Cell-Scaffold Interactions:

Scaffold Degradation Cell Attachment Cell Morphology Cell Contractility Cell Migration Cell Differentiation

Cell scaffold interactions

· Scaffolds also being used to characterize cell-scaffold interactions e.g. how cell be haviow (attachment, migration, contraction, differentiation) is affected by substrate

Scaffold degradation

- native ECM enzymes produced by cells result ELM over time; cells also synthesize new ECM to replace it e.g. bone-rates of resorption + synthesis depend on loading
- · cells also degrade tissue engineering scaffolds
- · length of time scaffold remains moluble called "residence time"
- · require scattold degradation to occur in a manner that does not interfere with new ECM synthesis
- scaffold residence time must be approx. Equal to the time required to symphesize new ECM

- · degradation rate for scaffold depends on its chemical composition + Crosslinking and on relative density of scaffold
- · Synthetic polymers Can vary molecular variation of polymers + ratio of Co-polymers eq. PLGA higher GA: LA ratio polymers degrade quicker
- · collagen based scaffolds can control degree of coss-linking

physical methods: - dehydrothermal (DHT) treatment (105°C vacan 24hs) - revolves water, forms interchain bonds through candensation

- uv treatment

chemical methods - glutaraldehyde; carbodiimite treatments

Cell adhesion

- · cells attach to ECM at focal adhesion
- · at focal adhesian
 - · cell has integrins transmembrane proteins that bind to ligends an ECM other end of integrin connects to sub - membrane plaque that then connects to cell's cyto skeleton (eq. to actin filaments)
- · cell behaviours such as attachment, migratian, proliferation, contraction
- a ffected by interactions between focal adhesians + integrins - biological activity of scaffelds depends on density of ligands available for integrins to bind to
- · ligand density depends an composition of scaffold + surface area/volume
- · Diological polymers that are constituents of native Zof scafford ECM (e.g. collagen) have a range of native binding sites
- · synthetiz polymers don't have binding sites + need to be functionalized with adhesive proteins such as fibronectin + laminin

- · specific surface area (salvoi) of scaffold depends on pire size, + relative density:
- · for a fetra kai decahedra 1 unit cell

 $\frac{SA}{V} \times \frac{1}{d} \left(\frac{p^*}{p_s} \right)^{\frac{1}{2}} \qquad \begin{bmatrix} SA/V = \frac{2\pi r \ln \alpha}{\lambda^3} \times \frac{r}{\lambda^2} \times \frac{r}{\lambda^2} \end{bmatrix} \xrightarrow{r} \frac{1}{d} \left(\frac{p^*}{p_s} \right)^{\frac{1}{2}} \frac{1}{d}$

2 1r

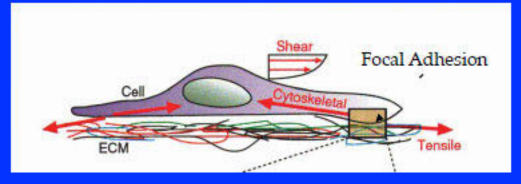
(4)

· dependence of cell attachment an specific surface area was measured by seeding cells (MC 3T3-ET mouse oskogenic) onto collagen-GAG scattoldo of constant relative density (p*ps=0.006) + Varying pore size

d= 96, 110, 121, 151 um

- . number of cells attached measured at 24, 48 hours
- · fraction of cells attached in created linearly with specific surface area <u>Cell Morphology</u>
- · cell orientation follows scafford pore orientation
- · cell morphology can depend an substrate stiffness
- <u>Cell contraction</u> Z se slides. <u>Cell misration</u> Z

Cell Adhesion



Gibson, L. J., M. Ashby, et al. *Cellular Materials in Nature and Medicine*. Cambridge University Press. © 2010. Figure courtesy of Lorna Gibson and Cambridge University Press.

Figure removed due to copyright restrictions. See Figure 9.1: Gibson, L. J., M. Ashby, et al. *Cellular Materials in Nature and Medicine*. Cambridge University Press, 2010. http://books.google.com/books?id=AKxIS4AKpyEC&pg=PA255

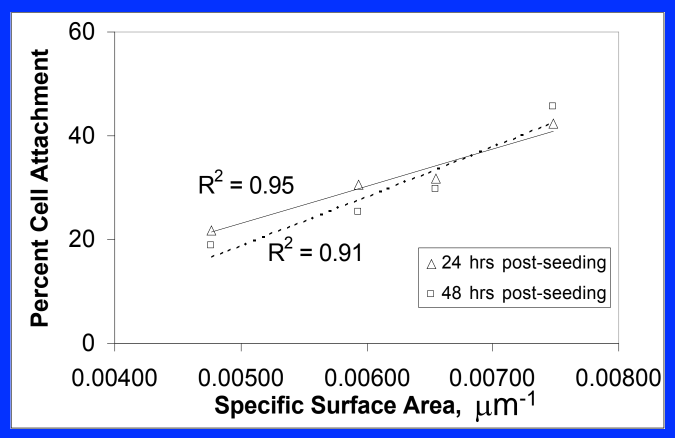
Gibson, Ashby and Harley, 2010

Cell Attachment

$$\frac{SA}{V} = \frac{3.65}{l} \left(\frac{\rho^*}{\rho_s}\right)^{1/2} = \frac{0.718}{d}$$

Open-cell tetrakaidecahedron Circular cross-section edges I = edge length d = pore size Collagen-GAG scaffold: $\rho^*/\rho_s = 0.005$, d = 96, 110, 121, 150µm

