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Plasma Kallikrein Mediates Retinal Vascular Dysfunction and Induces Retinal Thickening in Diabetic Rats

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OBJECTIVE—Plasma kallikrein (PK) has been identified in vitreous fluid obtained from individuals with diabetic retinopathy and has been implicated in contributing to retinal vascular dysfunction. In this report, we examined the effects of PK on retinal vascular functions and thickness in diabetic rats.

RESEARCH DESIGN AND METHODS—We investigated the effects of a selective PK inhibitor, ASP-440, and C1 inhibitor (C1-INH), the primary physiological inhibitor of PK, on retinal vascular permeability (RVP) and hemodynamics in rats with streptozotocin-induced diabetes. The effect of intravitreal PK injection on retinal thickness was examined by spectral domain optical coherence tomography.

RESULTS—Systemic continuous administration of ASP-440 for 4 weeks initiated at the time of diabetes onset inhibited RVP by 42% (P = 0.013) and 83% (P < 0.001) at doses of 0.25 and 0.6 mg/kg per day, respectively. Administration of ASP-440 initiated 2 weeks after the onset of diabetes ameliorated both RVP and retinal blood flow abnormalities in diabetic rats measured at 4 weeks' diabetes duration. Intravitreal injection of C1-INH similarly decreased impaired RVP in rats with 2 weeks' diabetes duration. Intravitreal injection of PK increased both acute RVP and sustained focal RVP (24 h postinjection) to a greater extent in diabetic rats compared with nondiabetic control rats. Intravitreal injection of PK increased retinal thickness compared with baseline to a greater extent (P = 0.017) in diabetic rats (from 193 ± 10 μm to 223 ± 13 μm) compared with nondiabetic rats (from 182 ± 8 μm to 193 ± 9 μm).

CONCLUSIONS—These results show that PK contributes to retinal vascular dysfunctions in diabetic rats and that the combination of diabetes and intravitreal injection of PK in rats induces retinal thickening.

Diabetic macular edema (DME) is a leading cause of vision loss attributed to diabetes. The 14-year incidence of this disease in individuals with type 1 diabetes followed in the Wisconsin Epidemiologic Study of Diabetic Retinopathy was 26% (1), and the progression to clinically significant macular edema was associated with increasing retinopathy severity (2). Although intensive glycemic and blood pressure control can reduce the incidence of DME (3) once this condition develops, the treatment options include laser and vascular endothelial growth factor (VEGF)-targeted therapies, which provide substantial improvement in visual acuity for ~50% of patients with DME (4). Thus, additional treatment options for this disease are needed.

DME is associated with a loss of blood-retinal barrier function, leading to increased diffusion of plasma components, thickening of the macula, and impairment in central vision (5,6). In addition to retinal thickening, increased retinal vascular permeability (RVP) alters the biochemical composition of the retinal interstitial fluid and the vitreous. Proteomic studies have begun to characterize the changes in the vitreous protein composition in people with diabetic retinopathy compared with nondiabetic subjects or diabetic subjects without diabetic retinopathy (7). We have previously reported an abundance of vasoactive plasma proteins, including components of the plasma kallikrein (PK)-kinin system (PKKS) in the vitreous of subjects with advanced diabetic retinopathy (7,8). These findings have suggested additional factors besides VEGF that may contribute to the decline in blood-retinal barrier integrity and vascular dysfunction in DME (9,10).

Plasma prekallikrein (PPK) is an abundant serine protease zymogen in blood that is converted to its catalytically active form, PK, by factor XIIa (11), contributing to the innate inflammatory response and intrinsic coagulation cascades (12). The mechanisms that lead to the activation of this pathway in vivo include interactions with polyphosphates released from activated platelets and deficiency of C1 inhibitor (C1-INH), the primary physiological inhibitor of the PKKS (13,14). PK-mediated cleavage of high-molecular weight kininogen generates the nonapeptide bradykinin (BK), which activates the BK2 (B2) receptor. Subsequent cleavage of BK by carboxypeptidases generates des-Arg9-BK, which activates the B1 receptor. Both B1 and B2 receptors are expressed by vascular, glial, and neuronal cell types, with the highest levels of retinal expression detected in the ganglion cell layer and inner and outer nuclear layers (15,16). Activation of B1 and B2 receptors causes vasodilation and increases vascular permeability (17–19). Previously, we have demonstrated that intravitreal injection of carboxic anhydrase-I (CA-1) increased RVP and that this response was blocked by the inhibition of PK and by BK receptor antagonists (8).

