Prevalence of diabetic retinopathy and diabetic macular edema in a primary care-based teleophthalmology program for American Indians and Alaskan Natives

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Abstract

Background

Diabetes and its complications are more common in American Indians and Alaska Natives (AI/AN) than other US racial/ethnic populations. Prior reports of diabetic retinopathy (DR) prevalence in AI/AN are dated, and research on diabetic macular edema (DME) is limited. This study characterizes the recent prevalence of DR and DME in AI/AN using primary care-based teleophthalmology surveillance.

Methods

This is a multi-site, clinic-based, cross-sectional study of DR and DME. The sample is composed of AI/AN patients with diabetes (n = 53,998), served by the nationally distributed Indian Health Service-Joslin Vision Network Teleophthalmology Program (IHS-JVN) in primary care clinics of US Indian Health Service (IHS), Tribal, and Urban Indian health care facilities (I/T/U) from 1 November 2011 to 31 October 2016. Patients were recruited opportunistically for a retinal examination using the IHS-JVN during their regular diabetes care. The IHS-JVN used clinically validated, non-mydriatic, retinal imaging and retinopathy assessment protocols to identify the severity levels of non-proliferative diabetic retinopathy (NPDR), proliferative diabetic retinopathy (PDR), DME, and sight threatening retinopathy (STR; a composite measure). Key social-demographic (age, gender, IHS area), diabetes-related health (diabetes therapy, duration of diabetes, A1c), and imaging technology variables were examined. The analysis calculated frequencies and percentages of severity levels of disease.

Results

Prevalence of any NPDR, PDR, DME, and STR among AI/AN patients undergoing DR teleophthalmology surveillance by IHS-JVN was 17.7%, 2.3%, 2.3%, and 4.2%, respectively.
Prevalence was lowest in Alaska and highest among patients with A1c $>/= 8\%$, duration of diabetes $>10$ years, or using insulin.

**Conclusions**

Prevalence of DR in this cohort was approximately half that in previous reports for AI/AN, and prevalence of DME was less than that reported in non-AI/AN populations. A similar reduction in diabetes related end-stage renal disease in the same population and time period has been reported by other researchers. Since these two diabetic complications share a common microvasculopathic mechanism, this coincident change in prevalence may also share a common basis, possibly related to improved diabetes management.

**Introduction**

American Indians and Alaska Natives (AI/AN) have an age-adjusted prevalence of diagnosed diabetes that is 2.0 times that of non-Hispanic whites. Prevalence varies by region from 6.0% among AN to 22.2% among AI in certain areas of the Southwest [1]. Diabetic retinopathy (DR) is the most common microvascular complication of diabetes [2]. The few published studies of DR in AI/AN populations have documented prevalence rates of 35% to 49% for non-proliferative diabetic retinopathy (NPDR) and 3% to 10% for proliferative diabetic retinopathy (PDR) [3–6]. These studies, conducted in the 1980s and 1990s individually included cohorts living in the US Southwest, Northern Plains, or Oklahoma. Similarly, in a Canadian First Nations Indian population, the prevalence of NPDR reported in 2007 was approximately 40% [7]. The prevalence of diabetic macular edema (DME) has not been reported among AI/AN.

DR [8] and DME [9] are leading causes of severe and moderate vision loss among working age adults in the US despite the effectiveness of timely diagnosis and treatment. Because approximately half the AI/AN population with diabetes fails to obtain the recommended annual retinal examination required to allow appropriate DR management, the Indian Health Service (IHS) implemented a primary care-based teleophthalmology program in 2000 to increase compliance with DR surveillance standard of care [10], and improve efficiencies [11].

This particular program, called the IHS-Joslin Vision Network Teleophthalmology Program (IHS-JVN) [12], uses a non-mydriatic retinal imaging protocol validated to American Telemedicine Association Category 3 [13–16]. The program has a diagnostic accuracy sufficient to determine levels of DR and DME commensurate with the Early Treatment Diabetic Retinopathy Study (ETDRS) clinical evaluation [15,16]. The IHS is a patient centered, public health-focused, federal health care organization serving AI/ANs in 35 states. AI/AN experience significant disparities in access to specialty care, in part due to their disproportionately rural distribution [17]. IHS-JVN has been integrated within existing IHS diabetes care programs to assess DR and DME without need for specialty referral or pupil dilation, thereby improving access to DR surveillance critical for timely management and prevention of avoidable vision loss [10].

The present study analyzed data from the IHS-JVN to characterize the prevalence of DR and DME among AI/AN undergoing teleophthalmology surveillance. Previous studies of DR prevalence in AI/AN populations were limited to only a few geographic areas, had limited tribal representation, and did not address DME. In addition, most were published two to three decades prior to the outcomes of clinical trials demonstrating reduction of diabetes related end-organ disease by improved diabetes management [18–21]. This study expands and
updates the literature on these retinal conditions among AI/AN, with implications for health care providers serving AI/AN, regulators, and for Indigenous populations globally.

Materials and methods

Sample

This was a retrospective data analysis of AI/AN persons with diabetes evaluated by the IHS-JVN. The IHS-JVN was started in 2000 and took its first images as a distributed clinical program in 2002. As of September 2017, the IHS-JVN was operational in 11 of the 12 IHS administrative areas spanning the United States, deployed in 142 IHS, Tribal, and Urban clinics, and conducted over 162,122 evaluations for DR (also known as “studies”). The IHS-JVN program is estimated to have a catchment population of 60–70% of AI/AN with known DM [22]. Thus, the IHS-JVN is an extensive program with a nationally geographic pattern that includes broad tribal representation and aligns with the frequency distribution of AI/AN people in the United States. This acts to minimize the impact of geographic and tribal variations in DR that is present in all previous reports of DR prevalence among AI/AN, as well as the impact of locations not served by the IHS-JVN.

This retrospective data analysis examines the cohort of patients evaluated by IHS-JVN from 1 November 2011 to 31 October 2016. The total number of facilities participating in the IHS-JVN in this timeframe was 96. Typically, these facilities are the predominate or only source of diabetes care available to AI/AN patients in the community. They are staffed by community health nurses and support staff who facilitate patient attendance to services, particularly patients with chronic disease like diabetes. In each of these facilities, patients with diabetes were recruited for retinal imaging consecutively during their primary care appointments, as a routine component of their standard diabetes care. Thus, evaluation by IHS-JVN did not require formal referral or a separate appointment, thereby mitigating barriers to access and the associated selection bias.

Ethics statement

This analysis was approved by the IHS National Investigational Review Board. The data were de-identified in accordance with Health Insurance Portability and Accountability Act (HIPAA) regulations.

Technology and protocol

The IHS-JVN uses two technical configurations for acquisition of retinal images. Both have been validated against the gold standard of ETDRS 7 standard fields photography. The first configuration uses a 45° (degree) field of view (FOV), low-illumination, nonmydriatic fundus photography (NMFP) digital imaging system (Topcon NW6S; Topcon Medical Systems, Inc., Paramus, NJ) with a custom digital camera back that is effective in low-light conditions (Mega Vision Retinal Image Capture; Santa Barbara, CA) [16]. Using the NMFP system, three 45° and two 30° FOV stereo pair digital images of the retina and one external image of the anterior segment for each eye are obtained by certified imagers as previously described [15]. Silva and colleagues found this configuration has near perfect agreement with the gold standard of ETDRS photography for diagnosis of DR severity levels ($\kappa = 0.81$, 95% confidence interval = 0.73–0.89) [16]. The second configuration uses nonmydriatic ultrawide-field (UWFI) scanning laser ophthalmoscopy (SLO) (Daytona, Optos, plc, Dunfermline, United Kingdom). It was first introduced in the IHS-JVN 1 October 2014 [14], and staged incrementally across 21 sites as upgrades to previous NMFP systems. The UWFI protocol includes a single macula-centered
200˚ FOV stereo pair image from each eye. Silva and colleagues found exact DR severity agreement between UWFI and ETDRS photography occurred in 84% of cases, and agreement within one level of severity occurred in 91% of cases (weighted $\kappa = 0.85$ and unweighted $\kappa = 0.79$) [23]. More IHS clinics will adopt UWFI over time, but currently the IHS-JVN uses both technologies because NMFP is less costly and relatively portable, allowing the program to access smaller and more remotely located populations that would otherwise not be represented here.

