The Effect of Lutein Supplementation on Blood Plasma Levels of Complement Factor D, C5a and C3d

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Abstract

Lutein is selectively taken up by the primate retina and plays an important role as a filter for harmful blue light and as an antioxidant. Recent studies have shown that lutein has systemic anti-inflammatory properties. Dietary lutein has been associated with reduced circulating levels of inflammatory biomarkers such as CRP and sICAM. Whether lutein also affects activation of the complement system has not yet been addressed and was the purpose of the study described here. Seventy-two subjects with signs of early macular degeneration were randomly assigned to receive either a 10 mg lutein supplement or a placebo during one year. EDTA blood samples were collected at 0, 4, 8 and 12 months. Complement factor D (CFD), a rate limiting component of the alternative pathway of complement activation and the complement activation products C5a and C3d were determined in the plasma samples by ELISA. A significant 0.11 µg/ml monthly decrease in plasma CFD concentration was observed in the lutein group (p<0.001), resulting in a 51% decrease from 2.3 µg/ml at baseline to 1.0 µg/ml at 12 months. The C5a concentration showed a significant 0.063ng/ml monthly decrease in the lutein group (p<0.001) resulting in a 36% decrease from 2.2ng/ml at baseline to 1.6ng/ml at 12 months. The C3d concentration showed a significant 0.19µg/ml monthly decrease in the lutein group (p=0.004) that gave rise to a 9% decrease from 15.4µg/ml at baseline to 14.4µg/ml at 12 months. In the placebo group we found a significant 0.04 µg/ml monthly decrease in plasma CFD concentration, whereas no changes were observed for C5a and C3d. Lutein supplementation markedly decreases circulating levels of the complement factors CFD, C5a and C3d levels, which might allow a simple method to control this inflammatory pathway of the innate immune system.

Introduction

Lutein selectively accumulates in the primate retina and as a major component of the macular pigment it plays an important role as a filter for harmful effects of blue light [1]. It also protects the retina from oxidative damage by quenching free oxygen radicals [2]. Various studies have recently shown that lutein also has anti-inflammatory properties (for review see Kijlstra et al. [3]). Lutein administration has led to beneficial effects in various models of experimental inflammation such as endotoxin induced uveitis [4], experimental age related macular degeneration [5,6], retinal ischemia [7,8] and diabetic retinopathy [9–11]. An inverse relationship has been shown between serum lutein concentration and markers of inflammation such as CRP and soluble ICAM-1 [12,13]. Whether lutein can also affect the complement pathway is not yet known and was the purpose of our study. The complement system plays an important role in the defense against microbial pathogens, clearance of apoptotic cells and chemotaxis of inflammatory cells [14]. A great deal of interest has recently been directed towards the role of the complement system in various eye diseases including age-related macular degeneration (AMD). This is based on the fact that drusen in the retina contain complement products [15,16] and because one of the strongest genetic associations with AMD has been found in genes encoding for a protein of this system (CFH gene) [17]. Furthermore, recent findings showed an increase in the level of various complement components in the circulation.
of AMD patients providing evidence for a systemic inflammatory component to the disease pathogenesis [18–22]. In view of these latter findings we chose individuals with early signs of AMD for our lutein intervention study.

Methods

Study design

The current study was a research project carried out with stored blood samples that had been collected during a randomized, double-blind, placebo-controlled two-centre intervention study (RCT) that was carried out in Manchester, UK and Maastricht, The Netherlands. This RCT was registered as a clinical trial NCT01042860 at clinicaltrials.gov and primary outcomes and results were recently published elsewhere [23]. Subjects were randomly assigned to receive either a 10 mg lutein supplement or a placebo based on soy bean oil. The UK arm of the study was approved by the South Manchester Regional Ethical Committee and The Netherlands arm of the study was approved by the Medical Ethical Committee at the University Hospital of Maastricht. The study was undertaken in accordance with the Declaration of Helsinki and all patients provided written informed consent. Patients were aware of the fact that blood samples were stored for future anonymous research questions. Uniform procedures and management strategies were used in the two centers. An advertising campaign was conducted within the universities and in local papers in both centers. Responding potential participants were contacted by letter and telephone.

Subjects

Men or women aged between 50 to 80 years who met the following inclusion criteria were eligible to participate in the study: AMD grade 0 to 4 in one eye, best corrected visual acuity (BCVA) of LogMAR of 0.5 or better and minimal cataract. Subjects were excluded if they had any ophthalmic disorder other than early signs of AMD. In particular diabetic retinopathy, optic atrophy, pigmentary abnormalities considered by a supervising ophthalmologist to be less typical of AMD and history of glaucoma. Subjects with a BMI over 30 or who were diabetic were excluded. Finally, any potential participant consuming supplements containing lutein, zeaxanthin or meso-zeaxanthin or who had taken any of these at least 12 months before the study was not enrolled.

