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The Application of Optical Coherence Tomography Angiography in Cerebral Small Vessel Disease, Ischemic Stroke, and Dementia: A Systematic Review

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Objective: To investigate the application of optical coherence tomography angiography (OCTA) in cerebral small vessel disease (SVD), ischemic stroke and dementia.

Methods: We conducted a systematic search in MEDLINE (from inception) and EMBASE (from 1980) to end 2019 for human studies that measured retinal parameters in cerebral SVD, ischemic stroke, and dementia using OCTA.

Results: Fourteen articles (n = 989) provided relevant data. Ten studies included patients with Alzheimer disease (AD) and mild cognitive impairment (n = 679), two investigated pre-symptomatic AD participants (n = 154), and two investigated monogenic SVD patients with cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy (n = 32) and Fabry disease (n = 124). Methods to reduce bias and risk factor adjustment were poorly reported. Substantial methodological variations between studies precluded a formal meta-analysis. Quantitative measurements revealed significant yet inconclusive changes in foveal avascular zone, perfusion density, and vessel density (VD) in AD, presymptomatic AD, and SVD patients. Two (n = 160) of three studies (n = 192) showed association between decreased VD and increased white matter hyperintensities. In three (n = 297) of seven studies (n = 563), better cognitive function was associated with increased VD. One study (n = 52) suggested increased VD was associated with increased ganglion cell–inner plexiform layer thickness in AD yet with no covariate adjustment.

Conclusions: Changes in retinal microvasculature identified using OCTA are associated with monogenic SVD and different stages of AD, but data are limited and partly confounded by methodological differences. Larger studies with risk factors adjustment and more consistent OCTA methods are needed to fully exploit this technology.

PROSPERO registration number: CRD42020166929.

Keywords: dementia, Alzheimer's disease, retinal vasculature, optical coherence tomography angiography, ischemic stroke, cerebral small vessel disease
INTRODUCTION

Cerebral small vessel disease (SVD) is an intrinsic disorder that affects the brain’s small perforating arterioles, capillaries, and probably venules, causing various lesions seen on pathological examination or brain imaging with magnetic resonance imaging (MRI) or computed tomography (1). Cerebral SVD is a common condition in older adults and causes a wide array of clinical syndromes, including 25% of ischemic strokes and 80% of intracerebral hemorrhages, and contributes to up to 50% of dementias worldwide (1–3).

However, the changes in the cerebral microcirculation are difficult to visualize in vivo. Because the retinal microvasculature and cerebral microvasculature share similar embryologic origins as well as anatomical and physiological properties (4), investigating the network of retinal vessels may provide new insights into cerebrovascular disease and the vascular contribution to pathologic features of dementia (5). Previous literature has mainly focused on retinal fundus photography or optical coherence tomography (OCT) (5, 6).

A newly presented extension of structural OCT, referred to as optical coherence tomography angiography (OCTA), represents a novel non-invasive, depth-selective modality that allows for visualization of retinal blood flow without dye injection (7). OCTA images are essentially motion-contrast images, which are based on the different backscattering of light between red blood cells and neurosensory tissue, because red blood cells are moving while neurosensory tissue is static (8). In many cases, these images are now approaching histology-level resolution (8), raising great interest in the possibility of detecting early microvascular pathological changes in people in vivo. A comprehensive review of the technology, its principles, limitations, and the clinical application in eye disease has been published (8), work that we extend here into the arena of neurological disease. OCTA may permit detection of a series of changes such as reduction in capillary vessels and perfusion density before they are visible on retinal photographs (9). Moreover, recent work suggests that high-resolution OCTA images combined with machine learning may provide great potential to automate detection and quantification of microvascular changes underlying common brain diseases such as SVD, stroke, and dementia, in the future (10, 11).

In this study, we aimed to conduct a systematic review of the literature to examine the application of OCTA in cerebral SVD, ischemic stroke, and/or any type of dementia and compared either (1) retinal parameters using OCTA vs. control, (2) OCTA parameters and brain imaging findings, (3) OCTA parameters and cognitive function, or (4) OCTA parameters and findings on fundus camera imaging or OCT.

The following studies were excluded: (1) duplicate publications or studies not meeting the inclusion criteria, (2) review studies, (3) single-case reports, (4) nonhuman studies, (5) non–English language studies, (6) conference presentations or summaries, (7) studies without details of diagnosis criteria or definitions, and (8) studies that despite examining retinal parameters did not conduct or report results on OCTA.

METHODS

This systematic review was based on a predefined protocol (PROSPERO registration no. CRD42020166929) following the Preferred Reporting Items for Systematic Reviews and Meta-analyses (12) guidelines. We refer to SVD MRI findings according to STRIVE (Standards for Reporting Vascular Changes on Neuroimaging) guidelines (13). As all analyses here are based on publicly available summary statistics and not individual-level data, no ethical approval or informed patient consent was required.

Search Strategy

We conducted a literature search of the Medical Literature Analysis and Retrieval System Online (MEDLINE, from inception) and the Excerpta Medica Database (EMBASE, from 1980) up to December 31, 2019, using the Ovid Web Gateway (detailed search strategy in Supplementary Material). We considered only studies published in English. References of relevant articles were hand-searched, and a forward citation search was performed to identify further studies.

Inclusion and Exclusion Criteria

The studies included patients with cerebral SVD, ischemic stroke, and/or any type of dementia and compared either (1) retinal parameters using OCTA vs. control, (2) OCTA parameters and brain imaging findings, (3) OCTA parameters and cognitive function, or (4) OCTA parameters and findings on fundus camera imaging or OCT.