The retinal images were securely transmitted to a central reading center for grading according to a standard validated protocol by certified optometrist tele-retinal readers with ophthalmologist supervision. The IHS-JVN reading software utilized computer assisted decision support based upon ETDRS criteria to further facilitate standardized grading of the retinal images. Pertinent health information for each patient was obtained from the IHS electronic health record (EHR). The reader renders a diagnosis of DR and DME severity and recommends a management plan in a report sent to the patient’s primary care provider (or designee) for further management of the patient. Ongoing quality assurance through a structured process of monthly administrative and clinical review provided an evidence basis for continued reader certification, and ongoing clinical performance consistent with the program’s original validation studies.

For the present study, diagnoses for both eyes were combined to create a person-level diagnosis representing the most severe level of DR and DME observed. If the diagnoses between a patient’s eyes differed, the diagnosis from the eye that indicated the most severe level of disease was selected. Each image underwent rule-based assessment for gradability [24]. If images for one eye could not be interpreted due to the technical quality of the images, but images for the fellow eye could be interpreted conclusively, the diagnosis from the interpreted images was used.

If a patient was imaged more than once in the five-year timeframe, the case indicating the most severe level of DR or DME was recorded. Otherwise, if the patient had no signs of DR or DME, or all records during the selected timeframe documented the same level of disease, the earliest case was recorded. Therefore, each patient is represented in the dataset only once.

**Measures**

**Outcomes.** The IHS-JVN identified the severity levels of DR and DME based on ETDRS criteria and consistent with the International Classification System for Diabetic Retinopathy and Macular Edema [25].

If the technical quality of the images was insufficient to make a conclusive determination of the level of DR or DME, the imager could not obtain all images for the patient, or there were technical difficulties resulting in an incomplete imaging protocol, the reader reported them as ‘ungradable’ for that condition. Because stereo imaging and overlapping retinal fields provided redundancy of data within a single retinal field, images may be ungradable for one condition, but sufficient to obtain a grade for others. Except for an initial examination of background characteristics, patients with ungradable images are excluded from the analyses, consistent with previous research [26,27]. A case designation of ‘ungradable’ resulted in an automatic referral for a conventional dilated retinal exam.

For DR, the possible outcomes were: 1) no apparent DR; 2) mild NPDR, meaning microaneurysms only; 3) moderate NPDR, meaning more than microaneurysms but less than severe NPDR; 4) severe NPDR, indicated by intra-retinal hemorrhage in each of the four quadrants or venous beading in two or more quadrants or intraretinal microvascular abnormalities (IRMAs) in one or more quadrants, but no PDR; and 5) evidence of PDR, indicated by neovascularization and/or vitreous preretinal hemorrhage.
For DME, the outcomes were: 1) absent; 2) not clinically significant, characterized by retinal thickening or hard exudates at or within 3,000 microns from the fovea or thickening in the posterior pole within the arcades that is outside the threshold for Clinically Significant Diabetic Macular Edema (CSDME); and 3) CSDME, characterized by retinal thickening at or within 500 microns of the fovea, hard exudates at or within 500 microns of the fovea with adjacent retinal thickening, one or more disc areas of retinal thickening any part of which is within 1,500 microns of the fovea or with center involvement [28].

We also determined the outcome for sight threatening retinopathy (STR), defined as present if severe NPDR and/or PDR and/or any DME was evident.

Health summary data. The IHS-JVN readers were presented with a five-year health summary that is obtained automatically from the IHS EHR and supplemented as needed by the imager. The summary includes evidence-based risk factors for the progression of DR [9,29] to guide the care plan. The health summary data included glycemic control, body mass index (BMI), smoking status, family history of diabetes, and presence of hypertension, hypercholesterolemia, nephropathy, and peripheral neuropathy. However, for health summary data accessible to this study, glycemic control was the most reliably recorded datum. Glycemic control was characterized as: A1c of less than 6%; A1c of 6% to 7.9%; A1c of 8% to 10%; A1c of greater than 10%; actual A1c not recorded, but ‘poor glycemic control’ is recorded; or missing.

The IHS-JVN software provides automated collection of diabetes duration from the EHR based upon time of diagnosis, rather than the patients’ recollection. Duration of diabetes is presented as a categorical variable. The software also collects information on the patients’ diabetes treatment, which includes diet only, oral diabetes medications, insulin only, insulin and oral medications, or unknown.

Social-demographics and technology used. The IHS-JVN software provides automated collection of demographic information from the patient’s EHR, such as age, gender, and imaging clinic name presented as categorical variables. Clinics were matched to the IHS’s twelve administrative areas and then consolidated into the following geographical areas: Southwest, Oklahoma, Northwest, Northern Plains, East of the Mississippi River, and Alaska. The type of technology used (NMFP or UWFI) was also documented.

Statistical analysis

First, the IHS-JVN cohort was characterized by calculating frequencies, column and row percentages (to show the conditional distributions) for the social-demographics, and health summary and technology variables. Although the focus of this analysis is patients with gradable images, consistent with previous research, the analyses compared the conditional distributions of whether images were gradable or ungradable using chi-square tests. Second, to obtain overall prevalence estimates, the analyses calculated the numbers and column percentages of the IHS-JVN population for each DR, DME, and STR severity level. Third, the analyses calculated the frequencies and percentages for each level of DR, DME, and STR by social-demographics, health summary data, and technology used, and conducted chi-square tests of independence. Lastly, although not the primary focus of this study, the analyses estimated multinomial logit models with all aforementioned variables (not shown, but available upon request) to document their net effects on DR and DME. All analyses were done using SAS 9.4 (Cary, NC).

Results

In the examined timeframe, 53,998 patients were imaged, 86.3% of which had gradable images (Table 1). Of those with gradable images, 40.3% were under age 50 and 8.8% were 70 years and older (mean age = 52.7 ± 12.8 years). The majority of patients were female (56.0%) and lived in
Table 1. Characteristics of the IHS-JVN patients, by gradable or ungradable images.