Cell Attachment

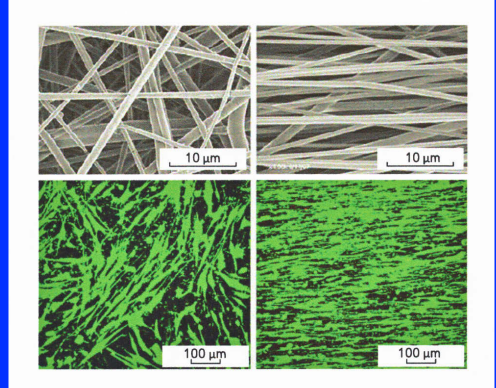


O'Brien, B. A. Harley, I. V. Yannas, et al. *Biomaterials* 26 (2005): 433-41. Courtesy of Elsevier. Used with permission. http://www.sciencedirect.com/science/article/pii/S0142961204002017

Mouse MC3T3 osteogenic cells on collagen-GAG scaffold

O' Brien

Cell Morphology



PLGA scaffolds

Seeded with rotator cuff fibroblasts

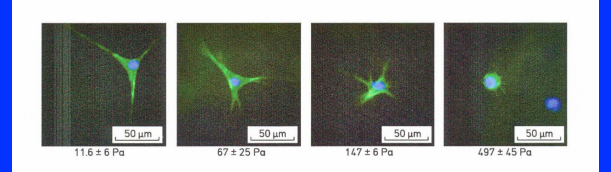
Random



Moffat, K. L., et al. *Clinics in Sports Medicine* 28 (2009): 157-76. Courtesy of Elsevier. Used with permission. http://www.sciencedirect.com/science/article/pii/S0278591908000707

Moffat et al, 2009b

Cell Morphology



E = 11.6 67 147 497 Pa

Dikovsky, D. H., et al. *Biophysical Journal* 94 (2008): 2914-25. Courtesy of Elsevier. Used with permission. http://www.sciencedirect.com/science/article/pii/S0006349508705411

Smooth muscle cells encapsulated in a PEG-fibrinogen hydrogels of varying modulus

Dikovsky et al., 2008

Cell Contractility: Wound Contraction and Scar Formation

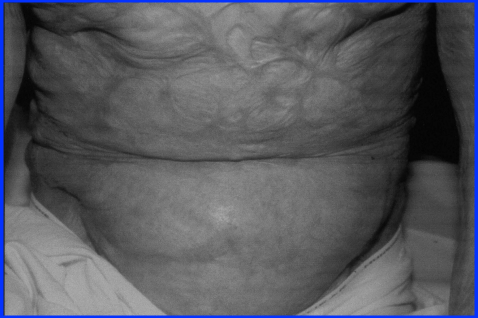


Image source unknown. All rights reserved. This content is excluded from our Creative Commons license. For more information, see http://ccw.mit.edu/help/faq-fair-use/.

Wound contraction associated with scar formation

Use of collagen-GAG matrix inhibits wound contraction and scar formation; results in synthesis of normal dermis

Photo courtesy of IV Yannas

This observation has led to interest in contractile response of cells on the scaffold

Contractility of Cells

Biological cells can contract a scaffold
Free-floating tests

Measure diameter change

Developed cell force monitor (CFM) to measure forces

Collagen-GAG Scaffold

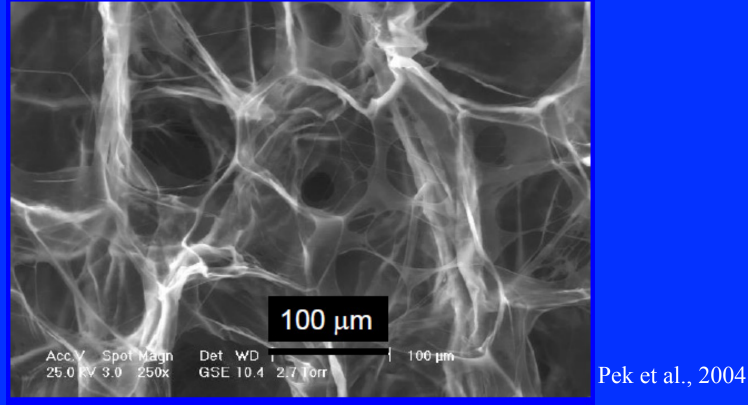
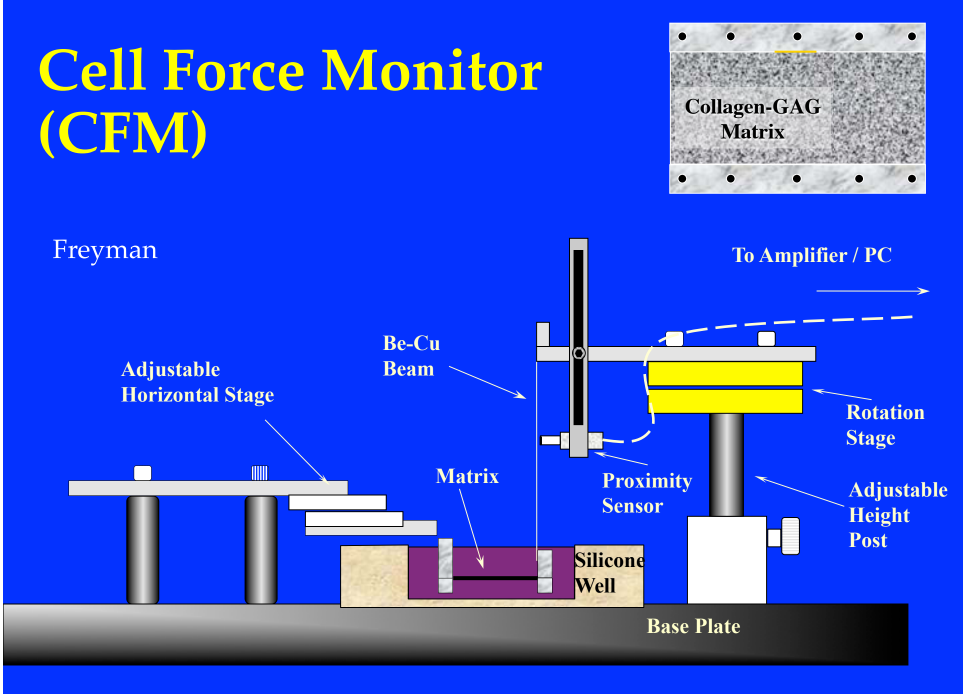


