Plasma Combination with Conventional Therapies for Cancer
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Abstract
Ambient gas plasmas are cytotoxic to cancer cells and bacteria at least in part because they create reactive oxygen and reactive nitrogen species (RONS). It is known that these reactive species can also play important roles in conventional therapies, including cancer chemotherapy and antibiotics. For example, anti-tumor synergy between NO-donating compounds and some redox-active chemotherapeutics has been documented [1]. We therefore explored the possibility that plasma-generated RONS could act synergistically against cancer and bacteria with conventional chemotherapeutic agents.

Motivation
• Anti-cancer therapies are generally delivered simultaneously, usually as drug cocktails. Synergies increase cocktail effectiveness.
• Combining therapies and taking advantages of synergies can prevent the development of resistance to chemotherapeutics.
• Simultaneous plasma treatment and chemotherapy may attenuate existing resistance to chemotherapeutics.

Background
Reactions of NO and O₂ produce RONS, causing oxidative and nitrosative stress in the cell. The effects of this stress include DNA strand breaks, and inhibition of DNA repair, including DNA ligation.

Sufficiently high concentrations of NO are known to be tumoricidal[2]. However, exposure to low doses of NO before treatment with chemotherapeutics such as doxorubicin and cisplatin increase the sensitivity of cancer cells to the chemotherapeutics. Notably, this effect has been observed for drugs known to have an RONS mechanism[1].

Materials and Methods
- Dielectric barrier discharge device
- Operated in indirect mode
- Operating conditions: 5 kV, 5 kHz
- Power: 0.15-0.2 W/cm²

Procedure:
• Plasma treat cells (MCF7 breast cancer line) drained of medium and add medium with doxorubicin 5 minutes after treatment
• Change medium after two days, incubate for three days more, and measure viability with Alamar blue fluorescence

Results
• Plasma treatment is fitted to exponential decay.

Conclusions
• For low concentrations of doxorubicin, plasma treatment and doxorubicin treatment are additive.
• More data are required to determine if synergy exists between ambient plasma treatment and doxorubicin, and if so, in what range of treatments.
• While studies on NO-donating compounds support the existence of synergy between doxorubicin and NO, our methodology may need to be changed to reveal it.
• Drug synergies are often cell-line specific, so testing multiple cell lines and multiple drugs may be needed to find plasma synergy.
• In particular, changing the order of plasma treatment and chemotherapy, and the plasma treatment conditions (power and gap height) may be essential.
• Similarly, glutathione depletion may allow for better resolution of any synergy, facilitating determination of statistical significance.

References

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