Peptide Derived from Cow’s Milk Kills Human Stomach Cancer Cells in Culture
Findings Reported in the Journal of Dairy Science® Show Promise for Treatment of Gastric Cancer


“Gastric cancer is one of the most common causes of cancer-related mortality worldwide, especially in Asian countries,” says Wei-Jung Chen, PhD, of the Department of Biotechnology and Animal Science of National Ilan University, Taiwan Republic of China. “In general, the main curative therapies for gastric cancer are surgery and chemotherapy, which are generally only successful if the cancer is diagnosed at an early stage. Novel treatment strategies to improve prognosis are urgently needed.”

Investigators evaluated the effects of three peptide fragments derived from lactoferricin B, a peptide in milk that has antimicrobial properties. Only one of the fragments, LFcinB25 reduced the survival of human AGS (Gastric Adenocarcinoma) cells in a dose-dependent and time-dependent manner.

Under a microscope the investigators could see that after an hour of exposure to the gastric cancer cells, LFcinB25 migrated to the cell membrane of the AGS cells, and within 24 hours the cancer cells had shrunken in size and lost their ability to adhere to surfaces. In the early stages of exposure, LFcinB25 reduced cell viability through both apoptosis (programmed cell death) and autophagy (degradation and recycling of obsolete or damaged cell parts). At later stages, apoptosis appeared to dominate, possibly through caspase-dependent mechanisms, and autophagy waned.
“This is the first report describing interplay between apoptosis and autophagy in LFcinB-induced cell death of cancer cells,” says Dr. Chen.

The research also suggested a target, Beclin-1, which may enhance LFcinB25’s cytotoxic action. Beclin-1 is a protein in humans that plays a central role in autophagy, tumor growth, and degeneration of neurons. In this study, the investigators found that cleaved beclin-1 increased in a time-dependent manner after LFcinB25-exposure, suggesting to the authors a new approach in drug development that may boost the anticancer effects of LFcinB25.

“Optimization of LFcinB using various strategies to enhance further selectivity is expected to yield novel anticancer drugs with chemotherapeutic potential for the treatment of gastric cancer,” concludes Dr. Chen.

NOTES FOR EDITORS


Full text of the article is available to credentialed journalists upon request. Contact Eileen Leahy at +1 732-238-3628 or jdsmedia@elsevier.com to obtain copies. Journalists wishing to set up interviews with Dr. Chen may contact him at +886 3-931-7623 or wjchen@niu.edu.tw.

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