Adherence to ophthalmology referral, treatment and follow-up after diabetic retinopathy screening in the primary care setting

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ABSTRACT

Introduction Telemedicine-based diabetic retinopathy screening (DRS) in primary care settings has increased the screening rates of patients with diabetes. However, blindness from vision-threatening diabetic retinopathy (VTDR) is a persistent problem. This study examined the extent of patients’ adherence to post-screening recommendations.

Research design/methods A retrospective record review was conducted in primary care clinics of a large county hospital in the USA. All patients with diabetes detected with VTDR in two time periods, differing in record type used, were included in the study: 2012–2014, paper charts only; 2015–2017, combined paper charts/electronic medical records (EMRs), or EMRs only. Adherence rates for keeping initial ophthalmology appointments, starting recommended treatments, and keeping follow-up appointments were determined.

Results Adequate records were available for 6046 patients; 408 (7%) were detected with VTDR. Only 5% completed a first ophthalmology appointment within recommended referral interval, 15% within twice the recommended interval, and 51% within 1 year of DRS. Patients screened in 2015–2017 were more likely to complete a first ophthalmology appointment than those in 2012–2014. Ophthalmic treatment was recommended in half of the patients, of whom 94% initiated treatment. A smaller percentage (41%) adhered completely to post-treatment follow-up. Overall, 28% of referred patients: (1) kept a first ophthalmology appointment; (2) were recommended for treatment; and (3) initiated the treatment. Most patients failing to keep first ophthalmology appointments continued non-ophthalmic medical care at the institution. EMRs provided more complete information than paper charts.

Conclusions Reducing vision impairment from VTDR requires greater emphasis on timely adherence to ophthalmology referral and follow-up. Prevention of visual loss from VTDR starts with retinopathy screening, but must include patient engagement, adherence monitoring, and streamlining ophthalmic referral and management. Revision of these processes has already been implemented at the study site, incorporating lessons from this investigation.

Significance of this study

What is already known about this subject?

► Regular diabetic retinopathy screening (DRS) is essential for the early detection of vision-threatening diabetic retinopathy (VTDR).
► The timely treatment of patients found to have VTDR can prevent severe vision loss in up to 90% of patients; delays in needed treatment can result in unnecessary and irreversible vision loss.

What are the new findings?

► A low proportion of patients detected with VTDR adhered completely to post-treatment follow-up. Most patients who did attend an ophthalmology visit were recommended for treatment, initiated that treatment, but a substantially smaller percent adhered completely to post-treatment follow-up.
► The majority of patients who failed to adhere to these recommendations were still receiving non-ophthalmic medical care within the same institution.

How might these results change the focus of research or clinical practice?

► The primary care team must be able to monitor efficiently the adherence of patients’ keeping a first ophthalmology appointment within the recommended time frame.
► Research is required to develop more efficient monitoring, and interventions to improve adherence to specialty referral, treatment and follow-up.

INTRODUCTION

Telemedicine-based diabetic retinopathy screening (DRS) programs in the primary care setting have become widespread,1 and have been successful in improving screening rates for retinopathy among patients with diabetes.2–4 Digital retinal imaging with a non-mydriatic camera can be effective in identifying individuals with vision-threatening
diabetic retinopathy (VTDR) who may need treatment of their eyes to prevent serious vision loss. However, to be most effective in preventing vision loss, screening must also be coupled with timely referral of patients with VTDR to eye specialists and with the administration of appropriate ocular therapy. To accomplish this, when DRS is conducted in the primary care setting, the non-specialist must educate the patient about the risks of VTDR and the importance of seeking timely eye specialist intervention. Extrapolation of data from randomized clinical trials of laser treatment for proliferative diabetic retinopathy (PDR) estimates that the rate of blindness from untreated severe PDR can be reduced by as much as 90% through early detection and prompt treatment before vision loss develops.

