A Quantitative Approach to Predict Differential Effects of Anti-VEGF Treatment on Diffuse and Focal Leakage in Patients with Diabetic Macular Edema: A Pilot Study

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Purpose: We use semiautomated segmentation of fluorescein angiography (FA) to determine whether anti-vascular endothelial growth factor (VEGF) treatment for diabetic macular edema (DME) differentially affects microaneurysm (MA)–associated leakage, termed focal leakage, versus non-MA–associated leakage, termed diffuse leakage.

Methods: We performed a retrospective study of 29 subjects treated with at least three consecutive injections of anti-VEGF agents for DME (mean 4.6 injections; range, 3–10) who underwent Heidelberg FA before and after anti-VEGF therapy. Inclusion criteria were macula center involving DME and at least 3 consecutive anti-VEGF injections. Exclusion criteria were macula center involving DME and at least 3 consecutive anti-VEGF injections. Exclusion criteria were macular edema due to causes besides DME, anti-VEGF within 3 months of initial FA, concurrent treatment for DME besides anti-VEGF, and macular photocoagulation within 1 year. At each time point, total leakage was semiautomatically segmented using a modified version of our previously published software. Microaneurysms were identified by an expert grader and leakage within a 117 μm radius of each MA was classified as focal leakage. Remaining leakage was classified as diffuse leakage. The absolute and percent changes in total, diffuse, and focal leakage were calculated for each subject.

Results: Mean pretreatment total leakage was 8.2 mm² and decreased by a mean of 40.1% (P < 0.0001; 95% confidence interval [CI], [−28.6, −52.5]) following treatment. Diffuse leakage decreased by a mean of 45.5% (P < 0.0001; 95% CI, [−31.3, −59.6]) while focal leakage decreased by 17.9% (P = 0.02; 95% CI, [−1.0, −34.8]). The difference in treatment response between focal and diffuse leakage was statistically significant (P = 0.01).

Conclusions: Anti-VEGF treatment for DME results in decreased diffuse leakage but had relatively little effect on focal leakage as assessed by FA. This suggests that diffuse leakage may be a marker of VEGF-mediated pathobiology. Patients with predominantly focal leakage may be less responsive to anti-VEGF therapy.

Translational Relevance: Fluorescein angiography can define focal and diffuse subtypes of diabetic macular edema and these may respond differently to anti-VEGF treatment.

Introduction

The prevalence of diabetes is increasing throughout the world and it is estimated that over 400 million people will be affected by the year 2025.¹ Diabetic retinopathy is considered the leading cause of vision loss in working age adults.² The main causes of vision loss due to diabetes are proliferative diabetic retinopathy (PDR) and diabetic macular edema (DME).³ Diabetic macular edema, which affects over 21 million people worldwide, is a major cause of central vision loss and results from complex pathobiology, including
breakdown of the blood retinal barrier and failure of Müller cell fluid pumping action. Currently, treatment options for DME include intravitreal anti-vascular endothelial growth factor agents (anti-VEGF), intravitreal corticosteroids, and thermal laser. While anti-VEGF agents are regarded by most as first line therapy for DME, they are not universally effective and corticosteroids and thermal laser also are effective therapies in a subset of patients with DME. Clinicians wishing to offer individualized treatment regimens for their patients are challenged to predict a priori which treatment is optimal. Thus, it is common practice to initiate treatment with anti-VEGF and then use alternate therapies for persistent or nonresponsive DME. This process of elimination approach is inefficient and increases the already substantial treatment burden for patients. In the RISE and RIDE studies, subjects treated with monthly ranibizumab for 24 months showed that complete resolution of leakage on FA occurred in only 17% to 30% of subjects. This suggests that some forms of leakage are not fully controlled by VEGF inhibition.

