

# Introduction of Gold Ion Solution into Living Cells: Uptake Mechanism and Applications



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## Introduction

•A better understanding of the relationship between living cells and nanotechnology is essential for successful clinical applications.

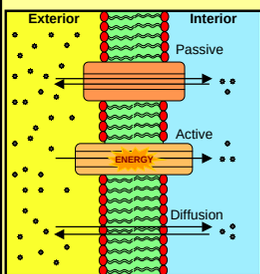
•By closely monitoring the response of a cell colony after the introduction and uptake of a gold ion solution, it may be possible to determine and predict the ideal conditions for a wide array of scientific techniques.

•Improvement in assorted imaging systems such as Raman spectroscopy

•Potent cancer treatment therapies.

•Advanced exploration of cellular biology.

## Fundamentals and Theory



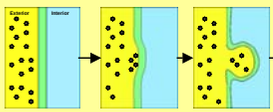
•The cell membrane controls the flow of nutrients in and out of the cell by using a number of different transportation pathways.

•Passive transport mediated by membrane proteins

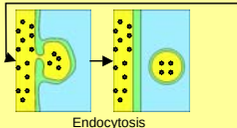
•Active transport (requires energy)

•Diffusion across selectively permeable membrane

•Endocytosis involves internalizing portions of the cell membrane, drawing external particles inside



•Gold ion solution uptake may occur through a combination of these pathways



## Questions

•Can living cells survive in gold ion solution, and, if so, what is the best combination of gold ion concentration and cellular medium for survival?

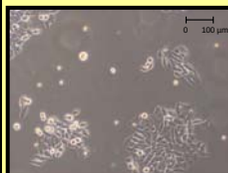
•Can gold ions be reduced inside the cell and form gold nanoparticles?

## Methods

•HeLa (human cervical cancer) cells were cultured in glass-bottom dishes using Dulbecco/Vogt Modified Eagle's Minimum Essential Medium (DMEM) for cellular nutrition, then incubated at 37 degrees Celsius and 5 percent CO<sub>2</sub> for approximately 48 hours.



Colony of HeLa cells and dishes



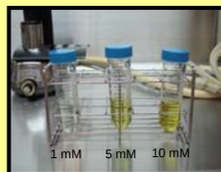
HeLa cells at approximately 30 percent confluency

•When dishes were 30-40 percent confluent (covered with cells), DMEM was exchanged with experimental media and gold (III) chloride (HAuCl<sub>4</sub>) was added.

•Media was either Phosphate Buffered Saline (PBS) with glucose, PBS without glucose, or DMEM without pH indicator (colorless).

•Gold (III) chloride solutions were in concentrations of 1 mM, 5 mM, or 10 mM.

•Control dishes without any gold (III) chloride were also prepared, to evaluate the effect of gold on cell viability.



Three different concentrations of gold (III) chloride solutions



Fluorescence Microscope

•After incubation for approximately 48 hours, dishes were monitored for 96 hours.

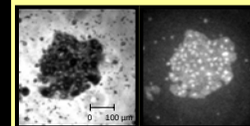
•Every 24 hours, cells were checked for viability (using propidium iodide (PI) indicator and fluorescence microscopy).

•In addition, time lapse photography was used in conjunction with dark field microscopy for periods of two or three hours to see whether gold uptake was visible over an extended period of time.

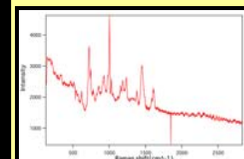
•Finally, after 48 hours of incubation with gold ion solution, Raman spectroscopy was used with the most successful results from above to confirm the formation of gold within the cells.



## Results



Confirming cell death by observing fluorescence during PI test



Raman spectra obtained from 1 mM HAuCl<sub>4</sub> in PBS without glucose

•In cases involving PBS with and without glucose, cells incubated with HAuCl<sub>4</sub> survived longer (approx. 24 hours) than controls.

•Although time lapse photography could not confirm the movement of HAuCl<sub>4</sub> into the cells, preliminary Raman spectra results suggest that gold ion solution was internalized by the cells incubated with 1 mM HAuCl<sub>4</sub> in PBS without glucose.

## Further Research and Applications

•The Laboratory for Scientific Instrumentation and Engineering plans to continue research on the effect that gold ion solution has on cellular viability.

•An expanded version of the trial will be conducted to facilitate easier and more accurate comparisons between experimental sets and controls.

•In addition, a wider variety of cell media and HAuCl<sub>4</sub> concentrations will be used to glean more information about the mechanisms behind experimental results.

•Transmission electron microscopy will be used to obtain additional images of gold particles within cells.

•Research may eventually result in an effective form of intracellular targeted photodynamic therapy.

## Acknowledgements

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