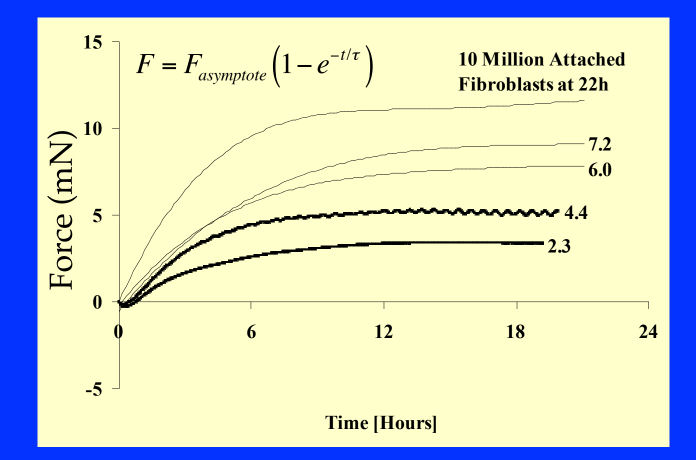
Fig. 1: Pek, Y. S., M. Spector, et al. *Biomaterials* 25 (2004): 473-82. Courtesy of Elsevier. Used with permission. http://www.sciencedirect.com/science/article/pii/S0142961203005416

Scaffold developed by IV Yannas (MIT)



Source: Freyman, T. M., et al. "Fibroblast Contractile Force is Independent of the Stiffness Which Resists the Contraction." *Experimental Cell Research* 272 (2002): 153-62. Courtesy of Academic Press/Elsevier. Used with permission.

CFM: Effect of Cell Number

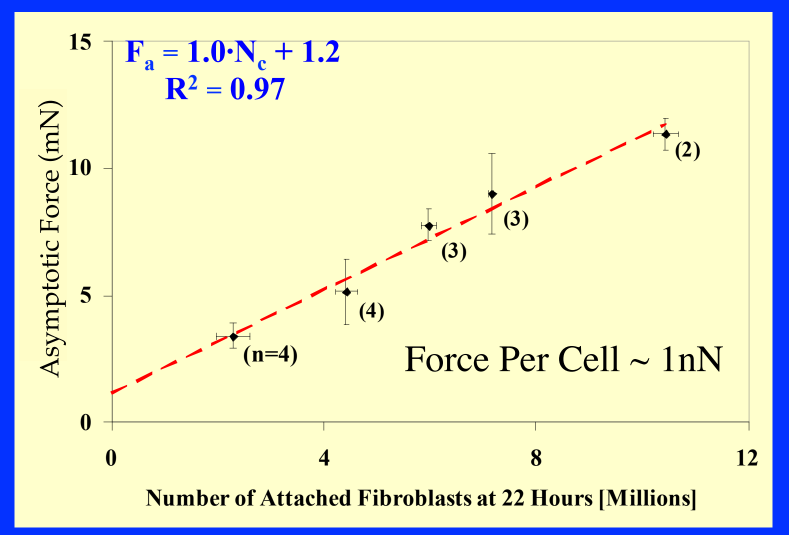


Time constant 5.7 hours



Freyman, T. M., I. V. Yannas, et al. "Fibroblast Contraction of a Collagen-GAG Matrix." *Biomaterials* 22 (2001): 2883-91. Courtesy of Elsevier. Used with permission.

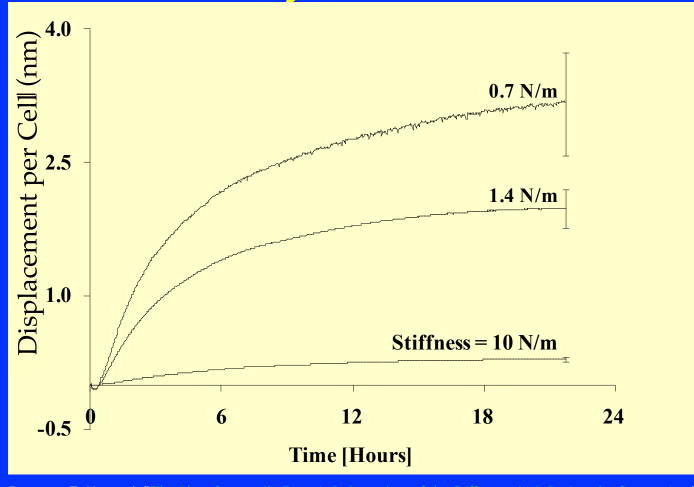
Effect of Cell Number



Freyman, T. M., I. V. Yannas, et al. "Fibroblast Contraction of a Collagen-GAG Matrix." *Biomaterials* 22 (2001): 2883-91. Courtesy of Elsevier. Used with permission.

