OPTICAL DEVICES FOR ADULTS WITH LOW VISION: A SYSTEMATIC REVIEW OF PUBLISHED STUDIES OF EFFECTIVENESS

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http://www.va.gov/vatap
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This summary form is intended as an aid for those who want to record the extent to which a HTA report meets the 17 questions presented in the checklist. It is NOT intended as a scorecard to rate the standard of HTA reports – reports may be valid and useful without meeting all of the criteria that have been listed.

### VA Technology Assessment Program Report
**Optical devices for adults with low vision:**
*A systematic review of published studies of effectiveness*  
(MAY 2003)

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Palo Alto, California
# Executive Summary

Purpose: The purpose of this review is to provide an overview of optical low vision devices and their effectiveness in improving the quality of life for individuals with vision loss. The review will cover various aspects such as definition, causes, VA services, demographics, uses, and methods of assessing the effectiveness of low vision devices.

## Background

### Definition of low vision
Low vision is defined as a visual impairment that limits daily activities and cannot be corrected by standard eyeglasses or contact lenses.

### Impact of vision loss on the visually impaired
Vision loss significantly affects the quality of life, independence, and employment opportunities for individuals with low vision.

### Causes of vision loss in the veteran population
Common causes of vision loss in the veteran population include macular degeneration, glaucoma, and diabetic retinopathy.

### VA services for visual impairment
The VA offers a range of services and resources to support individuals with visual impairment, including low vision devices.

### Demographics and trends in the veteran population
There is an increasing number of veterans with visual impairment, highlighting the need for effective low vision devices and services.

## Low Vision Devices Included in this Review

Special considerations for this review include the selection criteria for devices included in the review.

## Uses of Low Vision Devices

### Reading
Low vision devices enhance reading capabilities for individuals with low vision.

### Driving
Low vision devices support safe driving for individuals with visual impairment.

## Measuring the Effectiveness of Low Vision Devices

Methods of measuring effectiveness include clinical trials, patient satisfaction surveys, and quality of life assessments.

## Regulation and Reimbursement of Low Vision Devices

### Food and Drug Administration
The FDA regulates the safety and effectiveness of low vision devices.

### VHA Reimbursement
The VA provides reimbursement for low vision devices under its medical benefits.

### Federal Medicare Reimbursement
Low vision devices are reimbursable under Medicare for eligible individuals.

## Methods

### Scope of review
The scope of the review includes devices that are commonly used for low vision.

### Search strategy
The search strategy involves a comprehensive database search to identify relevant studies.

### Other data sources
Other sources of information include clinical guidelines and expert opinions.

### Inclusion criteria
Studies included in the review meet specific criteria based on methodology and relevance.

### Critical appraisal
A critical appraisal of the included studies is conducted to assess their quality.

## Results

### Primary data—reading
Results from primary studies on the effectiveness of reading devices are analyzed.

### Primary data—driving
Results from primary studies on the effectiveness of driving devices are analyzed.

### Prior reviews of effectiveness of low vision devices or low vision rehabilitation
Prior reviews of the effectiveness of low vision devices or low vision rehabilitation programs are evaluated.

### Limitations of this review
The limitations of the review, including potential sources of bias, are identified.

## Summary

Knowledge gaps and recommendations for future research are identified to improve the effectiveness of low vision devices.

## Conclusions and Recommendations

Conclusions based on the review findings are presented, along with recommendations for future research and clinical practice.

## End References

Articles included for critical appraisal, background articles, and excluded articles are listed at the end of the report.
EXECUTIVE SUMMARY

- Vision loss in adults is associated with many comorbidities, activity limitations and lower quality of life. The prevalence of vision loss both in the general and veteran populations is projected to increase dramatically over the next 20 years. With this trend is a growing need for low vision services targeted at detecting visual impairment and mitigating the functional consequences associated with age-related vision loss to improve quality of life.

- Provision of low vision services, including low vision devices, to veterans is a priority for VA. Popular among veterans and practitioners are electronic optical devices such as closed circuit TV (CCTV), computer assistive technologies, and non-electronic hand held models. However, advocates for visually impaired veterans have expressed concern over the quality of scientific evidence supporting the use of many low vision devices, particularly newly emerging electronic devices.

- A multidisciplinary task force of VA vision care experts has been charged with developing a process for evidence-based new technology evaluation and dissemination of information in VA. To inform the task force, this systematic review identified: 1) the existing evidence of effectiveness of optical low vision devices from the peer-reviewed published literature; 2) knowledge gaps, and; 3) evidence-based tools for assisting data collection and clinical decision-making.