The following studies were excluded: (1) duplicate publications or studies not meeting the inclusion criteria, (2) review studies, (3) single-case reports, (4) nonhuman studies, (5) non–English language studies, (6) conference presentations or summaries, (7) studies without details of diagnosis criteria or definitions, and (8) studies that despite examining retinal parameters did not conduct or report results on OCTA.

Data Extraction

We initially screened all studies identified in the systematic search by abstract and title for potentially relevant articles. Duplicate articles were removed, and the remaining articles were assessed for eligibility after full-text review. Data extracted from these studies included title; authors; publication year; study aim; study type; number of patients and controls; number of male and female; mean age, participant selection criteria, and diagnostic criteria; type of OCTA device; assessment of OCTA image quality; whether OCTA associations were adjusted for demographics, other risk factors, or other covariates; and outcomes. All data were cross-checked by a second reviewer. We report covariate-adjusted associations when available.

Quality Assessment

We used the STROBE checklist (www.equator-network.org) to score study methodology and assess study quality. We assigned up to 22 points using this checklist. This score was applied after study inclusion and did not influence whether a study was included in the review.

Owing to the small number of studies per category and the large methodological heterogeneity, it was not possible to combine study data by meta-analysis.
RESULTS

The search returned 2,611 articles. One additional article was found by hand-search. Among them, 621 duplicate articles were removed. The remaining 1,991 articles were screened by title and abstract only. Of those, 16 were considered potentially relevant and were assessed by full-text review. Figure 1 details numbers of articles excluded and the reasons.

Fourteen articles met the inclusion criteria. The population samples came from the United States (five articles), the Netherlands (two articles), Germany (two articles), Italy (two articles), Poland (one article), Turkey (one article), and China (one article). Multiple articles from the same study population were included only if different retinal parameters or outcomes were measured in separate articles.

Study Design and Patient Characteristics

Table 1 summarizes the study design and patient characteristics from the articles included. The 14 articles (14–27) included comprised 13 cross-sectional studies and 1 case–control study. Although two articles (18, 19) had overlapping samples, each study assessed different outcomes. Across the included studies (if two articles had an overlapping study population, the study with the larger sample was chosen), the total number of unique participants was 989. Participants in each study varied from 16 to 213 (median \( n = 55 \)).

Ten articles (14–23) included patients with AD and mild cognitive impairment (MCI) (AD, \( n = 221 \); MCI, \( n = 106 \); controls, \( n = 352 \)). Two studies (24, 25) investigated presymptomatic AD participants who had normal cognitive function with positive biomarker of amyloid-\( \beta \) (A\( \beta \)) (total \( n = 154 \)). No relevant studies were identified for other types of dementia. Two studies investigated monogenic SVD patients: one (26) studied cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy (CADASIL) patients (CADASIL, \( n = 11 \); controls, \( n = 21 \)), and the other (27) studied Fabry disease (Fabry disease, \( n = 54 \); controls, \( n = 70 \)). We did not find any study on

![FIGURE 1 | Search strategy flow chart.](image-url)
### Study design and patient characteristics.

| Article | Design of study | Number of subjects (eyes) | Eye selection | Mean age ± SD (years) | Sex (Male/Female) | MMSE (Mean) |
|---------|-----------------|---------------------------|---------------|-----------------------|-------------------|-------------|
| **AD studies** | | | | | | |
| Bulut et al. (14) | Cross-sectional study | AD 26 (26) | One eye from each subject was randomly selected | AD 74.2 ± 7.6 | 11/15 13/13 | AD 17 27 |
| Lahme et al. (15) | Cross-sectional study | AD 36 (36) | — | AD 68.0 ± 9.3 | 15/21 14/23 | AD 22 — |
| Haan et al. (16) | Cross-sectional study | AD 48 (NA) | Values from both eyes were averaged unless only one eye was suitable | AD 65.4 ± 8.1 | 25/23 24/14 | AD 23 29 |
| Querques et al. (17) | Cross-sectional study | AD 12 (12) | One eye for each subject was randomly selected | AD 72.9 ± 7.2 | 4/8 MCI 5/7 | AD 21 MCI 25 |
| Yoon et al. (18)* | Cross-sectional study | AD 39 (70) | Eye images with poor scan quality and motion artifact were excluded | AD 72.8 ± 7.7 | 13/26 MCI 17/20 | AD 20 MCI 23 |
| Yoon et al. (19)* | Cross-sectional study | AD 9 (17) | — | AD 75.2 ± 7.5 | 4/5 MCI 4/3 | AD 22 MCI 26 |
| Zabel et al. (20) | Cross-sectional study | AD 27 (27) | One eye of each patient was included | AD 74.1 ± 5.9 | 6/21 8/19 | AD 21 28 |
| Jiang et al. (21) | Cross-sectional study | AD 12 (NA) | Values from both eyes were averaged | AD 73.3 ± 9.6 | 7/14 | AD 20 MCI 26 |
| Wu et al. (22) | Cross-sectional study | AD 18 (28) | Eye images with poor scan quality and motion artifact were excluded | AD 69.9 ± 6.4 | — | AD 20 MCI 25 |
| Zhang et al. (23) | Cross-sectional study | aMCI/eAD 16 (16) | Chose right eyes. If the image quality failed, then chose left eyes. | aMCI/eAD 73.03 ± 8.24 | 3/13 MoCA 20 | MoCA 27 |
| van de Kreeke, et al. (24) | Cross-sectional study | Aji+ 13 Aji- 111 | Values from both eyes were averaged | ALL 68.6 ± 6.3 | ALL 58/66 | ALL 29 |
| O’Bryhim et al. (25) | Case-control study | ALL 30 (58) | Eye images with poor scan quality and motion artifact were excluded | ALL 74.5 ± 5.6 | ALL 14/16 (normal) | |
| **Presymptomatic AD studies** | | | | | | |
| Nelis et al. (26) | Cross-sectional study | 11 (21) | Values from both eyes were used. | 53.5 ± 10.7 | 3/8 | — |
| **Fabry disease study** | | | | | | |
| Cennamo et al. (27) | Cross-sectional study | 54 (54) | One eye for each subject was randomly selected | 44.1 ± 15.6 | 20/34 | 34/36 |

