



Oncotripsy: Targeting cancer cells selectively by means of tuned ultrasound

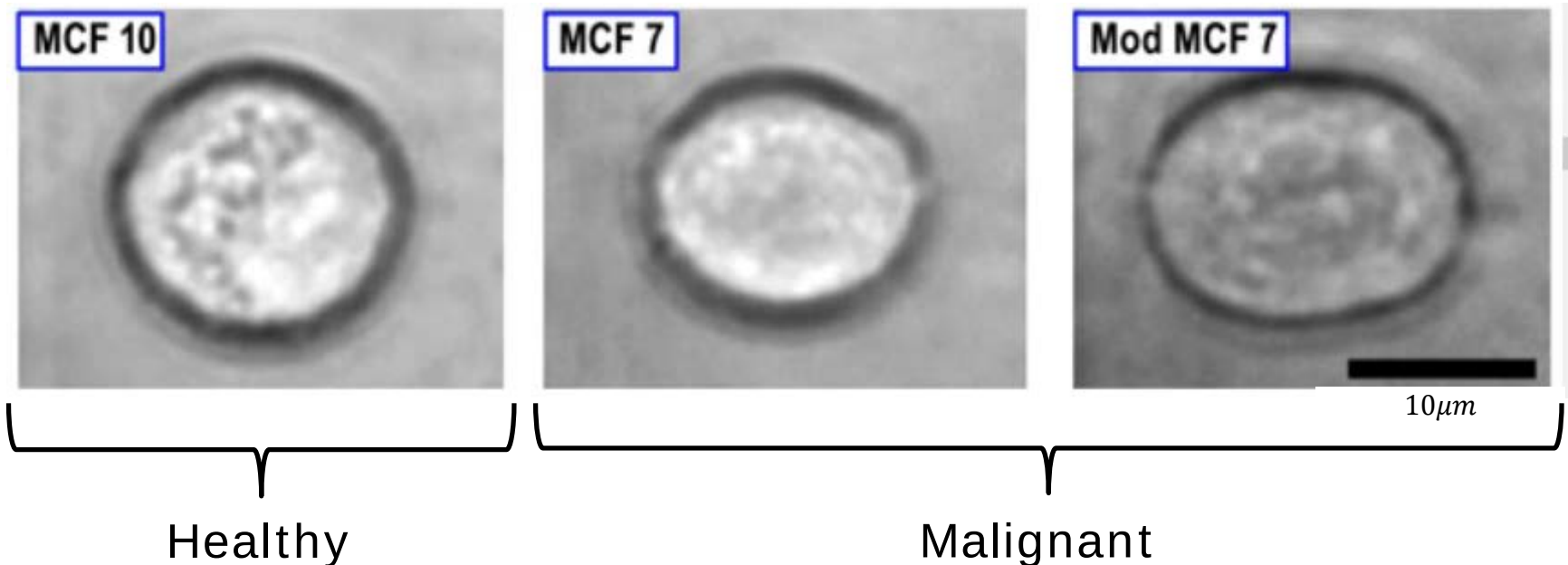
Michael Ortiz
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With: M. Gharib, D. Mittelstein, E.F. Schibber, M. Shapiro
(Caltech) and P. Lee, J. Ye (City of Hope)

Inaugural SEMTA Colloquium
Madrid, Spain, March 28, 2019

Oncotripsy: The key observation

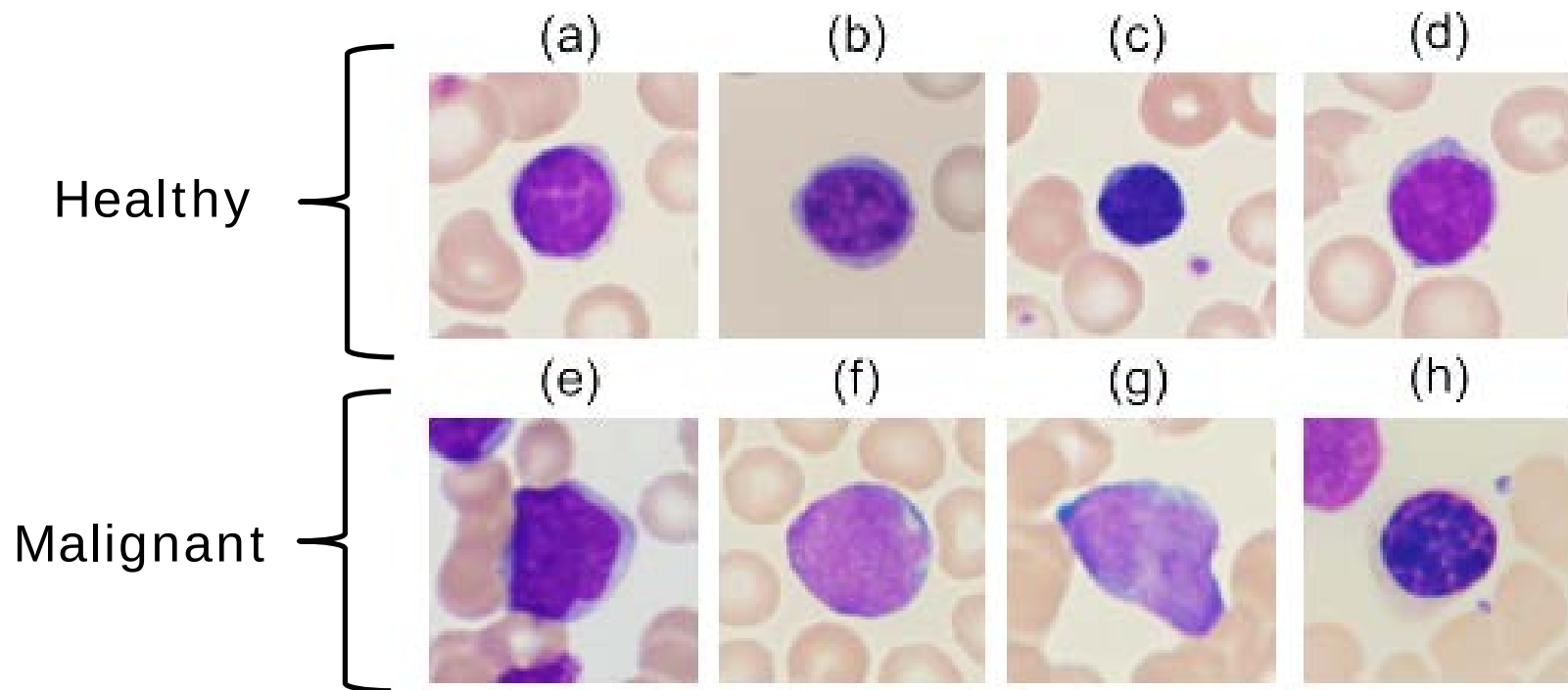
- Cancer cells have markedly different geometry, mechanical properties from healthy cells



Cells from MCF-7 breast cancer cell line
Morphological changes induced by malignancy

Oncotripsy: The key observation

- Cancer cells have markedly different geometry, mechanical properties from healthy cells



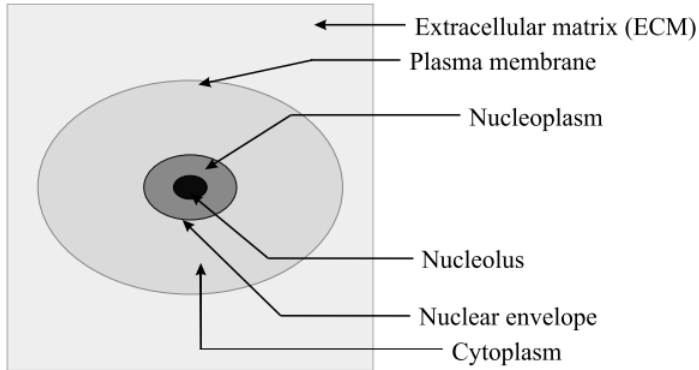
(a-d) Healthy lymphocyte cells

(e-h) Acute Lymphoblastic Leukemia (ALL) cells

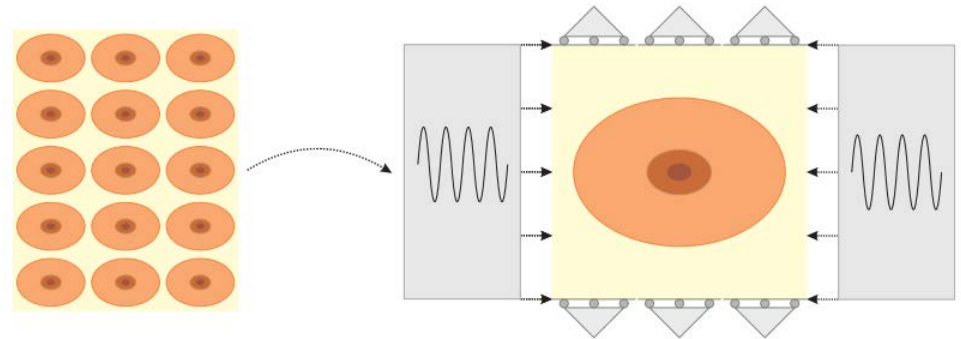
Oncotripsy: The key observation

- Numerous studies suggest that aberrations in both *cellular morphology* and *mechanical properties* of different cell constituents are typical of cancerous tissues
- Criterion for malignancy: *Size difference* between normal nuclei (average diameter of 7 to 9 microns) and malignant nuclei (can reach a diameter of over 50 microns)
- *Mechanical stiffness* of various cell components are found to vary significantly in healthy and diseased tissues [Berman, 2011]
- *Question*: Can cancer cells be *selectively targeted* by harmonic excitation at their resonance frequency? (*oncotripsy*) What are the *therapeutic ranges* of frequency, intensity?

