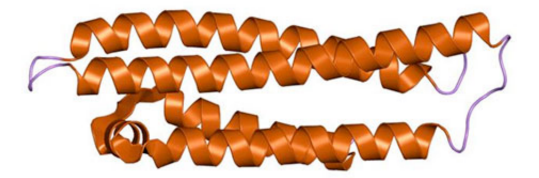


Drug sharply lowers cholesterol in animal tests

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Crystal structure of the receptor-binding domain of Apolipoprotein E, residues 1-165. Credit: European Bioinformatics Institute

An Arizona drug company will patent a small peptide—developed through research at the University of Alabama at Birmingham—that shows significant ability to lower blood cholesterol in animal models. Capstone Therapeutics Corp. and its joint venture affiliate, LipimetiX



Development Inc., recently announced the U.S. Patent application, claiming novel, more potent analogs of its lead peptide, AEM-28.

LipimetiX, based in Tempe, Arizona, has supported UAB research by G.M. Anantharamaiah, Ph.D., a professor in the UAB School of Medicine'sDivision of Gerontology, Geriatric and Palliative Care, to identify novel analogs of AEM-28 that have greater efficacy than the parent peptide. The new Apo E mimetic peptide, AEM-28-14, showed a 400 percent greater cholesterol-lowering efficacy and a several-fold increase in drug tolerability. Increased efficacy and tolerance have the potential to increase the safe and effective dose range compared with other AEM-28 analogs. Anantharamaiah's research at UAB focuses on the use of apolipoprotein mimetic peptides to treat atherosclerosis and cardiovascular disease.

Capstone and LipimetiX last year announced that the parent drug, AEM-28, showed a generally acceptable safety profile and some significant alterations in blood lipid biomarkers in a human Phase 1a/1b/2a clinical trial.

"We believe that the profound cholesterol-lowering effect of a single injection of AEM-28-14, accompanied by the enhanced toleration, will allow us to expand upon the statistically significant VLDL cholesterol and triglyceride reductions seen in our recently completed AEM-28 human clinical studies," said Dennis Goldberg, Ph.D., president of LipimetiX. "Subject to funding, LipimetiX plans to begin development of AEM-28-14 for cholesterol and triglyceride reduction in patients refractory to available therapeutic modalities."

AEM stands for chimeric Apolipoprotein E Mimetic peptides. UAB researchers created these small peptides to mimic the function of apolipoprotein E (Apo E), a class of protein essential for metabolism of cholesterol and triglycerides. After a meal, Apo E targets cholesterol-



and triglyceride-rich lipoproteins to specific receptors in the liver, decreasing the levels in the blood. Elevated plasma <u>cholesterol</u> and triglycerides are independent risk factors for atherosclerosis, the major cause of cardiovascular, peripheral artery and cerebral artery disease. These diseases can lead to heart attacks, loss of limbs and strokes. Faulty lipid metabolism also contributes to adult onset diabetes, and diabetics are very vulnerable to atherosclerosis and to heart and peripheral artery diseases.

UAB scientists patented the first chimeric Apo E mimetic peptide in 1999, reducing the 299-amino acid native Apo E to a 28-amino acid peptide that has two parts. One part of the peptide attaches to the surface of lipoproteins; the other domain binds to Apo E receptors in the liver.

In 2010, the founding scientist of LipimetiX obtained worldwide right to patents for Apo E mimetic peptides from the UAB Research Foundation, now a part of the UAB Institute for Innovation and Entrepreneurship. LipimetiX has an exclusive license with the UAB Research Foundation for AEM-28 and its analogs.

Provided by University of Alabama at Birmingham

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