FREE ENERGY MEASUREMENT DISTINGUISHES NORMAL FROM CANCER CELL, OFFERING A NEW PERSPECTIVE FOR CURING CANCER

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ABSTRACT

The interplay of the second law of thermodynamics with the normal state and the methodology needed for the measurement of free energy of normal and malignant cells and its practical implications has not been clearly addressed in current literature. The second law of thermodynamics is one of the most fundamental laws governing the known universe at all levels. A normal cell has an exceptional ability to minimize the speed of rise in entropy to saturation of the limits of the second law. By virtue of this law, any normal resting cell is at a maximum allowable free energy. In this regard mitosis could be viewed as an attempt to maximize the lowered cellular free energy. Here we present the result of our first series of measurements, which show a significant measurable difference between the free energy of normal and malignant cells using an Olympus 510 Argon laser to calculate a diffusion correlation as well as direct visualization of motion of malignant and normal cells cultured overnight in collagen mesh. We found a significantly higher vibratory motion of the normal cells after correction for confounding factors. We also propose a new way to increase the free energy of the malignant cell to match that of its normal counterpart. This could offer hope for cure by conversion of distorted energetics of the malignant cell.

Keywords: Free Energy, Cancer Cell, Warburg Effect, Entropy and Cancer

1. INTRODUCTION

Warburg originally described the “Warburg effect” several decades ago (Warburg, 1956). He discovered that the glycolytic pathway is the main source of ATP generation of malignant cell (Zhou et al., 2012; Heiden et al., 2009). Warburg thought that the dysfunction of the enzymes of the Krebs cycle is responsible for the disconnection between the Krebs cycle and the electron transport system and decreased energy production (Bensinger and Christofk, 2012). However, he lacked proper measurements as well as an in-depth explanation. Consequently, a cancer cell has significantly fewer number of ATP molecules available to it at any time (Jain et al., 2012). Furthermore recent findings define the critical role of glycine in purine and pyrimidine synthesis in cancer cells (Jain et al., 2012; Hsu and Sabatini, 2008). It is thus clear that cancer cells can use alternate pathways for its metabolic needs. Survival and rapid division of cancer cells, given their low supply of energy, has been one of the biggest puzzles of cancer biology. To date there has not been any report on how to measure this difference in a way that could be replicated by others. In order to achieve this goal, we utilized Cho-K1 cells, a commercially available normal epithelial cell line and compared its motion to malignant HT29 cells, a microsatellite-stable sporadic colon cancer cell line. Previously we had also come to realize that an increase in the number of available ATP molecules in a normal cell relative to a cancer cell, would translate into an instantaneous increase in free energy of its constituents through convection.
This homogeneous intracellular level of high free energy is maintained because the closed environment of the cell guarded by lipid bilayer membrane would protect against the dissipation of the net energy of the cell to the outside environment. In other words, promotion to a higher free energy level happens following availability of more ATP molecules and is maintained at all compartments of the cell instantaneously as a unit. The thinking as of this writing has been that when energy in the form of ATP becomes available to a specific compartment of the cell and for a specific function, it is an isolated event and does not diffuse to its surroundings. A corollary of homogeneous intracellular energetics is that a decrease in free energy of cancer cell would also affect its constituents. Any change in the intracellular free energy would essentially translate into a conformational change of its constituents as well (Canchi and Garcia, 2013). This conformational change has a major bearing on functionality of cell (Wand et al., 2013). We hypothesized that the decrease in free energy of cancer cell could translate into a decrease in vibratory in situ motion, which we could measure and potentially modify.

2. MATERIALS AND METHODS

2.1. HT29 Spheroid Formation and Culture of Chok1 Cells

HT29 cells were grown in standard stem cell medium including B27, N2, FGF, EGF, non-essential amino acids, glutamine as well as heparin and penicillin + streptomycin. Spheroids, which are the hallmark of stem cells, were formed within several days.

Cho-K1 cells were obtained commercially and grown in conventional Cho-K1 medium in 10 cm culture dish at 37°C. Confluence was achieved within 2-3 days.

2.2. Collagen Preparation

Rat tail collagen type 1 was purchased from Bd Biosciences at 3.37 mg mL⁻¹. Using the standard protocol, collagen was prepared in an eight-well chamber slide.