| Characteristic                         | Patients with gradable images for DR or DME (n = 46584) | Patients with ungradable images for both DR and DME (n = 7414) | Hypothesis tests of independence, gradable vs. ungradable (n = 53998) |
|----------------------------------------|---------------------------------------------------------|---------------------------------------------------------------|---------------------------------------------------------------------|
|                                        | n  | Column % | Row %  | n  | Row % | Chi-Sq (df) | P-value |
| Social-demographics                    |    |          |        |    |        |             |         |
| Age                                    |    |          |        |    |        |             |         |
| Less than 50 years                     | 18793 | 40.3     | 93.2   | 1381 | 6.8    | 2844.7 (3) | < 0.0001 |
| 50 to 59 years                         | 13977 | 30.0     | 88.6   | 1805 | 11.4   |             |          |
| 60 to 69 years                         | 9737  | 20.9     | 80.7   | 2324 | 19.3   |             |          |
| 70 years and older                     | 4077  | 8.8      | 68.2   | 1904 | 31.8   |             |          |
| Gender                                 |    |          |        |    |        |             |         |
| Male                                   | 20476 | 44.0     | 85.0   | 3627 | 15.0   | 64.3 (2)   | < 0.0001 |
| Female                                 | 26105 | 56.0     | 87.3   | 3786 | 12.7   |             |          |
| Area                                    |    |          |        |    |        |             |         |
| Southwest                              | 26829 | 57.6     | 87.0   | 4008 | 13.0   | 191.2 (5)  | < 0.0001 |
| Alaska                                 | 499   | 1.1      | 88.3   | 66   | 11.7   |             |          |
| East of Mississippi River              | 1743  | 3.7      | 79.0   | 462  | 21.0   |             |          |
| Northern Plains                        | 4401  | 9.4      | 82.4   | 941  | 17.6   |             |          |
| Northwest                              | 5326  | 11.4     | 86.8   | 807  | 13.2   |             |          |
| Oklahoma                                | 7786  | 16.7     | 87.3   | 1130 | 12.7   |             |          |
| Health summary data                    |    |          |        |    |        |             |         |
| Duration of diabetes                   |    |          |        |    |        |             |         |
| Less than 1 year                       | 5780  | 12.4     | 91.9   | 511  | 8.1    | 1015.2 (4) | < 0.0001 |
| 1 to 5 years                           | 12862 | 27.6     | 90.5   | 1347 | 9.5    |             |          |
| 6 to 10 years                          | 9237  | 19.8     | 88.3   | 1220 | 11.7   |             |          |
| More than 10 years                     | 14769 | 31.7     | 80.1   | 3668 | 19.9   |             |          |
| Unknown/missing                         | 3936  | 8.4      | 85.5   | 668  | 14.5   |             |          |
| Diabetes therapy                       |    |          |        |    |        |             |         |
| Diet only                              | 4287  | 9.2      | 87.8   | 593  | 12.2   | 373.3 (4)  | < 0.0001 |
| Oral medications                       | 23990 | 51.5     | 88.4   | 3152 | 11.6   |             |          |
| Insulin only                           | 5467  | 11.7     | 80.5   | 1324 | 19.5   |             |          |
| Insulin & oral medications             | 8638  | 18.5     | 83.5   | 1710 | 16.5   |             |          |
| Unknown/missing                         | 4202  | 9.0      | 86.9   | 635  | 13.1   |             |          |
| Glycemic control                       |    |          |        |    |        |             |         |
| A1c of less than 6% (< 42 mmol/mol)     | 4393  | 9.4      | 87.3   | 638  | 12.7   | 77.9 (5)   | < 0.0001 |
| A1c of 6 to 7.9% (42 to 63 mmol/mol)    | 16499 | 35.4     | 86.9   | 2488 | 13.1   |             |          |
| A1c of 8 to 10% (64 to 86 mmol/mol)     | 8788  | 18.9     | 86.2   | 1409 | 13.8   |             |          |
| A1c of greater than 10% (> 86 mmol/mol) | 8980  | 19.3     | 87.2   | 1320 | 12.8   |             |          |
| Missing but poor glycemic control noted| 797   | 1.7      | 82.6   | 168  | 17.4   |             |          |
| Unknown/missing                         | 7127  | 15.3     | 83.7   | 1391 | 16.3   |             |          |
| Technology                              |    |          |        |    |        |             |         |
| NMFP                                   | 30049 | 64.5     | 81.2   | 6948 | 18.8   | 2529.8 (1) | < 0.0001 |
| UWFI**                                 | 16535 | 35.5     | 97.3   | 466  | 2.7    |             |          |

Column percentages are shown to characterize the analytic sample, patients with gradable images only.
Row percentages are shown to examine differences in the likelihood of having gradable images, based on patient characteristics.
All chi-square tests examining likelihood of having gradable versus ungradable images (by characteristic) indicate the 2 groups differ along these characteristics.

* 3 people were ‘other’ for gender and were excluded.
** UWFI became available in this program Sept 2014.

IHS-JVN = Indian Health Service-Joslin Vision Network Teleophthalmology Program; DR = diabetic retinopathy; DME = diabetic Macular Edema; NMFP = nonmydriatic flash photography; UWFI = ultrawide-field imaging; df = degrees of freedom.

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the Southwest (57.6%). About 31.7% of patients had a diagnosis of diabetes for more than 10 years. Most patients were on oral medications alone for their diabetes therapy (51.5%). The most recent A1c was less than 6.0% (42 mmol/mol) for 9.4% of patients, 6.0 to 7.9% (42 to 63 mmol/mol) for 35.4% of patients, and 8% (64 mmol/mol) or greater for 38.2% of patients. For the remainder of patients, ‘poor glycemic control’ was noted or A1c data were missing.

Patients with gradable images tended to be younger, female, had shorter duration of diabetes, not taking insulin, and had a recorded value for A1c (i.e., not missing data). Because the clinics east of the Mississippi River were more recent additions to the IHS-JVN and were smaller, the imagers were less experienced and the area’s percentage of gradable images was lower. All chi-square tests comparing the characteristics of patients with gradable images versus the characteristics of patients with ungradable images were statistically significant, rejecting the null hypothesis of no differences between groups.

The total percentage of patients with any DR was 20.0% (Table 2), with 17.7% having NPDR and 2.3% having PDR. Prevalence of any DME was 2.3%. Prevalence of STR was 4.2%.

Compared with other age groups, a higher percentage of patients aged 60 years and older had mild NPDR (Table 3), and a higher percentage of patients less than age 60 years had moderate NPDR. A slightly higher percentage of males had moderate NPDR. With respect to geography, the highest percentage of patients with no DR was in Alaska, whereas the highest percentage of patients with PDR was in the Southwest. The percentages of people with any level of DR greater than ‘no apparent’ increased in expected ways when risk factors were considered; i.e., percentages were higher among patients with longer duration of diabetes and patients taking insulin alone or with oral medications. Higher percentages of mild and moderate NPDR were found using UWFI than NMFP, but there was no difference in percentage of severe NPDR. UWFI identified PDR twice as frequently as did NMFP.

Table 2. Numbers (n) and percentages (%) of IHS-JVN patients by level of DR and DME.

| Severity level | n   | %    |
|----------------|-----|------|
| DR             |     |      |
| No Apparent DR | 36381 | 80.0 |
| Mild NPDR      | 4284 | 9.4  |
| Moderate NPDR  | 3698 | 8.1  |
| Severe NPDR    | 67   | 0.1  |
| PDR            | 1052 | 2.3  |
| Total          | 45482| 100.0|
| DME            |     |      |
| Absent         | 44806| 97.7 |
| Not Clinically Significant | 653 | 1.4 |
| CSDME          | 394  | 0.9  |
| Total          | 45853| 100.0|
| STR            |     |      |
| Absent         | 43055| 95.8 |
| Present        | 1904 | 4.2  |
| Total          | 44959| 100.0|

Excludes patients with ungradable images.
The different total n for DR and DME is due to differing ungradable rates.

IHS-JVN = Indian Health Service-Joslin Vision Network; Teleophthalmology Program; DR = diabetic retinopathy; NPDR = nonproliferative diabetic retinopathy; PDR = proliferative diabetic retinopathy; DME = diabetic macular edema; CSDME = clinically significant DME; STR = sight threatening retinopathy.

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Table 3. Numbers (n) and percentages (%) of DR severity level among IHS-JVN patients, by their characteristics.