AD, Alzheimer disease; MCI, mild cognitive impairment; HC, healthy control; eAD, early Alzheimer-type dementia; aMCI, amnestic type of MCI; MMSE, Mini-Mental State Examination; MoCA, Montreal Cognitive Assessment; CADASIL, cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy.

*The two studies were from the same study population.
sporadic SVD or ischemic stroke that met the inclusion criteria (Figure 2).

Quality Assessment
The median study quality score as per STROBE checklist (www.equator-network.org) was 16/22 (Supplementary Table 1). Scores were mainly affected by inadequate description of study design in title or abstract, not explaining rationale for study size, inadequate description of efforts made to control potential sources of bias, and inadequate discussion of generalizability of results.

OCTA Devices, Parameters, and Calculation Method
Table 2 summarizes the methods used by each study to assess retinal parameters. Eight articles (14, 15, 20, 22, 23, 25–27) used AngioVue™ system of Optovue, and the other six articles (16–19, 21, 24) used AngioPlex™ of Carl Zeiss. Different manufacturers use different hardware, software, and algorithms for vessel segmentation, which complicates interstudy comparisons, and indeed, OCTA parameters with the same name may be defined differently across studies. Furthermore, different studies used different image quality assessment methods or cutoffs (Supplementary Table 2). Therefore, the results from those studies are not all interchangeable (28). See Table 3 for detailed explanation of these retinal parameters.

Main Analysis
OCTA Parameters in Patients vs. Controls
Figure 3 and Table 4 summarize the main results from the 14 included studies. Eight studies (14, 16, 18, 20, 22, 24–26) compared the foveal avascular zone (FAZ) size between patients and controls. Four studies \( n = 196, \) median \( n = 53, \) three AD and MCI studies (14, 20, 22); one presymptomatic AD study (25) found that patients had enlarged FAZ compared with controls. Among these studies, two (14, 22) were age- and sex-matched; another (25) adjusted for age, sex, scan quality, and diabetes; and the other (20) did not adjust for any covariates. Four other studies \( n = 455, \) median \( n = 105, \) two AD and MCI studies (16, 18); one presymptomatic AD study (24); one CADASIL study (26) showed no difference between patient and control groups. Among these studies, one study matched for age and sex (18). Two studies (16, 24) adjusted for age, one (16) of which also adjusted for spherical equivalent, and quality factor. The other study (26) did not adjust for any covariates.

Twelve studies (14–18, 20–24, 26, 27) compared the vascular density parameters (vessel density, vessel length density, perfusion density, etc.) between patients and controls. From these, eight studies \( n = 569, \) median \( n = 53, \) seven AD and MCI studies (14, 15, 18, 20–23); one CADASIL study (26) suggested decreased vascular density parameters in patients compared with controls. Among them, two studies (20, 21) did not adjust for any covariates, one study (18) adjusted the results for age and sex, and in four studies (14, 15, 23, 26), sample groups were matched for age, three of which (14, 22, 23) also matched for sex and one of them (23) also matched for race. However, two studies \( n = 56 \) and \( n = 86, \) involving AD and MCI patients (16, 17) showed no difference between patient and control groups. Interestingly, one study in presymptomatic AD \( n = 124 \) (24) found Aβ+ participants had higher vessel density than Aβ- participants after adjusting for age. Another study \( n = 124 \) (27) found Fabry disease patients had decreased vascular density in superficial capillary plexus and increased in deep capillary plexus compared with controls after adjusting for age and sex.

OCTA Parameters in Relation to Brain Imaging Findings
Only four studies \( n = 208, \) median \( n = 53, \) three in AD and MCI (15, 16, 19) and one in CADASIL (26) investigated the relationship between OCTA findings and brain imaging findings. Two of them \( n = 74 \) and \( n = 86 \) (15, 16) involving AD and MCI participants found increased vessel density (VD) was associated with having fewer white matter hyperintensities (WMHs), as per Fazekas scores. Between these two studies, one (15) matched for age, and the other (16) adjusted for age and sex. One study \( n = 16, \) (19) involving AD and MCI patients suggested a potential association between decreased VD and decreased perfusion density with increased volume of the inferior lateral ventricle (temporal horn), but it did not control for covariates, and patients and controls were not matched by age and/or gender. The study \( n = 32 \) (26) in CADASIL patients found no association between OCTA parameters and the number of small infarcts or WMHs, the latter estimated using Fazekas scores.