Numerous investigators have subtyped DME as diffuse, or focal according to various criteria and using variable tools, which include biomicroscopy, fundus photography, fluorescein angiography (FA), and optical coherence tomography (OCT). The use of nonangiographic methods of classifying DME has been reviewed previously. With regard to the use of FA to subtype DME, focal leakage has been defined most frequently as leakage originating from microaneurysms (MAs), while diffuse leakage has been defined as leakage without a clear source. The utility of classifying DME based on FA has been limited by inherent subjectivity, multiple definitions, and challenges in reproducibility. Additionally, classifying an entire eye as focal or diffuse does not take into consideration the fact that both types of leakage frequently coexist in most patients. For this study, focal leakage was defined as originating from MAs, while diffuse leakage was defined as leakage without a clear source. We hypothesize that the pathobiology of focal leakage is distinct from that driving diffuse leakage. Seeking to explore this hypothesis, we have developed custom MATLAB-based software (MathWorks, Natick, MA) to semi-automatically segment total fluorescein leakage and MAs. This software permits quantification of the diffuse and focal components of leakage. To probe whether one subtype is more responsive to anti-VEGF treatment, we have performed a retrospective analysis of 29 subjects who received FA before and after anti-VEGF therapy, and quantified changes in total, focal, and diffuse leakage.

**Methods**

**Subjects**

This study was approved by the Duke University Medical Center Institutional Review Board, was conducted in compliance with the Health Insurance Portability and Accountability Act (HIPAA), and adhered to the tenets of the Declaration of Helsinki. A total of 29 eyes of 29 subjects were included in the study. Subjects with visually significant DME who also received FA transiting the treated eye before and after anti-VEGF as part of their routine care between January 1, 2013 and November 3, 2016 were identified. Inclusion criteria were macula center involving DME and at least 3 consecutive anti-VEGF injections. Exclusion criteria were macular edema due to cause besides DME, anti-VEGF within 3 months of initial FA, concurrent treatment for DME besides anti-VEGF, and macular photocoagulation within 1 year. To test the use and generalizability of our image analysis software in a real world setting, image quality was not used as an exclusion criteria.

**Image Acquisition**

All images for this study were obtained using the Heidelberg Retina Angiograph 2 (HRA2; Heidelberg Engineering, Heidelberg, Germany) device and either 30° or 55° field of view. The transit phase of the study was captured in Movie mode using the high resolution setting (4.7 frames per second) and subsequent middle and late phase images were captured as single images in ART mode (averaging 9 images). Each grayscale image in the sequence was composed of 768 × 768 pixel images. Following acquisition, image files were deidentified and exported in E2E format for further analysis.

**Image Analysis**

To determine the amount of leakage attributable to MAs, we reviewed images from an independent dataset, which includes equal proportions of predominantly focal, predominantly diffuse, and mixed leakage patterns (as judged by expert clinicians MJA, PM, and SC), and was previously reported and is publicly available. An expert grader (MJA) identified MAs that were sufficiently isolated from
other leaking structures as to allow measurement of a radius of leakage. Fifty MAs were identified from this dataset and the radius of leakage was manually measured using the Heidelberg measurement tool. The mean radius of leakage was normally distributed with a mean of 117 μm (standard deviation [SD] 30.8; 95% confidence interval [CI], [105, 128]). Based on these data, leakage within a radius of 117 μm surrounding the center point of each MA was attributed to that MA. For each FA image, MAs were segmented by an expert retina-trained clinician grader (MJA) using both individual frames of the movie capturing the transit phase as well as subsequent mid and late phase still images. Microaneurysms were identified as punctate hyperfluorescent lesions that increased in intensity after their initial appearance and persisted over the course of the study.