| Characteristic                  | No Apparent DR | Mild NPDR | Moderate NPDR | Severe NPDR | Evidence of PDR | Hypothesis tests of independence |
|---------------------------------|----------------|-----------|---------------|-------------|-----------------|----------------------------------|
|                                 | n   | %   | n   | %   | n   | %   | n   | %   | Chi-Sq (df) | P-value |
| Age                             |     |     |     |     |     |     |     |     |             |         |
| Less than 50 years              | 14826 | 80.5 | 1440 | 7.8 | 1692 | 9.2 | 38   | 0.2 | 413 | 2.2 | 292.2 (12) | < 0.0001 |
| 50 to 59 years                  | 10868 | 79.6 | 1260 | 9.2 | 1195 | 8.7 | 20   | 0.1 | 315 | 2.3 |            |         |
| 60 to 69 years                  | 7576  | 79.9 | 1031 | 10.9| 630  | 6.6 | 6    | 0.1 | 233 | 2.5 |            |         |
| 70 years and older              | 3111  | 79.0 | 553  | 14.0| 181  | 4.6 | 3    | 0.1 | 91  | 2.3 |            |         |
| Gender                         |     |     |     |     |     |     |     |     |             |         |
| Male                           | 15525 | 77.9 | 1921 | 9.6 | 1953 | 9.8 | 27   | 0.1 | 495 | 2.5 | 147.4 (4)  | < 0.0001 |
| Female                         | 20853 | 81.6 | 2363 | 9.2 | 1745 | 6.8 | 40   | 0.2 | 557 | 2.2 |            |         |
| Area                           |     |     |     |     |     |     |     |     |             |         |
| Southwest                      | 20508 | 78.0 | 2516 | 9.6 | 2559 | 9.7 | 35   | 0.1 | 674 | 2.6 | 317.4 (20) | < 0.0001 |
| Alaska                         | 452  | 92.2 | 25   | 5.1 | 10   | 2.0 | 0    | 0.0 | 3   | 0.6 |            |         |
| East of Mississippi River      | 1357  | 82.3 | 154  | 9.3 | 100  | 6.1 | 1    | 0.1 | 37  | 2.3 |            |         |
| Northern Plains                | 3455  | 80.6 | 399  | 9.3 | 324  | 7.6 | 9    | 0.2 | 99  | 2.3 |            |         |
| Northwest                      | 4279  | 82.9 | 458  | 8.9 | 337  | 6.5 | 8    | 0.2 | 78  | 1.5 |            |         |
| Oklahoma                       | 6330  | 83.2 | 732  | 9.6 | 368  | 4.8 | 14   | 0.2 | 161 | 2.1 |            |         |
| Duration of diabetes           |     |     |     |     |     |     |     |     |             |         |
| Less than 1 year               | 5362  | 94.5 | 148  | 2.6 | 133  | 2.3 | 3    | 0.1 | 31  | 0.5 | 5901.4 (16) | < 0.0001 |
| 1 to 5 years                   | 11713 | 92.9 | 475  | 3.8 | 367  | 2.9 | 4    | 0.0 | 52  | 0.4 |            |         |
| 6 to 10 years                  | 7508  | 83.2 | 786  | 8.7 | 610  | 6.8 | 13   | 0.1 | 103 | 1.1 |            |         |
| More than 10 years             | 8569  | 59.8 | 2593 | 18.1| 2351 | 16.4| 43   | 0.3 | 777 | 5.4 |            |         |
| Unknown/missing                | 3229  | 84.1 | 282  | 7.3 | 237  | 6.2 | 4    | 0.1 | 89  | 2.3 |            |         |
| Diabetes therapy               |     |     |     |     |     |     |     |     |             |         |
| Diet only                      | 3971  | 94.5 | 124  | 3.0 | 72   | 1.7 | 0    | 0.0 | 34  | 0.8 | 4588.9 (16) | < 0.0001 |
| Oral medications               | 20393 | 87.0 | 1543 | 6.6 | 1263 | 5.4 | 17   | 0.1 | 232 | 1.0 |            |         |
| Insulin only                   | 2994  | 56.3 | 1017 | 19.1| 906  | 17.0| 20   | 0.4 | 382 | 7.2 |            |         |
| Insulin & oral medications     | 5466  | 65.0 | 1358 | 16.2| 1230 | 14.6| 25   | 0.3 | 328 | 3.9 |            |         |
| Unknown/missing                | 3557  | 86.6 | 242  | 5.9 | 227  | 5.5 | 5    | 0.1 | 76  | 1.9 |            |         |
| Glycemic control               |     |     |     |     |     |     |     |     |             |         |
| A1c of less than 6% (< 42 mmol/mol) | 3974 | 92.7 | 155  | 3.6 | 101  | 2.4 | 0    | 0.0 | 55  | 1.3 | 2416.0 (20) | < 0.0001 |
| A1c of 6 to 7.9% (42 to 63 mmol/mol) | 14077 | 87.2 | 1180 | 7.3 | 619  | 3.8 | 9    | 0.1 | 265 | 1.6 |            |         |
| A1c of 8 to 10% (64 to 86 mmol/mol) | 6195 | 72.4 | 1191 | 13.9| 883  | 10.3| 18   | 0.2 | 269 | 3.1 |            |         |
| A1c of greater than 10% (> 86 mmol/mol) | 5934 | 67.5 | 1108 | 12.6| 1428 | 16.2| 29   | 0.3 | 293 | 3.3 |            |         |
| Missing but poor glycemic control noted | 597  | 77.4 | 87  | 11.3 | 72  | 9.3 | 3    | 0.4 | 12  | 1.6 |            |         |
| Unknown/missing                | 5604  | 80.9 | 563  | 8.1 | 595  | 8.6 | 8    | 0.1 | 158 | 2.3 |            |         |
| Technology                     |     |     |     |     |     |     |     |     |             |         |
| NMFP                           | 24557 | 84.6 | 2345 | 8.1 | 1590 | 5.5 | 50   | 0.2 | 480 | 1.7 | 1214.8 (4) | < 0.0001 |
| UWFI                           | 11824 | 71.8 | 1939 | 11.8| 2108 | 12.8| 17   | 0.1 | 572 | 3.5 |            |         |

Excludes patients with ungradable images; All chi-square tests examine the likelihood of having a certain severity level of DR.

* Row percentage;
**3 people were ‘other’ for gender and were excluded.
IHS-JVN = Indian Health Service-Joslin Vision Network Teleophthalmology Program; DR = diabetic retinopathy; NPDR = nonproliferative diabetic retinopathy; NMFP = nonmydriatic flash photography; UWFI = ultrawide-field imaging; df = degrees of freedom.

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The percentages of patients with DME (not clinically significant) were higher in patients less than 60 years, male, had diabetes for more than 10 years, taking insulin, and with higher A1c or poor glycemic control (Table 4). The percentages of patients with CSDME were higher in patients who had diabetes for more than 10 years, were taking insulin only, and had an A1c of greater than 10%. Rates of DME (not clinically significant) and CSDME were lowest in Alaska; otherwise, there was no meaningful difference by area. There was no difference between NMFP and UWFI in the percentage of patients found to have any DME.

STR was highest for patients less than 70 years of age, males, patients with longer duration of diabetes, patients taking insulin, patients with higher A1c or missing A1c data in the IHS-JVN record, and patients imaged with UWFI (Table 5). It was lowest in people who live in Alaska.

Multinomial logistic regression analyses documented that the net, statistical associations of all the variables included were as expected given prior research on the progression of DR and DME [9,29].

**Discussion**

This study provides data on prevalence and severity of DR and DME in AI/AN who had undergone primary care based retinal imaging by the IHS-JVN from 1 November 2011 to 31 October 2016. The IHS-JVN program is validated, standardized, has robust quality assurance to ensure ongoing fidelity with validation studies, and provides recent retinal imaging data and pertinent medical record information for a geographically representative population of AI/AN, which predominately has type 2 diabetes. The expansive geographic scope of the IHS-JVN, federal funding of IHS health care, and opportunistic recruitment of patients from diabetes primary care workflow mitigates bias in DR prevalence due to local factors such as diet, difference in socio-economic status, health care utilization, and patient selection.

There are few studies of DR in AI/AN populations. Many AI/AN populations live in remote areas, making clinical studies of broad geographical scope difficult. Most of the previous studies are over two decades old. The results from these prior studies are summarized in Table 6.

The prevalence of DR in these earlier studies was higher than that reported here, i.e., in Pima Indians with type 2 diabetes [3], Navajo and Hopi Indians [4], and Sioux Indian tribes [5], the prevalence of NPDR was 37.8%, 40%, and 45.3% respectively. The prevalence of PDR was 2.7% (Pima), and prevalence of vision-threatening retinopathy was 8.2% (Navajo and Hopi). A longitudinal (12.7 years) study in AI examined in Oklahoma found that the overall incidence of PDR among survivors was 18.6%, and 45% of those with background DR (NPDR) at baseline later developed PDR [30]. A 2005 study in Oklahoma showed a DR prevalence of 20.1% [31], which is more consistent with our 2011 to 2016 results (16.7%, Table 3) from that region.

These earlier AI/AN NPDR prevalence rates (ranging between 20% and 49%) are similar to non-native groups documented in the Wisconsin Epidemiologic Study of Diabetic Retinopathy (WESDR), where DR prevalence varied from 29% (duration of diabetes less than 5 years) to 78% (duration of diabetes greater than 15 years), and PDR varied from 2% (duration less than 5 years) to 15.5% (duration greater than 15 years) [32].