OCTA Findings in Relation to Cognitive Function
Seven (14–16, 18, 20, 21, 23) studies investigated the relationship between OCTA parameters and cognitive function in AD and MCI participants. Three studies \( n = 297 \) (14, 18, 23) suggested better cognitive function was associated with increased VD. One study \( n = 52 \) (14) found enlarged FAZ area was associated with decreased Mini-Mental State Examination (MMSE) scores. Of these three studies, two (14, 23) were age- and sex-matched, one of which (23) was also race-matched. One study (18) did not adjust or match for covariates. However, four other studies \( n = 266, \) median \( n = 64 \) (15, 16, 20, 21) found no association between OCTA parameters and cognitive functions. Among them, one
| Article | OCTA device | Macular scan size (mm) | ONH scan size (mm) | OCTA parameters | Additional adjustments |
|---------|-------------|------------------------|-------------------|----------------|------------------------|
| Bulut et al. | Optovue | 6 | — | VD (%): Whole/fovea/parafovea superficial VD | None (age and sex matched) |
| Lahme et al. | Optovue | 3 | 4.5 | VD (%): Whole/fovea/parafovea superficial VD | None (age matched) |
| Haan et al. | Zeiss | 6 | — | VD (mm²): inner ring (Ø 1–3 mm around fovea) VD | Age, spherical equivalent, and quality factor |
| Querques et al. | Zeiss | 3 | 6 | PD (%), 3 and 6 mm | None (age and sex matched) |
| Yoon et al. | Zeiss | 3 | 6 | Superficial vascular plexus of VD (mm²) and PD (%) in: 3-mm circle, 3-mm ring (Ø 1–3 mm around fovea), and 6-mm circle | Age and sex |
| Yoon et al. | Zeiss | 3 | 6 | VD (mm²) | None |
| Zabel et al. | Optovue | 6 | 4.5 | VD (%): Whole superficial/deep-VD, Whole radial peripapillary capillaries VD | None |
| Jiang et al. | Zeiss | 3 | — | VD (Dbox): Area: ring (Ø 0.6–2.5 mm around fovea), Further separate (a), into four quadrantal sectors (b), into six concentric rings Segmentation: total, superficial and deep layers | None |
| Wu et al. | Optovue | 6 | — | VD (%): 1.5-mm ring (Ø 0.3–1.5 mm around fovea) 3-mm ring (Ø 1.5–3 mm around fovea) Each ring was separated into four quadrantal sectors | None (age and sex matched) |
| Zhang et al. | Optovue | 3 | 4.5 | VD (%) and VLD (%); Parofoveal superficial capillary plexus VD/VLD Peripapillary radial peripapillary capillary VD/VLD Peripapillary superficial vascular plexus VD/VLD Adjusted flow index: Parofoveal superficial capillary plexus | None (age, sex and race matched) |
| van de Kreeke, et al. | Zeiss | 6 | — | VD (mm²): inner ring (Ø 1–3 mm around fovea) VD outer ring (Ø 3–6 mm around fovea) ONH ring (Ø 3–6 mm around the ONH) VD | Age |
| O’Bryhim et al. | Optovue | 6 | — | VD (mm²) | Age, gender, scan quality, and diagnosis of diabetes |
| Nelis et al. | Optovue | 3 | 4.5 | VD (%): Parofoveal superficial/deep VD and OND VD Choriocapillaris [CC] parameters: Choriocapillaris (CC) decorrelation index (%) Choriocapillaris flow area (%) Analysis of signal voids | None (age matched) |
| Cennamo et al. | Optovue | 6 | — | VD (%): Superficial AND Deep capillary plexus: Whole image/parofoveal/fovea VD | Age and sex |

AD, Alzheimer disease; MCI, mild cognitive impairment; HC, healthy control; eAD, early Alzheimer-type dementia; aMCI, amnestic type of MCI; CADASIL, Cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy; OCTA, optical coherence tomography angiography; ONH, optic nerve head; VD, vessel density; VLD, vessel length density; PD, perfusion density; FAZ, foveal avascular zone; ANOVA, analysis of variance.
study (16) found no association, both adjusted and unadjusted for age and sex, whereas the other three studies (15, 20, 21) did not match or adjust for covariates.

**OCTA Parameters and Other OCT/Fundus Imaging Parameters**

Only one study (n = 52) (21) explored the relationship between OCTA features and OCT findings in patients with AD and found that increased VD was associated with increased ganglion cell–inner plexiform layer (GC-IPL) thickness. However, this study did not adjust or match for any covariates. None of the studies included investigated the relationship between OCTA parameters and traditional fundus imaging.

**DISCUSSION**

This systematic review found 14 studies investigating OCTA in patients with cerebral SVD or dementia, published between 2017 and 2020. These studies mainly focused on AD and MCI, pre-symptomatic AD, and monogenic SVD including CADASIL and Fabry disease. No other dementias, ischemic stroke, or sporadic SVD studies met the inclusion criteria. There were no longitudinal studies. Collectively, these studies do not provide a clear conclusion of whether the retinal microvasculature is impaired in cerebral SVD, ischemic stroke, and dementia patients or not.

FAZ is a capillary-free area in the foveal zone susceptible to ischemia (29). Enlargement of FAZ is a sign of ischemia (14). The analysis of the FAZ measurements yielded inconclusive results. Four studies (n = 196) (14, 20, 22, 25) found increased FAZ in patients compared with controls, whereas four other studies (n = 455) (16, 18, 24, 26) showed no difference between groups. It is possible that the relatively small sample size of some studies precludes detection of group differences. Another possibility is that some studies included patients in the earlier stage of the disease process, which might explain why no FAZ difference was found between groups (24). However, as ophthalmological confounders, quality factors, and other possible risk factor confounders including age were not always taken into account, studies reporting group differences may represent an overestimation of true disease effects (16). Furthermore, significant variation in FAZ area in normal eyes has been reported that may be associated with gender, central retinal thickness, etc. (30). Therefore, studies with larger sample size and good matching of potential confounders are needed to further verify these findings. Also, the lack of longitudinal studies means that it is difficult to know if the early OCTA changes could predate worsening neurological state, and longitudinal studies are urgently needed.