Recently, we reported that intravitreal injection of PK increased RVP in nondiabetic rats, and systemic administration of a small-molecule PK inhibitor, ASP-440, decreased RVP in rats subjected to angiotensin II (AngII)-induced hypertension (19). In the current study, we investigated the effects of PK on retinal vascular functions and retinal thickness in diabetic rats.
Diabetes induction. Male Sprague-Dawley rats (250–300 g) were obtained from Taconic Farms (Hudson, NY). Diabetes was induced by intraperitoneal injection of streptozotocin (STZ, Sigma-Aldrich, Milwaukee, WI) in 50 mmol/L sodium citrate at 55 mg/kg after a 12-h overnight fast. Blood glucose was measured by tail sampling using a One Touch Ultra glucometer 24 h after injection. Injections of blood glucose were >250 mmol/L, which was considered diabetic for the study. Anesthesia used for these experiments was an intramuscular injection of ketamine (50 mg/kg; Bioniche Pharma, Lake Forest, IL) and xylazine (10 mg/kg; Sigma-Aldrich). At the conclusion of the studies, animals were killed by inhalation of carbon dioxide. The experimental protocols were approved by the Joslin Diabetes Center Institutional Animal Care and Use Committee. All animals were handled and cared for in accordance with the Association for Research in Vision and Ophthalmology Resolution on the Use of Animals in Research.

AngII-induced hypertension model. Under anesthesia, each rat was implanted with a subcutaneous Alzet osmotic pump (model 1007D, 0.5 μL/h; Durect Corporation, Cupertino, CA) containing saline or AngII (18 μg/kg/h) and a second Alzet osmotic pump containing either saline or the PK inhibitor ASP-440 at 4 mg/mL (model 1007D, 0.5 μL/h or model 2001, 1 μL/h). Five days after pump insertion, blood pressure was measured using a noninvasive blood pressure–heart-rate–monitoring system (UR-5000; Ueda Electronic, Tokyo, Japan).

Primary and secondary intervention studies with the PK inhibitor ASP-440. For the primary intervention study, diabetic and age-matched control rats were randomly assigned on day 2 after STZ injection, and diabetic rats received a subcutaneous Alzet osmotic pump (either model 2002 [0.5 μL/h] containing AngII-440 at 5 mg/mL, model 2004 [0.25 μL/h] containing ASP-440 at 24 mg/mL, or vehicle [50% glycerol/50% saline]). For the secondary intervention study, a second set of rats with STZ-induced diabetes and age-matched control rats was randomly assigned on day 14 after confirmation of diabetes and implanted with a subcutaneous Alzet osmotic pump (model 2002 [0.5 μL/h] containing AngII-440 at 12 mg/mL, or vehicle [10% polyethylene glycol90% saline]).

C1-INH intervention. After the onset of diabetes was confirmed, nondiabetic and diabetic rats that were under anesthesia received intravital injections of human C1-INH (200 ng/kg; EMD Chemicals, Gibbstown, NJ) in one eye and balanced salt solution (BSS) in the contralateral eye on days 2, 7, and 12. Under sterile conditions, a 31-gauge needle was inserted into the vitreous pars plana, and a 10-μL bolus of solution was slowly infused.

RVP measurements by Evans blue-dye permeation. The Evans blue-dye permeation technique was used to quantify RVP (20). In brief, under anesthesia, each animal was infused with Evans blue dye (45 mg/kg) through an indwelling jugular catheter. The dye was allowed to circulate for 2 h prior to the time the rats were killed. After tissue fixation, the eyes were enucleated. Retinas were extracted with dimethyl formamide, and the resultant supernatant was used to determine Evans blue-dye content. Results are expressed as the mean ± SD.