Analyses of DR prevalence in diabetic populations, both globally [29] and in the US [26,33], provide a benchmark, though none provide data on AI/AN populations. A meta-analysis on global prevalence of any DR (NPDR and/or PDR and/or DME) [29]—including a total of 35 studies (1980–2008) and data from 22,896 non-AI/AN individuals with diabetes (both type 1 and 2)—showed an overall prevalence of 34.6% for any DR, 7.0% for PDR, 6.8% for DME and 10.2% for STR (PDR and/or DME). For only individuals with type 2 diabetes, the prevalence for
Table 4. Numbers (n) and percentages (%) of level of DME among IHS-JVN patients, by their characteristics.

| Characteristic                        | Absent       | Not Clinically Significant | CSDME         | Hypothesis tests of independence |
|---------------------------------------|--------------|----------------------------|---------------|----------------------------------|
|                                       | n  | %  | n  | %  | n  | %  | Chi-Sq | (df) | P-value |
| Age                                   |    |    |    |    |    |    |        |      |         |
| Less than 50 years                    | 1821 | 97.7 | 284 | 1.5 | 139 | 0.7 | 48.6   | (6)  | < 0.0001 |
| 50 to 59 years                        | 1341 | 97.2 | 242 | 1.8 | 141 | 1.0 |        |      |         |
| 60 to 69 years                        | 9343 | 98.1 | 106 | 1.1 | 76  | 0.8 |        |      |         |
| 70 years and older                    | 3836 | 98.5 | 21  | 0.5 | 38  | 1.0 |        |      |         |
| Gender                                |    |    |    |    |    |    |        |      |         |
| Male                                  | 19585 | 97.4 | 336 | 1.7 | 187 | 0.9 | 17.8   | (2)  | 0.0001   |
| Female                                | 25218 | 98.0 | 317 | 1.2 | 207 | 0.8 |        |      |         |
| Area                                  |    |    |    |    |    |    |        |      |         |
| Southwest                             | 25787 | 97.6 | 402 | 1.5 | 221 | 0.8 | 14.3   | (10) | 0.1617   |
| Alaska                                | 492  | 99.4 | 2  | 0.4 | 1   | 0.2 |        |      |         |
| East of Mississippi River             | 1685 | 98.1 | 18 | 1.0 | 15  | 0.9 |        |      |         |
| Northern Plains                       | 4209 | 97.7 | 66 | 1.5 | 35  | 0.8 |        |      |         |
| Northwest                             | 5123 | 97.6 | 72 | 1.4 | 52  | 1.0 |        |      |         |
| Oklahoma                              | 7510 | 97.9 | 93 | 1.2 | 70  | 0.9 |        |      |         |
| Duration of diabetes                  |    |    |    |    |    |    |        |      |         |
| Less than 1 year                      | 5696 | 99.1 | 29 | 0.5 | 21  | 0.4 | 556.5  | (8)  | < 0.0001 |
| 1 to 5 years                          | 12669 | 99.2 | 63 | 0.5 | 36  | 0.3 |        |      |         |
| 6 to 10 years                         | 8965 | 98.3 | 101| 1.1 | 53  | 0.6 |        |      |         |
| More than 10 years                    | 13703 | 95.4 | 414| 2.9 | 253 | 1.8 |        |      |         |
| Unknown/missing                       | 3773 | 98.0 | 46 | 1.2 | 31  | 0.8 |        |      |         |
| Diabetes therapy                      |    |    |    |    |    |    |        |      |         |
| Diet only                             | 4225 | 99.4 | 14 | 0.3 | 11  | 0.3 | 504.3  | (8)  | < 0.0001 |
| Oral medications                      | 23388 | 98.6 | 205| 0.9 | 134 | 0.6 |        |      |         |
| Insulin only                          | 4995 | 94.7 | 173| 3.3 | 107 | 2.0 |        |      |         |
| Insulin & oral medications            | 8108 | 95.9 | 231| 2.7 | 120 | 1.4 |        |      |         |
| Unknown/missing                       | 4090 | 98.7 | 30 | 0.7 | 22  | 0.5 |        |      |         |
| Glycemic control                      |    |    |    |    |    |    |        |      |         |
| A1c of less than 6% (< 42 mmol/mol)   | 4296 | 99.3 | 16 | 0.4 | 14  | 0.3 | 420.9  | (10) | < 0.0001 |
| A1c of 6 to 7.9% (42 to 63 mmol/mol)  | 16086 | 98.9 | 89 | 0.5 | 85  | 0.5 |        |      |         |
| A1c of 8 to 10% (64 to 86 mmol/mol)   | 8376 | 97.2 | 156| 1.8 | 86  | 1.0 |        |      |         |
| A1c of greater than 10% (> 86 mmol/mol)| 8415 | 95.3 | 279| 3.2 | 134 | 1.5 |        |      |         |
| Missing but poor glycemic control     | 760  | 96.0 | 21 | 2.7 | 11  | 1.4 |        |      |         |
| Unknown/missing                       | 6873 | 97.8 | 92 | 1.3 | 64  | 0.9 |        |      |         |
| Technology                             |    |    |    |    |    |    |        |      |         |
| NMFP                                  | 29014 | 97.8 | 416| 1.4 | 241 | 0.8 | 2.49   | (2)  | 0.2876   |
| UWFI                                  | 15792 | 97.6 | 237| 1.5 | 153 | 0.9 |        |      |         |

Excludes patients with ungradable images; All chi-square tests examine the likelihood of having a certain severity level of DME.

* Row percentage;
** 3 people were 'other' for gender and were excluded.
IHS-JVN = Indian Health Service-Joslin Vision Network Teleophthalmology Program; DME = diabetic macular edema; CSDME = clinically significant diabetic macular edema; NMFP = nonmydriatic flash photography; UWFI = ultrawide-field imaging; df = degrees of freedom.

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any DR was 25.2%, for PDR was 3.0%, and for STR was 6.9%. In the US, an analysis of 2005–2008 National Health and Nutrition Examination Survey (NHANES) data [26] showed a prevalence of DR (NPDR plus PDR) of 28.5%, PDR of 1.5%, CSDME of 2.7% and STR (Severe NPDR, PDR, and CSDME) of 4.4% in a non-AI/AN diabetic patient population over the age of 40.

Table 5. Numbers (n) and percentages (%) of STR among IHS-JVN patients, by their characteristics.

| Characteristic                        | No STR Apparent | STR Apparent | Hypothesis tests of independence |
|---------------------------------------|-----------------|--------------|----------------------------------|
|                                       | n       | %    | n       | %    | Chi-Sq (df) | P-value |
| Age                                   |         |      |         |      |             |         |
| Less than 50 years                    | 17585   | 95.9 | 759     | 4.1  | 8.7        | (3)     | 0.0337  |
| 50 to 59 years                        | 12914   | 95.4 | 627     | 4.6  |            |         |         |
| 60 to 69 years                        | 8928    | 95.9 | 377     | 4.1  |            |         |         |
| 70 years and older                    | 3628    | 96.3 | 141     | 3.7  |            |         |         |
| Gender**                              |         |      |         |      |             |         |
| Male                                  | 18754   | 95.4 | 913     | 4.6  | 14.3       | (1)     | 0.0002  |
| Female                                | 24298   | 96.1 | 991     | 3.9  |            |         |         |
| Area                                  |         |      |         |      |             |         |
| Southwest                             | 24816   | 95.5 | 1174    | 4.5  | 23.6       | (5)     | 0.0003  |
| Alaska                                | 481     | 98.8 | 6       | 1.2  |            |         |         |
| East of Mississippi River             | 1561    | 96.0 | 65      | 4.0  |            |         |         |
| Northern Plains                       | 4041    | 95.6 | 184     | 4.4  |            |         |         |
| Northwest                             | 4912    | 96.4 | 186     | 3.6  |            |         |         |
| Oklahoma                              | 7244    | 96.2 | 289     | 3.8  |            |         |         |
| Duration of diabetes                  |         |      |         |      |             |         |
| Less than 1 year                      | 5580    | 98.7 | 72      | 1.3  | 556.5      | (8)     | < 0.0001 |
| 1 to 5 years                          | 12388   | 98.9 | 137     | 1.1  |            |         |         |
| 6 to 10 years                         | 8688    | 97.3 | 243     | 2.7  |            |         |         |
| More than 10 years                    | 12775   | 90.7 | 1305    | 9.3  |            |         |         |
| Unknown/missing                       | 3624    | 96.1 | 147     | 3.9  |            |         |         |
| Diabetes therapy                      |         |      |         |      |             |         |
| Diet only                             | 4114    | 98.7 | 53      | 1.3  | 1358.1     | (4)     | < 0.0001 |
| Oral medications                      | 22712   | 97.8 | 515     | 2.2  |            |         |         |
| Insulin only                          | 4610    | 88.5 | 599     | 11.5 |            |         |         |
| Insulin & oral medications            | 7677    | 92.5 | 621     | 7.5  |            |         |         |
| Unknown/missing                       | 3942    | 97.1 | 116     | 2.9  |            |         |         |
| Glycemic control                      |         |      |         |      |             |         |
| A1c of less than 6% (< 42 mmol/mol)   | 4149    | 98.2 | 78      | 1.8  | 439.4      | (5)     | < 0.0001 |
| A1c of 6 to 7.9% (42 to 63 mmol/mol)  | 15569   | 97.6 | 389     | 2.4  |            |         |         |
| A1c of 8 to 10% (64 to 86 mmol/mol)   | 7972    | 94.5 | 467     | 5.5  |            |         |         |
| A1c of greater than 10% (> 86 mmol/mol) | 8060    | 92.6 | 643     | 7.4  |            |         |         |
| Missing but poor glycemic control noted | 727    | 94.3 | 44      | 5.7  |            |         |         |
| Unknown/missing                       | 6578    | 95.9 | 283     | 4.1  |            |         |         |
| Technology                             |         |      |         |      |             |         |
| NMFP                                  | 27782   | 96.4 | 1040    | 3.6  | 77.7       | (1)     | < 0.0001 |
| UWFI                                  | 15273   | 94.6 | 864     | 5.4  |            |         |         |