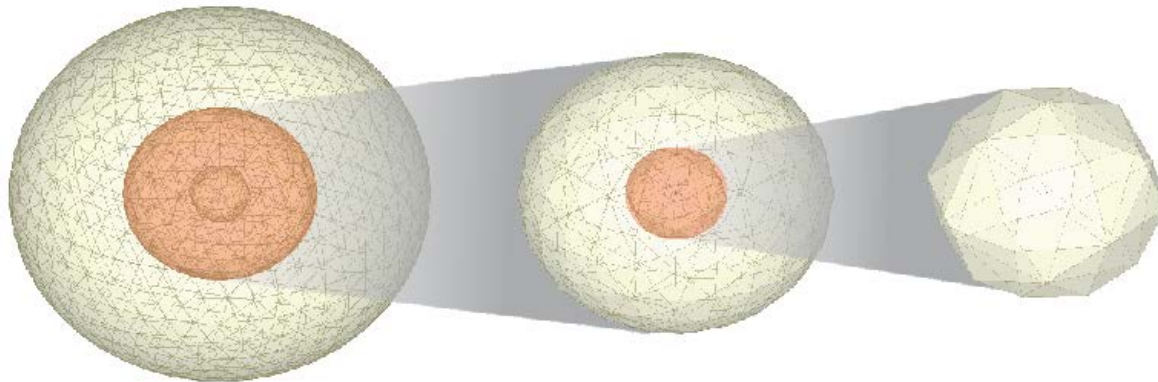
Oncotripsy: The spectral gap



Cell in extracellular matrix



Periodic (Bloch) model of tissue



Finite-element model of
cytoplasm, nucleoplasm and nucleolus

Heyden, S. and Ortiz, M., *JMPS*, **92**:164-175, 2016.

Heyden, S. and Ortiz, M., *CMAME*, **314**, 09 2016.

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Oncotripsy: The spectral gap

Hepatocellular Carcinoma (HCC)

Malignant	κ [kPa]	μ_1 [kPa]	μ_2 [kPa]
Plasma membrane	39.7333	0.41	0.422
Cytoplasm	39.7333	0.41	0.422
Nuclear envelope	239.989	2.41	2.422
Nucleoplasm	239.989	2.41	2.422
Nucleolus	719.967	7.23	7.266
ECM	248.333	5.0	5.0
Healthy	κ [kPa]	μ_1 [kPa]	μ_2 [kPa]
Plasma membrane	71.5199	0.738	0.7596
Cytoplasm	71.5199	0.738	0.7596
Nuclear envelope	431.98	4.338	4.3596
Nucleoplasm	431.98	4.338	4.3596
Nucleolus	1295.94	13.014	13.0788
ECM	198.666	4.0	4.0

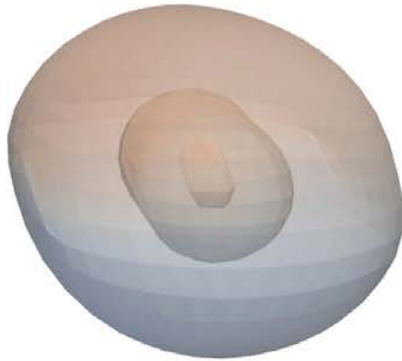
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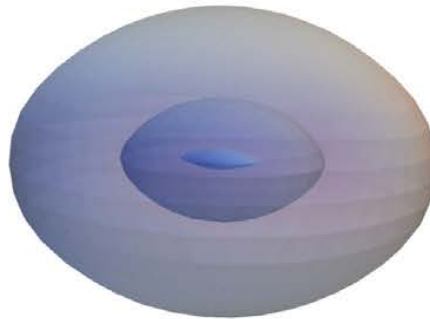
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Oncotripsy: The spectral gap

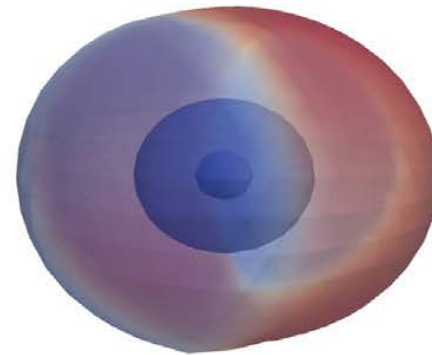
Hepatocellular Carcinoma (HCC)



$$\omega = 558031 \text{ rad/s}$$



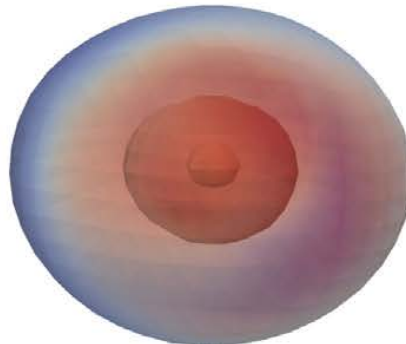
$$\omega = 576073 \text{ rad/s}$$



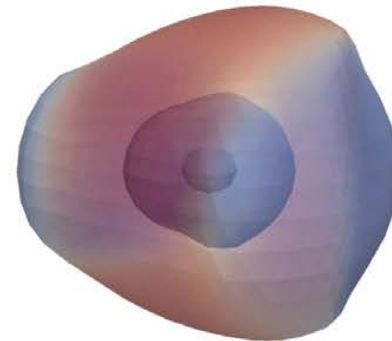
$$\omega = 816846 \text{ rad/s}$$



$$\omega = 849764 \text{ rad/s}$$



$$\omega = 979926 \text{ rad/s}$$



$$\omega = 991430 \text{ rad/s}$$

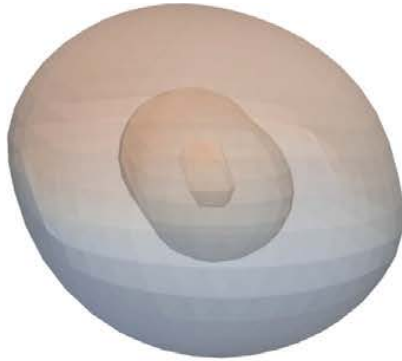
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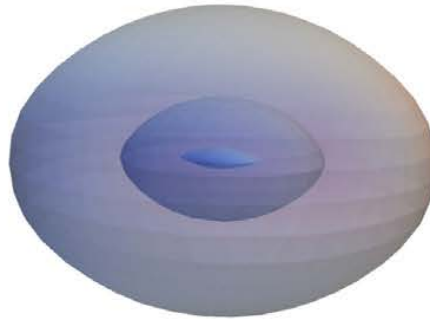
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Oncotripsy: The spectral gap

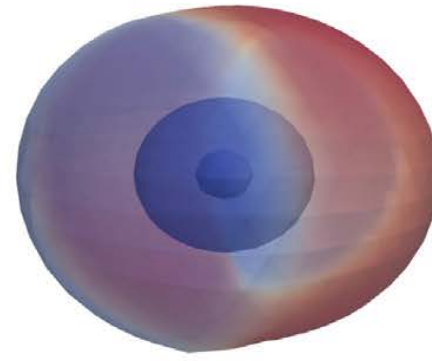
Hepatocellular Carcinoma (HCC)



$$\omega = 501576 \text{ rad/s}$$



$$\omega = 502250 \text{ rad/s}$$



$$\omega = 508795 \text{ rad/s}$$

	ω_1 [rad/s]	ω_2 [rad/s]	ω_3 [rad/s]	ω_4 [rad/s]	ω_5 [rad/s]
Cancerous	501576	502250	508795	532132	537569
Healthy	271764	274141	364259	364482	367413
	ω_6 [rad/s]	ω_7 [rad/s]	ω_8 [rad/s]	ω_9 [rad/s]	ω_{10} [rad/s]
Cancerous	538512	557291	667107	678287	678771
Healthy	375570	376000	380063	424226	425327

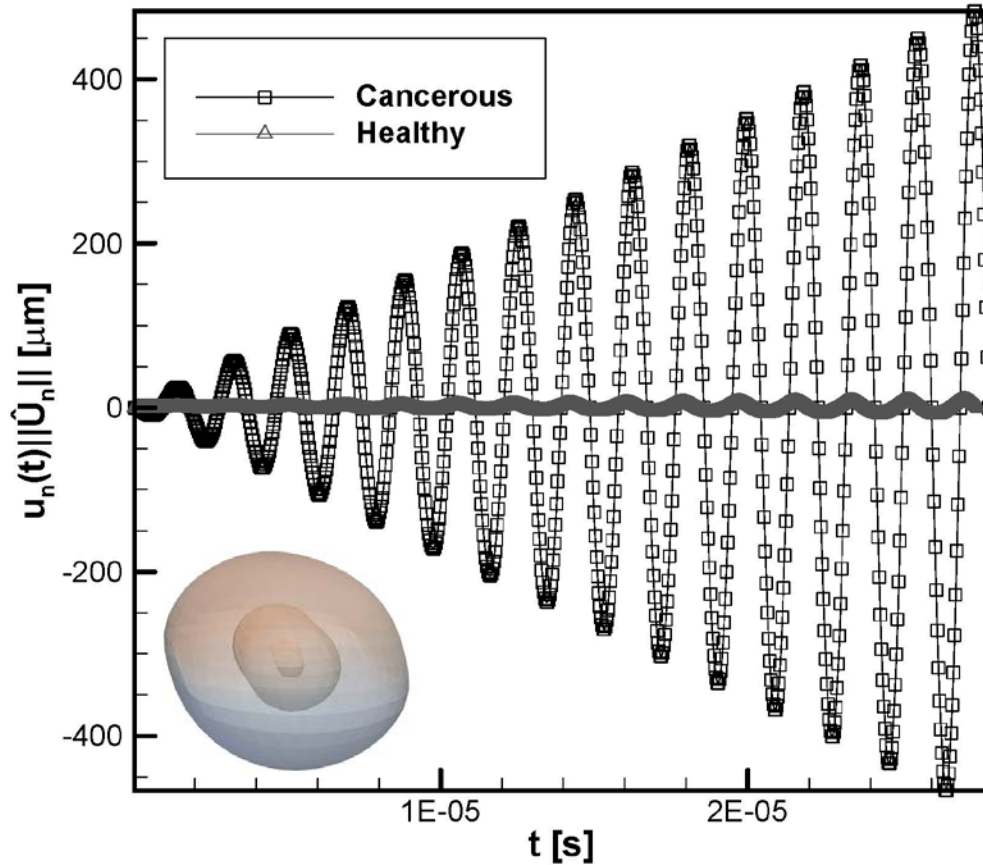
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Oncotripsy: The spectral gap

Hepatocellular Carcinoma (HCC)



Modal displacements of HCC and healthy cells excited at HCC resonant frequency showing vastly different growth rates: Malignant cells can be brought to lysis first!

