Overcoming the Crabtree Effect is Critical to Mitochondrial Toxicity Testing

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Introduction

Potential drug toxicity testing has now been moved further upstream in the overall drug discovery process. This has been done in the hopes of limiting, or eliminating potential adverse toxicological effects or interactions. This translates to lower overall costs and risks to a manufacturer than if the adverse effects were discovered further downstream or after market launch.

Toxicity testing is subsequently performed on cell-based models, and even though primary cells are ideal for testing drug interactions, they are expensive and short-lived. Many researchers instead turn to less expensive, immortalized tumor-derived cells. The downside to using in vitro tumor-derived cells is that they may not be truly indicative of in vivo primary cell behavior in the presence of the drug compound.

For example, energy is normally produced by mitochondria in eukaryotic cells. These intracellular organisms use oxidative phosphorylation to create adenosine triphosphate (ATP), used for metabolism, signal transduction and other molecular processes. In this pathway, also known as an electron transport chain, five distinct protein complexes oxidize, reduce and interact with other intracellular biochemicals to create a proton gradient across the mitochondria’s inner membrane. This gradient is then used to form ATP from adenosine diphosphate, which the cell uses as energy.

In highly proliferative cells, such as many cancer cells, ATP is generated via glycolysis instead of oxidative phosphorylation. Although there are ten steps involved in converting glucose to pyruvate and releasing ATP, it is a simpler process than oxidative phosphorylation and does not require oxygen. Favoring glycolysis and reducing oxygen uptake is known as the Crabtree Effect. First described by Herbert Crabtree in 1926, this effect can significantly alter toxicology testing results as potential mitotoxicants have little effect on cell growth or viability.

For example, substituting glucose with galactose in culture will force the cancer cells to revert back to oxidative phosphorylation for ATP production, thus avoiding the Crabtree Effect and offering a truer indication of toxic potential in vivo.

In the following example, we use a multiplex assay to show how the Crabtree Effect can adversely affect cancer cell test results, and how to revert their ATP production back to oxidative phosphorylation.

The multiplex assay measures cell membrane integrity as a function of cytotoxicity, and mitochondrial function via ATP production concurrently, thus distinguishing between compounds that exhibit mitochondrial toxicity versus overt cytoxicity. General cytotoxicity is characterized by a decrease in ATP production and a loss of membrane integrity.
Methods

Cryopreserved hepatocytes were thawed and resuspended in InVitroGro HT Medium. The cultures were centrifuged, supernatant was removed, and the cells were resuspended in serum-free, glucose-free media consisting of No Glucose DMEM, 5 mM HEPES, 10 mM galactose, 2 mM glutamine, 1 mM Na-Pyruvate and 1% Penicillin/Streptomycin.

HepG2 cells were propagated in media consisting of High Glucose DMEM, 10% FBS and 1% Penicillin/Streptomycin. After removal from the growth flask, the cells were resuspended in the previously described serum-free, glucose-free medium.

All compounds were titrated in 100% DMSO to create an 11-point dose-response curve. The compounds were then diluted from the 1000X stock in non-serum/non-glucose medium to a final 2X concentration before addition to the assay plates.

Per each assay, 10 μL of hepatocytes or HepG2 cells were added to each well in a 384-well microplate. Compound, 10 μL, was added to each well, and the plates were placed in a 5% CO2/37°C incubator for 1, 2, 3, 4, or 6 hours. After incubation, 5 μL of 5X cytotoxicity reagent was added to each well, and the plate was incubated using the same environmental conditions for thirty minutes. The fluorescent signal was then read using the aforementioned Synergy H4 settings.

The microplate was then equilibrated to room temperature for approximately 15 minutes. 25 μL of ATP Detection Reagent was added to each well, and the plate was incubated for five minutes at room temperature. The resulting luminescent signal was then read on the Synergy H4.

Data Analysis

Luminescent or fluorescent values from wells containing media, treatments, and assay reagents were subtracted from raw values detected from cell containing wells. % Unstimulated Control was then computed using the following formula:

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\% \text{ Unstimulated Control} = \left( \frac{\text{Value(T)}}{\text{Avg Value(U)}} \right) \times 100
\]

Where Value(T) equals the background subtracted value from wells containing compound, and Avg Value(U) equals the average value from background subtracted basal wells containing no compound.
Results

Per Figure 3, ATP Detection Assay data, ATP production, as a function of the detected luminescent signal, substantially decreases in the hepatocyte cells in the presence of antimycin. The lack of significant change in cytotoxicity data proves that toxicity from antimycin is due to the compound impeding mitochondrial function, and not a result of primary necrosis. ATP production and cytotoxicity data are similar in hepatocytes grown with and without glucose.

ATP production in the HepG2 cells (Figure 4) reveals a sharp contrast between cells grown in glucose media and those in non-glucose media. Mainly, those HepG2 cells grown in non-glucose media were forced to revert to produce ATP via mitochondrial oxidative phosphorylation, and therefore behaved similarly to the hepatocyte cells, while HepG2 cells grown in glucose media produced ATP via glycolysis, therefore the known antimycin toxic properties on mitochondria were masked in a classic Crabtree Effect.

Summary

Because of their ease of use, and availability, highly proliferative tumor-derived immortalized cell lines have become the model of choice for mitochondrial toxicity studies. This has led to possible false conclusions regarding lead compound toxicity due to the cells propensity to derive energy from glycolysis rather than through oxidative phosphorylation in the mitochondria. In order to circumvent this phenomenon, and attain the correct data, it is necessary to force these cell lines to use mitochondrial generated ATP through the use of a non-glucose media. Here we have demonstrated how this effect can be monitored through the use of a multiplexed cytotoxicity and ATP detection assay. The homogeneous format allows for automation of the assay using simple, yet robust instrumentation. The results show how accurate titration data for cytotoxicity and ATP production can be attained, and the Crabtree Effect overcome, when using the correct media or cell model conditions.

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