Micro & nanofabrication methods for nanoscience applications

Subtitle: toward 3D fabrication, manipulation and characterization

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Tilted stage for 3D exposures

3D building block for microfluidic

NanoImprinting

3D Metal Structures

LEO 1540XB CrossBeam**®** Workstation

System Features:

- Super Eucentric 6-axis stage X 102 mm, Y 102 mm
- Gas injection system
- 4" Airlock (optional)
- Automated aperture change on FIB column
- Enhanced vacuum system

Options:

- EDS
- CAD
- Lithography
- SIMS

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Beam deposition and gas assisted etch

FIB on AFM tip

Focused Ion Beam - microsculpture

3D manipulation et al:Optical Tweezers, Two Photon lithography, two photon microscopy

L'uso combinato di forbici e pinze laser

consente di effettuare delicate manipolazioni a livello subcellulare. Nella procedura illustrata, che dovrebbe essere realizzabile entro una decina d'anni, due fasci che fungono da pinze (in rosa) trattengono strettamente la cellula. Un fascio forbice (in blu chiaro) penetra

nella cellula per eliminare un gene difettoso (in rosso). Un secondo fascio forbice (in blu scuro) crea nella membrana cellulare un foro attraverso il quale potrà entrare una sequenza genica appropriata (punti neri). Si potranno poi ottenere cloni della cellula modificata che saranno trapiantati a scopo terapeutico. **KINT National Nanotechnology Laboratories ICTP winter college 7-18 Febbraio 2005 Trieste TASC - INFM Optical tweezers - background Two regimes of operation:**

- \triangleright Rayleigh regime (size of particle $\lt\lt\lambda$)
- \triangleright Mie regime (size of particle $> \lambda$)

Ashkin, A.; Dziedzic, J. M.; Bjorkholm, J. E.; and Chu, S., "Observation of a Single-Beam Gradient Force Optical Trap for Dielectrical Particles", Optics Letters 11, pp 288-290 (1986) - ray optics model

Implementation of multiple optical tweezers using diffractive optical elements

Dynamic optical tweezers – setup at LILIT

Experimental results

2 D – Array of optical tweezers

Experimental results

3D displacement (2)

3 microspheres are trapped and moved in X-Y-Z

Experimental results

Transfer of orbital angular momentum with doughnut beams

External Force Gradients and Cell Polarity (Collaboration with V. Emiliani- Jacques Monod Institute)

...a key role is played by the cell capability to sense mechanical gradients and tensions in the environment

• Durotaxis: cell movement can be guided by physical interactions at the cell-substrate interface

> Lo CM, Wang HB, Dembo M, Wang YL, Biophys. J 79, 144 (2000)

• Tensile stress: stimulate Microtubule outgrowth, RAC (GTPAses) activity, axons growth

via micro-needles and flexible substrates

I. Kaverina et al. J. Cell Science 115, 2283 (2002) S. Chada et al. J. Cell Science 110, 1179 (1997)

equi-biaxial stretch device

Akira Katsuni et al. J Cell Biol. 158, 153 (2002)

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Force gradient

The trapping force is proportional to the laser intensity: by making an array of laser spots with a gradient in intensity we can mimic a gradient in the substrate rigidity

Mechanical stress

The all array can be deformed in a symmetric or asymmetric way:

Imaging: Microtubules and Actin network organization

Optical manipulation of liposomes as microreactors

Figure 2: Schematic representation of the procedure rigure 2: Schematic representation of the procedure
for the trapping and fusion of two liposomes, con-
taining different chemicals. a) Two liposomes, one taining different chemicals. a) Two liposomes, one
containing reagent A and the other one containing reagent B, are identified in the sample. b) The two
liposomes are trapped in separate optical tweezers
and translated su the contact point. d) The membranes repair spontathe contact point of the inclusion of the contact of the point of the reagents A and B mix.

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Figure 3: Fusion of two liposomes. The images are recorded by video microscopy. The fusion was initiated by the UV laser at the time when the first image scatted The next two images capture the proposition gress concled.

Figure 6: Fusion of two liposomes, one containing fluo-3 dye and the other one containing calcium ions. The bright field video microscopy images and the fluorescence images recorded simultaneously before and after the fusion was initiated (upper and lower images, respectively). After the fusion the fluorescence increases as a consequence of the reaction in which fluo-3 chelates the calcium ions.

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