2.3. Culturing HT29 Spheroids and Cho-K1 Cells in Collagen

Cho-K1 cells were scraped off the 10 cm dish and counted. 50,000 cells were separated and put in 300 mcL of Cho-K1 medium and added to the well containing collagen and allowed to grow over night. An equal volume of stem cell medium containing Ht29 spheroids was added to another well of the 8 well chamber slide and allowed to grow overnight.

2.4. Imaging

Using an Olympus 510 Argon laser, serial images were obtained from HT29 spheroids and Cho-K1 cells over a time span of 4-8 h. Their motion was recorded by using a t and z scale at wavelength of 488 nm with a pixel ratio of 256×256 and pinhole of 112 um. Data was analyzed by both direct visualization and by using a software program invented in the fluorescence live microscopy and Nano imaging division of UCI.

3. RESULTS

We noticed a significant increase in motion of Chok1 cells (Fig. 1), as compared with HT29 cells (Fig. 1).

Fig. 1. Cho-K1 cells (left) and HT29 spheroid (right) visualized using an argon laser at 488 nm
Computational analysis of the motion of Cho-K1 cells in collagen mesh demonstrating a greater diffusion rate (movement) designated by D of 0.0005, compared to its HT29 counterpart. The motion was rapid and jerky and mostly in situ. Using fluctuation spectroscopy, fluctuations in the fluorescence signal allows for the determination of diffusion between cells. Cho-K1 cells demonstrated a greater diffusion rate compared to its malignant HT29 counterpart, 0.0005 versus 0.00001 respectively (Fig. 2 and 3).

4. DISCUSSION

For the first time we have come to discover the physical representation of the free energy of normal and cancer cell and present the methodology to measure it in a reproducible way. Warburg originally discovered the decrease in energy supply of cancer cell several decades ago. However the inability to define and measure it in a quantitative and reproducible way had plagued the biomedical field until this writing. Warburg could not describe why components of Krebs cycle become dysfunctional in cancer cell. We now know that under the low free energy state of cancer cell the quaternary structures of enzymes, which decide about their functionality become perturbed. Restoration of the free energy of cancer cell to the level of its normal counterpart could potentially reverse the dysfunction of critical proteins. Such reversal could open the window of opportunity for potential cure of cancer. Without precise and reproducible measurement, such reversal is not possible. Our findings of a significant decrease in physical in situ motion of cancer cell using the t and z score, would allow for precise quantitative measurement of such difference with normal cell. This would open the way to design the methodology for closing the gap in free energy of cancer cell.
We propose Nano technology as a reasonable way for delivery of necessary modifying measures such as nanotubes with inherent vibratory potential to the site of action (Caraglia et al., 2012; Grossman and McNeil, 2012; Medhe et al., 2013). We also suggest that it is critically important to identify the exact location of the pathological lesion inside the cell. We think that the “master regulator complex” of the cancer cell is the potential source for the decrease in free energy as a result of molecular lesions that lead to an irreversible decrease in their free energy. Some examples in this regard are the MITF protein in melanocyte development (Li et al., 2012), the IkB kinase complex regulation of NF-kB (Abu-Amer, 2012) and the role of the miR-200 family in determining epithelial phenotype of cancer cells (Castilla et al., 2012). We also strongly believe that until the time that we come up with such delivery technology; cancer would continue to remain an incurable disease simply because none of the available and popular methodologies such as chemotherapy, radiotherapy, immunotherapy, small molecule tyrosine kinase inhibitors and even gene modifying and microRNA related therapies of today are addressing the fundamentals and deeply seated energetics perturbances of cancer cell.

5. CONCLUSION

Our discovery and findings open the way on a new and revolutionary approach to cancer therapeutics. By using our methodology through the use of Argon Laser microscopy for measurement of free energy of cancer cell and its normal counterpart, we could design a precise, reproducible and customized therapy for our cancer patients. Conversion of deeply seated aberrancies in bioenergetics of cancer cell would replace our current treatment modalities which are mostly based on destruction. Employment of Nanotechnology and epigenetics for fine tuning these aberrancies would make chemotherapy, radiation and immuno therapy something of the past.
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