Cisplatin resistance reversal by disulfiram and caffeine

Sir,

Therapy resistance and relapse are the major issues associated with the current management of cancer. Cancer stem cells (CSCs) have inbuilt mechanisms to develop resistance. The most commonly overexpressed markers of CSCs are aldehyde dehydrogenase (ALDH) enzymes and ATP-binding cassette transporters (e.g., ABCG2). Moreover, cells have the capability to induce checkpoint responses when exposed to genotoxic agents (i.e., chemotherapy or radiation therapy), causing arrest of cells cycle. Resistance followed by chemotherapy administration as observed clinically can be divided into two phases. The first phase involves conversion of actively dividing cells to a completely dormant stage where cells do not divide though remain metabolically active. This stage can be defined as therapy-induced senescence. At this stage, chemotherapeutic agents fail to produce desired effect because cells are not dividing. The second phase is conversion of these dormant cells to actively dividing cells after longer duration. These cells may be even more resistant than parent cells and hence, require higher concentration of chemotherapeutic agents to arrest cell division. Disulfiram, an antabuse drug, is reported to inhibit ALDH, efflux pump, and some other potential cellular targets in cancer cells. Caffeine is reported to inhibit the checkpoint responses that are responsible for development of senescence. Hence, both disulfiram and caffeine simultaneously can be used to target CSCs and senescent cells, which are major contributors of therapy resistance.

A549 cell line was procured from the National Centre for Cell Science (Pune, India). It was cultured in Dulbecco’s modified Eagle’s media supplemented with 10% fetal bovine serum and antibiotics (100 µg/ml streptomycin and ampicillin). When the culture became 70–80% confluent, media were discarded and replaced with fresh media containing cisplatin and incubated for 3 days. After that, the media were replaced with cisplatin-free media, and survived cells were allowed for recovery for 4 days. These procedures were repeated with increasing concentrations of cisplatin (1.66, 3.33, 6.66, and...
After exposure to cisplatin, the cells appeared bigger (2–4 times), flattened, and elongated with bigger and clearly visible nuclei [Figure 1d]. Vacuoles were seen in the cytoplasm of some cells. The cisplatin IC$_{50}$ got almost doubled on A549-R cells. Both the curves presented in [Figure 1a] were significantly different ($P < 0.0001$) from each other. Both caffeine and disulfiram pretreatment reduced

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Table 1: Curve fitting analysis for different curves of Graph b and Graph c

| Graph b [A549] | Cisplatin + caffeine (14.31) | Cisplatin + disulfiram (1.43) | Cisplatin + disulfiram + caffeine (1.43) | Graph c [A549-R] | Cisplatin + caffeine (2.53) | Cisplatin + disulfiram (2.44) | Cisplatin + disulfiram + caffeine (1.13) |
|----------------|-------------------------------|-------------------------------|--------------------------------|-------------------|-----------------------------|-------------------------------|--------------------------------|
| Cisplatin (15.14) | NS                            | ****                          | ****                          | Cisplatin (30.09) | ****                        | ****                          | ****                          |
| Cisplatin + caffeine (14.31) | -                            | ****                          | ****                          | Cisplatin + disulfiram (2.53) | -                           | NS                            | ****                          |
| Cisplatin + disulfiram (1.43) | -                            | NS                            | -                             | Cisplatin + disulfiram (2.44) | -                           | -                             | ****                          |

****Significantly different curves ($P<0.0001$). NS=Not significant. The values in parentheses indicate IC$_{50}$ in µM.
the IC$_{50}$ of cisplatin in A549-R cells [Figure 1b]. Both when combined, a further reduction was observed [Figure 1c]. A similar study on A549 cells showed reduced IC$_{50}$ in only disulfiram pretreated cells. Further, how A549 and A549-R cells responded to given therapies can be depicted from graphs. The outcome of statistical treatment that was given to the different curves is shown in Table 1.

Both CSC-related markers expressing cells$^{[1]}$ and senescent cells$^{[4]}$ have been individually reported to be the major cause of resistance. Targeting both of them together can be expected to improve disease prognosis. The present study was conducted to determine benefits of such combined targeting. We observed a reduced toxicity of disulfiram in senescent cells than that observed in naive A549 cells. However, its pretreatment reversed the cisplatin resistance. Checkpoint responses can be inhibited by caffeine but at higher concentrations.$^{[7]}$ In this study, we used relatively lower concentration of caffeine to improve cisplatin cytotoxicity. Cisplatin efficacy was not altered by caffeine pretreatment to A549 cells. However, against senescent cells, it completely reversed the cisplatin resistance. The results indicated that caffeine might be acting by inhibiting the checkpoint responses in the senescent cells and forcing the cells forward in the cell cycle. The presence of cisplatin at this juncture becomes more meaningful in producing cytotoxicity. Exposure of A549-R cells to combination of caffeine and disulfiram caused reduction in IC$_{50}$ of cisplatin even more than that observed in naive A549 cells.

Thus, simultaneous use of caffeine and disulfiram as adjuvant therapy to chemotherapeutic agent is likely to improve the prognosis of the disease. However, a clinical study in this direction can throw more light.

Acknowledgment
Disulfiram powder, for the research purpose, was provided as a gift sample by Tripada Healthcare Pvt. Ltd., Ahmedabad, India.

Financial support and sponsorship
Nil

Conflicts of interest
There are no conflicts of interest.

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Received: 02-05-2016
Revised: 31-05-2016
Accepted: 15-07-2016

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