Calculating retinal thickness. Retinal thickness was measured from the inner limiting membrane (ILM) to the retinal pigment epithelium (RPE) using spectral domain optical coherence tomography (SD-OCT) (S805DOCT System; Bioptigen, Durham, NC). Rectangular volumes of each retina were obtained comprising of 1000 A-scans by 100 B-scans over a 1.5 × 1.5 mm area centered upon the ONH. Retinal thickness was measured at 5 μm relative to the ONH at points defined by calipers on the OCT-derived en face image. At the intersection of the distance calipers and the corresponding B-scan, retinal thickness was measured using calibrated calipers from the retinal pigment epithelium (RPE) to the internal limiting membrane (ILM). Four caliper measurements were taken at each site, corresponding to the temporal, nasal, superior, and inferior quadrants of the retina. The measurements were averaged to produce a single thickness value for each retina. Retinal thickness false-color topographic maps were generated using semi-automated image segmentation software based on a smoothing filter and a Sobel operator method for edge detection in Matalab from exported retinal OCT images. Before segmentation, a median filter was applied to reduce speckle noise and smooth the image data in the transverse direction. Edges were detected using a modified Hough transform. Results were visualized by enhanced chemiluminescence (Cell Signalling, Danvers, MA). Quantification of autofluorescence was performed using ImageQuant 5.1 (Molecular Dynamics, Sunnyvale, CA).

Statistics. Statistical analysis was performed with SigmaStat statistical software (Jandel, San Raphael, CA). Multiple group comparisons for statistical significance used Bonferroni correction for one-way ANOVA. Unpaired comparisons were made with the Student paired t test. Results were considered statistically significant when P < 0.05. Results are expressed as mean ± SD.

RESULTS

Effect of PK inhibition by ASP-440 on AngII-induced RVP. We examined the effective dose of AngII-440 on RVP to Evans blue dye using an AngII-induced hypertension (19). Infusion of AngII increased the systolic blood pressure (SBP) of rats compared with vehicle controls (200 ± 32 vs. 152 ± 9 mmHg, P < 0.001). The SBP of rats coinfused with AngII and ASP-440 at 0.2 and 0.4 mg/kg per day was 193 ± 14 and 220 ± 24 mmHg, respectively, which was not significantly different than the SBP in rats infused with AngII alone (Supplementary Table 1). AngII infusion for 7 days increased RVP by 97% compared with vehicle-treated control rats (39.6 ± 12.3 vs. 20.1 ± 3.3 μL/g/h, P < 0.001). Cotreatment with ASP-440 at 0.2 and 0.4 mg/kg per day decreased the AngII-mediated increase in RVP in a dose-dependent manner by 61% (27.6 ± 9.1 μL/g/h, P = 0.015) and 90% (22.0 ± 11.0 μL/g/h, P = 0.002) compared with AngII plus vehicle-treated rats (Supplementary Fig. 1). RVP was similar in vehicle- and ASP-440–treated normotensive control rats (20.1 ± 3.3 vs. 21.4 ± 8.7 μL/g/h).

Effect of PK inhibition on RVP in diabetic rats. We examined the effect of systemic administration of AngII-440 on RVP in rats with STZ–induced diabetes. Subcutaneous injection of AngII-440 was repeated 2 days after STZ injection and delivered continuously for 4 weeks. Blood glucose and body weights were similar compared with rats receiving vehicle and AngII-440 (Supplementary Fig. 2). Diabetes increased RVP to Evans blue by 2.6-fold (P < 0.001) compared with age-matched control nondiabetic rats.
rats. Treatment with ASP-440 decreased RVP in diabetic rats in a dose-dependent manner by 42% ($P = 0.013$) at 0.25 mg/kg per day and 83% ($P < 0.001$) at 0.6 mg/kg per day compared with diabetic rats that received vehicle alone (Fig. 1A). Next, we examined the effect of intervention with subcutaneous osmotic pumps containing ASP-440 implanted after 2 weeks’ diabetes duration on RVP measured at 4 weeks’ diabetes duration. RVP in diabetic rats that received ASP-440 (0.6 mg/kg per day) was decreased by 56% ($P = 0.011$) compared with diabetic rats that received vehicle (Fig. 1B).