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Heyden, S. and Ortiz, M., *CMAME*, **314**, 09 2016.

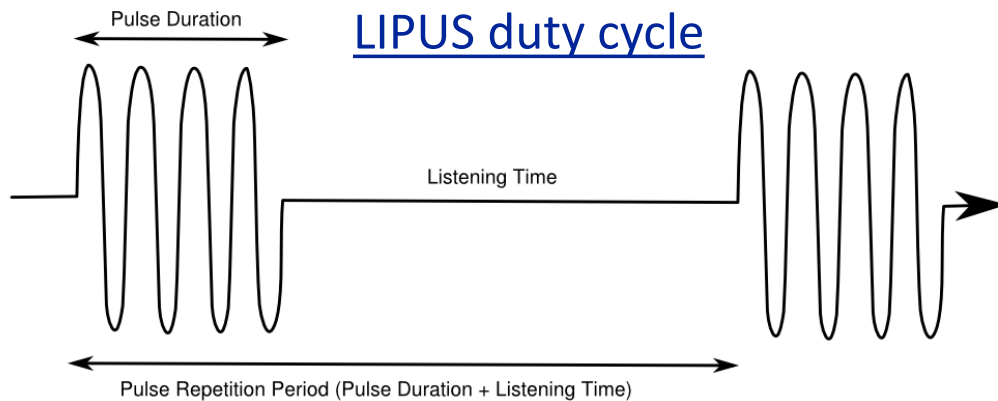
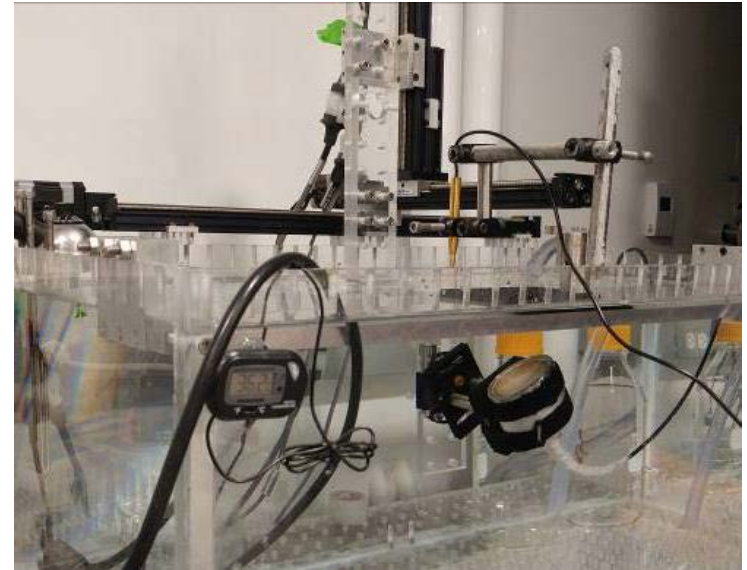
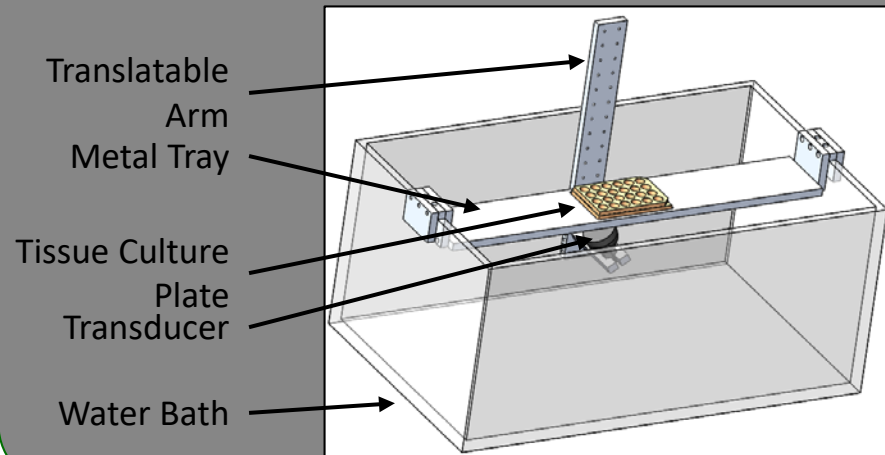
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Oncotripsy: The spectral gap

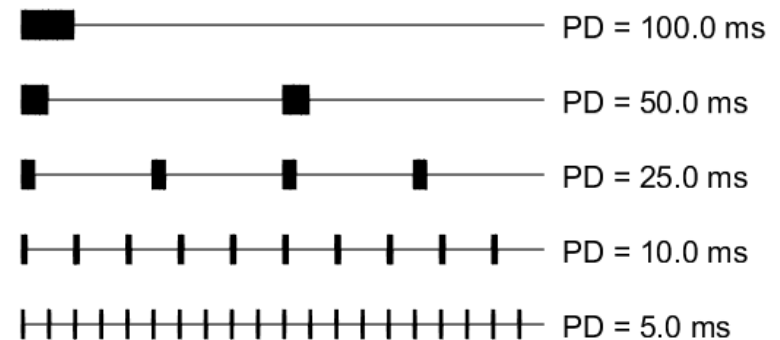
- Computational studies of HCC give natural frequencies of ~ 80 kHz (malignant) and ~ 43 kHz (healthy): *Ultrasound range*
- Spectral gap of ~ 37 kHz: Window for *selective targeting* of malignant cells (*oncotripsy*)
- Energy deposition rates ~ 1 W/m²: Low-intensity pulsed ultrasound (*LIPUS*)
- LIPUS is widely used in clinical applications. New non-invasive cancer therapies?
- *Is oncotripsy observed in the laboratory? (in vitro, in vivo, models, humans...)*

In vitro testing of cells in suspension

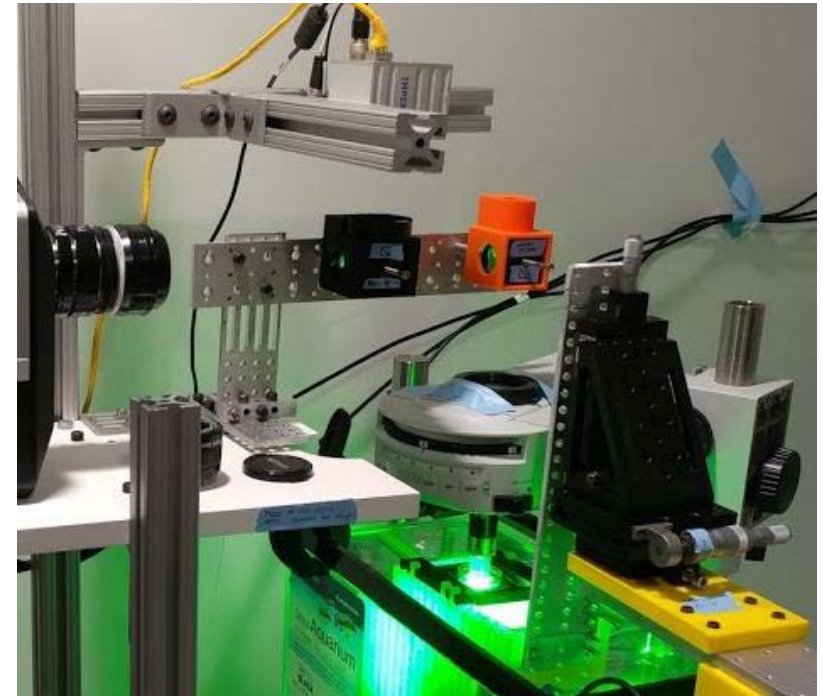
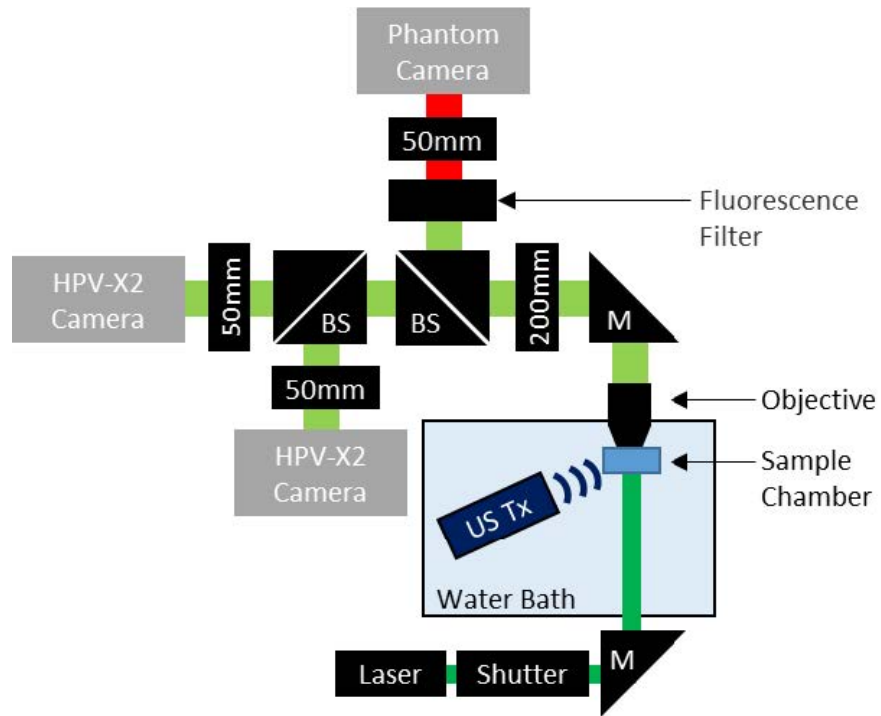
In-vitro experimental setup:



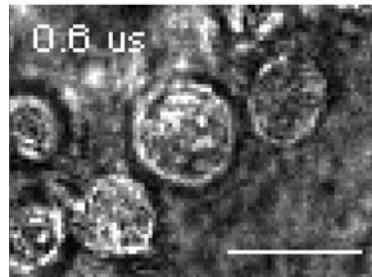
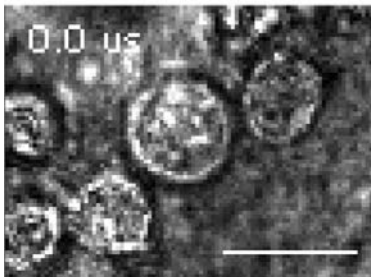
LIPUS duty cycle



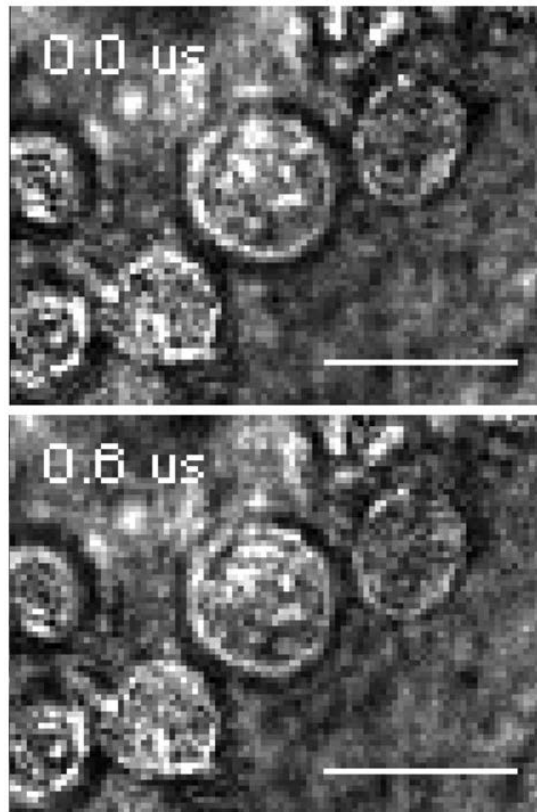
In vitro testing of cells in suspension



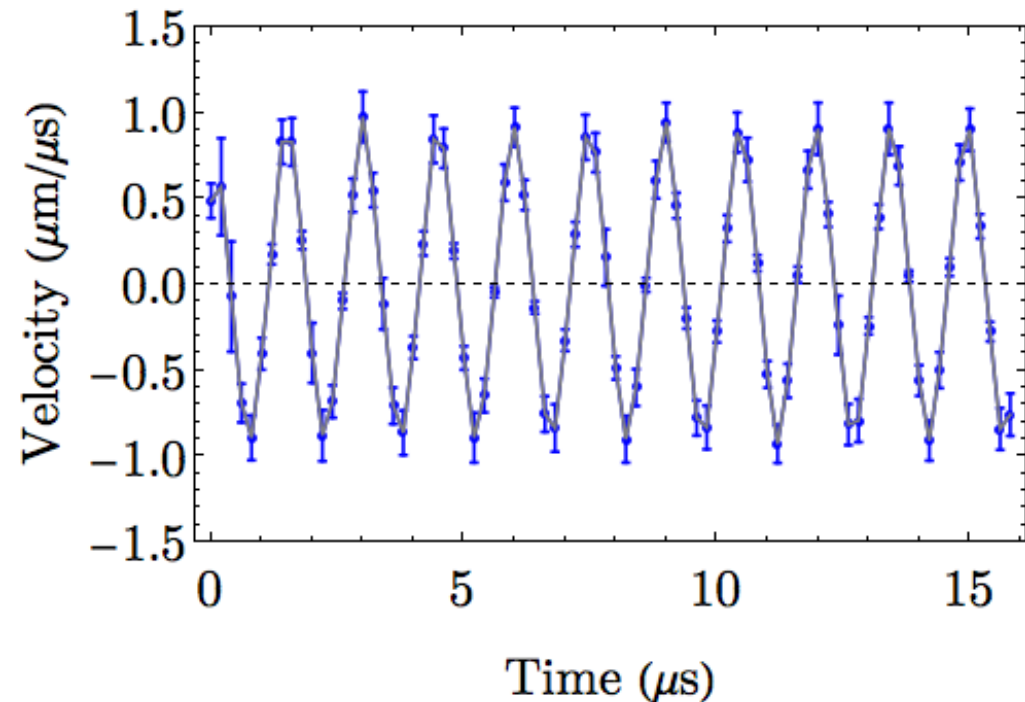
High-speed camera setup



In vitro testing of cells in suspension



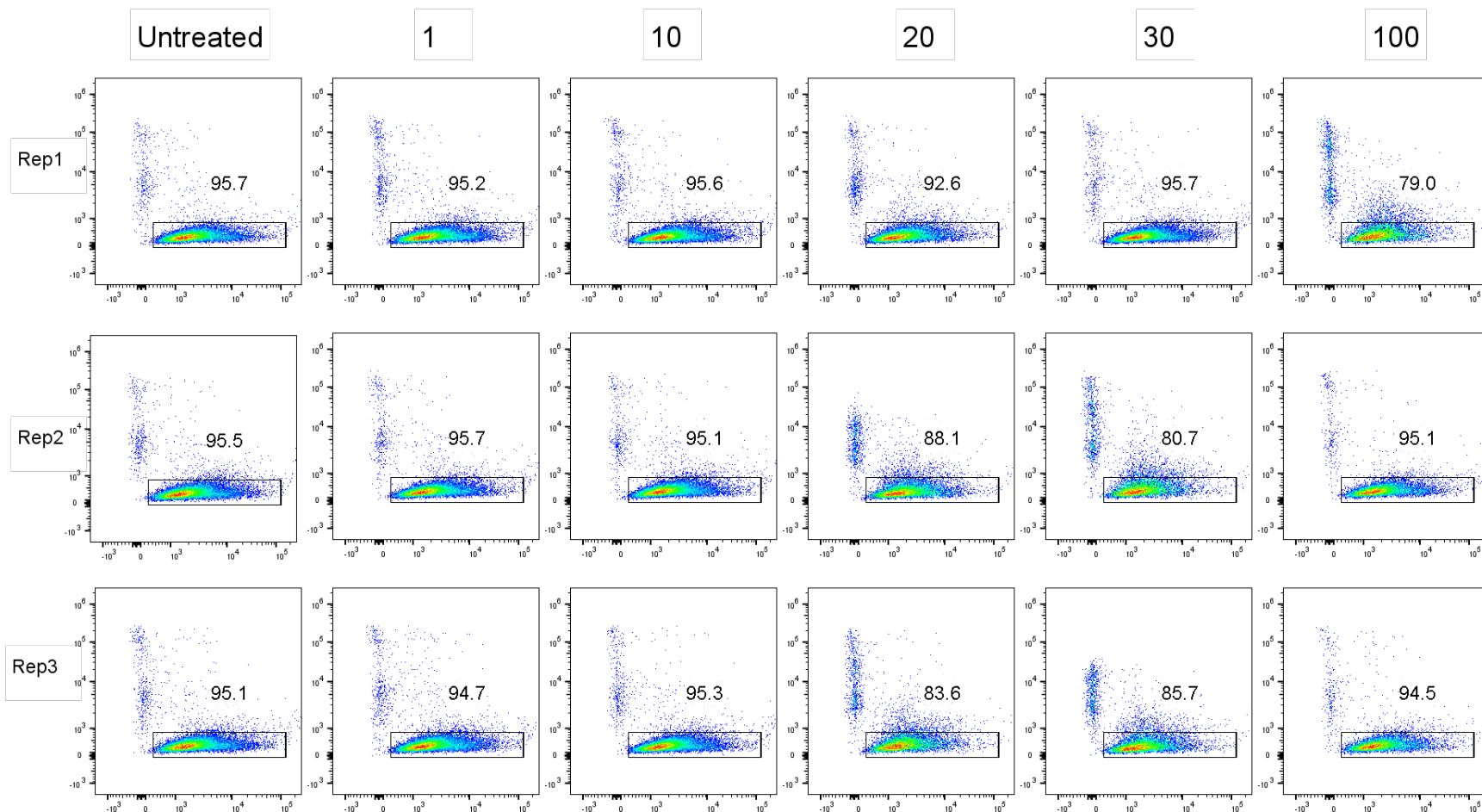
High frame-rate camera recordings showing minimal K-562 cell distortion after 100 ms of 670 kHz ultrasound exposure (scale bar 20 microns)



