktail (Calbiochem, San Diego, CA), with a final concentration of 0.5 μM. Tyrosine phosphatases have been known to play an important role in force sensing in cells, but there have been no quantitative studies on the changes in the mechanical properties of the cell due to the inhibition of PTPases.

Figure 13 compares the compliance of treated and untreated cells, as measured from bead displacements. A spring constant for a cell can be found by fitting the data to a modified Kelvin–Voigt model. The spring constant for untreated Drosophila cells was $756 \pm 67 \times 10^{-6}$ N/m, SR2+ cells that were treated with the PTPase inhibitor showed greater compliance. The spring constant for cells treated with the PTPase inhibitor was found to be $240 \pm 53 \times 10^{-6}$ N/m.

Using the MHTS technology, a single experiment enabled mechanical measurements on cells in four independent cell cultures. The PTPase-treated SR2+ Drosophila cells exhibited thrice the compliance of control conditions; this difference is outside the uncertainty in the calibration of each well. Studies using the MHTS to study cell mechanoresponse under varying biochemical conditions are ongoing.

V. CONCLUSIONS

We have presented critical technology for a MHTS based on commercial HTS standards and scalable to 96 wells. Our prototype applies independently controlled forces to 16 specimen wells. Force calibrations demonstrate forces in excess of 1 nN and reveal the saturation behavior of the magnetic pole material and expected force field dependence that approaches $F \sim r^{-3}$. We have demonstrated that MHTS technology can be used to perform cell manipulation experiments on independently prepared and treated cell cultures.

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