**Effects of PK inhibition on retinal hemodynamics.** Retinal hemodynamics were assessed by measurements of MCT from retinal arteries to veins, measurements of primary retinal vessel diameters, and calculated RBF in rats with 4 weeks’ diabetes duration in the absence or presence of treatment from week 2 to 4 with ASP-440. We show that MCT was prolonged ($P < 0.001$) and RBF was decreased ($P = 0.001$) in diabetic rats compared with nondiabetic control rats (Fig. 2), as described previously (22). Intervention with ASP-440 resulted in partial normalization of the diabetes-induced retinal hemodynamic changes, decreasing MCT ($P = 0.015$) and increasing RBF ($P = 0.001$) compared with diabetic rats that received vehicle alone. ASP-440 treatment in diabetic rats was also associated with increased diameter of retinal arteries ($9.7 ± 0.7$ vs. $8.7 ± 0.7$ pixels, $P = 0.011$) and veins ($13.3 ± 0.8$ vs. $12.3 ± 0.7$ pixels, $P = 0.03$) compared with diabetic rats that received vehicle alone.

**Circulating PPK levels in diabetes.** PPK levels in plasma from both diabetic and nondiabetic rats were quantified. We show that PPK protein levels measured by immunoblot were increased by 74% ($P = 0.005$) in diabetic rats and 64% ($P = 0.027$) in ASP-440–treated diabetic rats compared with nondiabetic rats (Supplementary Fig. 3).

**Effect of C1-INH on RVP in diabetic rats.** C1-INH is the primary physiological inhibitor of PK and plays a critical role in suppressing plasma extravasation in areas of local inflammation (14,23). We examined the effect of intravitreal injection of C1-INH on RVP in both diabetic and nondiabetic rats. RVP assessed by Evans blue-dye permeation was increased after 2 weeks of diabetes compared with nondiabetic controls (Fig. 3). We show that eyes that received an intravitreal injection of C1-INH (200 ng) at
days 2, 7, and 12 after diabetes onset displayed a 61% (P = 0.003) decrease in RVP compared with contralateral eyes that received control injections with saline vehicle. Intravitreal injection of C1-INH did not significantly affect RVP in nondiabetic rats.

**Effects of intravitreal injection of PK on RVP.** We examined the effects of intravitreal injection of PK, compared with vehicle, on RVP in diabetic and nondiabetic rats. We estimated the vitreous volume to be ~100 μL and injected 50 ng hPK per eye. This concentration of PK in the vitreous is comparable with 0.5 ng/eye and injected 50 ng hPK per eye. This concentration of PK in the vitreous is comparable with 0.5 ng/μL PK levels that we have detected in human diabetic retinopathy vitreous (data not shown). We demonstrate that intravitreal injection of hPK increased RVP in both nondiabetic and diabetic rats, compared with eyes that received saline vehicle injections (Fig. 4). It is noteworthy that we found that the magnitude of RVP increase induced by intravitreal injection of hPK in diabetic rats was significantly greater than that observed in nondiabetic rats (67 ± 30 vs. 34 ± 19%, P = 0.037).

We next examined the effect of intravitreal injections of ASP-440 or vehicle on RVP in nondiabetic rats induced by subsequent intravitreal injections with hPK, CA-1, and VEGF. Intravitreal injection with ASP-440 (20 μmol/L) pretreatment decreased PK- and CA-1-induced RVP by 75 and 73% (P < 0.001), respectively, compared with PK and CA-1 alone (Fig. 5). In contrast, ASP-440 did not alter VEGF-mediated RVP. These findings confirm that PK inhibition blocks CA-1-mediated RVP (8) but not VEGF-mediated RVP, which has been shown to be inhibited by >95% after intravitreal injection of the protein kinase Cβ inhibitor, LY333531 (24). These results suggest that ASP-440’s inhibition of either AngII-induced RVP or diabetes-induced RVP proceeds via inhibition of endogenous PK and not via a direct effect on endogenous VEGF or protein kinase Cβ.