Quantification of total leakage used our previously published segmentation algorithm with changes and additions detailed below. To permit segmentation of MAs, grader correction of errors in leakage segmentation, and the quantification of focal and diffuse leakage, a graphical user interface (GUI) was developed (Fig. 1). The GUI was designed to allow: (1) uploading the transit stage video and the single late phase image, (2) registering early video frames and the late phase image, (3) manual correction of automated leakage segmentation, (4) manual marking of MAs, and (5) automatic quantification of focal and diffuse leakage. To reduce bias in manual segmentation, the GUI allowed segmentation of leakage and MAs to be performed independently; that is, the grader was masked to detected leakage while segmenting MAs and vice versa. The leakage in the 117 μm radius circle centered at each MA was classified as focal leakage. Remaining leakage was classified as diffuse leakage. After completion of segmentation the total, focal, and diffuse leakage areas were estimated automatically. Representative examples of clinical imaging and segmentation results are shown in Figure 2.
Figure 2. Fluorescein angiograms from three representative subjects are shown. For each subject, the top row shows pretreatment imaging and the bottom row shows post-treatment imaging. The left column is the composite early frame image, the middle column is the late image, and the right column shows the segmentation result. In the segmentation image, the region of interest is outlined by the yellow circle, leakage is outlined in red and shaded, and MAs are marked with green dots with the radius of attributable leakage appearing as a yellow circle around each MA.
Statistical Analysis

Descriptive and comparative statistics were performed using JMP, Version 12 (SAS Institute, Inc., Cary, NC). Normality was tested using histogram plot and Shapiro-Wilks test. Wilcoxon signed rank testing was used to compare total, diffuse, and focal leakage before and after anti-VEGF treatment.

Results

Subjects received an average of 4.6 injections between initial and follow up FAs. Ten subjects received bevacizumab, 6 received ranibizumab, 6 received aflibercept, and 7 received more than one anti-VEGF agent. Absolute total leakage decreased from a mean of 8.2 mm$^2$ (median, 7.6 mm$^2$) to a mean of 5.0 mm$^2$ (median, 3.9 mm$^2$) and this was statistically significant ($P = 0.001$). There also was a statistically significant reduction in absolute diffuse leakage after treatment; mean absolute diffuse leakage was 7.0 mm$^2$ (median, 6.5 mm$^2$) pretreatment and 4.0 mm$^2$ (median, 2.9 mm$^2$) post-treatment ($P = 0.0006$). Mean absolute focal leakage was 1.3 mm$^2$ (median, 1.2 mm$^2$) pretreatment and 1.0 mm$^2$ (median, 0.90 mm$^2$) post-treatment, but this change was not statistically significant ($P = 0.14$). Change in leakage also was analyzed by percent change in total, diffuse, and focal leakage. Total leakage decreased by a mean of 40.1\% ($P < 0.0001$; 95\% CI, [−28.6, −52.5]) following treatment. Diffuse leakage decreased by a mean of 45.5\% ($P < 0.0001$; 95\% CI, [−31.3, −59.6]). Focal leakage was 1.1 mm$^2$ (median, 1.0 mm$^2$) and decreased by a mean of 17.9\% ($P = 0.02$; 95\% CI, [−1.0, −34.8]). The difference in treatment response between focal and diffuse leakage was statistically significant for absolute ($P < 0.0001$) and percent ($P = 0.01$) change. The percent change in diffuse and focal leakage for each subject is displayed in Figure 3. As expected, there was heterogeneity in clinical response. However, most subjects have improvement in both forms of leakage. Notably diffuse leakage improved more than focal leakage in 22 of 29 subjects.