The decreased vascular density (including parameters such as VD, perfusion density, vessel length density, etc.) is another sign of microvasculature impairment. Eight studies (n = 569, seven with AD and MCI and one with CADASIL) (14, 15, 18, 20–23, 26) suggested decreased vascular density parameters in patients compared with controls. However, two studies (n = 56–86) (16, 17) showed no difference between groups. Different sample size, disease stage, and confounder adjustment might explain the disparity in results. Interestingly, van de Kreek et al. (n = 124) (24) found Aβ+ participants had higher VD than Aβ- participants in pre-symptomatic AD. Here, early amyloid accumulation in the retina might induce an inflammatory reaction with hypoxia, leading to increased retinal blood flow (24). Larger studies including those following patients longitudinally will help explore retinal vessel changes. Moreover, Cennamo et al. (n = 124) (27) found Fabry disease patients had decreased VD in superficial capillary plexus and increased VD in deep capillary plexus compared with the control group. The authors considered the increased VD in deep capillary plexus as a compensatory mechanism to support the reduced VD in superficial capillary plexus (27).
We further explored the relationships between OCTA and brain imaging findings. Currently, cerebral small vessels are hard to visualize in vivo. Other imaging features are provided as SVD biomarkers such as WMHs, microbleeds, enlarged perivascular spaces, and lacunes (1). Because the retinal vasculature is a proxy of cerebral vessels (4), establishing the relationship between retinal microvasculature and cerebral imaging findings could aid in our understanding of disease mechanisms. This systematic review found only four studies (15, 16, 19, 26) that investigated the relationship between OCTA parameters and brain imaging findings of SVD. Two studies (n = 74–86) (15, 16) found decreased VD was associated with more WMHs, whereas another study (n = 32) (26) found no association between OCTA parameters and WMHs or the number of small infarcts, which might be due to small sample size. MRI changes, such as atrophy of the brain and consequential ventricular enlargement, can be seen in AD or SVD patients (31). A pilot study (n = 16) (19) suggested the potential association between decreased VD and perfusion density with increased volume of inferior lateral ventricle (temporal horn). These small sample studies show a promising role of OCTA to investigate SVD in vivo. A prospective cross-sectional study (32) published recently when our work was under review investigating the amyloid-positive AD-related cognitive impairment (n = 28), subcortical vascular cognitive impairment (n = 18) patients, and amyloid-negative cognitively normal subjects (n = 14) found that VD was negatively correlated with SVD scores and suggested OCTA as a potential imaging tool to screen for the degree of SVD. Further studies in larger samples comparing a range of SVD brain imaging biomarkers with OCTA parameters are needed to confirm relationships and explore the underlying pathophysiological mechanisms in SVD.

Previous studies reported that the changes in retinal vessel caliber on fundus imaging were associated with cognitive decline (5). This systematic review found seven studies investigated the relationship between OCTA findings and cognitive function. Among them, three studies (n = 297) (14, 18, 23) suggested better cognitive function was associated with higher VD. One study (n = 52) (14) found enlarged FAZ area was associated with decreased MMSE score. However, four other studies (n = 266) (15, 16, 20, 21) found no association between OCTA parameters and cognitive function. A cross-sectional study (33) in 27 amnestic mild cognitive impaired patients and 29 controls published after completing this review did not find any correlation between MMSE scores and OCTA parameters. Contradictory results, relatively small samples with limited or no adjustment for covariates, and no long-term follow-up of the studies that suggest OCTA parameters as potential predictors of cognitive decline point to the necessity of larger, longitudinal studies in this area of research.

We found only one study (n = 52) (21) investigating the relationship between OCTA features and OCT findings that suggested increased VD was associated with increased GC-IPL thickness in AD patients. However, this study did not adjust for covariates. The optic nerve and retina develop as a direct extension of the diencephalon during embryonic development (4). OCT is a non-invasive method and offers high-resolution images of the retinal structure, including the neuronal layers (6). Exploring the relationship between OCT and OCTA findings especially in longitudinal studies will not only provide evidence of the temporal changes between retinal nerve layers and retinal microvasculature, but also offer potential predictors in disease progression.

We did not find any study investigating the relationship between OCTA parameters and fundus imaging. Retinal arterioles and venules typically measure 15 to 150 µm in size, whereas the capillaries measure 5 to 15 µm (34). There is an extensive literature studying the retinal vessel changes in SVD, ischemic stroke, and dementia using fundus imaging (5, 35). However, recent fundus imaging can only measure relatively large retinal vessels, whereas OCTA provides high-resolution, depth-resolved information that has been available only since 2016 (8). The relationships between
TABLE 4 | Results of included studies.

