

Protective effects of ADM-RAMP2 system make it a new therapeutic target for retinal vein occlusion

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A: Topical endoscopy fundus imaging showing the ocular fundus of untreated and central retinal vein occlusion (CRVO)-treated mice (day 7). In the CRVOtreated mice, note the retinal hemorrhage and venous dilatation with tortuousness resembling clinical CRVO. B: Retinal flat-mount images of untreated and CRVOtreated mice (day 7). In the CRVO-treated mice, retinal hemorrhage is prominent. C: Fluorescein isothiocyanate (FITC)-perfused retinal flat-mount images analyzed using AngioTool software in wild-type (WT), Adm knockout (ADM KO), and Ramp2 knockout (RAMP2 KO) mice after CRVO treatment

(day 7). Blood vessels are depicted in red, and vascular branch points are depicted in blue. D: Percent vascular area in FITC-perfused retinal flat-mounts quantified using AngioTool. E: FITC-perfused retinal flat-mounts analyzed using AngioTool software in control (Ctrl) and ADM-administered (ADM pump) mice after CRVO treatment (day 7). Blood vessels are depicted in red, and vascular branch points are depicted in blue. F: Percent vascular area in FITC-perfused retinal flat-mounts quantified using AngioTool. Credit: *American Journal of Pathology*

A clot in the retinal vein can lead to severe and irreversible loss of vision. In a report in the *American Journal of Pathology* investigators utilize a newly developed model of central retinal vein occlusion (CRVO) in mice that mimics many of the clinical features of CRVO in humans to study the pathologic effects of retinal occlusion and demonstrate the retinoprotective effects of the peptide adrenomedullin (ADM) and its receptor activity-modifying protein RAMP2.

"CRVO is the second most common retinal vascular disease next to diabetic retinopathy. In recent years, intravitreal administration of antivascular endothelial growth factor (VEGF) agents has been used to treat CRVO by suppressing retinal vascular permeability. However, recurrence of macular swelling and decreasing drug effectiveness are limitations of this approach. Therefore, identification of other therapeutic agents that enhance vascular integrity is desirable. In this study, we proved that the ADM-RAMP2 system is one of the novel therapeutic targets for treatment of CRVO," explained Takayuki Shindo, MD, Ph.D., of the Department of Cardiovascular Research at Shinshu University Graduate School of Medicine, Matsumoto, Japan.

The CRVO model was generated by combining intraperitoneal injection of Rose Bengal, a photo-activator dye that enhances thrombus formation, with laser photocoagulation. The most prominent changes

noted were retinal hemorrhage and venous dilatation with tortuosity. Measurements of retinal thickness revealed prominent edema early after CRVO was induced, followed by gradual atrophy.

There are several possible ways ADM works in CRVO. ADM was originally identified as a vasodilating peptide expected to improve blood flow in CRVO. ADM also exerts anti-coagulating effects, which may suppress the hypercoagulability in CRVO. The researchers showed that ADM suppresses inflammation and oxidative stress associated with CRVO, both of which promote endothelial injury. ADM also suppresses vascular hyperpermeability, which is the cause of retinal edema, and increases vascular reperfusion. Thus, endogenous ADM likely exerts protective effects against the retinal damage caused by CRVO.

Based on these findings, investigators propose that the retinoprotective effects of the ADM-RAMP2 system make it a novel therapeutic target for treatment of CRVO.

"As the biological functions of ADM are likely different from VEGF antibodies, co-administration of ADM and VEGF antibodies may be a choice in some diseases. It may also be used as secondary therapy when patients become resistant to VEGF antibodies," commented Dr. Shindo.

Retinal vein occlusion, which affects an estimated 16.4 million individuals worldwide, can occur as a complication of hypertension, diabetes, dyslipidemia, or aging. Hallmark features are capillary loss, hemorrhage, macular edema, inflammation, and vascular complications. CVRO can be ischemic or non-ischemic depending upon whether <u>blood</u> flow has been interrupted; the ischemic type is irreversible and resistant to treatment. Anti-VEGF agents have been used to treat CVRO, but their use is limited by troublesome side effects, temporary benefit, and high costs.

More information: "Development of a Novel Model of Central Retinal Vascular Occlusion and the Therapeutic Potential of the Adrenomedullin-Receptor Activity-Modifying Protein 2 System," *American Journal of Pathology* (2019). DOI: <u>10.1016/j.ajpath.2018.10.021</u>

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