Freyman

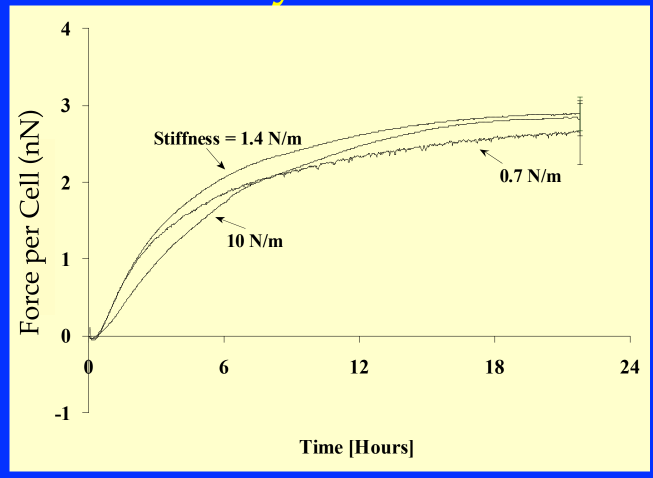
Effect of System Stiffness



Freyman, T. M., et al. "Fibroblast Contractile Force is Independent of the Stiffness which Resists the Contraction." *Experimental Cell Research* 272 (2002): 153-62. Courtesy of Elsevier. Used with permission.



Effect of System Stiffness



Freyman, T. M., et al. "Fibroblast Contractile Force is Independent of the Stiffness which Resists the Contraction." *Experimental Cell Research* 272 (2002): 153-62. Courtesy of Elsevier. Used with permission.

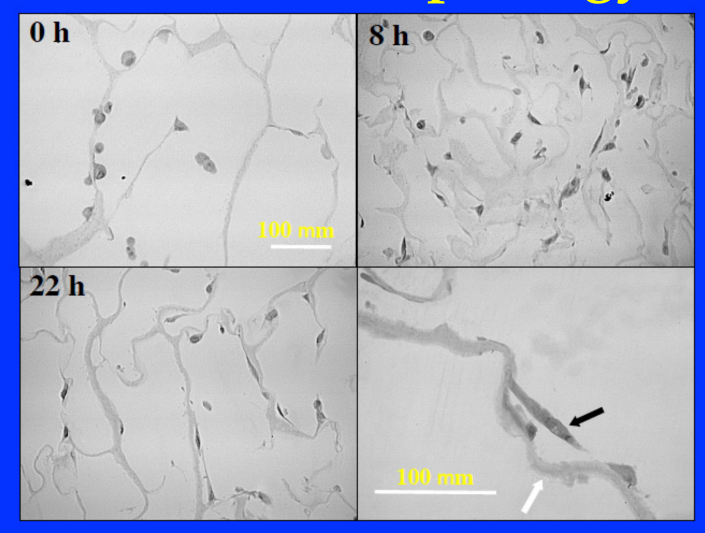
Freyman

Methods: Cell Elongation

Average aspect ratio of cells

- Time points 0, 4, 8, 15, 22, and 48 h (n=3)
- Hematoxylin & eosin (H&E) stained glycomethacrylate (GMA) sections (5mm)
- Digital image analysis (~200 cells per sample)

Fibroblast Morphology





Source: Freyman, T. M., et al. "Micromechanics of Fibroblast Contraction of a Collagen-GAG Matrix." *Experimental Cell Research* 269 (2001): 140-53. Courtesy of Academic Press/Elsevier. Used with permission.

Fibroblast Morphology

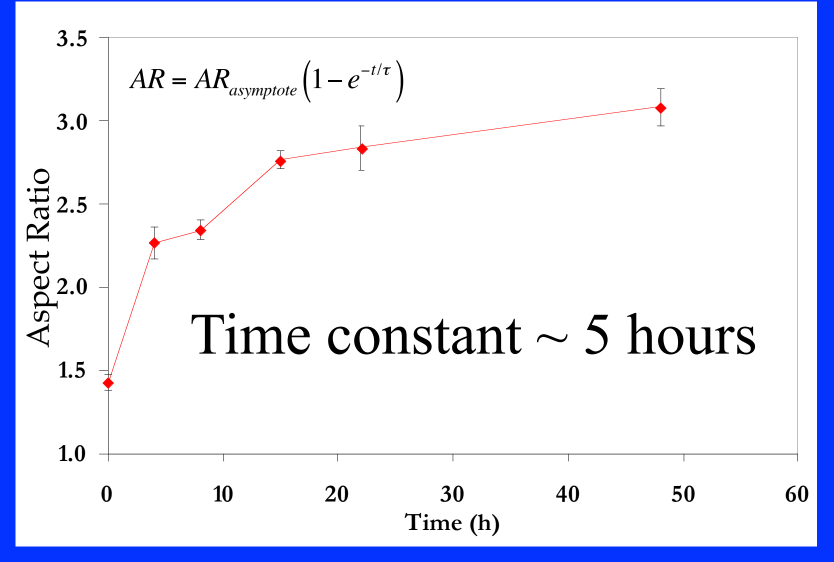


Image after Freyman, T. M., et al. "Micromechanics of Fibroblast Contraction of a Collagen–GAG Matrix." *Experimental Cell Research* 269 (2001): 140-53.