Excludes patients with ungradable images; All chi-square tests examine the likelihood of having STR.

* Row percentage;

** 3 people were ‘other’ for gender and were excluded.

IHS-JVN = Indian Health Service-Joslin Vision Network Teleophthalmology Program; STR = sight threatening retinopathy; NMFP = nonmydriatic flash photography; UWFI = ultrawide-field imaging; df = degrees of freedom.

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The design of this study prevents identification of factors contributing to the lower prevalence of DR as compared with prior studies of AI/AN. However, evidence from studies of non-AI/AN populations suggest that improved diabetes management has resulted in a general decline in diabetes related end-organ disease [34]. In this context, it is noteworthy that the reduction in DR prevalence observed in this study is temporally associated with the IHS Special Diabetes Program for Indians (SDPI), initiated in 1997 [35,36]. The specific character of SDPI programs varied among health care facilities to suit local needs, but was otherwise

| Lead Author [Reference Number] | Rate [4] | Newell [6] | Nagi [3] | Berinstein [5] | Lee [31] | Ross [7] |
|--------------------------------|----------|------------|----------|----------------|----------|----------|
| Study Type                     | Cross-sectional and clinic-based | Cross-sectional and clinic-based | Cross-sectional population based | Cross-sectional | Cross-sectional | Cross-sectional |
| Population/ Sample             | Sample Size | Hopi (77) and Navajo (60) | Predominantly Cheyenne—Arapaho | Pima | Cheyenne River Sioux Tribe and the Oglala Sioux Tribe | Seven Oklahoma tribes | First Nations people of Canada |
| Tribe(s)                       | Location  | North-eastern Arizona, US | Clinton, Oklahoma, US | Gila River Indian Community, South central Arizona, US | Northern Plains, US | Oklahoma, US | Southern Alberta, Canada |
| Study Duration                 | Study Duration | Jul 1979—Jun 1980 | Oct 1985 –July 1987 | Apr 1982-Dec 1990 | 1991 | Sep 1995 –Mar 1998 | 1986 |
| Background Characteristics     | Age (years) | Range: >5 = 20 | Average: 55.8; Range: 27 to 87 | Average: 47; Range: 15 to 88 | Range: 45–75. Average not reported. | Range: 48 to 82. 42.5%; aged 48–59; 36.5% aged 60–69; 21.0% aged 70–82 | Not reported |
| % Male                         | 48.2 | 39.4 | 37 | Not reported | 39.8 | Not reported |
| Diabetes Duration              | Average: 7.5 years (SD not reported) | 0–5 years duration = 30.3%; 6–10 years = 26.1%; 11 + years = 33.8%; the rest unknown | 9.1 years | Averages: 6.4 ± 7.9 years if no DR; 12.3 ± 7.5 if NPDR; 14.2 ± 10.3 if PDR | Not reported | Average: 8.26 ± 7.13 years |
| Medication                     | 40% taking oral medications; 37% taking insulin | 50.7% taking oral medications only; 35.2% taking insulin with or without oral medications | 28% taking oral medications; 34% taking insulin | Not reported | Not reported | Not reported |
| A1C (%)                        | Not reported | Averages: 11.8 ± 3.8 if taking insulin; 12.4 ± 2.8 if not taking insulin | 9.9 ± 2.6 years | Averages: 7.7 ± 2.6 if no DR; 8.9 ± 2.0 if NPDR; 9.3 ± 2.5 if PDR | Not reported | Average: 7.52 ± 2.0 |
| Prevalence                     | DR Any DR = 36% | Any DR = 49.3%, PDR = 21.4% | Any DR = 37.8%, PDR = 2.7% | NPDR = 40.2%; PDR = 5.1% | Any DR = 20.1% | Any DR = 39.7% PDR = 5.5% |
| DME                            | Not reported | Not reported | Not reported | Not reported | DME not reported. CSDME = 2.6% (MD exam) or 2.4% (Reading Center exam) | Not reported |
| Other Comments                  | No difference in prevalence between the 2 tribes. | Data for people with diabetes only reported here. | Risk factors reported by presence or level severity of retinopathy. | DR prevalence not the primary outcome. Number of subjects with diabetes and factors pertinent to diabetes/DR not reported. | 2247 was total sample size, with 232 being natives. Data for natives only reported here. |

DR = diabetic retinopathy; NPDR = nonproliferative diabetic retinopathy; PDR = proliferative diabetic retinopathy; DME = diabetic macular edema; CSDME = clinically significant diabetic macula edema; Sight threatening retinopathy (STR) was reported in studies on nonnative populations, but not for the studies of native populations.

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uniformly available across all facilities serving AI/AN for the purpose of improving diabetes management, and in many instances included the IHS-JVN [36]. This systematic program of public health and population management approaches for diabetes care doubled, or in some cases tripled access to organizational diabetes health services, including diabetes clinics, diabetes clinical teams, nutrition services, culturally tailored diabetes education, physical activity specialists, and weight management programs [36]. It was associated with clinically significant improvements in A1c levels, blood lipid levels, and blood pressure [37], which are risk factors for onset and progression of DR and DME. For example, average A1c among AI/AN dropped from 9.0% in 1996 to 8.1% in 2010 [3]. These and other improvements in diabetes management coincident with the implementation of SDPI have been related to a 54% reduction in diabetes related end stage renal disease (dESRD) among AI/AN [37,38]. This is particularly relevant since dESRD and DR share a similar microvasculopathy etiology. Thus, it is not surprising to see a parallel improvement in prevalence of diabetes-related retinal disease in this population. Similar significant improvements in DR and other diabetes-related complications have recently been observed for non-AI/AN populations in the US and Europe [34,39].

To our knowledge, no other studies have investigated the prevalence of DME in the AI/AN population. Comparisons between this study and previous non-AI/AN studies are difficult because the criteria for DME and CSDME differ across the studies or are not defined. In the US, two separate studies using 2005 to 2008 NHANES data showed a DME prevalence rate of 3.8% in patients aged over 40 years [9] and a CSDME prevalence of 2.7% [24]. Further, in the Wisconsin epidemiologic study of diabetic retinopathy. IV [40], which included patients who were over 30 years of age at diagnosis, the prevalence rates of DME ranged from 3.0% (duration of diabetes less than 5 years) to 28.0% (duration greater than 20 years). In contrast, the Multi-Ethnic Study of Atherosclerosis (MESA) reported a 9.0% DME prevalence [27]. The AI/AN prevalence of DME reported here is lower than that reported in the published literature for non-AI/AN populations. In general, these studies depended upon monoscopic, surrogate indicators of macular edema rather than directly observed retinal thickening by stereoscopic viewing as used by the IHS-JVN. This has the effect of under-reporting DME determined by surrogate indicators in the absence of micro aneurysms or hard exudates, and under-reporting CSDME vs DME. However, studies have shown that this technique is reliable for identification of any level of DME [41,42]. Although a basis for the lower rate of DME observed in this study cannot be established, this difference, in part, may be due to the aforementioned improvements in diabetes management, a potential protective effect of the retinal pigmentation in AN/AI [43], and/or other racial/ethnic differences in the nature of DME.