Despite overall improvement in screening rates nationwide, diabetic retinopathy remains the leading cause of blindness among working age adults in the USA. A factor contributing to this may be a breakdown and inefficiency in the process for eye specialty referrals, treatment recommendations and follow-up appointments. There is generally no comprehensive system for tracking patients referred from primary care settings to ophthalmology, and very few studies have been published on this topic. From those that have been published, it appears that many patients with VTDR are not being optimally treated.

To understand the relationship between DRS-generated ophthalmology referral and ocular treatment for VTDR, a retrospective record review study was conducted in a large county medical center, where primary care-based DRS had been successfully implemented since 2011. Data from 2012 to 2017 were collected and analyzed.

**RESEARCH DESIGN AND METHODS**

**Patient identification and record reviews**

The study was conducted in the Family Medicine Department at the Riverside University Health System-Medical Center (RUHS-MC), Riverside, California. Following Institutional Review Board approval, all patients who had received telemedicine-based DRS in the Family Care Center (FCC) at RUHS-MC, over the time periods specified below, were identified in the screening database. The patients had been referred for DRS either from the FCC itself, from the Internal Medicine Clinic at RUHS-MC, or from satellite community health centers. All patients with a diagnosis of diabetes mellitus or pre-diabetes were recommended for an annual DRS, unless they had seen an eye care specialist within the previous 2 years.

The retinal images had been evaluated remotely by two experienced, certified retinal graders using a standardized structured grading protocol. The same consultants evaluated the retinal images over the entire study period. All patients with a screening diagnosis VTDR were identified in the DRS database. VTDR was defined as one or more of the following in at least one eye: (1) PDR; (2) severe non-proliferative diabetic retinopathy (SNPDR); (3) clinically significant macular edema (CSME). Only patients recommended for referral to ophthalmology were included in the record review. The referral included a recommended referral interval, the time duration within which the patient should be seen, as determined by the retinopathy severity; the recommended referral intervals were consistent with published clinical guidelines. The default referral intervals were: SNPDR, 3 months; PDR, 1 month; CSME, 1 month. Certified graders could over-ride the defaults, but rarely did, except for patients with advanced PDR who were referred within 2 weeks or less. All patients were referred internally to the ophthalmology clinic in the same building as the FCC. Most were initially evaluated in the general eye service and referred to the retina specialty service if treatment was anticipated.

The medical record reviews were divided into two time periods, based on the type of medical records in use during the screening and subsequent follow-up periods. (The RUHS switched from paper charts to an electronic medical record (EMR) system in October 2016.) These were designated time period A: 2012–2014, paper charts only; time period B: 2015–2016, combined paper charts and EMRs, and 2017, EMRs only. Data were analyzed separately for the two time periods, to look for: (1) trends in screening and referral adherence over time; (2) differences in the information obtainable from the two record types; and (3) evidence that the EMR improved adherence with referral fulfillment, as has been reported elsewhere.

The following data were extracted for each patient: (1) date of keeping first ophthalmology appointment; (2) ocular treatment recommendation status; (3) date of initiating recommended treatment; and (4) dates of keeping subsequent treatment episodes and/or follow-up visits.

**Chart review interval and definition of adherence indicators**

1. The ‘chart review interval’ was the time between the date of a given event (eg, DRS) and the last date the records were reviewed for each patient. The ‘overall chart review interval’ was calculated from the date of the DRS. The minimum chart review intervals required to include a specific adherence rate in the study analysis were: (1) keeping first ophthalmology appointment, 12 months after DRS; and (2) treatment and post-treatment follow-up, 12 months after first ophthalmology appointment. Patients with shorter chart review intervals for a given measure were designated ‘indeterminate’ for that measure.

2. ‘Adherence to keeping a first ophthalmology appointment’ was the proportion of patients showing a kept appointment within a specified ‘adherence interval’. The ‘adherence interval’ was the length of time between the DRS and first visit to the ophthalmology clinic.

3. ‘Adherence to initiating treatment’ was the proportion of patients initiating treatment after the recommendation was made.