Freyman

Time Constants

- Time constant for contraction ~ 5.7 hours
- Time constant for elongation ~ 5 hours
 Suggests a link between the average elongation of the cell population and the macroscopic contraction of the population

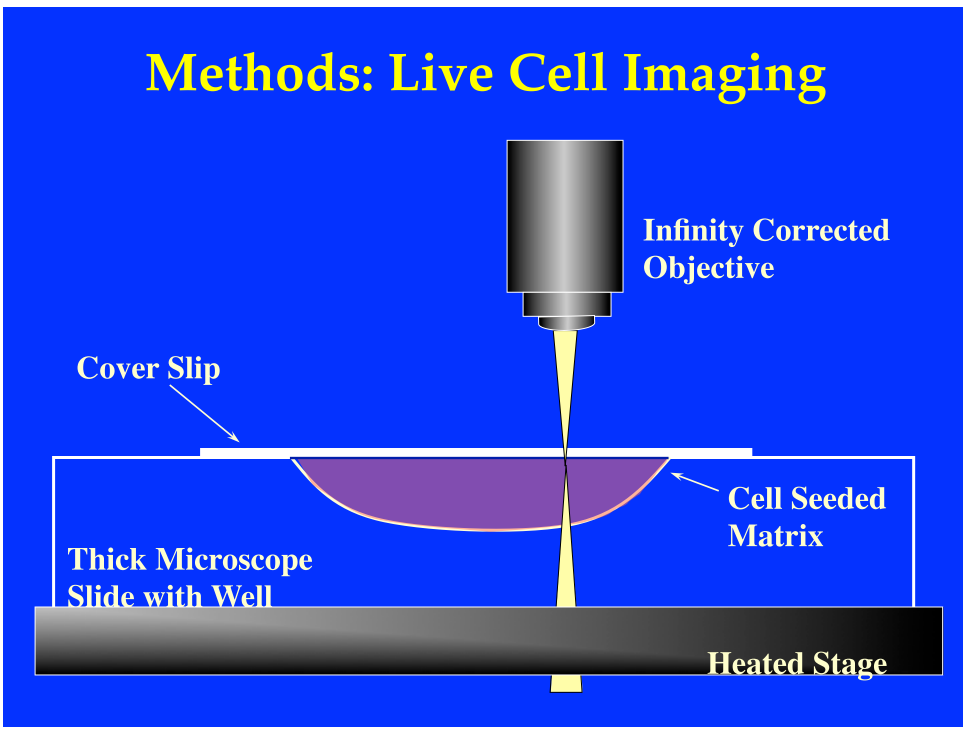
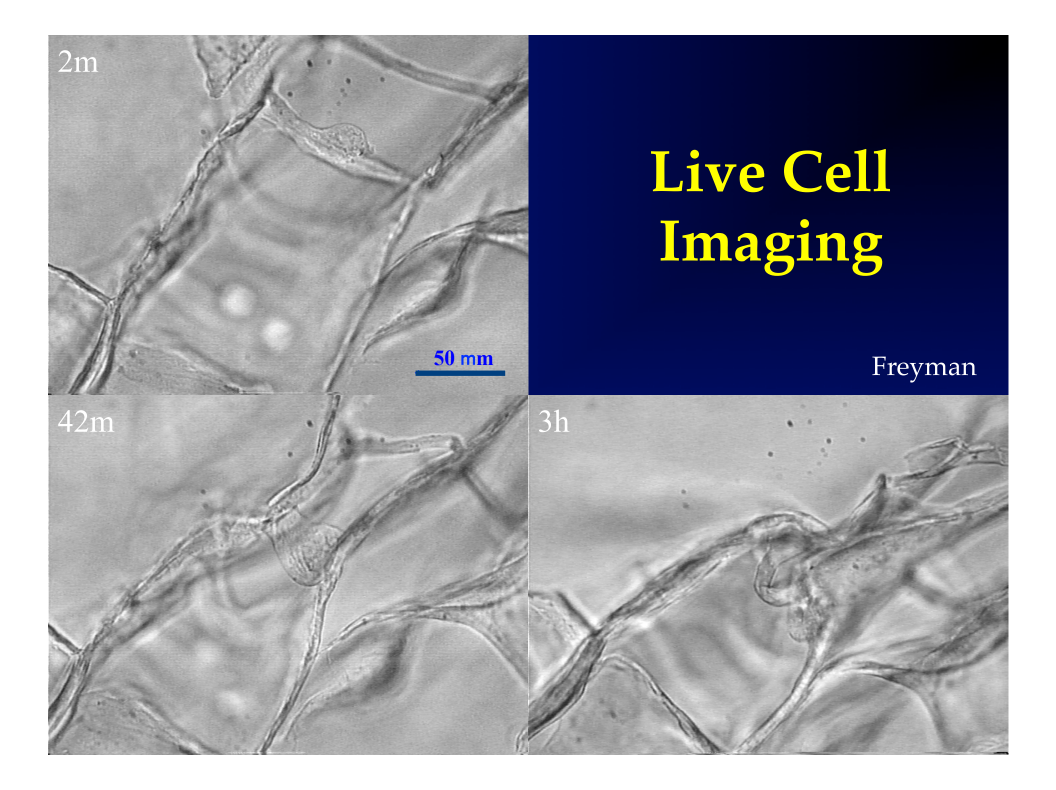


Image after Freyman, T. M., et al. "Micromechanics of Fibroblast Contraction of a Collagen–GAG Matrix." *Experimental Cell Research* 269 (2001): 140-53.

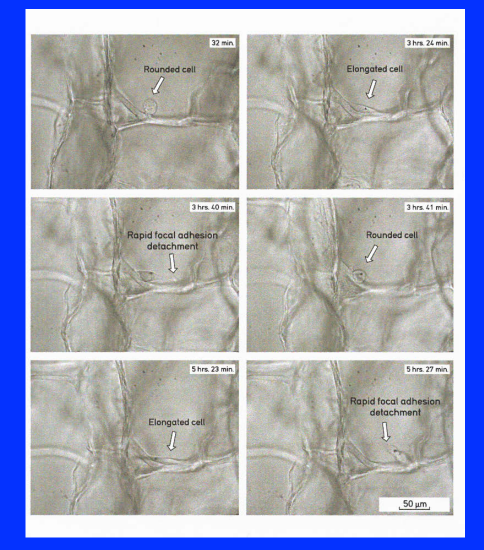


Live Cell Imaging

Figure removed due to copyright restrictions. See Figure 7: Freyman, T. M., et al. "Micromechanics of Fibroblast Contraction of a Collagen–GAG Matrix." *Experimental Cell Research* 269 (2001): 140-53.