A potential limitation of this study is selection bias. First, patients with more advanced retinal conditions may already be under specialty ophthalmological care and may have chosen to defer IHS-JVN as a result. Also, patients with diabetic retinal disease may also have poor glycemic control (A1c >7%) and would be at higher risk of mortality [44]. Both possibilities could result in some underreporting of DR and/or DME prevalence. Second, the sample may be biased in that the IHS-JVN accesses people who go to primary care clinics in facilities that participated in the teleophthalmology program. However, due to the aforementioned reach and geographic distribution of the program, and the fact that its services are offered to all known patients with diabetes receiving care at the facility, we believe risk of this type of bias is low. Indeed, unlike many previous reports of DR and DME prevalence, these patients were not recruited in specialty eye clinics, so this powerful source of selection bias was mitigated.

Another potential limitation of this study is that images for some patients were ungradable. Except for glycemic control, factors associated with the outcome of ‘ungradable’ paralleled those for risk of DR and DME [42, 45]. Thus, presence of a risk factor is more likely to result in ungradable images and referrals. But a designation of ungradable does not necessarily mean...
disease is visible or clinically overt. Images may be ungradable because of smaller pupil size and media opacity, both of which are more common with increasing age and duration of diabetes [45]. To address the question of DR and DME prevalence among patients who had ungradable images in the present analysis, we extracted the diagnoses codes from the medical records for a sample of 799 unique patients whose images were ungradable for DR and/or DME in 2013 and 2014 and who got a dilated eye exam within 365 days of that original ungradable finding. Of these patients 1.5% had NPDR “not otherwise specified” (i.e., level unknown), 2.4% had mild NPDR, 2.5% had moderate NPDR, 0.6% had severe NPDR, 2.6% had PDR, and 2.9% had DME. Another 24.0% had a diagnosis code indicating ‘background diabetic retinopathy’ only, with no NPDR severity level specified. These percentages indicate that the rate of sight-threatening diabetic eye disease is not substantially higher among patients who had ungradable images during the period of this study, whereas levels of DR that do not (yet) threaten sight might be. Further analysis of the outcomes for people who had ungradable images is a substantial, separate undertaking to be addressed in a future project.

To further address the question of DR and DME prevalence among patients with ungradable images, several of the tables herein reported findings from the NMFP and the UWFI separately. The ungradable rate using the NMFP was 18.8%, compared with 2.7% using UWFI, both of which are consistent with other reported studies [46]. Despite differences in ungradable rates, the NMFP and UWFI technologies yielded similar findings for prevalence and rates for severe NPDR and any DME in the present analysis, likely due to the more central occurrence of the condition, but the rate for PDR was twice as high with UWFI than with NMFP, likely resulting from the larger aggregate FOV that UWFI provides [46]. Other reports of UWFI imaging for DR suggest that it may be a more sensitive measure of DR in both native and non-native populations. [24,46] This is being explored by the Diabetic Retinopathy Clinical Research Network (DRCRnet) protocol AA [47], and its results may impact future standards for DR identification and risk stratification. Until then, the current standard for DR diagnosis remains ETDRS 7 standard fields photography, with which both technologies used by the IHS-JVN have high levels of agreement. Since all previous reports of DR prevalence were based upon central fields, direct comparisons of prevalence using UWFI is somewhat difficult, so comparison with earlier reports is best aligned technically with our NMFP data. Since our data includes more sensitive UWFI, this report may slightly overstate the prevalence as compared to legacy central field methods. This report shows a substantial decrease in the prevalence of DR using either UWFI or NMFP as compared to previous reports using central fields.

Conclusions
To our awareness, this is the only systemic report of DR and DME among AI/AN, and updates older reports, which were limited to specific regions and AI/AN tribes. The AI/AN prevalence of NPDR, PDR, DME, and STR among AI/AN patients undergoing DR teleophthalmology surveillance by the IHS-JVN was 17.7%, 2.3%, 2.3%, and 4.2% respectively. This represents a decrease in DR prevalence of at least 50% compared to previous reports in the 1980s and 1990s. Identifying trends in DME prevalence is more problematic since comparators are not available for AI/AN, and reporting methods for DME is less standardized than DR. Nonetheless, the DME prevalence shown herein is lower than that in recent reports of non-AI/AN population. This study provides a benchmark for future research on DME in AI/AN.

The reduction in DR prevalence is temporally coincident with the implementation of the IHS SDPI, which has been similarly coincident with improvements in diabetic outcome metrics and dESRD. A simultaneous reduction in risk for these two major diabetes complications
in this at risk population following the implementation of systemic programmatic changes in diabetes care provides clinicians, regulators, and federal funders possible further evidence for adherence to best practices of diabetes care. This information may be particularly helpful for clinicians, chronic disease program managers, and public health decision makers in the IHS and Tribal health care since it better defines a major chronic disease in AI/AN and facilitates the development of effective programs and provides a reliable baseline for evaluating the effects of programs over time.

Further research is required to clarify the impact of ungradable images on the prevalence of DR determined by the IHS-JVN, and to investigate links between improved diabetes care in the IHS population and reduced rates of retinal complications using IHS-JVN data available since 2000.

Supporting information
S1 Table. (XLSX)

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References
1. CDC National Diabetes Statistics Report, 2017. Estimates of diabetes and its burden in the United States. https://www.cdc.gov/diabetes/pdfs/data/statistics/national-diabetes-statistics-report.pdf.
2. Fong DS, Aiello LM, Gardner TW, King GL, Blankenship G, Cavallerano JD, et al. for the American Diabetes Association. Diabetic retinopathy. Diabetes Care. 2003; 26: s99–s102. PMID: 12502630
3. Nagi DK, Pettitt DJ, Bennett PH, Klein R, Knowler WC. Diabetic retinopathy assessed by fundus photography in Pima Indians with impaired glucose tolerance and NIDDM. Diabet Med. 1997; 14: 449–456. https://doi.org/10.1002/(SICI)1096-9136(199706)14:6<449::AID-DIA367>3.0.CO;2-D PMID: 9212309
4. Rate RG, Knowler WC, Morse HG, Bonnell MD, McVey J, Chervenak CL, et al. Diabetes mellitus in Hopi and Navajo Indians. Prevalence of microvascular complications. Diabetes. 1983; 32: 894–899. PMID: 6618018

5. Berenstein DM, Stahn RM, Welty TK, Leonardson GR, Herlihy JJ. The prevalence of diabetic retinopathy and associated risk factors among Sioux Indians. Diabetes Care. 1997; 20: 757–759. PMID: 9135938

6. Newell SW, Tolbert B, Bennett J, Parsley TL. The prevalence and risk of diabetic retinopathy among Indians of southwest Oklahoma. J Okla State Med Assoc. 1989; 82: 414–424. PMID: 2769467

7. Ross SA, McKenna A, Mozejko S, Fick GH. Diabetic retinopathy in native and nonnative Canadians. Exp Diabetes Res. 2007; 2007: 76271. https://doi.org/10.1155/2007/76271 PMID: 18317512

8. Ferris FL, Davis MD, Aiello LM. Treatment of diabetic retinopathy. N Engl J Med. 1999; 341: 667–678. https://doi.org/10.1056/NEJM199908263410907 PMID: 10460819

9. Varma R, Bressler NM, Doan OV, Gleson M, Danese M, Bower JK, et al. Prevalence of and risk factors for diabetic macular edema in the United States. JAMA Ophthalmol. 2014; 132: 1334–1340. https://doi.org/10.1001/jamaophthalmol.2014.2854 PMID: 25125075

10. Wilson C, Horton M, Caravallano J, Aiello LM. Addition of primary care–based retinal imaging technology to an existing eye care professional referral program increased the rate of surveillance and treatment of diabetic retinopathy. Diabetes. 2005; 28: 318–322.