4. ‘Adherence to post-treatment follow-up’ was defined in four categories: (1) ‘non-adherent’: no follow-up...
visits after initial treatment session; (2) ‘partially adherent’: returned for some follow-up visits, but failed to return for one or more visits within 6 months of the recommended follow-up interval; (3) ‘completely adherent’: returned for all follow-up visits within 6 months of the recommended intervals; and (4) ‘indeterminate’: could not determine due to missing data or to an inadequate chart review interval.

5. ‘Adherence to follow-up Ophthalmology Clinic visits among patients not recommended for treatment’ was defined using the same categories as adherence to post-treatment follow-up.

Additional analyses
The screening database was searched for repeat screening to determine whether patients failing to keep a first ophthalmology appointment within 1 year of the DRS were ‘recaptured’ for re-evaluation and possible re-referral. The date of repeat DRS, diagnosis, ophthalmological referral and adherence to first ophthalmology appointment were recorded. Also, the records were searched for patients who died after the DRS.

Statistical analyses
Patients’ demographic variables including age, gender, ethnicity and insurance type were collected from paper charts and/or EMRs. Clinical characteristics such as years of diabetes, comorbid conditions, diabetes type and HbA1c value were also collected at or close to baseline screening.

Descriptive statistics of the study population are listed in table 1. χ² tests were used to test differences of characteristics in two time periods. Kaplan-Meier curves and log rank test were performed to compare two time periods for keeping ophthalmology appointments. Outcome of time-to-event analysis was the time between screening until the date of the earliest of the following events: having a first ophthalmology appointment, or last chart review date. Patients who did not have ophthalmology appointments were included as censored observations.

Additionally, bivariate and multivariate analyses were performed to test whether patient-related factors correlated with the adherence rate for keeping a first ophthalmology appointment. The results are reported in online supplementary appendix B. A multivariate Cox proportional hazards model was fitted to estimate the HRs of keeping first ophthalmology appointments for the various patient characteristics. Analyses were performed using SAS V.9.4 (SAS Institute). Statistical significance was set at p<0.05 with two-tailed tests.

RESULTS
Some of the major findings in the study are illustrated in the flow chart (figure 1) and are described in detail below.

Baseline characteristics of study population
A total of 3064 patients were screened in time period A, of whom six were eliminated because medical records were not available. Among those remaining, 147/3058 (5%) were recommended for referral for VTDR; a greater proportion was referred in time period B, 2991 screened, 3 eliminated, and 261/2988 (9%) referred (p<0.001). Since the retinal image grading criteria did not change over time, the reason for the increase in referrals for time period B was inapparent. The same readers graded the images over the entire duration of the study.

The distribution of the characteristics of the referred study population, subdivided by time period, is shown in table 1. Differences of distributions between time periods were tested with χ² test or Fisher’s exact test.

The two time periods showed statistically significant differences as follows:

Age group: Time period B had a higher proportion of patients in the oldest age group (60+) and fewer in the youngest group (18–49).

Ethnicity: Hispanic patients formed the largest ethnic group, and this preponderance was greater in time period B.

Retinopathy severity: There was a higher percentage of patients with PDR in time period A, and a higher percentage of the milder retinopathy categories in time period B.

CSME: CSME was present in the majority of patients (81.1%), and was more prevalent in time period B.

Primary care site: Of patients referred for DRS from the Internal Medicine Clinic, a higher percentage was referred in time period B, and of the satellite clinics a higher percentage was referred in time period A. There was little difference in the percentage of patients referred from the FCC between each time period.

Insurance type: The most common types of insurance in descending order were MediCal, Medicare, and private; close to 10% of patients were covered by the Medically Indigent Services Program (MISP). (This is a safety net program sponsored by Riverside County, and designed to cover ‘medical care to prevent disability’. It is not, strictly speaking, insurance, but is treated as such for the purpose of this analysis.) Other types included: uninsured, charity patients, and self-pay. For slightly more than 25% of patients, the insurance status could not be determined from the records.