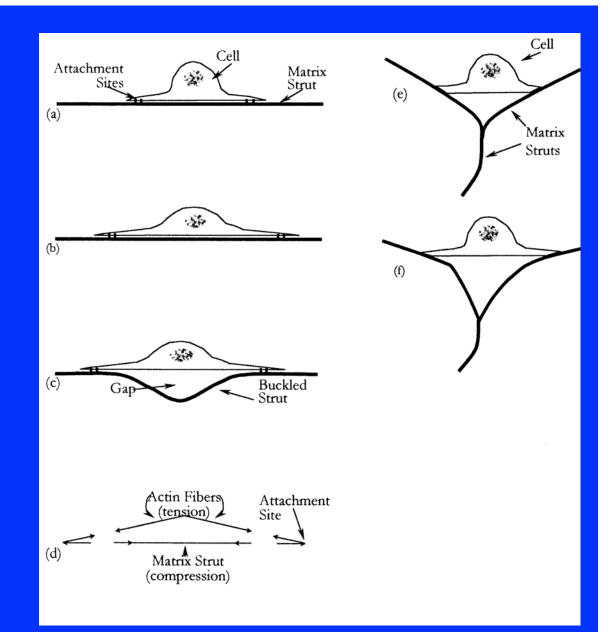


Live Cell Imaging



Source: Freyman, T. M., et al. "Micromechanics of Fibroblast Contraction of a Collagen–GAG Matrix." *Experimental Cell Research* 269 (2001): 140-53. Courtesy of Academic Press/Elsevier. Used with permission.

Schematic of cell elongation and matrix contraction



Freyman

Source: Freyman, T. M., et al. "Micromechanics of Fibroblast Contraction of a Collagen–GAG Matrix." *Experimental Cell Research* 269 (2001): 140-53. Courtesy of Academic Press/Elsevier. Used with permission.

Discussion

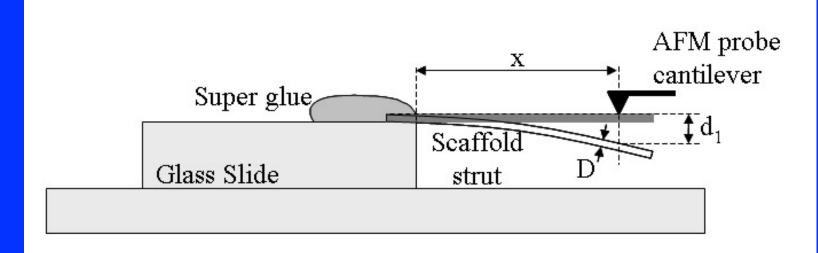
• Cell elongation linked to contraction

- time constants for cell elongation and contractile force development similar (τ ~ 5h)
- as cell elongates, observe gap between central portion of cell and matrix
- adhesion points at periphery of cell
- tensile forces in actin filaments induce compression in the matrix => buckling

Single Cell Contractile Force

- Contraction: cell buckling
- Measure E_s from AFM bending test
- Allows calculation of contractile force of single fibroblast

Single Cell Contractile Force



E_s = 762 MPa (dry)

$E_s = 5.28 \text{ MPa}$ (wet)

Source: Harley, B. A., et al. *Acta Biomaterialia* 3 (2007): 463-74. Courtesy of Elsevier. Used with permission. http://www.sciencedirect.com/science/article/pii/S1742706107000025

Harley, Silva

Single Cell Contractile Force

• Euler buckling:

$$F = \frac{n^2 \pi^2 E_s}{l^2}$$

$$I = \frac{\pi d^4}{64}$$

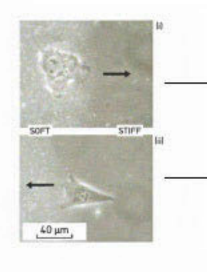
 $n^2 = 0.34$ (hydrostatic loading of tetrakaidecahedral cells (Triantafillou) $d = 3.9 + - 0.8 \mu m$; I from live cell imaging

$$F_c = 11$$
 to 41 nN (average 26 nN)

Harley, Wong

Cell Migration

Figure removed due to copyright restrictions. Figure 3: Cornwell, K. G., et al. Journal of Biomedical Material Research A 80 (2007): 362-71. http://onlinelibrary.wiley.com/doi/10.1002/jbm. a.30893/abstract



Source: Lo, et al., Biophysical Journal 79 (2000): 144-52.

Courtesy of Elsevier. Used with permission. http://www.sciencedirect.com/science/article/pii/S0006349500762795

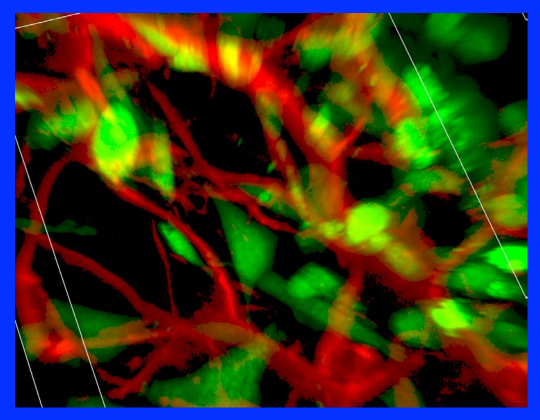
Top: Cornwell et al., 2007; Bottom: Lo et al, 2000

Migration speed on onedimensional fiber constructs

NIH 3T3 cells on 2D flat substrate: Cells on soft substrate cross to stiff substrate

Cells on stiff substrate will not cross onto soft substrate; instead spread out at boundary

Cell Migration: Fibroblasts in CG Scaffold



Courtesy of Brendan Harley. Used with permission.