The effects of PK on RVP were also assessed 24–48 h postinjection by visualizing the leakage of FITC-dextran (molecular mass 2 × 10^6 Da) conjugate. We show that eyes subjected to intravitreal injection of PK displayed focal areas of leakage to FITC-dextran that were not observed in retinas from eyes that received saline vehicle injection (Fig. 6). At least one area of focal leakage was observed on retinal whole mounts in five of eight eyes injected with PK compared with none of eight saline vehicle–injected eyes in diabetic rats, indicating a sustained effect of intravitreal PK on focal leakage. In nondiabetic rats, eyes injected with PK had at least one area of focal leakage in two of eight eyes compared with none of eight eyes injected with saline vehicle. These findings show that intravitreal injection of PK induced sustained focal leakage in both nondiabetic and diabetic rats, with a trend for more leakage in diabetic rats.

**Effect of intravitreal injection of PK on retinal thickness.** The effects of diabetes and intravitreal PK injection on retinal thickness were examined using SD-OCT. OCT scans were obtained for each retina at baseline and at 24 h postinjection. Retinal thickness measurements were registered using en face images relative to the center of the ONH (Figs. 7A–D). Representative en face images and a corresponding B-scan at 500 μm superior to the ONH are shown in Fig. 7. At 24 h postinjection for nondiabetic and diabetic rats that received BSS (Fig. 7A and C) or PK (Fig. 7B and D). B-scans illustrate a single caliper measurement from RPE to ILM at the 500-μm intersection point.
Observations from the en face images indicated that PK injection induced vascular dilation and tortuosity that were not evident in the BSS control retina. Baseline retinal thickness in nondiabetic and diabetic rats were 182 ± 8 and 193 ± 10 μm, respectively (P = 0.024) (Fig. 7E). The reproducibility of measurements from consecutively acquired SD-OCT images was 1.2 ± 1.0 μm. Retinal thickness in nondiabetic rats 24 h postintravitreal injection of BSS or PK was 189 ± 4 and 193 ± 9 μm, respectively. Retinal thicknesses in diabetic rats following BSS or PK intravitreal injection was higher at 196 ± 8 and 223 ± 13 μm (P < 0.01), respectively. Results are summarized in Supplementary Table 2.

To further evaluate the regions of retinal thickness, an edge-detection algorithm was applied to each B-scan to identify the RPE and ILM boundaries. Point differences between layers were measured along each B-scan, and the results were compiled to create a retinal-thickness map. Representative thickness maps for corresponding baseline and 24-h post-PK injection are shown for nondiabetes (Fig. 8A and B) and diabetes (Fig. 8C and D), respectively. Automated segmentation was not possible near the ONH because of the absence of the RPE as a reference position and the presence of hyaloid remnants. This region is indicated in white. Thickness false-color topographic maps illustrate diffuse PK-induced thickening throughout the retina in both nondiabetic and diabetic rats. These thickness topographic maps also show increased retinal thickness coincident with primary vessels, which appeared elevated above the plane of the nerve fiber layer and approximately uniform thickness between these vessels on the 500-μm annulus in the inferior-superior and nasotemporal axes, corresponding to the locations of caliper measurements indicated in Fig. 7.

DISCUSSION

This report provides the first characterization of the effect of PK inhibition on retinal vascular dysfunction in diabetes and the effect of intravitreal injection of PK on retinal thickness. We show that systemic administration of a selective small-molecule PK inhibitor, ASP-440 (19), ameliorated both RVP and RBF abnormalities in diabetic rats. Using a prevention protocol, continuous delivery of ASP-440 for 4 weeks inhibited diabetes-induced increased RVP by >80% without altering blood glucose or body weight. Using an intervention protocol, we show that ASP-440 inhibited diabetes-induced RVP when treatment was initiated 2 weeks after diabetes onset, although this did not affect RVP compared with baseline in nondiabetic normal rats. The efficacious dose range of ASP-440 in diabetic rats was comparable with that observed in rats with 7 days of AngII-induced hypertension. Although ASP-440 inhibited PK-induced RVP, it did not affect VEGF-induced RVP, suggesting that its beneficial effects on pathological RVP are not a result of a direct inhibition of VEGF-induced blood-retinal barrier permeability. These findings suggest that PK activity contributes to the elevated levels of RVP associated with both hyperglycemia and hypertension, both of which are major risk factors for diabetic retinopathy progression and DME (1,3), and also that systemic PK inhibition does not alter RVP in normal, healthy animals. We also show that intravitreal injection of C1-INH decreased RVP in diabetic rats, suggesting that local intracellular activation of the PKKS contributes to increased RVP in diabetic rats. We have previously shown that intravitreal injection of C1-INH blocked CA-1–induced RVP (8), suggesting that PK inhibition may also reduce RVP associated with retinal hemorrhages, which often increase in number and severity during diabetic retinopathy progression.