- This review reveals a paucity of high quality evidence in the peer-reviewed, published literature to inform choices about provision of optical low vision devices in VA. The best evidence consisted of seven small, prospectively controlled clinical studies comparing the performance of optical low vision devices for reading tasks at various distances in a controlled indoor setting. Evidence of preferences and use were anecdotal. Sustained use of these devices in the subject’s life setting, resources required in terms of costs and training associated with each alternative, and the link between device use and health related quality of life were unknown.

- In the absence of compelling evidence from published research and a standard taxonomy of desired outcomes, clinicians must continue to rely on industry data, clinical observations, patient self-reporting, and real-world trials in determining appropriate provision of low vision aids.

- Future research is needed to determine the appropriate candidacy for low vision devices, suitable prescription of these devices, and outcome measures that define the quality of life in subjects with age-related visual impairment along the continuum of visual impairment and disability. TAP encourages using an evidence-based framework in evaluating evidence of effectiveness, partnering with industry and consumers to carry out evidence-based technology evaluation, and seeking out systematic information that can improve and standardize current prescription practices.
OPTICAL DEVICES FOR ADULTS WITH LOW VISION: A SYSTEMATIC REVIEW OF PUBLISHED STUDIES OF EFFECTIVENESS

PURPOSE

The VA Technology Assessment Program (TAP) generated this systematic review in response to a request from VA Office of Patient Care Services (PCS) to evaluate the effectiveness of low vision aids for the visually impaired veteran.

VA is committed to rehabilitation of visually impaired veterans because of its own mission and on behalf of stakeholders. The Blinded Veterans Association (BVA)\(^1\) expressed concerns over the quality of the scientific evidence supporting the use of many low vision aids, particularly electronic devices that have emerged in recent years. Within VA, PCS and VA’s Visual Impairment Advisory Board\(^2\) identified timely evaluation of technology as a priority area of service delivery. Finally, the VA Under Secretary for Health identified visual impairment, including provision of low vision services, as an important subject area for the Veterans Health Initiative.\(^3\)

VA clinicians and managers involved in care for the visually impaired veteran face requests from veterans for new devices that are marketed without evidence of effectiveness, evaluation criteria or application. To develop a process for new technology evaluation and dissemination of information in VA, a multidisciplinary task force of VA vision care experts is being formed.

Proponents of newly marketed products often advocate improvements over existing technologies. Choices about which technologies to purchase and use require evidence of effectiveness relative to available alternatives. To that end, this systematic review will inform the task force in a number of ways:

- By identifying the current knowledge base in the peer-reviewed published literature that addresses the relative effectiveness of a wide range of optical low vision aids;
- By identifying gaps in the existing knowledge base from which recommendations for future research can be made;
- By recommending instruments and techniques to assist evidence-based data collection and clinical decision-making.

\(^1\) BVA is a veteran service organization that provides access to information about new technologies, including education and awareness regarding efficacy, and advocates for all visually impaired individuals who rely on technology to function in their work and personal lives.

\(^2\) VIAB is a multidisciplinary panel of subject matter experts in eye care and blind rehabilitation with representation from the field, Central Office, Veteran Integrated Service Network administrative and clinical leaders, and stakeholders.

\(^3\) Veterans Health Initiative is “a comprehensive program in VA to recognize the connection between certain health effects and military service, to allow veterans to document their military medical history, and to prepare health care providers to better serve their veteran patients, and to establish a database for further study.” [http://www.va.gov/VHI/]

VA OPCS Technology Assessment Program \hspace{1cm} http://www.va.gov/vatap
BACKGROUND

Definition of low vision
While there is no universal consensus on a definition of low vision, generally low vision is thought of as vision loss that cannot be corrected medically, surgically or with conventional eyeglasses, and interferes with daily activities. The World Health Organization’s definitions of the terms “disorder”, “impairment”, “disability”, and “handicap” are used frequently in defining low vision and remedial services (Leat 1999). Disorder refers to the disease in terms of ocular pathology and etiology. Impairment refers to sensory deficit in the visual system, while disability refers to the inability to perform activities that are important to the person. Handicap relates to the economic, social or psychological changes incurred as a result of a visual impairment.