Insurance category: There was a much higher percentage of insured patients in time period B and a higher percentage of missing insurance data in time period A. The higher insured rate in time period B was mainly due to an increase in MediCal-covered patients.

The distribution of recommended referral intervals was similar in the two time periods: on average, immediate to 2 weeks, 22/408 (5%); 1 month, 358/408 (88%); 2–3 months, 23/408 (6%); and 6 months, 5/408 (1.2%).
Table 1  Baseline patient characteristics by time period

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Continued
Clinical Care/Education/Nutrition

Table 1 Continued

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*P value from $\chi^2$ test or Fisher’s exact test for difference of patient characteristics in two time periods.

MISP, Medically Indigent Services Program; PDR, proliferative diabetic retinopathy; SNPDR, severe non-proliferative diabetic retinopathy.

Adherence rates

Adherence rates for keeping a first ophthalmology appointment

Very few patients kept their first ophthalmology appointment within the recommended referral interval: 8/147 (5%) in time period A; 12/261 (5%) in time period B. For keeping first appointments within twice the recommended interval, only 19/147 (14%) in period A and 42/261 (16%) in period B adhered. When the latter results are stratified by the length of the recommended interval, those recommended for <2 weeks and those for 3 months had a higher average adherence rate, 10/22 (45%) and 6/23 (26%), respectively, with those recommended for 1 month interval, 44/358 (11%) (p<0.0001).

For keeping a first ophthalmology appointment within 1 year of the DRS, the adherence rates were 70/147 (48%) in time period A and 147/261 (56%) in time period B (p<0.05). An additional 13/147 (9%) in time period A and 33/261 (13%) in time period B kept a first ophthalmology appointment beyond 1 year of the DRS.

Figure 2 shows the adherence data over time plotted as time-to-event estimates (Kaplan-Meier curves), where the ‘event’ was keeping a first ophthalmology appointment. The upper curve (2015–2017) shows a more rapid development of the event than the lower curve (2012–2014), log rank p=0.0109.

Only 17/195 (9%) of patients failing to keep an ophthalmology appointment within 1 year of the initial DRS had a repeat screening within 2 years of the DRS. Among the patients with repeat screening, 14/17 (82%) were again referred to ophthalmology for VTDR, and of

Figure 1 Diagram showing the original study sample and the major findings of the study, app’t, appointment; DRS, diabetic retinopathy screening; OPTH, ophthalmology; VTDR, vision-threatening diabetic retinopathy.

Figure 2 Time-to-event estimates: Kaplan-Meier curves comparing time period A, 2012–2014 (lower curve), with time period B, 2015–2017 (upper curve). The ‘event’ was keeping a first ophthalmology appointment. The number of subjects at risk for each time period is shown above the x-axis.
Clinical Care/Education/Nutrition

Figure 3  (A) Proportion of patients starting treatment who adhered to post-treatment follow-up recommendations after treatment was started. (B) Proportion of patients keeping a first ophthalmology appointment and NOT recommended for treatment, who adhered to follow-up recommendations in the eye clinic.

Adherence to initiation of treatment

One hundred and sixteen out of 123 (94%) patients recommended for treatment initiated the treatment. Treatment was usually started in the ophthalmology clinic the same day that it was recommended.

Adherence to post-treatment follow-up

The rate of complete adherence to post-treatment follow-up was higher in time period A, 27/48 (56%), than in time period B, 21/68 (31%) (p=0.006), although it was suboptimal in both (figure 3A). The rates of partial adherence were not statistically significantly different by time period: time period A, 16/48 (33%), and time period B, 34/68 (51%). Non-adherence was relatively low in both time periods (8%, 10%) as were the proportion of patients for whom adherence was indeterminate (4%, 7%).

