# **ARIC Manuscript Proposal # 1134r**

PC Reviewed: 03/_21/06	<b>Status:</b>	Priority:
SC Reviewed:	<b>Status:</b>	Priority:

- **1.a. Full Title**: Associations of Negative Emotions with Retinal Microvascular Disease and Age-Related Maculopathy
  - b. Abbreviated Title (Length 26 characters): Negative Emotions and Retinal Diseases
- 2. Writing Group (list individual with lead responsibility first):

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#### 3. Timeline:

Manuscript proposal to Publication's Committee:

Data analysis completed:

Completed manuscript to Publication's Committee:

Jan / 2006

June / 2006

Oct / 2006

# 4. Rationale:

Negative emotional states (anger, hostility and depression) have been linked with development and progression of cardiovascular diseases<sup>1</sup>. Much of this research has focused on the impact of negative emotion on macrovascular disease. For example, numerous studies have shown that anger and hostility predict incident coronary heart disease <sup>2,3</sup>, incident stroke<sup>4</sup> and cardiovascular mortality<sup>5</sup>. Anger also appears to predict risk of hypertension<sup>6</sup>. In the ARIC cohort, Williams and colleagues found that high trait anger was associated with a 2.7-times greater risk of CHD morbidity and death in normotensives<sup>7</sup>, and a two-fold greater risk of incident stroke among participants aged 60 or younger<sup>8</sup>.

Similarly, depression has also been linked to various cardiovascular risk factors and cardiovascular diseases<sup>8, 9</sup>, including progression of carotid atherosclerosis<sup>10-12</sup> as well as incident stroke<sup>13-16</sup>. Vital exhaustion is a closely related construct and overlaps considerably with measures of depression, and has also been linked with incident cardiovascular disease<sup>17-19</sup>. Preliminary findings from ARIC also revealed significant associations between vital exhaustion and metabolic syndrome<sup>20</sup>.

Although the pathophysiological basis linking these negative emotions with CVD have not been fully elucidated, several hypotheses have been proposed. First, individuals with higher levels of negative emotions tend to have poorer cardiovascular risk profiles (e.g., more likely to be smokers and to have less physical activity), although the excess cardiovascular risk associated with negative emotions persists even when studies controlled for these risk factors. Second, negative emotions may interact with other psychosocial risk factors (e.g., individuals high in hostility may engender stressful interpersonal environments and ultimately reduced social support). Finally, there is increasing evidence that negative emotions may influence CVD risk via increased sympathetic and neuroendocrine responses, resulting in alterations in the hypothalamic-pituitary-adrenal axis and sympatho-adreno-medullary axis<sup>21</sup>.

There have been few studies on the possible impact of negative emotions on small vessel diseases. "Microvascular angina", or cardiac syndrome X, a condition caused by coronary microvascular dysfunction<sup>22</sup>, has been linked with measures of anxiety and depression in both men<sup>23</sup> and women<sup>24</sup>. It has also been associated with trait anger and neuroticism<sup>25</sup>. In addition, previous meta-analysis has reported statistically significant associations of depression with diabetes complications including diabetic retinopathy  $(r = 0.17)^{26}$ . However, whether negative emotions, including vital exhaustion and trait anger, also impact on retinal microvascular changes is unclear.

Furthermore, the data addressing the effect of negative emotions on age-related macular degeneration (AMD) are also scarce. A review of the recent clinical studies proposed AMD as a risk factor for depression<sup>27</sup>. Nevertheless, there is a lack of population-based data to support this notion. In addition, whether there are associations between AMD and other measures of negative emotions such as vital exhaustion and trait anger still remains undetermined.

In the current study, we propose to examine the association between negative emotions (trait anger and vital exhaustion) with retinal microvascular disease and AMD in the ARIC cohort.

# **Main Hypothesis/Study Questions:**

We hypothesize that trait anger and vital exhaustion will be positively associated with retinal microvascular signs and AMD, independent of standard cardiovascular risk factors.

- Data (variables, time window, source, inclusions/exclusions):
  - **A.** Retinal microvascular variables at **Visit 3**: retinal arteriolar diameter, retinal venular diameter. arteriovenous nicking, focal arteriolar narrowing, blot hemorrhages, soft exudates (cotton wool spots), and microaneurysms
  - **B.** AMD variables at **Visit 3**. Any AMD, early AMD, late AMD and specific AMD lesions (drusen, RPE de-pigmentation, any pigmentary changes)
  - C. Vital exhaustion at Visit 2: the Maastricht questionnaire
  - **D.** Trait anger at **Visit 2** and **Visit 4**: the Spielberger Trait Anger Scale
  - E. Other variables:
    - Cardiovascular risk factors at Visit 1, 2 and 3: hypertension, blood pressure, diabetes, cigarette smoking, plasma total cholesterol, HDL cholesterol, LDL cholesterol, triglyceride, glucose, BMI
    - ii. Medications with known psychotropic effects

### Plan of analysis

Participants with a history of stroke or TIA prior to Visit 3 or missing data will be excluded from the

analysis. We will analyze the association	n of vital ext	naustion and t	rait anger characterist	tics at Visit 2 with
retinal variables at Visit 3, adjusting for	potential co	<mark>nfounders at V</mark>	Visit 2. In sub-group a	<mark>analyses, we will</mark>
also (1) examine the association of trait	anger at Vis	it 4 with retin	al signs at Visit 3, and	d (2) adjust for
confounders (systolic blood pressure, gl	<mark>lucose, total</mark>	<mark>cholesterol, tr</mark>	iglyceride and BMI)	with values that ar
the average of Visits 1 to 3. Limitations	that this is n	ot a pure cros	s-sectional analysis v	vill be discussed in
<mark>the paper.</mark>				
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8.b. If yes, is the author aware that either DNA of be used, or the file ICTDER02 must be used to exuse/storage DNA"?						exclude	·						
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10. V	What ar	e the	most	related	d manuscr	ipt propo	sals in	ARIC (a	utho	rs are	encou	raged to	

10. What are the most related manuscript proposals in ARIC (authors are encouraged to contact lead authors of these proposals for comments on the new proposal or collaboration)?

ARIC MS #666, Williams et al. (Stroke, 2002) demonstrated an association between trait anger and incident stroke risk.

We are aware of no ARIC manuscripts or proposals related to negative emotions and retinal disease.

11. Manuscript preparation is expected to be completed in one to three years. If a manuscript is not submitted for ARIC review at the end of the 3-years from the date of the approval, the manuscript proposal will expire.

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