Applying this construct to service delivery, Massof (1995) states that medical and surgical interventions treat disorders to limit or reverse impairments, whereas rehabilitation enhances impairment to limit or reverse disabilities. Human services assist, accommodate, and educate persons with disabilities to limit or reverse handicaps. The goal of low vision rehabilitation, including provision of low vision devices discussed in this review, is to ameliorate vision disabilities and improve the quality of life of these patients.

In the US impairment-based definitions classified by performance on visual tests such as visual acuity and visual field are commonly used to define low vision. According to the International Classification of Diseases, 9th revision (ICD-9-CM), abnormal vision includes five levels ranging from moderate low vision (20/80) to total blindness (no light perception). The term “legal blindness” is defined in the US as visual acuity of 20/200 or worse in the better eye with best corrective lens or visual field restricted to 20 degrees or less in diameter in the better eye (Public 2002). In practice, visual impairment can begin to have a functional affect at approximately 20/50, which is roughly the size of newsprint, and many who are classified as legally blind may have residual vision and can benefit from low vision services.

Impact of vision loss on the visually impaired
A growing body of research suggests a correlation between vision loss and a variety of patient outcomes. An analysis of data from the second Supplement on Aging, 1994 (SOA-II) using an International Classification of Functioning, Disability and Health (ICIDH-2) framework showed that vision impairment among the elderly was associated with a variety of medical conditions (diabetes, arthritis, hypertension, heart conditions, stroke, osteoporosis, disorientation and confusion, depression, broken hips and history of falls), activity limitations and participation restrictions (Crews 2001). Recent

4ICIDH-2 is a conceptual framework that systematically groups functional states associated with health conditions. It identifies three domains of human experience (body functions and structures, activity, and participation) and recognizes the role of the environment as a factor that enables and disables people (WHO 2001).
observational studies reported an association between visual impairment and increased risk of mortality (Wang 2001; McCarty 2001), falls and hip fracture (McCarty 2002; Ramrattan 2001; Lord 2001; Ivers 2000 (a); Klein 1998), depression (Rovner 1998; Brody 2001), and memory decline (Anstey 2001).

Visual impairment in patients with glaucoma, age-related macular degeneration, and diabetes was associated with decreased functional status, decreased self-reported quality of life, and increased emotional distress (Stelmack 2001). Among patients with permanent visual acuity loss due to age-related macular degeneration or diabetic retinopathy, utility values in terms of time trade-off were directly dependent on the level of visual acuity in the better seeing eye caused by the disease, and not on the disease itself.

Dual sensory impairment (vision and hearing loss) is a particular challenge for the elderly and their caregivers. To note, approximately 52% of veterans enrolled in VA Blind Rehabilitation programs have some degree of hearing loss (VA BRS 2002). Recent cross-sectional research suggests a correlation between dual sensory impairment (visual and hearing) and a variety of health outcomes in the elderly, such as comorbidities and greater difficulty performing activities (Campbell 1999), as well as functional status independent of either mental status or comorbid illness (Keller 1999; Reuben 1999). Evidence on the association between dual sensory impairment and mortality is inconclusive (Reuben 1999; Appollonio 1995; Laforge 1992).

According to the Veterans Health Initiative Independent Study Course on Vision Impairment, the impact of vision loss on an individual’s activities of daily living can be dramatic and affect the individual’s social, familial, occupational, and recreational life (VHI 2002). Vision loss can change a person’s ability to perform ordinary but important tasks such as reading, driving, preparing meals, taking prescription medications, and maintaining personal finances. Hobbies may end, and tasks such as correspondence and facial recognition may be compromised. As a result, persons with vision loss can experience less mobility and functional independence, and greater psychological stress, isolation from family and friends, risk of poverty, and potential loss of their homes.

While the reasons underlying the correlations between vision loss and patient outcomes require further examination, the evidence suggests potentially far-reaching effects of visual impairment on the health and lives of the aged. A reasonable rationale can be made for provision and evaluation of a full range of rehabilitation services to veterans using frameworks that take into account the individual’s age, overall health, performance of activities, social circumstances, and environment.

**Causes of vision loss in the veteran population**

Causes of vision loss in the veteran population are primarily age-related (VA BRS 2002). They are macular degeneration, glaucoma, diabetic retinopathy, and cataract. Of these, cataracts are commonly remediated through surgery. Other conditions frequently leave the veteran with visual impairment, which can be severe. Less common causes found in the veteran population are cerebrovascular accidents, optic
nerve disease (eg. multiple sclerosis), retinitis pigmentosa, and service-related injuries or complications such as trauma and long-term effects of inhumane treatment of prisoners of war.