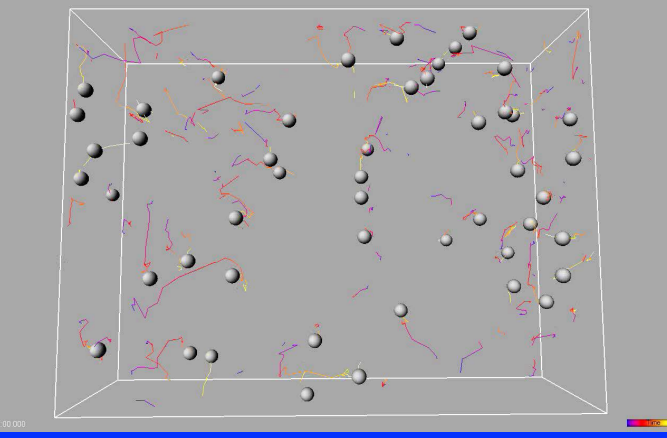
Confocal Microscopy

NR6 Fibroblasts CMFDA Live Cell Tracker

CG Scaffold Alexa Fluor 633 Stain

Harley

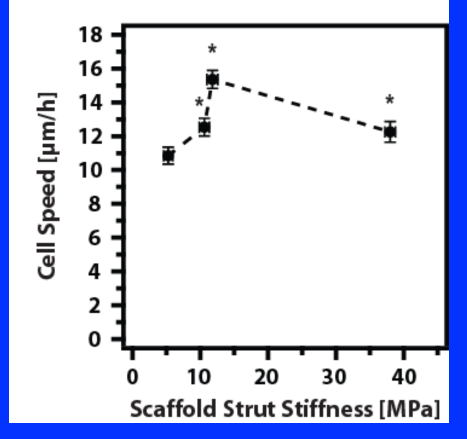
Fibroblast Migration: Spot Tracking



Courtesy of Brendan Harley. Used with permission.

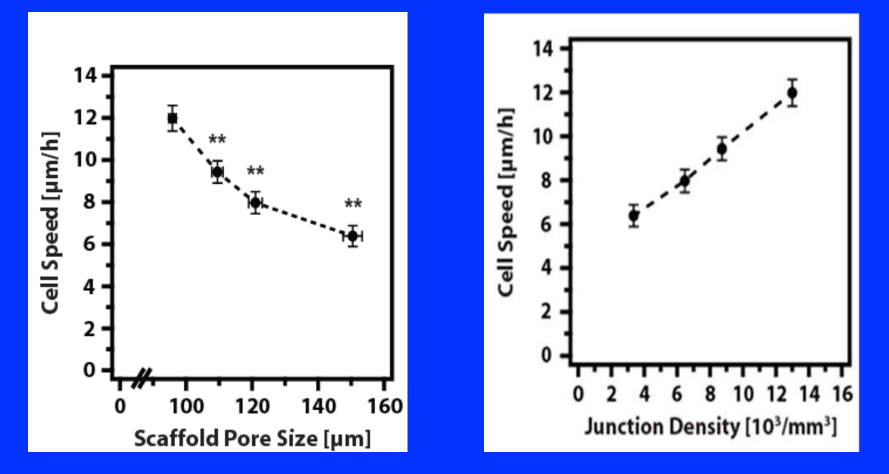


Migration Speed vs Strut Stiffness



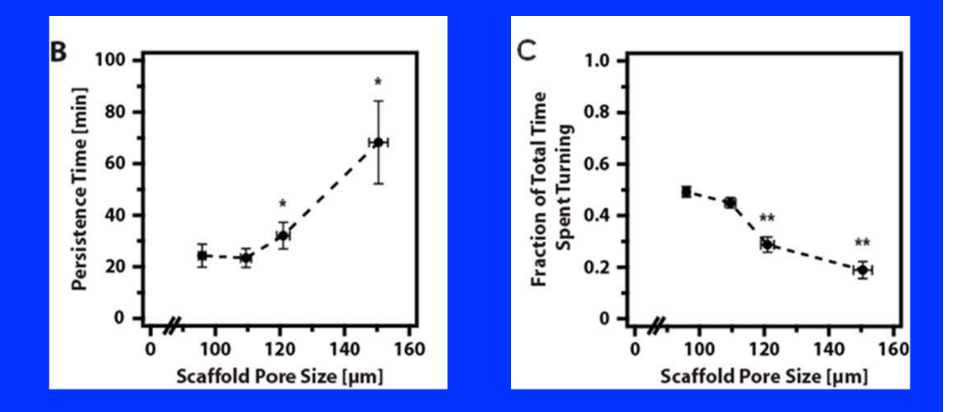
Source: Harley, B. A. C., et al. *Biophysical Journal* 95 (2008): 4013-24. Courtesy of Elsevier. Used with permission. http://www.sciencedirect.com/science/article/pii/S0006349508785394