Randomization and masking

The supplement was manufactured for the study by Cognis GmbH (now BASF SE; Rheinpromenade 1, 40789 Monheim, Germany) in accordance with Good Manufacturing Practice. A randomization code was generated by the sample manufacturer and capsules were distributed in accordance with this list. Each patient was assigned a specific treatment number. Treatment numbers were allocated in ascending order using the next available consecutive number. If a discontinued patient was replaced, the next available treatment number was used. The code linking treatment number with capsule allocation remained with the manufacturer until the end of the intervention trial. The study personnel were unaware of which patients were assigned to which groups. The study products were packaged and delivered under the responsibility of Cognis. Each participant was allocated a box (a treatment unit) in which there were 14 white plastic cartons. Each of these held 36 capsules, sufficient for 1 month (31 days plus 5 extra). The placebo and lutein containers were indistinguishable. Each treatment unit represented the complete study product for one participant. It contained a total of 504 capsules, enough capsules for 12 months of treatment (365 days plus 139 more). The supplementation products were stored safely according to manufacturer’s guidelines at temperatures below 25 °C and not exposed to light. Four 3 monthly visits were scheduled at which four cartons (144 capsules) were given to each participant. The safety of the lutein supplement has been addressed elsewhere [24,25].

Blood sampling and complement factor analysis

Blood samples were taken using EDTA containing tubes at the time of the scheduled visits. They were centrifuged within one hour and stored in aliquots at -80 degrees Celsius. CFD was measured at a 1/4000 dilution using a commercially available development kit (DuoSet) for human complement factor D (R&D Systems, Minneapolis, USA) according to the manufacturer’s instructions. C5a was measured at a 1/10 dilution using a commercially available development kit (DuoSet) for human complement component C5a (R&D Systems, Minneapolis, USA). C3d was measured at a 1/10 dilution using a development kit for human complement fragment C3d (MyBioSourceSan Diego, USA).

Statistical analysis

Statistical analysis was performed using SPSS 20.0.0. Differences in gender distribution over the experimental groups were tested using the Pearson Chi-square test, while differences in age and plasma concentrations between groups was evaluated using ANOVA. To quantify the course in time, a Linear Mixed Models analysis (LMM) was performed with subject ID as grouping factor, and gender, supplementation time, and the interaction term of the latter two as covariates. All statistical tests were done at a two-sided significance level of 0.05.

Results

Seventy-two subjects were randomly assigned to the two intervention groups. An extensive flow diagram of the included volunteers is shown in the paper published of the original RCT [23]. Table 1 shows the baseline characteristics of the 72 subjects. Age en gender as well as baseline plasma concentrations of CFD, C5a, and C3d were not statistically different among the two groups.

Following supplementation with a daily dose of 10 mg lutein, all measured complement factors (CFD, C5a and C3d) decreased over time in the lutein group (Table 2). In the placebo group, C3d and C5a levels remained constant over
time, whereas the CFD levels showed a decrease in time. Separate analysis of the values from each individual relative to the baseline concentration showed a decrease of 51%, 36% and 8% for CFD, C5a and C3d respectively in the lutein group. In the placebo group, CFD showed a 14% decrease, whereas no significant changes were observed for C5a and C3d (see Figure 1).

Linear Mixed Model (LMM) analysis revealed a significant effect for time (p<0.001), as well as the interaction between supplementation and time (p=0.006) on CFD plasma levels. Levels of significance for the time effect for plasma C5a and C3d were p<0.001 and p=0.057. For the interaction between supplementation and time these values were p<0.001 and p=0.010 for C5a and C3d, respectively.

Stratifying for supplementation we found a significant 0.11 µg/ml monthly decrease in plasma CFD concentration in the lutein group (p<0.001), resulting in a decrease from 2.3 µg/ml at baseline to 1.0 µg/ml at 12 months. The C5a concentration showed a significant 0.063ng/ml monthly decrease in the lutein group (p<0.001) resulting in a decrease from 2.2ng/ml at baseline to 1.6ng/ml at 12 months. The C3d concentration showed a significant 0.19µg/ml monthly decrease in the lutein group (p=0.004) that gave rise to a decrease from 15.4µg/ml at baseline to 14.4µg/ml at 12 months. In the placebo group we found a significant 0.04 µg/ml monthly decrease in plasma CFD concentration. No significant changes were observed for C5a and C3b. We incorporated gender as covariate into the LMM analysis, but did not find a significant effect of gender on the effect of supplementation on plasma concentrations of the complement factors. We also found no difference between the two participating centers.

Figure 2 shows the change in serum C5a and C3d concentration from baseline to 12 months as a function of the change in CFD concentration from baseline to 12 months (r=0.37, p = 0.04 and r=0.19, p=0.16 respectively). A LLM analysis whereby either C5a or C3d plasma levels were taken as dependent, subject ID as grouping factor and CFD plasma levels as covariates yielded a significant association between plasma C5a and CFD concentration (β=0.090, p=0.006). For C3d no significant association was observed.

**Discussion**

This study shows for the first time that daily supplementation with lutein can decrease the circulating levels of the complement factors CFD, C3d and C5a in an elderly population.