**VA services for visual impairment**

Since World War II VA has provided the only nation-wide comprehensive rehabilitation service to American’s blinded veterans and is recognized internationally as a leader in rehabilitation of the blind. The Blind Rehabilitation Service provides:

“…a **continuum of care** for blinded veterans extending from their home environment to the local VA facility and to the appropriate rehabilitation setting. These services include adjustment to blindness counseling, patient and family education, benefits analysis, comprehensive residential inpatient training, outpatient rehabilitation services, the provision of assistive technology, and research.” (VA BRS 2002)

VA offers a variety of low vision services to eligible veterans depending on the degree of visual impairment. VA enrollees or veterans who are legally blind and eligible for non-vision related VA health care services are eligible for a comprehensive array of assistance through the VA Blind Rehabilitation Service. Visually impaired veterans who experience functional difficulties but are not eligible for blind rehabilitation may be eligible for assistance from low vision clinics at VA facilities, the Visual Impairment Services Outpatient Rehabilitation (VISOR) program, or the Vision Impairment Center to Optimize Remaining Sight (VICTORS) inpatient/outpatient programs.

The Prosthetic and Sensory Aids Service (PSAS) furnishes blind aids and prosthetic equipment related to sight loss to eligible veterans with visual impairment. For veterans with visual impairment not caused as a result of active military service or with an income and net worth above established thresholds, co-payment for services is generally required (DVA eligibility 2002).

**Demographics and trends in the veteran population**

A VA national survey described the typical veteran receiving low vision services through VA Blind Rehabilitation Services:

“He is an elderly Caucasian male, who is married and lives in a home with his wife. His visual impairment is due to macular degeneration, and his visual acuity is approximately 20/200. This typical veteran had a high school education, and has now retired on an income that would place his family in the lower middle class. The veteran is active, and participates in such activities as socializing with friends, shopping for groceries, and participating in clubs or organizational activities.” (Watson 1997(a))

Between 2000 and 2020 the total population of veterans is projected to decline, but the **proportion** of veterans age 65 and older is projected to increase from 38% to 51%,
respectively (DVA 2000). Even more dramatic is a projected eight-fold increase in the veteran population age 85 and older during the same time period.

There are currently over 93,000 legally blind veterans. Applying prevalence estimates of legal blindness for specific chronological ages\(^5\) to the VetPop2001\(^6\) veteran population projections, by the year 2007, over 161,000 will meet the criteria for legal blindness, and over 1.2 million will have visual impairments including but not limited to legal blindness (De l'Aune 2002).

The VA Blind Rehabilitation Service has added outpatient rehabilitation services in the form of Visual Impairment Service Team (VIST) coordinators and Blind Rehabilitation Outpatient Specialists (BROS) to meet the special vision needs of an increasingly blinded geriatric veteran population. From Fiscal Year (FY) 1997 to FY 2000, the number of veterans served by VIST coordinators and BROS increased 16% (from 30,313 patients to 35,172 patients) and 152% (from 873 patients to 2,199 patients), respectively (Beck 2001).

Data from the PSAS National Prosthetic Patient Database from FY 1999 to FY 2001 show a 7% increase in the number of discharges at Blind Rehabilitation Centers with a corresponding 23% increase in the number of aids furnished for the blind (Table 1).

Table 1. National Prosthetic Patient Database: Blind aids furnished by PSAS from FY 99 to FY 01

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<td>BRC discharges</td>
<td>1,760</td>
<td>1,817</td>
<td>1,885</td>
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<td>Aids for the Blind</td>
<td>61,477</td>
<td>72,560</td>
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* Projected based on 1\(^{st}\) & 2\(^{nd}\) quarters data

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5 Center for Health Statistics, Health Interview Survey, Disability Supplement (1994-95) under funding from DVA Rehabilitation Research and Development (C-2704-I “Secondary Data Analysis Relevant to Low Vision Rehabilitation”) and the DVA Visual Impairment Advisory Board.