| Article | OCTA parameters in subjects vs. controls | OCTA and brain imaging findings | OCTA and cognitive function | OCTA and fundus or OCT images |
|---------|--------------------------------------|---------------------------------|-----------------------------|-----------------------------|
| Bulut et al. (14) | The group of patients with AD had lower VD and enlarged FAZ area compared with the control group | Not assessed | Increased MMSE score associated with all increased VD parameters and decreased FAZ area | Not assessed |
| Lahme et al. (15) | The group of patients with AD had lower VD compared with control group | Increased VD associated with decreased Fazekas score | No association between MMSE score and VD | Not assessed |
| Haan et al. (16) | No difference between AD and control groups | Increased VD associated with decreased Fazekas score | No association between MMSE score and OCTA parameters | Not assessed |
| Querques et al. (17) | No differences were found among groups | Not assessed | Not assessed | Not assessed |
| Yoon et al. (18) | The AD group showed reduced VD and PD compared with MCI group and control group | No difference between MCI and control groups | Increased MMSE score associated with increased VD and PD parameters | Not assessed |
| Yoon et al. (19) | Not assessed | Decreased VD and PD with increased volume of inferolateral ventricle | No association between MMSE score and FAZ | Not assessed |
| Zabel et al. (20) | AD group showed reduced VD and increased FAZ compared with control group | Not assessed | No association between MMSE score and OCTA parameters | Not assessed |
| Jiang et al. (21) | AD group showed reduced VD compared with control group | Not assessed | No association between MMSE score and VD | Increased VD associated with increased GC-IPL thickness in AD patients |
| Wu et al. (22) | The patient groups had lower VD and enlarged FAZ area compared with the control group | Not assessed | Not assessed | Not assessed |
| Zhang et al. (23) | The aMCI/eAD group showed decreased VD and adjusted flow index compared with control group | Not assessed | Increased MoCA score associated with increased VD and vessel length density | Not assessed |
| van de Kreeke et al. (24) | Aβ+ participants had higher VD than Aβ- participants | No differences in vessel length density and FAZ between two groups | Not assessed | Not assessed |
| O’Bryhim et al. (25) | The FAZ was increased in the biomarker-positive group compared with controls | Not assessed | Not assessed | Not assessed |
| Nelis et al. (26) | CADASIL group showed decreased VD compared with control group | No association between OCTA parameters and Fazekas score and number of small infarcts | Not assessed | Not assessed |
| Cennamo et al. (27) | Fabry disease group showed decreased VD in superficial capillary plexus compared with control group | Not assessed | Not assessed | Not assessed |

AD, Alzheimer disease; MCI, mild cognitive impairment; HC, healthy control; MMSE, Mini-mental State Examination; MoCA, Montreal Cognitive Assessment; OCTA, Optical coherence tomography angiography; OCT, Optical coherence tomography; FAZ, Foveal avascular zone; VD, vessel density; PD, perfusion density; VLD, Vessel length density; CADASIL, Cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy; GC-IPL, ganglion cell–inner plexiform layer.

Changes in the relatively larger vessels and smaller vessels seen only on OCTA are not fully understood. Further studies investigating the relationship between these two measurements are required. It should also be noted that longitudinal studies assessing fundus images and future neurological outcomes are also limited pointing to the need for more longitudinal studies of OCTA and fundus images in SVD.
The methodological differences limit between-study comparisons and interpretation. The variations in the different device types, software and hardware, macular scan size, segmentation methods, and retinal parameters all add to heterogeneity. Other potentially confounding factors such as age, race, and baseline status including diagnosis of diabetes are inevitable. The asymmetry between eyes of the same individual should be considered because it might cause potential misleading results when we choose one eye as a proxy for both eyes (36). On the contrary, studies utilizing both eyes of participants should address the statistical challenges cautiously because the pair of eyes are non-independent (36). OCTA scan quality has impacts on derived parameters, and researchers should perform strict quality control to ensure good data quality, and thus results are reliable and comparable. Patients with more severe disease status may have fixation problems, which may cause poor scan quality (14). Lastly, studies reviewed here were cross-sectional, and no conclusions can be drawn on the value of OCTA measurements for evaluation of disease progression.

CONCLUSIONS

The findings of this review show promise that changes in retinal microvasculature identified using OCTA are associated with monogenic SVD and different stages of AD. Larger studies with risk factors adjustment and more consistent OCTA methods are needed to fully exploit this technology.

DATA AVAILABILITY STATEMENT

All datasets generated for this study are included in the article/Supplementary Material.