Data reduction from video showing nearly harmonic motion of the cell

Flow cytometry measurements

K-562 (leukemia bone marrow)

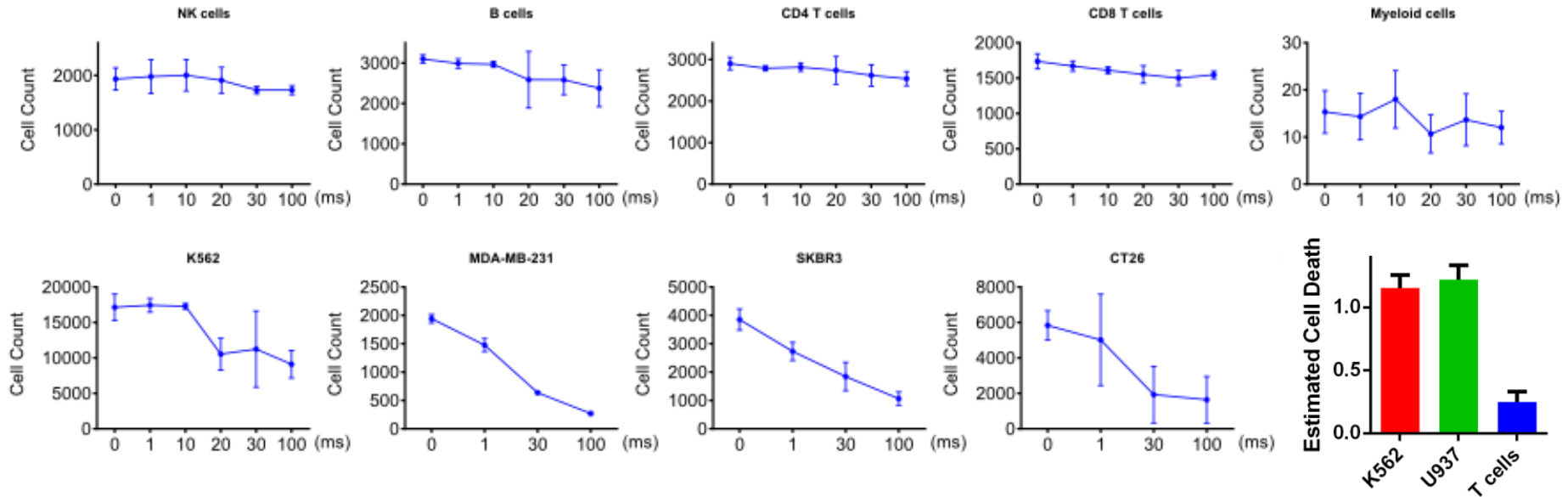


Double fluorescence dot plots from cytometry analysis
Dead-cell fractions as a function of exposure and duty cycle

Source: Lee, P. and Ye, J., *City of Hope*, 2019.

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Flow cytometry measurements

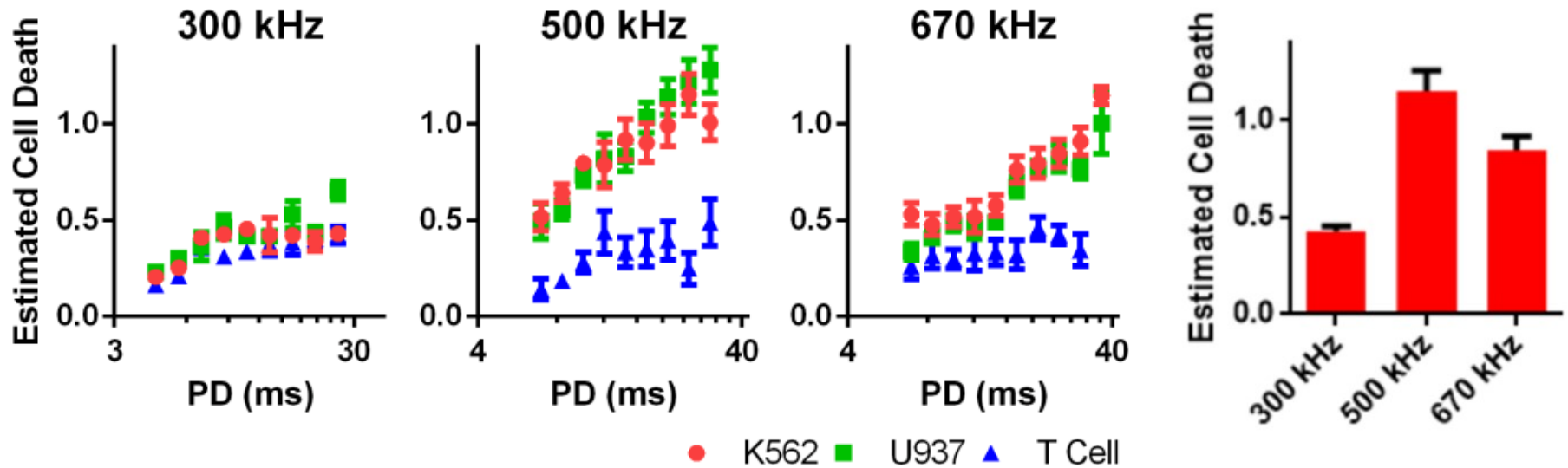


Cell counts at 500 kHz, 20 ms PD,
on healthy cell models (row 1) and cancer cell models (row 2),
demonstrate significant therapeutic index

Sources: D. Mittelstein *et al.* (2019) (manuscript),
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Flow cytometry measurements



Cell death patterns for different cell types
assessed in high-throughput screen
as a function of insonation frequency and pulse duration

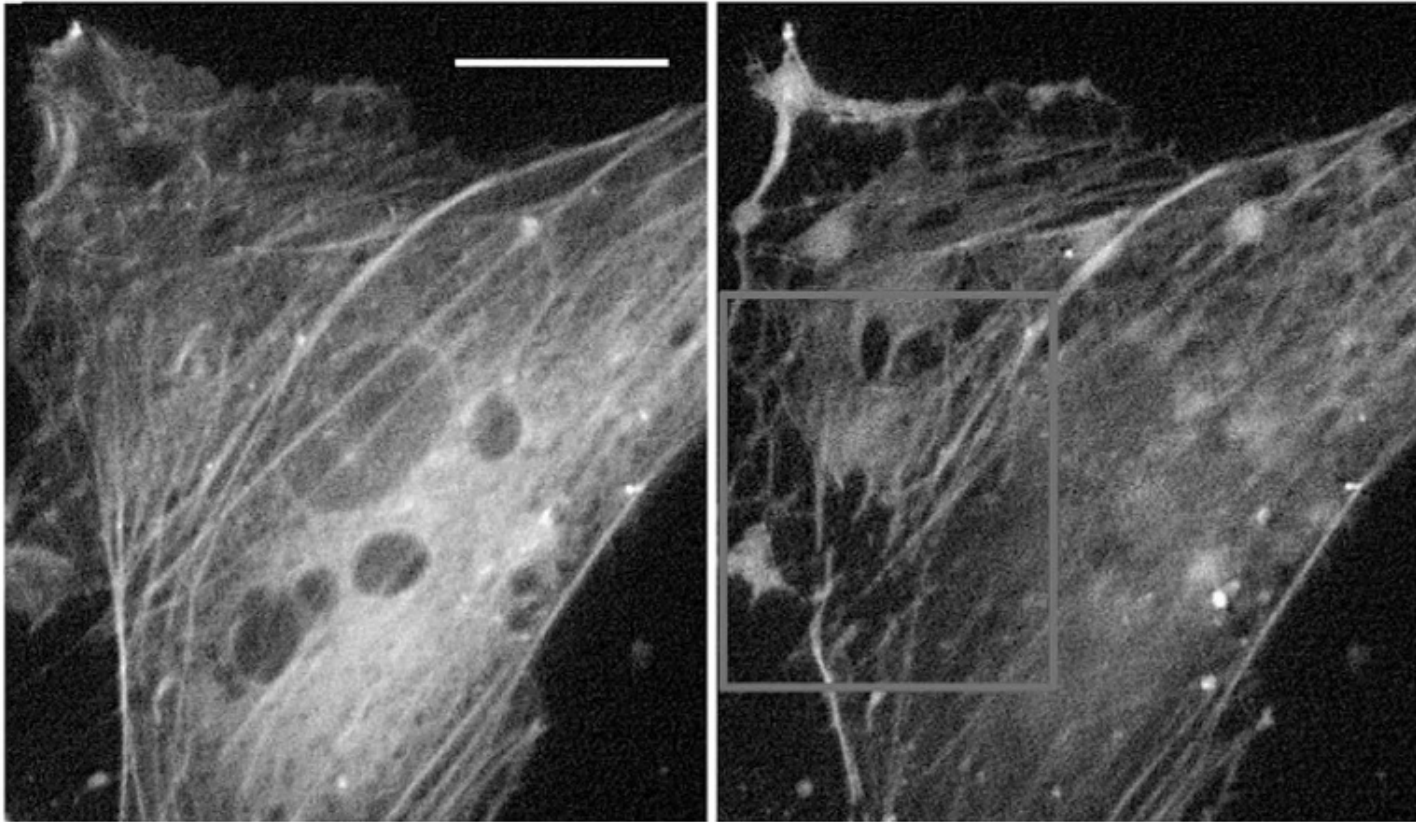
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In vitro testing of cells in suspension