Adherence to follow-up visits in the ophthalmology clinic for patients not recommended for treatment

Most patients not recommended for treatment were advised to return to the ophthalmology clinic for repeat evaluations. The rates of adherence with follow-up were generally low (figure 3B). In time period A, almost half of the patients were either non-adherent, 6/35 (17%), or partially adherent, 9/35 (26%); in time period B, a

injections with anti-vascular endothelial growth factor (anti-VEGF agents), and vitrectomy surgery. Sixty-seven of 123 (54%) patients had a combination of treatment types recommended, and 56/123 (46%) a single treatment type. PRP and intravitreal anti-VEGF injections comprised the most common combination, 35/67 (52%), as well as being the most common single treatment types, 25/56 (45%) and 24/56 (43%), respectively.

Overall, 137/263 (52%) of the referred patients keeping a first ophthalmology appointment were thought not to need treatment. Of these, 55/137 (40%) were diagnosed with VTDR by ophthalmological examination, but were not recommended for treatment. The decision not to treat at that time had been based on one or more factors: (1) SNPDR, and no diabetic macular edema; (2) prior treatment with PRP, and no additional treatment needed; (3) macular edema present, but not involving or threatening the central macula. The rest, 79/137 (58%), were found to have retinopathy less severe than VTDR by ophthalmological examination.

Among the referred patients with VTDR who kept a first ophthalmology appointment who kept a first ophthalmology appointment, the ophthalmological examination found a greater proportion of patients with moderate non-proliferative diabetic retinopathy and no CSME in time period B, 63/180 (37%), compared with 18/83 (22%) in time period A; none of these patients were recommended for treatment. Conversely, there were more patients with PDR in time period A (37/83 (45%)) than in time period B (58/180 (32%)); 92% of these were recommended for treatment in time period A and 72% in time period B.

Adherence to follow-up visits in the ophthalmology clinic for patients not recommended for treatment

Most patients not recommended for treatment were advised to return to the ophthalmology clinic for repeat evaluations. The rates of adherence with follow-up were generally low (figure 3B). In time period A, almost half of the patients were either non-adherent, 6/35 (17%), or partially adherent, 9/35 (26%); in time period B, a
greater proportion of patients were either non-adherent, 47/108 (44%), or partially adherent, 27/108 (25%) (χ² test p value=0.0108). The proportion of patients for whom the adherence was indeterminate was relatively high in both time periods (34% and 14%).

Comparison of adherence to follow-up among all patients recommended for treatment (Rx) with all those not recommended (no Rx) showed that the complete compliance rate in the Rx group, 48/109 (44.0 %), was significantly greater than in the no-Rx group, 27/116 (23.3%) (p=0.0010).

Multivariate Cox proportional hazards model
The HRs for keeping a first ophthalmology appointment, derived from the multivariate Cox proportional hazards model, are reported in online supplementary appendix B. Patients in time period B were more likely to be adherent compared with patients in time period A (HR=1.40, 95% CI 1.05 to 1.87, p=0.023). Other covariates that were significantly associated with greater adherence were Hispanic patients, patients with PDR (compared with less than SNPDR), patients with one or more comorbid conditions (compared with no comorbid condition) and diabetes duration 0–5years (compared with 6–10years). Health insurance status was not significantly related to adherence.

DISCUSSION
While telemedicine-based DRS can dramatically increase the screening rate by moving the screening locus from the specialty eye care setting to the primary care setting, the management of patients identified with VTDR through DRS still requires a referral out to a specialist. Therefore, one would expect that many of the same barriers that impeded DRS by referral would also impede treatment by referral. The major difference from the healthcare delivery and quality improvement viewpoints is that with successful DRS, at least we know who the patients are who need to be ‘moved’, and the challenge becomes how to encourage them to move, not how to find them.

Although several published studies have reported adherence data for ophthalmic follow-up after DRS in the primary care setting, they have usually been restricted to tracking the first ophthalmology visits and not the subsequent treatment and post-treatment visits.410–18 There are a number of studies conducted entirely in the ophthalmology setting that track the adherence of patients with VTDR to various modalities of treatment.19–21 The present study appears to be the first to report the results of tracking patients from the primary care screening setting all the way through the specialty referral and subsequent treatment and post-treatment phases.