REFERENCES

1. Wardlaw JM, Smith C, Dichgans M. Small vessel disease: mechanisms and clinical implications. Lancet Neurol. (2019) 18:684–96. doi: 10.1016/S1474-4422(19)30079-7
2. Bos D, Wolters FJ, Darweesh SKL, Vernooij MW, de Wolf F, Ikram MA, et al. Cerebral small vessel disease and the risk of dementia: a systematic review and meta-analysis of population-based evidence. Alzheimers Dement. (2018) 14:1482–92. doi: 10.1016/j.jalz.2018.04.007
3. Pantoñi L. Cerebral small vessel disease: from pathogenesis and clinical characteristics to therapeutic challenges. Lancet Neurol. (2010) 9:689–701. doi: 10.1016/S1474-4422(10)70104-6
4. Patton N, Aslam T, Macgillivray T, Pattie A, Deary IJ, Dhillon B. Retinal vascular image analysis: as a potential screening tool for cerebrovascular disease: a rationale based on homology between cerebral and retinal microvasculatures. J Anat. (2005) 206:319–48. doi: 10.1111/j.1469-7580.2005.00395.x
5. McGroty S, Cameron JR, Pellegrini E, Warren C, Doublan FN, Deary IJ, et al. The application of retinal fundus camera imaging in dementia: a systematic review. Alzheimers Dement. (2017) 6:91–107. doi: 10.1016/j.dadm.2016.11.001
6. Chan VTT, Sun Z, Tang S, Chen IJ, Wong A, Thom CC, et al. Spectral-domain OCTA measurements in Alzheimer’s disease: a systematic review and meta-analysis. Ophthalmology. (2019) 126:497–510. doi: 10.1016/j.ophtha.2018.08.009
7. Jia Y, Bailey ST, Hwang TS, McClintic SM, Gao SS, Pennesi ME, et al. Quantitative optical coherence tomography angiography of vascular abnormalities in the living human eye. Proc Natl Acad Sci USA. (2015) 112:E2395–402. doi: 10.1073/pnas.1500185112
8. Kashani AH, Chen CL, Gahm JK, Zheng F, Richter GM, Rosenfeld PJ, et al. Optical coherence tomography angiography: a comprehensive review of current methods and clinical applications. Prog Retin Eye Res. (2017) 60:66–100. doi: 10.1016/j.preteyeres.2017.07.002
9. Rosenfeld PJ, Durbin MK, Roisman L, Zeng F, Miller A, Robbins G, et al. ZEISS angioplex spectral domain optical coherence tomography angiography: technical aspects. Dev Ophthalmol. (2016) 56:18–29. doi: 10.1159/000442773
10. The Lancet. Artificial intelligence in health care: within touching distance. Lancet. (2018) 390:2739. doi: 10.1016/S0140-6736(17)31540-4
11. Sandhu HS, Elmogy M, El-Adawy N, Eltanboly A, Shalaby A, Keynton R, et al. Automated diagnosis of diabetic retinopathy using clinical biomarkers, optical coherence tomography (OCT), and OCT angiography. Am J Ophthalmol. (2020) 216:201–6. doi: 10.1016/j.ajo.2020.01.016
12. Moher D, Liberati A, Tetzlaff J, Altman DG. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. PLoS Med. (2009) 6:e1000097. doi: 10.1371/journal.pmed.1000097
13. Wardlaw JM, Smith EE, Biessels GJ, Cordonnier C, Fazekas F, Frayne R, et al. The Lancet. Artificial intelligence in health care: within touching distance. Lancet. (2018) 390:2739. doi: 10.1016/S0140-6736(17)31540-4
14. Sandhu HS, Elmogy M, El-Adawy N, Eltanboly A, Shalaby A, Keynton R, et al. Automated diagnosis of diabetic retinopathy using clinical biomarkers, optical coherence tomography (OCT), and OCT angiography. Am J Ophthalmol. (2020) 216:201–6. doi: 10.1016/j.ajo.2020.01.016
15. Moher D, Liberati A, Tetzlaff J, Altman DG. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. PLoS Med. (2009) 6:e1000097. doi: 10.1371/journal.pmed.1000097
16. Wardlaw JM, Smith EE, Biessels GJ, Cordonnier C, Fazekas F, Frayne R, et al. Neuroimaging standards for research into small vessel disease and its contribution to ageing and neurodegeneration. Lancet Neurol. (2013) 12:822–38. doi: 10.1016/S1474-4422(13)70124-8
17. Bulut M, Kurtulus F, Gozkaya O, Erol MK, Cengiz A, Akidan M, et al. Evaluation of optical coherence tomography angiographic findings in Alzheimer’s type dementia. Br J Ophthalmol. (2018) 102:233–7. doi: 10.1136/bjophthalmol-2017-310476