6 VA’s official estimate and projection of the number and characteristics of veterans based on Census 1990 data, as of 9/30/01. Prepared by the Office of the Actuary, Office of Policy and Planning, Department of Veterans Affairs, April 2002. [http://www.va.gov/vetdata/demographics/index.htm](http://www.va.gov/vetdata/demographics/index.htm). Prevalence estimates using VetPop2001Adj, VA’s official adjusted population estimates based on Census 2000 data were not available at the time this report was released.
Figures 1 and 2 show that in FY 2002, several optical low vision devices were among the top 12 blind aids furnished to veterans in terms of either high unit costs (eg. 2,047 CCTVs prescribed at an average unit cost of $2,047) or high prescribed volume (eg. 21,524 hand held devices prescribed at an average unit cost of $75).

Figure 1. National Prosthetic Patient Database: Top 12 Blind Aids by Total Cost (Raw Data) for FY 2002

Figure 2. National Prosthetic Patient Database: Top 12 Blind Aids by Total Quantity (Raw Data) for FY 2002
LOW VISION DEVICES INCLUDED IN THIS REVIEW

Most low vision devices (LVD) are designed to enhance access to information and ensure safe mobility (Scherer 1996). A range of adaptive technologies for low vision exists in electronic and non-electronic forms from optical devices that incorporate refractive correction and require prescription to non-optical devices.

Optical devices are the subjects of this review because of their popularity among VA patients and practitioners and their associated costs. An optical LVD is any device that alters the image focus, size, contrast, brightness, color or directionality of an object through the use of lenses or other technology. Optical LVDs are intended: 1) to improve visual acuity by enlarging images or by clarifying images through improved illumination, color, or contrast enhancement, or 2) to enhance the field of view.

In VA, such devices include, but are not limited to, spectacles with or without tint, microscopic spectacles, hand held magnifiers, stand magnifiers, telescopes (monocular or binocular), head mounted lenses, minifiers, prisms, and closed-circuit televisions (CCTV) (DUSH 2000). This review will consider the full range of optical devices available to VA patients.

Special considerations for this review

Intraocular lens. Intraocular lens (or “implants”) are implanted surgically in patients with cataract surgery to restore lost vision due to diseased tissue. Since this review focuses on low vision aids used to optimize irreversible vision loss, these devices were not included. However, individuals with implants may be included in studies in this review if they have persistent, irreversible low vision after surgery and are candidates for low vision rehabilitation.

Contact lens. Occasionally, contact lenses are prescribed as an “add-on” technology for veterans with vision loss (G. Mancil: personal communication July 18, 2002). The major use of contact lenses is to correct myopia, but other uses of contact lenses are: to correct hyperopia, astigmatism, presbyopia, aphakia, and irregular corneal surfaces; to provide therapeutic protection for certain conditions, e.g. bullous keratopathy and recurring corneal erosion; and to improve comfort, vision, and wound healing during the postoperative period immediately following photorefractive keratectomy (Yanoff 1999). Contact lenses may be preferred purely for cosmetic reasons (e.g. to avoid wearing spectacles or to change iris color). The two types of contact lenses are scleral and corneal. Scleral contact lenses were developed first, but corneal lenses have superceded their use except for rare clinical conditions (Foss 1994).

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7 spectacle-mounted lenses that exceed the upper power limit of standard reading glasses (+3.50 diopters) and generally range in power from +4.00 to +80.00 diopters; also called high-plus lenses, strong reading lens or high adds
8 devices that increase the amount of information in the visual field in proportion to minification (e.g. a 2x minifier provides 2x more information in the visual field); visual scanning is improved but usually at the expense of visual acuity.
Contact lenses may be prescribed along with other low vision aids, for example with a spectacle-mounted objective lens to create a contact lens telescope. However, improved visual acuity through the use of a telescope comes at the expense of a reduced visual field. The rationale for using a contact lens telescope is: 1) to increase the visual field (relative to that provided by spectacle-mounted telescopes) by reducing the distance between the eye and the contact lens, and 2) to improve cosmesis and comfort by creating a single spectacle-mounted lens. For this review, any contact lens that is used alone or with other low vision aids to correct refractive abnormalities in a patient who would be a candidate for low vision rehabilitation will be considered.

**Visual field enhancers.** Approaches to rehabilitation for patients with visual field deficits employ a variety of optical aids and adaptive strategies to make effective use of residual vision. The patient with a significantly restricted field has only a limited range of easy and comfortable eye movement scanning in the direction of the field loss beyond which scanning requires undue effort or head movement (Bailey 1978). Commonly called “field expanding devices”, they do not increase the usable visual field. Rather they bring awareness to the area of visual deficit by displacing images toward the residual field to make it easier for the patient by reducing the extent of ocular and head rotation.