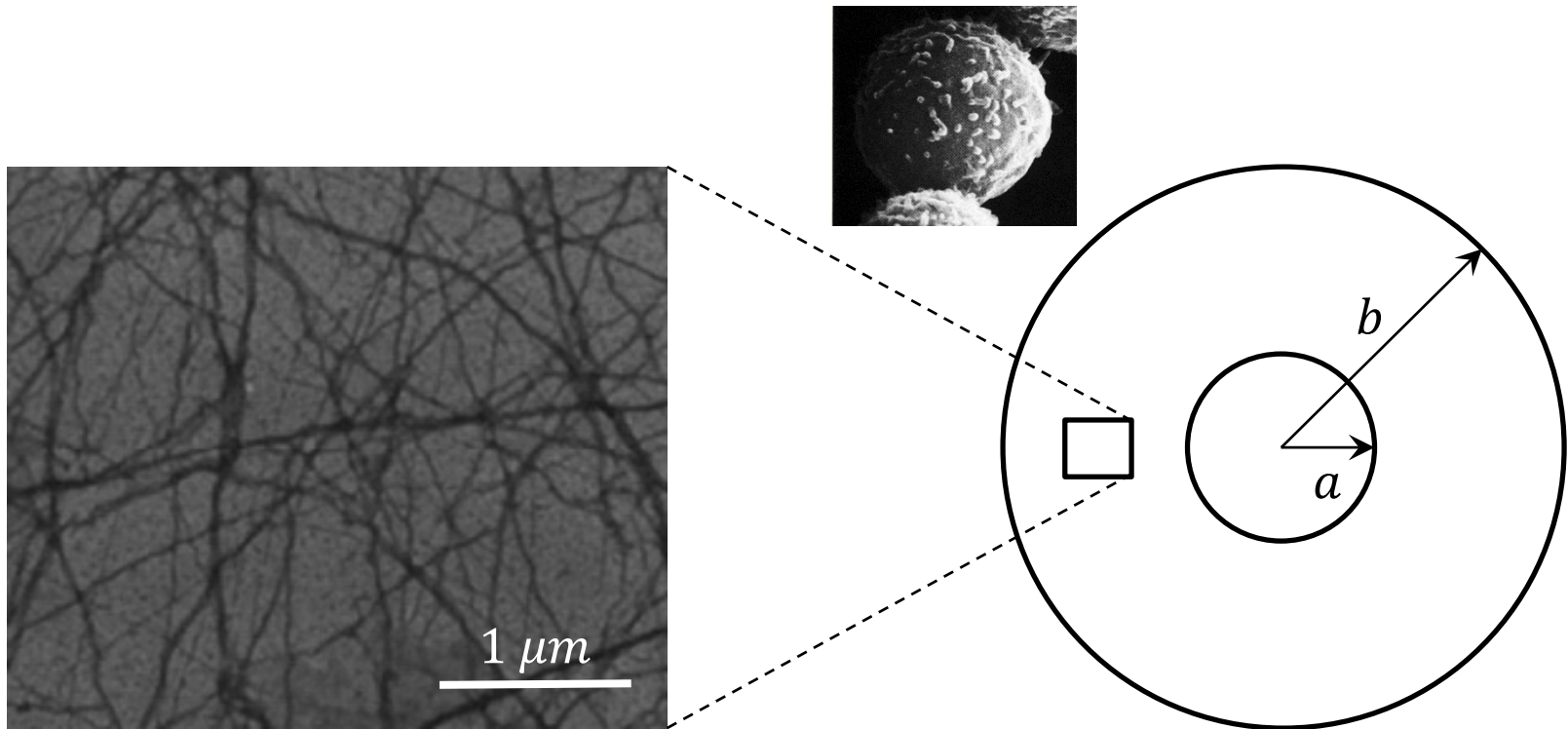
- Cell death in response to ultrasound exhibits *frequency-dependence, peak frequency*
- Targeted US induces *highly selective cell death*, demonstrating significant therapeutic index
- These observations bear out *oncotripsy*
- *But*: Cell death dependent on pulse duration, despite constant energy deposited
- Cell death requires a *large number of pulses*
- *Hypothesis*: Cells in aqueous suspension behave as *internal resonators*, die by slow accumulation of *damage to the cytoskeleton* (fatigue)

Response of actin network to US



Cell actin network subjected to high-intensity US, progressively disassembles within 3 min exposure (Mizrahi et al., 2012)

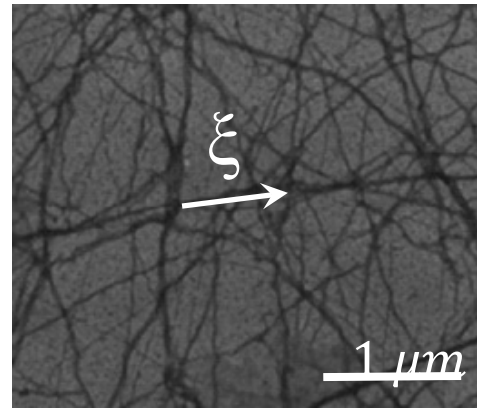
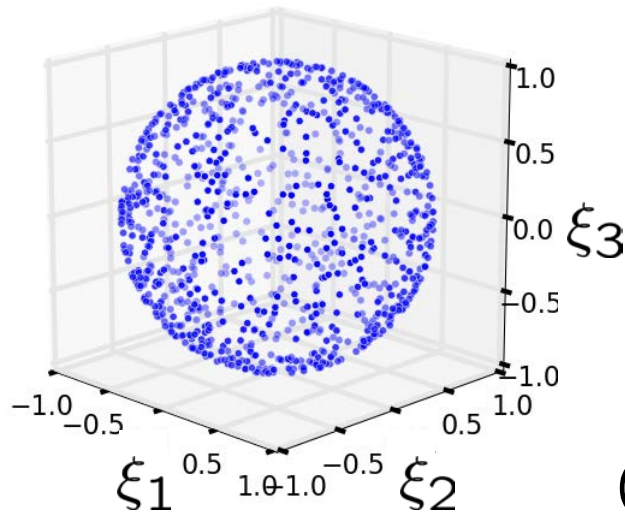
High-cycle fatigue model



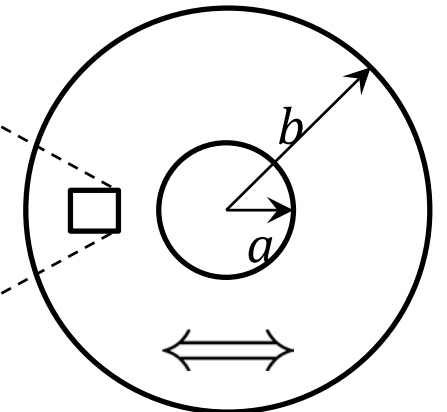
Electron micrograph of F-actin
(Gardel et al., 2008)

Globular cell moving with fluid particle velocity

High-cycle fatigue model



(Gardel *et al.*, 2008)

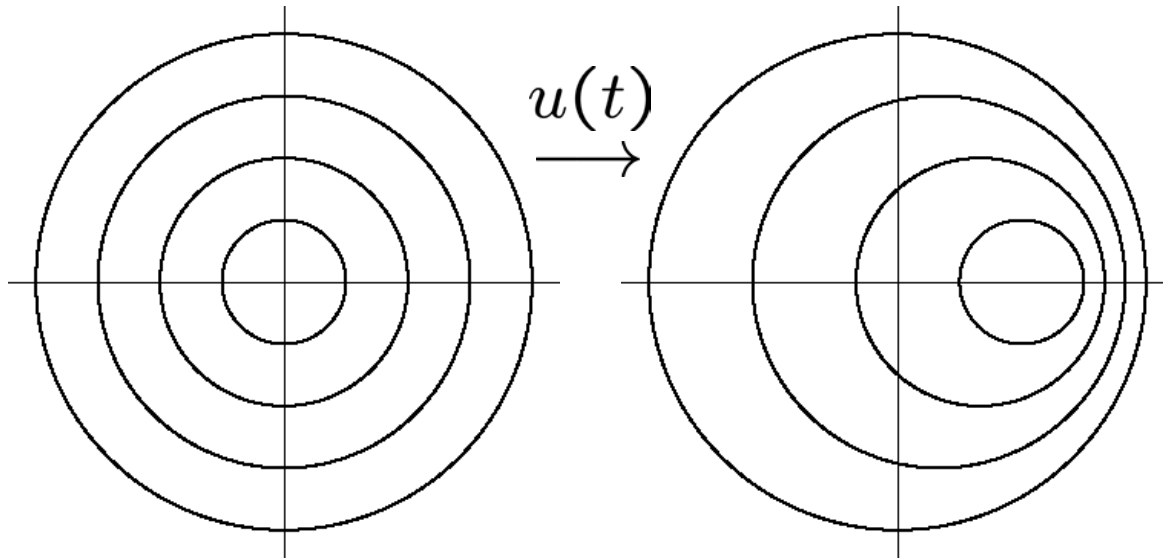


- Network theory of elasticity: $A(F, T, q) =$

$$\int_{S^2} p(\xi) \left(\frac{\mu(T)}{2} (1 - q(\xi))^2 (\lambda^2(\xi) + \lambda^{-2}(\xi)) + \frac{\beta}{2} q^2(\xi) \right) d\Omega$$

- Linear kinetics: $\alpha \dot{q}(\xi) + \frac{\partial A}{\partial q(\xi)} = 0$