At baseline we found a mean CFD plasma concentration of 2.09± 1.43 µg/ml. This is in between the median values of...
2.31 µg/ml for AMD subjects and 2.08 µg/ml in controls as recently reported by others using the same ELISA kit [21]. Using a different kit to determine CFD, two other studies found somewhat lower values: 1.26 µg/ml for AMD subjects and 0.95 µg/ml for controls [18] and 1.50 µg/ml for AMD subjects and 1.16 µg/ml for controls [22]. The mean C5a concentration at baseline was 2.21 ng/ml in this study. This is in between the median values of 1.85 ng/ml and 4.28 ng/ml in AMD subjects and 1.67 ng/ml and 4.06 ng/ml in controls as recently reported by others using a different kit [18,22]. The mean C3d concentration at baseline was 15.4 µg/ml in our study. This is lower than median values of 55.2 µg/ml and 40.7 µg/ml in AMD subjects and 46.9 ng/ml and 35.8 µg/ml in controls as recently reported by others using a different kit [18,22]. Note that these studies all showed raised levels of the complement factors in AMD subjects as compared to controls pointing to a role in AMD pathogenesis.

A deregulated control of the alternative pathway of the complement system has been shown to play an important role in the pathogenesis of AMD [17]. CFD is a key enzyme in this pathway, resulting in the assembly of the alternative pathway C3 convertase and controlling the subsequent steps leading to important inflammatory activation products such as C3a and C5a [26,27]. CFD is the rate limiting enzyme in the activation sequence of the alternative pathway and its level in blood are quite low. The limitation is overcome at 9-10 times the normal physiological level [28]. Small alterations of the CFD level will thus have profound effects on the activation of the alternative pathway of complement. In this study we found a 51% decrease in CFD and an accompanying 36% decrease in C5a, in accordance with this concept. It was further confirmed by the fact that we found a positive correlation (r=0.37, p=0.004) between the change in CFD and the change in C5a (see Figure 2) and the fact that a LMM analysis showed a significant association between plasma CFD and C5a concentration (β=0.090, p=0.006). C3d only showed a small but significant 9% decrease, and the association with CFD changes was less clear. This may be due to the fact that C3d, which is formed following decay of C3b, is mainly present on the activating surface and only small amounts are generated in the fluid phase.

We thus hypothesize that lutein supplementation decreases CFD levels, which as a result will lower complement activation products downstream in the complement cascade. CFD, also known as adipsin, is mainly produced by adipocytes [28,29]. Its low blood level is caused by an extremely high catabolic rate and loss by glomerular filtration. Whether lutein would affect production or catabolism and whether other carotenoids have the same effect remains an open question. Of interest is the fact that adipocytes are the main storage site for carotenoids including lutein [30]. A literature search failed to find studies linking lutein in adipocytes and whether it could affect the production of CFD.

A small but significant decrease in CFD levels was also found in the placebo group. Before entering the trial subjects were informed about the nature and background of the study. This may have led to a change in their dietary habits. It could also be due to a seasonal effect, but analysis of the dates that subjects entered the trial did not support this explanation. We also investigated CRP levels in plasma over time but did not find a significant change either in the treatment or placebo group (data not shown). A phase I/II clinical trial was recently started to lower CFD in the eye itself. In this trial the effect of an intravitreal injection of a monoclonal antibody against CFD (FCFD4514S) is being evaluated for the treatment of geographic atrophy (Genentech, San Francisco, USA, ClinicalTrials.gov NCT01229215). A much more appealing option would be to control CFD levels via the diet, as put forward by our study.

The role of nutrition in the prevention and progression of AMD has received a great deal of attention over the past decades. Studies on nutritional modulation of AMD were
recently reviewed by Weikel and Taylor [31]. They suggested that 10 mg of lutein, the amount also provided in this study, confers the highest benefit to the health of the retina.

There are a number of limitations in our study. It had a relatively small sample size and we did not equalize for possible confounders such as dieter health of the participants. We did exclude individuals with diabetes or with a BMI above 30. Other possible confounders such as weight changes during the study, the presence of chronic (inflammatory) disease in the study group which has an age over 65, use of anti-inflammatory medication, total energy intake, physical activity or smoking were not recorded. In the trial we replaced discontinuing subjects with new ones. The reasons for discontinuation and those lost to follow up were described in detail in our earlier paper on this study [23]. It might have resulted in unequal distributions between treatment and control groups, but this could not be evaluated because data on diet, lifestyle or medication which might have an influence on blood complement levels were not collected and compared. Larger trials are needed in the future to confirm the present observations and to rule out the possible confounding factors listed above.

In conclusion, due to the key role of CFD in the activation pathway of the alternative pathway of the complement system and the potentially profound effect of lutein on circulating levels of CFD, this study suggests that lutein supplementation may be a powerful tool in the prevention and treatment of AMD, although further studies to replicate the significance of these findings are now clearly warranted.

Author Contributions
Conceived and designed the experiments: AK IM TB. Performed the experiments: YT RvdV MM. Analyzed the data: YT AK TB. Wrote the manuscript: YT AK TB.

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