11. Maa AY, Wojciechowski B, Hunt KJ, Dismuke C, Shyu J, Janjua R, et al. Early experience with technology-based eye care services (TECS): A novel ophthalmologic telemedicine initiative. Ophthalmol. 2017; 124: 539–546.

12. Carroll M, Cullen T, Ferguson S, Hogge N, Horton M, Kokesh J. Innovation in Indian Healthcare: Using health information technology to achieve health equity for American Indian and Alaska Native populations. Perspect Health Inf Manag. 2011; 8: 1d.

13. Li HK, Horton M, Bursell SE, Caravallano J, Zimmer-Galler I, Tennant M, et al. American Telemedicine Association Diabetic Retinopathy Telehealth Practice Recommendations Working Group. Telehealth practice recommendations for diabetic retinopathy, second edition. Telemed J E Health. 2011; 17: 814–837. https://doi.org/10.1089/ tmj.2011.0075 PMID: 21970573

14. Silva PS, Caravallano JD, Sun JK, Noble J, Aiello LM, Aiello LP. Nonmydriatic ultrawide field retinal imaging compared with dilated standard 7-field 35-mm photography and retinal specialist examination for evaluation of diabetic retinopathy. Am J Ophthalmol. 2012; 154: 549–559.e2. https://doi.org/10.1016/j.ajo.2012.03.019 PMID: 22626617

15. Bursell SE, Caravallano J, Caravallano A, Clermont AC, Birkmire-Peters D, Aiello LP, et al. Stereo nonmydriatic digital-video color retinal imaging compared to ETDRS 7-field 35-mm stereo color photos for determining level of diabetic retinopathy. Ophthalmol. 2001; 108: 572–585. PMID: 11237913

16. Silva PS, Walia S, Caravallano JD, Sun JK, Dunn C, Bursell SE, et al. Comparison of low-light nonmydriatic digital imaging with 35-mm ETDRS seven-standard field stereo color fundus photographs and clinical examination. Telemed J E Health. 2012; 18: 492–499. https://doi.org/10.1089/tmj.2011.0232 PMID: 22627402

17. US Department of Agriculture, Economic Research Service. American Indians remain disproportionately rural. 2016. https://www.ers.usda.gov/data-products/chart-gallery/gallery/chart-detail/?chartId=77893.

18. Diabetes Control and Complications Trial Research Group, Nathan DM, Genuith S, Lachin J, Cleary P, Crofford O, et al. The effect of intensive treatment of diabetes on the development and progression of long-term complications in insulin-dependent diabetes mellitus. N Engl J Med. 1993; 329: 977–986.

19. Nathan DM, Cleary PA, Backlund JY, Genuith SM, Lachin JM, Orchard TJ, et al; Diabetes Control and Complications Trial/Epidemiology of Diabetes Interventions and Complications (DCCT/EDIC) Study Research Group. Intensive diabetes treatment and cardiovascular disease in patients with type 1 diabetes. N Engl J Med. 2005; 353: 2643–2653. https://doi.org/10.1056/NEJMoa052187 PMID: 16371630

20. UK Prospective Diabetes Study (UKPDS) Group. Intensive blood-glucose control with sulphonylureas or insulin compared with conventional treatment and risk of complications in patients with type 2 diabetes (UKPDS 33). Lancet. 1998; 352: 837–853. PMID: 9742976

21. UK Prospective Diabetes Study (UKPDS) Group. Effect of intensive blood-glucose control with metformin on complications in overweight patients with type 2 diabetes (UKPDS 34). Lancet. 1998; 352: 854–65. PMID: 9742977

22. Indian Health Service: Find Health Care. 2016. https://www.ihs.gov/findhealthcare/.

23. Silva PS, Caravallano JD, Sun JK, Noble J, Aiello LM, Aiello LP. Nonmydriatic ultrawide field retinal imaging compared with dilated standard 7-field 35-mm photography and retinal specialist examination for evaluation of diabetic retinopathy. Am J Ophthalmol. 2012; 154: 549–559. https://doi.org/10.1016/j. ajo.2012.03.019 PMID: 22626617
24. Silva PS, Cavallerano JD, Tolls D, Omar A, Thakore K, Patel B, et al. Potential efficiency benefits of nonmydriatic ultrawide field retinal imaging in an ocular telehealth diabetic retinopathy program. Diabetes Care. 2014; 37: 50–55. https://doi.org/10.2337/dc13-1292 PMID: 23939541
25. International Council of Ophthalmology. ICO guidelines for diabetic eye care. 2017. http://www.icoph.org/downloads/ICOGuidelinesForDiabeticEyeCare.pdf.
26. Zhang X, Saaddine JB, Chou CF, Cotch MF, Cheng YJ, Geiss LS, et al. Prevalence of diabetic retinopathy in the United States, 2005–2008. JAMA. 2010; 304: 649–656. https://doi.org/10.1001/jama.2010.1111 PMID: 20699456
27. Wong TY, Klein R, Islam A, Cotch MF, Folsom AR, Klein BE, et al. Diabetic retinopathy in a multi-ethnic cohort in the United States. Am J Ophthalmol. 2006; 141: 446–455. https://doi.org/10.1016/j.ajo.2005.08.063 PMID: 16490489
28. Early Treatment Diabetic Retinopathy Study Research Group. Photocoagulation for diabetic macular edema. Early Treatment Diabetic Retinopathy Study Report number 1. Arch Ophthalmol. 1985; 103: 1796–1806.
29. Yau JW, Rogers SL, Kawasaki R, Lamoureux EL, Kowalski JW, Bek T, et al. Meta-Analysis for Eye Disease (META-EYE) Study Group. Global prevalence and major risk factors of diabetic retinopathy. Diabetes Care. 2012; 35: 556–564. https://doi.org/10.2337/dc11-1909 PMID: 22301125
30. Kempen JH, O’Colmain BJ, Leske MC, Haffner SM, Klein R, Moss SE, et al. Eye Diseases Prevalence Research Group. The prevalence of diabetic retinopathy among adults in the United States. Arch Ophthalmol. 2004; 122: 552–563. https://doi.org/10.1001/archophthalm.123.12.1699 PMID: 16344442
31. Klein R, Klein BE. Are individuals with diabetes seeing better?: A long-term epidemiological perspective. Diabetes. 2010; 59: 1853–1860. https://doi.org/10.2337/db09-1904 PMID: 20668290
32. Horton MB, Paolo PS, Cavallerano JD, Aiello LP. Clinical Components of Telemedicine Programs for Diabetic Retinopathy. Curr Diab Rep. 2016; 16: 129–139. https://doi.org/10.1007/s11892-016-0813-8 PMID: 27796779
33. Wolfe JA, Horton MB, McAteer MB, Szuter CF, Clayton T. Race, macular degeneration, and diabetic maculopathy. Arch Ophthalmol. 1993; 111: 1603–1604. PMID: 7512329
34. Zhang Y, Hu G, Yuan Z, Chen L. Glycosylated hemoglobin in relationship to cardiovascular outcomes and death in patients with type 2 diabetes: a systematic review and meta-analysis. PLoS ONE. 2012; 7: e42251. https://doi.org/10.1371/journal.pone.0042251 PMID: 22912709
35. Klein BE, Klein R, Wang Q, Moss SE. Older-onset diabetes and lens opacities. The Beaver Dam Eye Study. Ophthalmic Epidemiol. 1995; 2: 49–55. PMID: 7985233
46. Silva PS, Horton MB, Clary D, Lewis DG, Sun JK, Cavallerano JD, et al. Identification of diabetic retinopathy and ungradable image rate with ultrawide field imaging in a national teleophthalmology program. Ophthalmol. 2016; 123: 1360–1367.

47. DCRR.net Policies, Procedures, Protocols and Protocol Idea Forms. 2017. http://dcrnet.jaeb.org/ViewPage.aspx?PageName=Investig_Info.