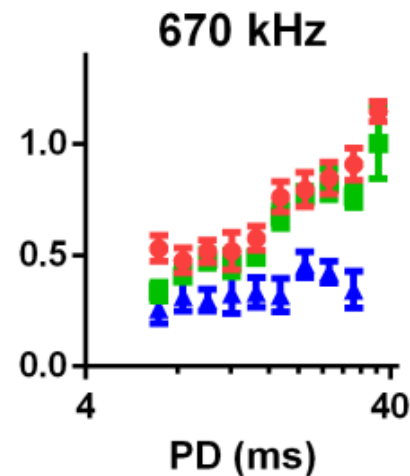
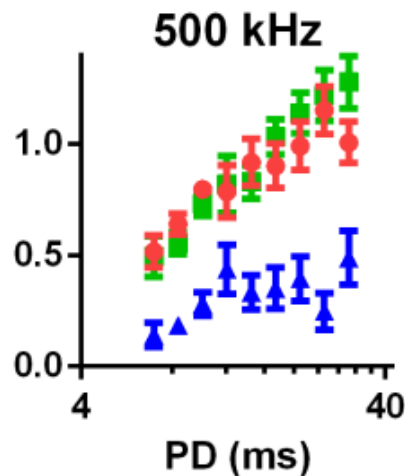
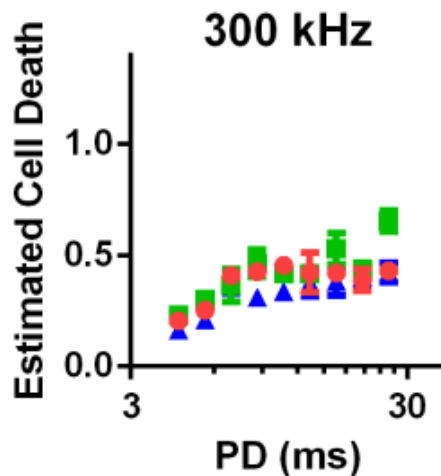
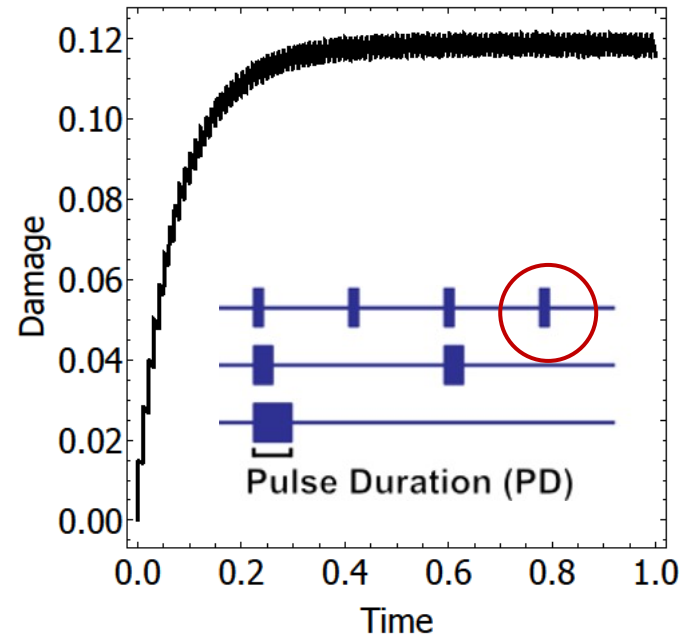
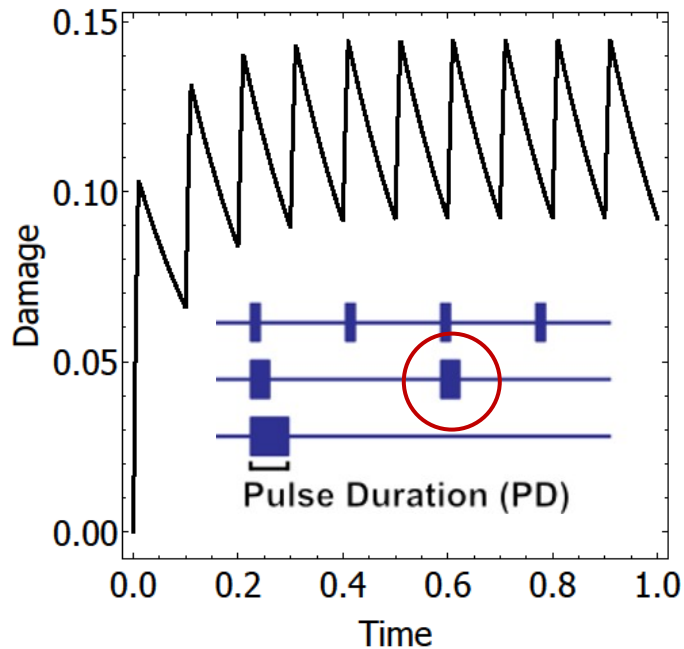
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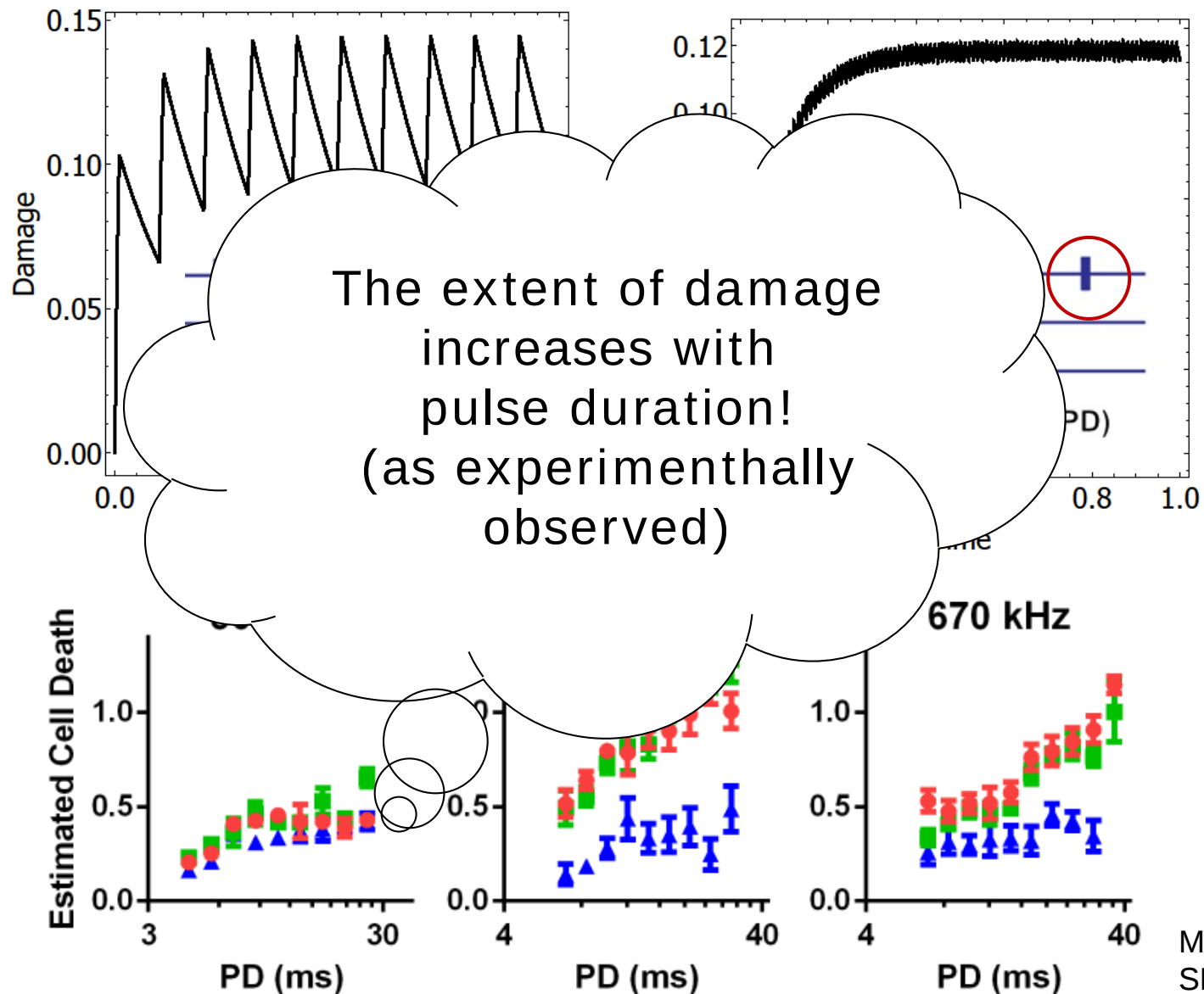
- Reduced (two d.o.f.) dynamical model:

$$\left. \begin{aligned} m\ddot{u}(t) + c\dot{u}(t) + (1 - q(t))^2 ku(t) &= -m\dot{v}(t) \\ \alpha\dot{q}(t) + \beta q(t) &= (1 - q(t))\frac{k}{2}u^2(t) \end{aligned} \right\}$$

