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Okayama University research: Simplified boron compound may treat brain tumours

(Okayama, 16 September) **Effective and targeted uptake among malignant cells demonstrates the promise of a simplified boron compound for neutron radiation therapy, report researchers at Okayama University and Kinki University.**

“Glioblastoma multiforme is the most common malignant central nervous system primary tumour that has been incurable for decades,” explain Hideki Matsui and colleagues at Okayama University Graduate School of Medicine and Kinki University in their recent report. Treatment with ‘boron neutron capture therapy’ (BNCT) has been found to double the duration of patient survival, but finding effective compounds for it has been challenging. Now Michiue and colleagues have demonstrated - both in cell lines and mice - a readily made boron compound that targets and penetrates tumour cells for effective BNCT [1].

The compound BSH is high in boron content and has the advantage of accumulating primarily in tumour tissue rather than healthy tissue. However BSH does not penetrate inside the tumour cells limiting the therapeutic effects of BNCT using it.

Previously Michiue and a collaboration of researchers in Japan demonstrated they could enhance cell penetration by fusing BSH with a large peptide comprising up to 11 amino acid ‘arginine’ groups [2]. However with the large added peptide it was difficult both to produce the compound and track its effects in the body.

Studies of protein transcription aided by a smaller version of the peptide suggested that a compromise in size may allow effective BSH cell penetration without the need for prohibitively complicated synthesis techniques. [3]. The latest results from Michiue and colleagues have now shown that while peptides with just one or two arginine groups were ineffective, three arginine groups were enough to improve cell uptake.

The BSH fused with a peptide of three arginine groups was simpler to produce and they could also track the drug in the body using positron emission tomography. They conclude, “In our present study, we discovered that BSH-3R overcame many obstacles to develop a BSH-peptide suitable for future clinical use.”
Background

Boron neutron capture therapy
One of the challenges with treating glioblastoma multiforme is removing all cancer cells without causing damage to the brain. The destruction of nearby healthy cells is a significant side effect for traditional radiation therapy.

BCNT has attracted particular interest because the radiation caused is highly localised. When boron is delivered to malignant cancer cells, irradiation with low-energy thermal neutrons then triggers boron fission into a high energy alpha particle and lithium atom. While lethal to cells in their direct path the high-energy alpha particle and lithium atom produced cannot travel further than roughly a cell length before their energy is absorbed leaving nearby healthy cells intact.

BNCT has already been used clinically for treating brain tumours, malignant melanoma, head and neck cancers, and hepatoma. Hospital sources for low-thermal-energy neutrons to trigger the fission are in continued development. However clinical use so far has highlighted shortcomings in the boron delivery drugs currently available.

Boron delivery drugs
Previous clinically used drugs for BNCT include an essential amino acid analog p-borono-Lphenylalanine (BPA) and disodium mercaptoundecahydrododecaborate ([B12HnSH]^{2-2Na^+}, BSH). BPA readily delivers high boron content to cells but it does not target malignant cells alone and is not taken up by slowly proliferating malignant cells.

In contrast accumulation of BSH is greatly enhanced in tumour regions but it is present between the cells making it less effective for BNCT. Simulations in previous work have highlighted the limitations of BNCT from outside the cell.

Arginine
Arginine is a type of amino acid. Previous work has shown that with an 11-arginine-strong peptide the cancer killing effect of the BSH compound was greater than with a dose 100 times stronger of BSH without the added peptide. However it is difficult to synthesise BSH with the 11-arginine peptide and it cannot be readily tracked to monitor its effects in the body. The results of this latest work resolve some of the practical issues around using arginine groups to improve BSH cell penetration for cancer treatment using BNCT.
Figure caption

Chemical structures of the readily synthesised BSH fused with comparatively short ‘nR’ peptides, where R is the arginine group and n is the number of arginine groups. When fused with 3R the BSH penetrates the cell for effective BNCT. The chemical is additionally readily fused with a fluorescent moiety, Tmr (a), and DOTA for positron emission tomography scans (b) to track the drug in the body.

References

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Correspondence to
Assistant Professor Hiroyuki Michiue, M.D., Ph.D.
Department of Physiology, Okayama University
Graduate School of Medicine, Dentistry and Pharmaceutical Sciences, 2-5-1 Shikata-Cho,
Okayama, 700-8558, Japan.
E-mail: hmichiue@md.okayama-u.ac.jp

Further information
Okayama University
1-1-1 Tsushima-naka , Kita-ku , Okayama 700-8530, Japan
Planning and Public Information Division, Okayama University
E-mail: www-adm@adm.okayama-u.ac.jp
Website: http://www.okayama-u.ac.jp/index_e.html
Okayama Univ. e-Bulletin: http://www.okayama-u.ac.jp/user/kouhou/ebulletin/
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