Discussion

We used semiautomated segmentation software to quantify total as well as the focal and diffuse
components of leakage before and after anti-VEGF therapy. Anti-VEGF therapy effectively reduced total leakage as well as diffuse and focal leakage. However, the absolute and percent reduction in diffuse leakage was significantly greater than the reduction in focal leakage. These findings suggested that diffuse leakage is significantly more responsive to a moderate course of anti-VEGF therapy (approximately 5 injections on average). Numerous investigators have subtyped DME as diffuse or focal according to various criteria and using variable tools, which include biomicroscopy, fundus photography, FA, and OCT. To date, the utility of classifying DME based on FA has been limited by inherent subjectivity, multiple definitions and challenges in reproducibility. Fluorescein angiography has been used to distinguish diffuse and focal DME in numerous studies and in randomized clinical trials. Most of these studies examined various treatment modalities in the management of diffuse DME. The RESTORE study performed a subgroup analysis of subjects classified as having diffuse, focal, or mixed DME, and failed to identify differences in treatment response among groups. However, the methodology used in RESTORE differs from our study in several important ways, which may explain the differences in results. In RESTORE, trained graders categorized each study eye as “focal,” “diffuse,” or “mixed” based on the proportion of leakage in the central subfield associated with MAs (0%–33% of leakage from MAs classified as diffuse, 33%–66% of leakage from MAs classified as mixed, and 66%–100% of leakage from MAs classified as focal). Because most eyes have components of diffuse and focal leakage, using a global classification for the entire eye could obscure differential responses to treatment between focal and diffuse leakage types. By contrast, our study examined the diffuse and focal components of leakage in each eye independently and, therefore, may have been able to detect differences in response to treatment in focal and diffuse leakage types. In addition, using a quantitative, semi-automated segmentation approach to classify leakage may be more sensitive and less subjective than the qualitative assessment of expert graders. Finally, it must be stated that RESTORE is a randomized, prospective study, while ours is a small, retrospective case series. Clearly, our results must be confirmed in the context of a larger prospective study. This study currently is underway.

Our study suggested that focal and diffuse leakage are, indeed, differentially responsive to anti-VEGF therapy. From a mechanistic standpoint, preclinical studies characterizing the effects of VEGF on the retinal vasculature support the concept that diffuse leakage may be a sign of VEGF-driven pathobiology. Vascular endothelial growth factor has been well characterized as a mediator of vascular permeability in many diseases, including diabetic retinopathy. A single intravitreal injection of VEGF has been shown to cause vessel dilation, tortuosity, and extensive diffuse leakage in a primate model. By contrast, only repeated intravitreal injection of VEGF induced formation of MAs. While this finding suggests a role for VEGF in MA formation, it does not inform the effects of VEGF inhibition on the biology of MAs. Observations from large clinical trials of anti-VEGF agents do not clearly address this question either. For example, reduced progression or even regression of nonproliferative diabetic retinopathy, which includes MAs in its definition has been reported with long-term administration of anti-VEGF in the RISE and RIDE trials. However, the specific effects of anti-VEGF therapy on MA turnover or resolution have not been specifically explored. Our results suggested that focal leakage is slow to respond to anti-VEGF when compared to diffuse leakage, but it is unknown what the findings would be after a longer continuous regimen of anti-VEGF therapy. This is a topic worthy of future study.

Our study is limited by its retrospective nature and small sample size. Because this is a retrospective study of subjects who received a second FA as part of their care, our cohort may be biased in favor of more severe or recalcitrant disease. A larger prospective observational trial with a standardized imaging protocol currently is under way and will allow us to confirm and extend the findings of the current study as well as examine the impact of other treatment modalities, such as thermal laser and corticosteroids.

In addition, the emergence of OCT angiography (OCT-A), raises the possibility that noninvasive, nondye based imaging will be capable of discriminating among different subtypes of DME. It also will be informative to evaluate whether there is correlation between FA subtypes and specific findings on traditional OCT. We have found in this study that a single eye may contain variable amounts of diffuse and focal leakage, which suggests the need for localized rather than global comparisons between FA and OCT or OCT-A. These investigations are part of our ongoing work.

In conclusion, we have found that in eyes with a mixture of leakage subtypes, the diffuse component is
more responsive to anti-VEGF therapy. It is possible that focal leakage is less responsive to anti-VEGF and that, therefore, patients with predominantly focal leakage may benefit from early or adjunctive focal laser therapy targeting leaking MAs or possibly steroid based treatments. From a more broad perspective, there may be differences in the pathobiology driving different leakage subtypes. As additional therapeutics become available to treat DME, robust subtyping using multimodal imaging will become increasingly important to guide individualized treatment planning.

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