Migration Speed vs Pore Size



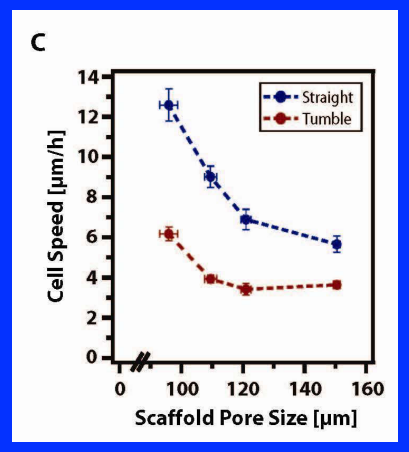
Source: Harley, B. A. C., et al. *Biophysical Journal* 95 (2008): 4013-24. Courtesy of Elsevier. Used with permission. http://www.sciencedirect.com/science/article/pii/S0006349508785394

Migration Speed vs Pore Size



Source: Harley, B. A. C., et al. *Biophysical Journal* 95 (2008): 4013-24. Courtesy of Elsevier. Used with permission. http://www.sciencedirect.com/science/article/pii/S0006349508785394

Migration Speed vs Pore Size



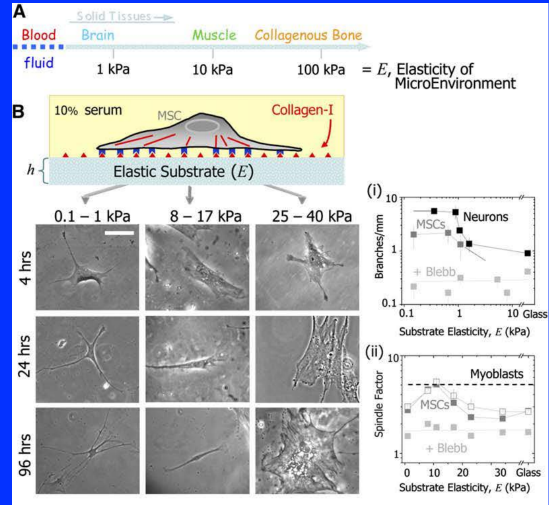
Cells on scaffolds with smaller pore sizes have a higher speed both along a strut and at a strut junction than cells in scaffolds with larger pores

As pore size decreases, specific surface area increases and # binding sites increases

Source: Harley, B. A. C., et al. *Biophysical Journal* 95 (2008): 4013-24. Courtesy of Elsevier. Used with permission.

http://www.sciencedirect.com/science/article/pii/S0006349508785394

Cell Differentiation

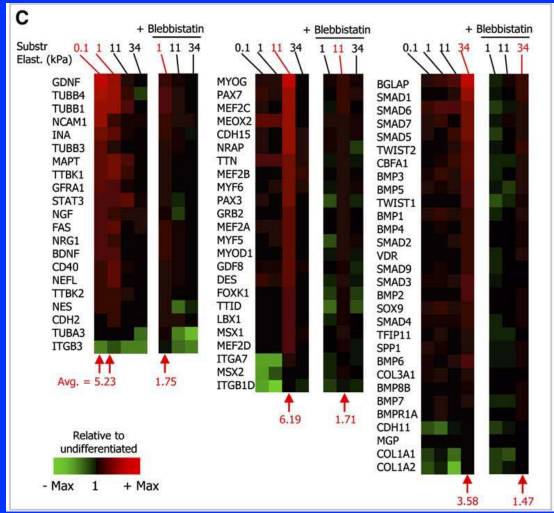


Engler et al., 2006

Neuron-like Myoblast-like Osteoblast-like

Source: Engler, A. J., et al. *Cell* 126 (2006): 677-89. Courtesy of Elsevier. Used with permission. http://www.sciencedirect.com/science/article/pii/S0092867406009615

Cell Differentiation



Source: Engler, A. J., et al. *Cell* 126 (2006): 677-89. Courtesy of Elsevier. Used with permission.

http://www.sciencedirect.com/science/article/pii/S0092867406009615

Engler et al, 2006

 Cell attachment increases linearly with specific surface area (binding sites)

 Cell morphology depends on orientation of pores in scaffold and on the stiffness of the scaffold

• Cell contractile behaviour:

- Cells bind at periphery of cells
- As they spread and elongate, unsupported length increases
- Compressive force in strut reaches buckling load
- For a population of cells in the cell force monitor, force per cell ~ 1nN
- Contractile force calculated from buckling of a strut by a single cell ~ 11-41 nN

- Cell migration speed increases with stiffness of 1D fibers
- Cells will not migrate from a stiff 2D substrate to a soft one
- In collagen-GAG scaffolds:
 - Cell migration speed increases at low scaffold stiffness and then decreases at higher scaffold stiffnesses
 - Cell migration speed increases at smaller pore sizes

Cell differentiation

- Mesenchymal stem cells differentiate to different morphologies, resembling different cell lineages (neuron, myoblast, osteoblast), depending on substrate stiffness
- Differentiated cells on substrates of different stiffness have cell markers associated with the different cell lineages (neurons, myoblasts, osteoblasts)

Acknowledgements

- Drs. TM Freyman, BA Harley, FJ O' Brien, M Zaman
- JH Leung, R Yokoo, Y-S Pek, MQ Wong, ECCM Silva, HD Kim, K Corin
- Profs. IV Yannas, D Lauffenburger, KJ Van Vliet
- Drs. Spector and Germaine
- NIH Training Grant, NIH grant (DE 13053), Matoula S. Salapatas Professorship, Cambridge-MIT Institute

3.054 / 3.36 Cellular Solids: Structure, Properties and Applications Spring 2014

For information about citing these materials or our Terms of Use, visit: http://ocw.mit.edu/terms.