Overall, about 50% of patients failed to keep any first ophthalmology appointment within 1 year of the DRS, a deficit that can lead to clinically significant vision loss, especially with PDR.22 No formal system was in place to identify and rescreen patients who failed to keep an ophthalmology appointment. Rather, there was a general policy in the primary care clinics to offer DRS to any patient who had not been screened for at least 12 months or who had not seen an eye care provider for at least 2 years. However, the rate of rescreening for the non-adherent patients was very low, underscoring the need for a more robust tracking and recall system for patients failing to keep a first ophthalmology appointment.

The records showed that only six patients died during the observed time frame after the DRS. The effect of death on adherence rates to keeping a first ophthalmology appointment within 1 year was negligible; four patients adhered, one patient did not adhere, but died beyond the 1 year post-DRS period, while only one patient died within the 1 year post-DRS period.

There were differences in adherence rates to keeping a first ophthalmology appointment depending on the recommended referral interval. Higher adherence rates for keeping a first ophthalmology appointment within twice the recommended interval were found both for patients recommended for referral within 2 weeks of the DRS (45%) or within 3 months (26%) compared with those with a 1 month interval (11%).

The higher adherence rate for the ≤2 weeks’ interval can be attributed to an informal ‘fast track workaround’, in which a primary care staff directly contacted the ophthalmology clinic, either by telephone or in person, to facilitate the referral of the most urgent patients.23 The higher adherence for the 3–6 months’ interval can probably be explained simply by the longer interval, giving the patients more time to adhere to the recommendation.

Overall, patients in time period B went sooner to their first ophthalmology appointment than those in time period A, according to the Kaplan-Meier time-to-event estimates. This is reinforced by finding an increased likelihood of keeping the first ophthalmology appointment among patients in time period B by multivariate regression analysis (online supplementary appendix B). These results could relate to more efficient referral processes with the EMR. However, finding that almost 30% of the year 2017 patients who failed to keep a first appointment within 1 year showed no evidence of a formal referral to ophthalmology in the EMRs (online supplementary appendix A) demonstrates that while the EMR can improve communications between primary care providers and specialists and improve the quality of care,4 fail-safe processes for making and tracking referrals are still needed.

Other patient characteristics significantly associated with a higher likelihood of adherence were Hispanic ethnicity, a comorbidity index of 1 or more, and the diagnosis of PDR. Having health insurance did not significantly correlate with adherence likelihood (online supplementary appendix B). The latter finding is consistent with reports elsewhere that the increased coverage with Medicaid, achieved after the introduction of the Affordable Care Act, did not, by itself, lead to increased utilization of certain screening services.24 25
A higher percentage of patients with kept first ophthalmology appointments were recommended for treatment in time period A than in time period B. This could be due to the higher prevalence of patients with PDR in the time period A sample, which, in turn, could indicate a backlog of previously undetected patients with PDR in the earlier phase of the retinal screening program.

Once treatment was recommended, a very high proportion (94%) of patients initiated the treatment. Post-treatment follow-up visits were always scheduled to evaluate the effects of treatment, and to consider adding treatment. This was especially important for patients receiving intravitreal anti-VEGF treatment, for whom at least three to four treatment sessions in a series were frequently planned over a period of 3–6 months.

The proportion of patients who initiated treatment and then adhered to all post-treatment follow-up visits (within 6 months of the recommended visit date) was significantly greater in time period A than in time period B. This could be an artifact relating to greater difficulty in identifying missed visits in the paper charts. While more missed visits have been reported among patients receiving PRP compared with anti-VEGF injections, our sample was not large enough to stratify the results by individual treatment types.

Among the patients not recommended for treatment, the non-adherence rates for follow-up were significantly higher in time period B compared with A. However, since the proportion of patients for whom adherence was indeterminate was significantly higher in time period A compared with B, the validity of comparing adherence rates by time period is questionable.

