Case Report

Nicotinamide as a treatment option of Age-Related Macular Degeneration

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Age related macular degeneration is a severe disease of mainly elderly people and leads to central vision loss because of the degeneration of the retinal pigment epithelium [1]. Genetic and environmental factors are responsible for the accumulation of extracellular material and deposit formation near the retinal pigment epithelial (RPE) layer, which leads to loss of photoreceptors and induction of chronic inflammation. The deposits are composed of lipids and proteins including many complement proteins, indicating the involvement of the complement system in the degenerative process and chronic inflammation [2]. So far there is no treatment for the dry form of AMD, except nutritional supplementation with antioxidants and vitamins [3]. Combined with a prolonged lifetime expectation in developed countries, AMD is developing to a social and economic burden. Therefore, there is an urgent need for a treatment of AMD that can delay disease manifestation and progression for several years.

In a recent manuscript, Saini and colleagues analyzed a vitamin B derivate, nicotinamide (NAM), as a potential treatment strategy of AMD [4]. The anti-inflammatory activity of NAM is not entirely new and may have inspired the investigators to look in detail at the effect of this drug for AMD treatment. In 2008, Temple and her group reported about the treatment of 3 x Tg-AD mice, a mouse model for Alzheimer’s disease (AD), with NAM. Their work suggested that NAM treatment selectively reduces a phosphorylated species of tau associated with microtubule depolymerization and implicated in AD. Treatment of mice restored cognitive decline related to AD pathology indicate a readily available therapeutic for the treatment [5]. Collectively NAM was identified as a cytoprotectant that blocks cellular inflammatory cell activation, early apoptotic phosphatidylserine exposure, and late nuclear DNA degradation [6]. Therefore, NAM supplementation was suggested for the treatment of chronic diseases with an immune system dysfunction as seen in diabetes and age-related diseases.

Saini et al. [4], investigated the effects of NAM in a human induced pluripotent stem cell (iPSC) model of AMD which they they previously established. The advantage of this model is evident. The investigators used cells from AMD patients with or without a specific ARMS2/HTRA1 risks haplotype as well as from control people to generate iPSCs and to compare the effects of NAM. The authors showed that iPSC cells express a broad panel of drusen like deposits and AMD associated molecules. Also, iPSC-RPE from all AMD patients studied, demonstrated an increased expression of inflammatory and complement factors as compared to controls. The expression of complement genes C3 and CFI was significantly enhanced in patient cell lines carrying the homozygous
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**ARMS2/HTRA1** risk alleles. Interestingly, the expression of the classical complement components C1s and C1r was enhanced in AMD derived iPSC-RPE, thus indicating a role of the classical pathway in AMD.

To determine the effect of NAM on the iPSC-RPE, the authors used a large panel of methods (ELISA, transcriptome analysis, qPCR) for analysis. Overall, the results revealed similar responses to NAM of all generated iPSC-RPEs, with enhanced inhibition of several markers including, drusen components ApoE,ApoJ, and VEGF in AMD lines. Furthermore, NAM inhibited transcription of complement and inflammatory factors. Moreover, NAM increased RPE survival by targeting aging-associated pathways.

Given the strong involvement of complement in AMD (genetic association [7], complement proteins in drusen [4,8,9], ARMS2 protein function in complement [10], factor H on microglial [11] and the dramatic reduction of C3 expression levels in iPSC-RPE lines upon NAM treatment, justified the question by the investigators whether the NAM effect was largely explained by the reduction of C3 transcription. To answer this question, the authors inhibited C3 expression in their iPSC cell models via shRNA. This treatment inhibited C5 transcription, but in contrast to NAM treatment, they did not observe any effects on complement regulator *CFH* expression or factor I (CFI) (Figure 1). Silencing of C3 expression also did not increase cell viability as seen with NAM. This is likely explained by intracellular C3 functions which were reported to be involved in signal transduction pathways directing survival, repair, and regeneration [12,13]. Based on these results the authors concluded that C3 inhibition does not have the multiple beneficial effects like NAM.

However, although C3 inhibition does not have a similar suppressive effect on gene transcription like NAM, the subsequent inhibition of C5 expression might have a specific impact on complement mediated inflammatory responses of microglia cells attraction and activation of blood monocytes, which was not tested here. Although NAM holds great potential for multiple disease entities, there are also sounds of caution based on complex biological roles NAM plays in cell survival pathways and unwanted detrimental effects in cellular proliferation that leads to cancer [14]. Clinical trials with NAM will clarify whether this readily available promising substance fulfills the expectations to reduce inflammation and damage in AMD patients.

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**Figure 1:** NAM-effect on complement in iPSC-RPE cells. Treatment of iPSC-RPE cells with NAM inhibits more complement genes compared to C3-silenced cells. Furthermore VEGF secretion is reduced in NAM treated RPE cells, but not in C3 -gene silenced RPE cells.
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References