The effects of the PKKS on vascular permeability at sites of vascular injury and inflammation have been attributed
FIG. 7. Representative en face OCT and B-scan images of rat retina obtained by spectral domain optical coherence tomography (SD-OCT) at 24 h after intravitreal injection and resultant retinal thickness measurements. After 4 weeks’ diabetes duration, baseline SD-OCT scans of the retina from each eye were obtained (100 B-scans of 1,000 A-scans over a 1.5 × 1.5 mm area). Rats were given 10-μL intravitreal injections of PK (50 ng/eye) and BSS in the contralateral eye. At 24 h after injection, repeat SD-OCT scans of the retina from each eye were obtained. A–D: The center of the optic nerve was identified, and a 500-μm radius was drawn on the en face image as shown in. The intersection of the radii terminus and the corresponding B-scan identified the retinal sites for thickness measurement. Retinal thickness was measured on the intersecting B-scan from the RPE to the ILM using calipers calibrated in microns. En face images were saved for retinal registration between timed SD-OCT scans. En face and B-scan slices are shown for nondiabetes (NDM) plus BSS (A), nondiabetes plus PK (B), diabetes (DM) plus BSS (C), and diabetes plus PK (D). B-scans correspond to the horizontal green line shown on each en face image. A single caliper is drawn on the corresponding B-scan at the 500-μm point from the ONH. B-scans of PK-treated nondiabetic and diabetic retina illustrate increased thickness compared with BSS alone. En face images...
to the production and actions of BK peptides (12,14). Both B1 and B2 receptors are expressed in retina, and their direct activation via intravitreal injections of BK and des-Arg^9-BK, respectively, has been shown to increase RVP (17,19). Abdouh et al. (17) have shown that retinal B1 receptor expression and des-Arg^9-BK–induced RVP were increased in diabetic rats compared with nondiabetic controls and that 7 days of systemic treatment with a B1 receptor antagonist significantly reduced diabetes-induced RVP. Likewise, Lawson et al. (25) reported that B1 receptor antagonism decreased RVP in Wistar rats with 4 weeks of STZ-induced diabetes. These reports suggest that activation of the PKKS can increase RVP via both B1 and B2 receptors and that diabetes appears to increase actions mediated via the B1 receptor. However, although upregulation of retinal B1 receptors has been demonstrated in rodent models with short duration of diabetes (6,17), there is no information currently available on retinal B1 receptor expression levels in diabetic retinopathy in humans. Kedzierska et al. (27) have reported that circulating PPK levels are elevated in people with diabetic retinopathy. In the current study, we show that PPK levels in the plasma of diabetic rats were elevated compared with nondiabetic control rats (Supplementary Fig. 3), which is consistent with previous findings using STZ-induced diabetic Wistar rats (28,29). Lawson et al. (25,30) have shown that systemic administration of the B1 antagonist R-954 in diabetic rats decreased vascular permeability in other organs, including aorta, heart, kidney, and skin. Taken together, these findings suggest that systemic activation of the PKKS may contribute to increased vascular permeability in multiple tissues that are associated with diabetic vasculopathies.