Comparison between patients recommended for treatment (Rx), with those not recommended for treatment (no Rx), showed significantly poorer adherence with recommended follow-up visits in the no-Rx group. Recommended follow-up intervals were usually longer in the no-Rx group, up to 1 year in many cases. With the longer intervals, a specific appointment may not have been made during the initial ophthalmology visit. In addition, there was probably a lesser sense of urgency imparted to these patients because of the low immediate threat to vision. Again, the relatively high indeterminate rate in the no-Rx group limits the validity of this comparison.

The results of this study indicate that the EMR provides more complete data on the referral process and on adherence characteristics than the paper charts. This included dates of formal referral to ophthalmology, attendance at other outpatient clinics, and missed follow-up visits, data for which were often not available or not reliable in paper charts. (See online supplementary appendix A.)

There are limits to the generalizability of the findings from this study. The present study was conducted in a primary care clinic of a safety net hospital serving a mostly low-income population. The insurance payer mix of the subjects in the study was weighted toward safety net, publicly funded insurance, specifically MediCal and MISP. In settings where there is a preponderance of private insurance payers and Medicare, the adherence characteristics would conceivably be better, as has been reported for the rates of DRS itself. The ophthalmology services at RUHS-MC were located in the same building as the primary care clinics. In primary care settings where ophthalmology is located off-site, the adherence characteristics might understandably be worse.

The present study was also limited by the lack of availability of data from patients who may have been evaluated and treated elsewhere for their VTDR. It is likely that at least some of the non-adherence occurred for that reason. As example, we found that 16/114 (14%) of patients referred for VTDR in time period B who failed to keep a first ophthalmology appointment within 1 year had no RUHS outpatient visit beyond 6 months after the date of the DRS. Tracking of patients like this in future studies would provide a more complete picture of the true adherence rates. Conversely, most patients who were non-adherent or partially adherent to eye care recommendations continued to receive non-ophthalmic medical care at RUHS. This suggests that with an enhanced tracking and recall system, improved adherence to eye care may be achievable through targeted interventions.

Our study did not look at adherence to treatment and follow-up in the ophthalmology clinic for patients with other chronic eye diseases such as glaucoma. It is possible that the adherence deficits found for patients with diabetes and VTDR are a more generalized problem for patients with chronic eye disease, or, as seems likely, for the safety net population as a whole. While addressing the multifactorial causes for adherence deficits in the safety net system is daunting, collecting adherence data, such as was done retrospectively in the present study, is an essential step in improving the quality of care. Otherwise, the problem cannot be defined or may not even be recognized.

As a result of our findings, the primary care clinics at RUHS-MC have redesigned the workflow for DRS and the referral process. They have begun to provide results of DRS immediately to the patients before they leave the clinic. This has been coupled with enhanced patient engagement to explain the risk of vision loss from untreated VTDR and how to prevent it, as well as provision of prompt appointments to the ophthalmology clinic for the more urgent referrals (≤1 month). An improved tracking and recall system for patients not keeping a first ophthalmology appointment has also been instituted. The adherence rates of patients enrolled in this new program will be compared with the results reported in the present study in an ongoing prospective study.

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Contributors  GB designed the study protocol, researched the data, contributed to discussion, drafted the manuscript, and reviewed and edited the manuscript. JAC helped design the study protocol, researched the data, contributed to discussion, and reviewed and edited the manuscript. PC and AL researched the data, contributed to discussion, and reviewed and edited the manuscript. SF, ERP, GW, and MK contributed to discussion, and reviewed and edited the manuscript. JC and LJ conducted the statistical analyses and reviewed and edited the manuscript. GB is the guarantor of this work, and as such had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis. All authors have read and approved the paper.

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Patient consent for publication  Not required.

Ethics approval  This study was approved by the Riverside University Health System IRB Committee (No. 824116-9). All patient data were deidentified, and patient consent for this retrospective study was not required by the IRB.

Provenance and peer review  Not commissioned; externally peer reviewed.

Data availability statement  Data are available upon reasonable request. The data are deidentified participant data, available from GB; visionforal2020@gmail.com.

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