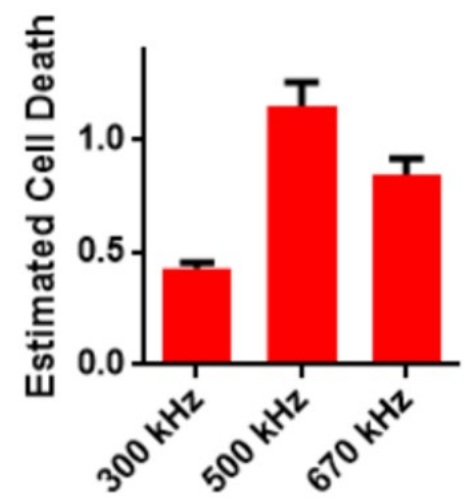
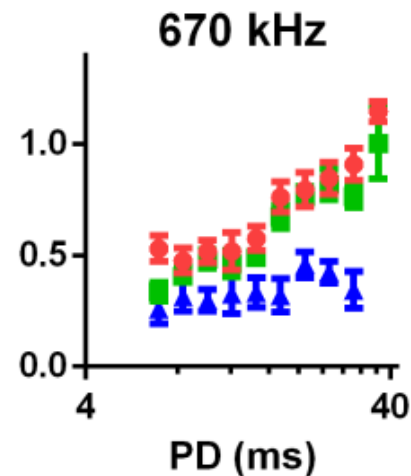
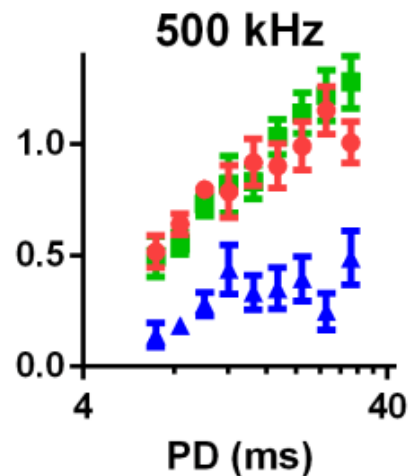
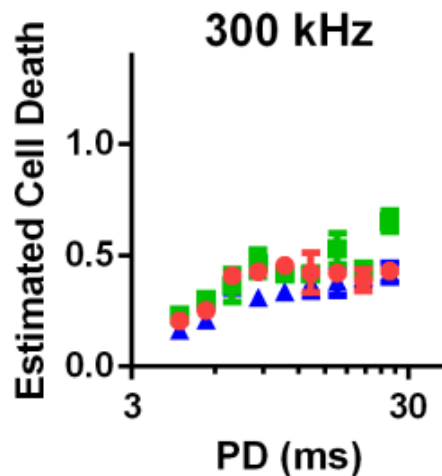
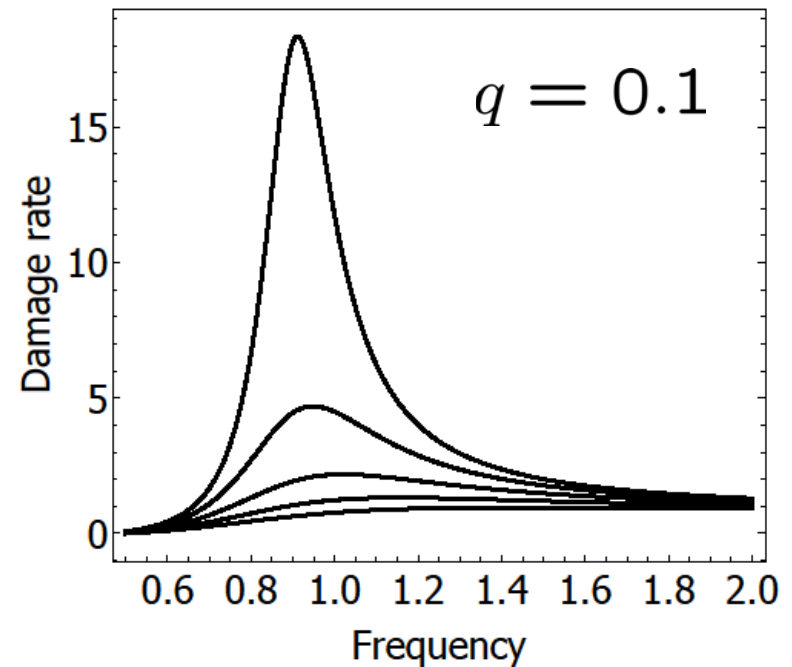
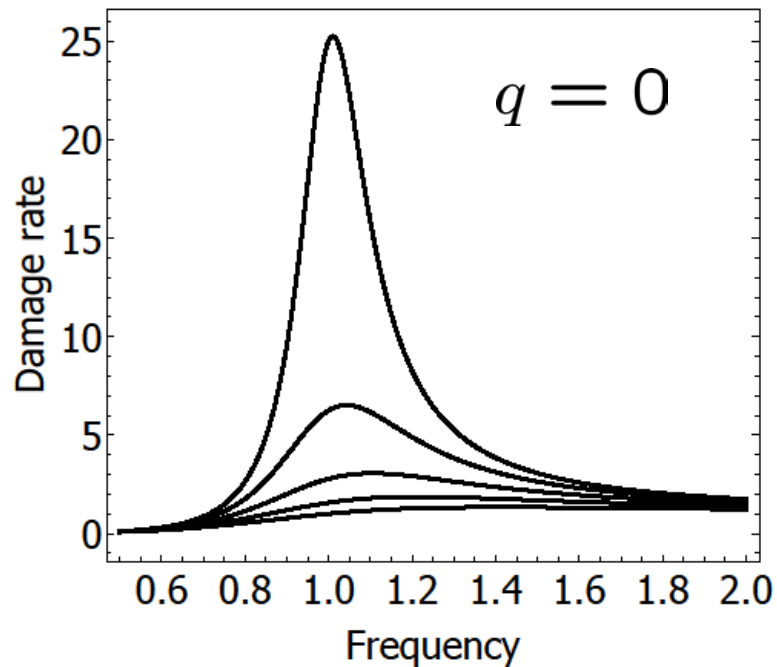
High-cycle fatigue model



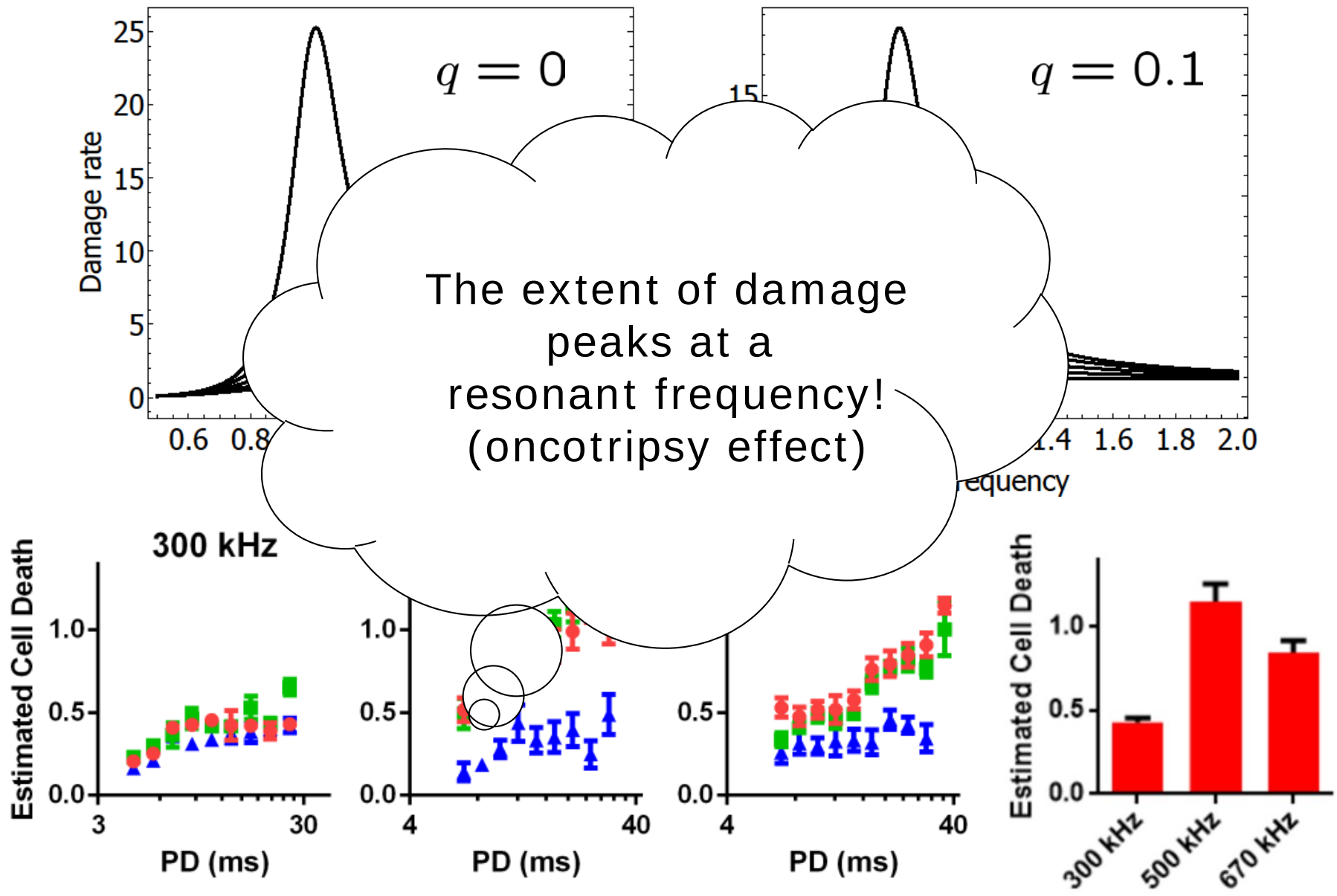
High-cycle fatigue model



High-cycle fatigue model



High-cycle fatigue model



Concluding remarks

- A simple dynamical model of *oncotripsy* captures and quantifies all the observed experimental trends pertaining to the response of cells in aqueous suspension to low-intensity pulsed ultrasound

Thank you!