We have previously shown that the retinal MCT, which measures the rate of hemodynamics across the retinal microvasculature, is prolonged, and the RBF is decreased in diabetic rats with 2 weeks of STZ-induced diabetes (22). In this report, we show that the abnormalities in MCT and RBF were significantly ameliorated in rats that received ASP-440 for 2 weeks in the intervention protocol. It is surprising that we found that ASP-440 increased retinal vessel diameters in diabetic rats, suggesting that activation of endogenous PK in diabetic rats leads to vasoconstriction of retinal vessels. Although acute administration of BK has been shown to increase retinal vessel diameters (26), the effects of chronic PK activation and increased RVP on retinal hemodynamics are not yet known. Although the role of retinal hemodynamics on the pathology of diabetic retinopathy is not fully understood, decreases in RBF have been described for diabetic patients with no diabetic retinopathy or mild nonproliferative diabetic retinopathy (31–33), suggesting that it is associated with early retinal vascular responses to diabetes and diabetic retinopathy. The reduction in RVP and the increase in RBF observed in diabetic rats that received ASP-440 compared with vehicle-treated diabetic controls show that PK inhibition ameliorates two early retinal vascular abnormalities caused by diabetes.

We have reported previously (7,8) that PK, high-molecular weight kininogen, and FXII are present at elevated levels in vitreous fluid obtained from people with diabetic retinopathy. The vitreous in diabetic retinopathy contains a number of proteins that have been shown to bind and activate components of the PKKS, including collagen and laminin (34,35), misfolded proteins (36), CA-1 (8), and heat shock protein 90 (7,37). We have shown a high ratio of PK relative to PPK in diabetic retinopathy vitreous (8), suggesting that PPK within the vitreous is efficiently activated to PK. Although the mechanisms contributing to PKKS expression in the vitreous are not fully understood, increased RVP in diabetes could enable diffusion of PKKS components from the blood to the retinal interstitial fluid and into the vitreous. In this report, we demonstrate that the magnitude of the acute increase in RVP to fluorescein and the prolonged increase in focal areas of RVP to FITC–dextran conjugate 24 h post-PK intravitreal injection were greater in diabetic rats compared with nondiabetic controls. Recently, using a proteomics approach, we have shown that PK induces the proteolysis of extracellular matrix proteins released from astrocytes, including collagen, fibronectin, laminin, and ntidogen (38). These findings suggest that PK-induced proteolysis of basement membrane may contribute to its sustained effects on the retina.

We demonstrate that intravitreal injection of PK in diabetic rats results in an increase in retinal thickness measured using SD-OCT. We used two methods to evaluate retinal thickness using SD-OCT data collected at baseline and at 24 h after intravitreal injection. The first method involved the generation of a 500-μm annulus on the fundus image centered on the ONH that was used to define four regions for retinal thickness measurements from B slices using calipers. By use of this approach, we were able to perform image registration for pre- and post-injection measurements of retinal thickness. This method enabled reproducibility of retinal-thickness measurements with a tolerance of 1–2 μm. In the second method, we used an algorithm to automate segmentation of B slices and compiled thickness data to generate thickness topographic maps. Thicknesses were visualized using a colored-scale heat map centered at 200 μm. This method provided the advantage of a wider field of visualization of retinal thickness compared with the caliper approach described above. These thickness maps indicated that PK-induced retinal thickening was relatively uniform in all quadrants of the retina. We show that PK intravitreal injection in diabetic rats resulted in a marked increase in thickness compared with both baseline measurements and to the response observed in nondiabetic rats, a response consistent with the PK-induced changes in RVP. Our results suggest that the level of RVP induced by diabetes alone may not be sufficient to induce a robust increase in retinal thickness in rats. This is consistent with clinical findings that have demonstrated a correlation between DME and the severity of RVP (39). Previous studies based on vitreous proteomics have suggested that RVP and hemorrhage can
activate pathways that are additive to those induced by diabetes alone and that this additive effect may overwhelm compensatory mechanisms that mediate retinal fluid balance (8). Our results suggest that increased PK levels in the vitreous may be one of the factors that contribute to retinal thickening in diabetes.

In conclusion, this report shows that inhibition of PK reduces RVP in diabetic rats and describes a new model of...
retinal thickening that is generated by the intravitreal injection of PK in diabetic rats. These findings suggest that either systemic administration of a small-molecule PK inhibitor, such as ASP-440, or intravitreal injection of C1-INH, a physiological PK inhibitor, may provide new pharmaceutical treatment options to reduce RVP associated with diabetic retinopathy and DME.

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