AUTHOR CONTRIBUTIONS

J-FZ, SW, and MV-H searched the literature, drafted the manuscript, collected the data, and performed statistical analyses. J-FZ, SW, MV-H, FD, BD, Y-CW, and JW revised the manuscript. J-FZ, SW, MV-H, Y-CW, and JW contributed to conception, design, and data interpretation of the study. SW, MV-H, FD, BD, Y-CW, and JW contributed to manuscript revision for critical intellectual content and supervision of the study. All authors read and approved the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fneur.2020.01009/full#supplementary-material
15. Lahme L, Esser EL, Mihailovic N, Schubert F, Lauermann J, Johnen A, et al. Evaluation of ocular perfusion in Alzheimer’s disease using optical coherence tomography angiography. J Alzheimers Dis. (2018) 66:1745–52. doi: 10.3233/JAD-180738
16. Haan JD, van de Kreeke JA, van Berckel BN, Barkhof F, Teunissen CE, Scheltens P, et al. Is retinal vasculature a biomarker in amyloid proven Alzheimer’s disease? Alzheimers Dement. (2019) 11:383–91. doi: 10.1016/j.jalz.2019.03.006
17. Querques G, Borrelli E, Sacconi R, De Vitiis L, Leocani L, Santangelo R, et al. Functional and morphological changes of the retinal vessels in Alzheimer’s disease and mild cognitive impairment. Sci Rep. (2019) 9:63. doi: 10.1038/s41598-018-13727-6
18. Yoon SP, Grewal DS, Thompson AC, Polascik BW, Dunn C, Burke JR, et al. Retinal microvascular and neurodegenerative changes in Alzheimer's disease and mild cognitive impairment compared with control participants. Ophthalmol Retina. (2019) 3:489–99. doi: 10.1016/j.iors.2019.02.002
19. Yoon SP, Thompson AC, Polasck BW, Caliote C, Burke JR, Petrella J, et al. Correlation of OCTA and volumetric MRI in mild cognitive impairment and Alzheimer disease. Ophthamlic Surg Lasers Imaging Retina. (2019) 50:709–18. doi: 10.3928/23258160-20191031-06
20. Zabel P, Kaluzny J, Wlkosc-Debczynska M, Gbeka-Toloczek M, Suwala K, Zabel K, et al. Comparison of retinal microvasculature in patients with Alzheimer’s disease and primary open-angle glaucoma by optical coherence tomography angiography. Invest Ophthalmol Vis Sci. (2019) 60:5447–55. doi: 10.1167/iovs.19-27028
21. Jiang H, Wei Y, Shi Y, Wright CB, Sun X, Gregori G, et al. Altered macular microvasculature in mild cognitive impairment and Alzheimer disease. J Neuroophthalmol. (2018) 38:292–8. doi: 10.1097/WNO.0000000000000580
22. Wu J, Zhang X, Azhati G, Li T, Xu G, Liu F. Retinal microvascular attenuation in mental cognitive impairment and Alzheimer’s disease by optical coherence tomography angiography. Acta Ophthalmol. (2020) 98:e781–7. doi: 10.1111/aos.14381
23. Zhang YS, Zhou N, Knoll BM, Samra S, Ward MR, Weintraub S, et al. Parafoveal vessel loss and correlation between peripapillary vessel density and cognitive performance in amnestic mild cognitive impairment and early Alzheimer’s disease on optical coherence tomography angiography. PLoS ONE. (2019) 14:e0214685. doi: 10.1371/journal.pone.0214685
24. van de Kreeke JA, Nguyen HT, Konijnenberg E, Tomassen J, den Braber A, Ten Kate M, et al. Optical coherence tomography angiography in preclinical Alzheimer’s disease. Br J Ophthalmol. (2020) 104:157–61. doi: 10.1136/bjo.2019-314127
25. O’Bryhim BE, Apte RS, Kung N, Coble D, Van Stavern GP. Association of preclinical Alzheimer’s disease with optical coherence tomographic angio findings. JAMA Ophthalmol. (2018) 136:1242–8. doi: 10.1001/jamaophthalmol.2018.3556
26. Nelis P, Kleffner I, Burg MC, Clemens CR, Alnawanishe M, Motte J, et al. OCT-angiography reveals reduced vessel density in the deep retinal plexus of CADASIL patients. Sci Rep. (2018) 8:8148. doi: 10.1038/s41598-018-26475-5
27. Cennamo G, Di Maio LG, Montorio D, Tranfa F, Russo C, Pontillo G, et al. Optical coherence tomography angiography findings in Fabry disease. J Clin Med. (2019) 8:528. doi: 10.3390/jcm8040528
28. Li XX, Wu W, Zhou H, Deng Jj, Zhao MY, Qian TW, et al. A quantitative comparison of five optical coherence tomography angiography systems in clinical performance. Int J Ophthalmol. (2018) 11:1784–95. doi: 10.18240/jio.2018.11.09
29. Snodderly DM, Weinzhaus RS, Choi JC. Neural-vascular relationships in central retina of macaque monkeys (Macaca fascicularis). J Neurosci. (1992) 12:1169–93. doi: 10.1523/JNEUROSCI.12-04-01169.1992
30. Fujiwara A, Morizane Y, Hosokawa M, Kimura S, Shiode Y, Hirano M, et al. Factors affecting foveal avascular zone in healthy eyes: an examination using swept-source optical coherence tomography angiography. Prog Retin Eye Res. (2017) 62:188572. doi: 10.1017/S0140670X17000583
31. Prioni GB, Fox NC, Jack CR Jr, Scheltens P, Thompson PM. The clinical use of structural MRI in Alzheimer disease. Nat Rev Neurol. (2010) 667–77. doi: 10.1038/nrneurol.2009.215
32. Lee JY, Kim JP, Jang H, Kim J, Kang SH, Kim JS, et al. Optical coherence tomography angiography as a potential screening tool for cerebral small vessel diseases. Alzheimers Res Ther. (2020) 12:73. doi: 10.1186/s13195-020-00638-x
33. Criscuolo C, Cennamo G, Montorio D, Carotenuto A, Strianese A, Salvatore E, et al. Assessment of retinal vascular network in amnestic mild cognitive impairment by optical coherence tomography angiography. PLoS ONE. (2020) 15:e0233975. doi: 10.1371/journal.pone.0233975
34. An D, Balаратnasimgam C, Heisler M, Francke A, Ju M, McAllister IL, et al. Quantitative comparisons between optical coherence tomography angiography and matched histology in the human eye. Exp Eye Res. (2018) 170:13–9. doi: 10.1016/j.exer.2018.02.006
35. Cheung CY, Ikram MK, Chen C, Wong TY. Imaging retina to study dementia and stroke. Prog Retin Eye Res. (2017) 57:89–107. doi: 10.1016/j.preteyeres.2017.01.001
36. Cameron JR, Megaw RD, Tatham AJ, McGrory S, MacGillivray TJ, Double BN, et al. Lateral thinking - interocular symmetry and asymmetry in neurovascular patterning, in health and disease. Prog Retin Eye Res. (2017) 59:131–57. doi: 10.1016/j.preteyeres.2017.04.003

Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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GLOSSARY

OCTA, optical coherence tomography angiography; SVD, small vessel disease; AD, Alzheimer disease; MCI, mild cognitive impairment; CADASIL, cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy; VD, vessel density, OCT, optical coherence tomography; FAZ, foveal avascular zone; WMHs, white matter hyperintensities; MMSE, Mini-Mental State Examination.