

POSTER PRESENTATION

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Targeting uroporphyrinogen decarboxylase for head and neck cancer treatment

Emma Ito^{1,2*}, Shijun Yue², Eduardo H Moriyama¹, Angela B Hui², Inki Kim², Wei Shi², Nehad M Alajez², Nirmal Bhogal², GuoHua Li³, Alessandro Datti^{4,5}, Aaron D Schimmer^{1,2}, Brian C Wilson¹, Peter P Liu³, Daniel Durocher⁴, Benjamin G Neel^{1,2}, Brian O'Sullivan^{6,7}, Bernard Cummings^{6,7}, Rob Bristow^{1,2,6,7}, Jeff Wrana⁴, Fei-Fei Liu^{1,2,6,7}

From São Paulo Advanced School of Comparative Oncology
Águas de São Pedro, Brazil. 30 September - 6 October 2012

Background

Head and neck cancer (HNC) is the 8th most common malignancy worldwide. Despite advances in therapeutic options over the last few decades, treatment toxicities and overall clinical outcomes have remained disappointing, underscoring a need to develop novel therapeutic approaches, particularly those that enhance tumor cell death, while minimizing damage to the surrounding normal tissues.

Materials and methods

An RNA interference (RNAi)-based high-throughput screen (HTS) was performed for the large-scale identification of novel genes that will selectively sensitize HNC cells to ionizing radiation. The Dharmacon Protein Kinase and Druggable Genome siRNA Libraries were screened using FaDu cells (human hypopharyngeal squamous cell cancer). Radiosensitizing targets were subjected to *in vitro* and *in vivo* characterizations.

Results

Sixty-seven target sequences with potential radiosensitizing effects were identified. Targets reducing the surviving fraction by >50% at 2 Gy relative to their un-irradiated counterparts were selected for further evaluation. A key regulator of heme biosynthesis, uroporphyrinogen decarboxylase (UROD), was thereby identified as a novel tumor-selective radiosensitizing target, demonstrating both *in vitro* and *in vivo* efficacy. Radiosensitization appeared to be mediated *via* enhancement of tumor

oxidative stress from perturbation of iron homeostasis and increased free radical production. UROD was significantly over-expressed in HNC patient biopsies, wherein lower pre-radiation mRNA levels correlated with improved survival, suggesting UROD could potentially predict radiation response. UROD down-regulation also radiosensitized several different human cancer models, while sparing normal cells.

Conclusions

An RNAi-based radiosensitizer HTS has revealed UROD as a potent tumor-selective sensitizer for radiation, with potential relevance to many human malignancies.

Financial support

Canadian Institutes of Health Research (CIHR; grant 69023); Elia Chair in Head and Neck Cancer Research; philanthropic support from Wharton Family, J. Finley, and G. Tozer; Campbell Family Institute for Cancer Research; Ministry of Health and Long-Term Planning; CIHR Resource Maintenance grant (PRG-82679).

Author details

¹Department of Medical Biophysics, University of Toronto, Toronto, Canada.
²Ontario Cancer Institute, Campbell Family Cancer Research Institute, University Health Network, Toronto, Canada. ³Toronto General Research Institute, University Health Network, Toronto, Canada. ⁴Samuel Lunenfeld Research Institute, Mount Sinai Hospital, Toronto, Canada. ⁵Department of Experimental Medicine and Biochemical Sciences, University of Perugia, Perugia, Italy. ⁶Department of Radiation Oncology, Princess Margaret Hospital, University Health Network, Toronto, Canada. ⁷Department of Radiation Oncology, University of Toronto, Toronto, Canada.

Published: 4 April 2013

* Correspondence: emma.ito@rmpuhn.on.ca

¹Department of Medical Biophysics, University of Toronto, Toronto, Canada
Full list of author information is available at the end of the article

doi:10.1186/1753-6561-7-S2-P19

Cite this article as: Ito et al.: Targeting uroporphyrinogen decarboxylase for head and neck cancer treatment. *BMC Proceedings* 2013 7(Suppl 2): P19.

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