Optical aids for visual field loss include spectacle-mounted mirrors, reverse telescopes\(^9\), amorphic lens\(^10\), cemented prisms, Fresnel press-on prisms, CCTV, as well as devices for simple magnification and illumination. Adaptive strategies comprise scanning therapy, eccentric viewing, orientation and mobility training, occupational therapy, cane techniques, and guide dogs. Proficiency with these devices and strategies for visual field enhancement usually requires extensive training and adaptation. The optical aids used for visual field loss are emphasized in this review, but they may be compared with training techniques to accomplish the same task—orientation, mobility and reading being the most common.

**USES OF LOW VISION DEVICES**

*Reading and driving are the uses covered in this review.* Reading encompasses many vital activities of daily living important for maintaining independence and is the most common objective of low vision rehabilitation, including prescription of LVDs (Tobin 1990; Elliott 1997; Hall 1987; Leat 1999). Driving is an essential component of American culture. Retaining licensure for driving is important for preserving independence and quality of life in a growing number of older adults with low vision, and a large volume of literature has been devoted to driving with visual impairment. The VA Physical Medicine and Rehabilitation Service offers a specialized Drivers Rehabilitation Program to eligible veterans at 39 VA Medical Centers nationwide, in which visual assessment is an important component (VA PM&RS 2003).

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\(^9\) a telescope that minimizes, as opposed to magnifies, images to allow more information in the visual field

\(^10\) a spectacle mounted cylindrical reverse telescope designed to expand the horizontal field
Reading
Difficulty with reading can be the most serious and important result of visual impairment. Evaluation of reading performance is important for recommending appropriate aids to patients with low vision. The literature suggests that a constellation of visual and nonvisual factors may influence reading performance (Mancil 1986; Legge 1992). Visual factors include nature of the ocular disease, reduction in visual field, visual acuity, or contrast, inadequate lighting, effects of magnification, and text and background colors. Nonvisual factors include motivation, attitude, educational background, age, cosmesis and type of optical system.

Generally, reading performance is measured using standardized tests of reading speed (e.g. Minnesota Low-vision Reading Test) at prescribed distances and letter sizes in a clinical setting. Reading comprehension and duration may also be factored in. However, these tests may be impractical for general practitioners to administer, and their results may not represent situations encountered in real world conditions. Research to identify accurate, simpler, resource-conserving methods using clinical factors or visual tests as predictors of reading performance has been inconsistent or inconclusive (Humphry 1986; Legge 1992; Turco 1994; Ahn 1995; Leat 1997; Watson 1997(b); Lowe 2000; Wolffsohn 2000; Brabyn 2001; West 2002). For now, direct measurement of reading performance is required to find the low vision aid that provides maximum performance and meets the patient’s needs, ideally under real world conditions.

Driving
Driving is a complex skill requiring integration of visual, cognitive and motor components. There is general agreement that individuals with severe visual impairment should not drive, but driving for moderately visually impaired individuals is controversial (Owsley 1999; Barron 1991). A variety of LVDs, such as bioptics\textsuperscript{11} and tinted lenses and filters, can be used to make driving possible for a portion of visually impaired individuals.

Standard tests for visual acuity and occasionally visual field and color vision are done to identify high-risk drivers, but minimum visual requirements for licensure of visually impaired individuals vary across states and usually involve several steps of evaluation and training to ensure safe driving (Peli 2002). To note, the use of bioptic devices as a visual aid for driving is permitted in 28 states.

Research into a causal relationship between visual factors (e.g. static and dynamic visual acuity, contrast sensitivity, visual field, glare, depth perception, useful field of view\textsuperscript{12} (UFOV), night vision, and color vision) and driving performance and safety in

\textsuperscript{11} A small spectacle-mounted telescope used to increase visual acuity and aid the driver in seeing objects at a distance

\textsuperscript{12} Useful field of view\textsuperscript{®} is a test of divided attention. UFOV\textsuperscript{®} is a proprietary trademark of Visual Resources, Inc. (Chicago, IL); its version is a computer-administered and computer-scored test of divided attention that uses both visual and cognitive skills to determine how an adult driver processes visual information.
older drivers has been inconclusive (Staplin 1998; Owsley 1999). Recent crosssectional analyses of subjects with cataracts suggest associations between decreased visual acuity and a variety of complex driving situations including driving at night, decreased contrast sensitivity and difficulty in making left turns, UFOV® and difficulty driving in the rain, and decreased contrast sensitivity in one or both eyes on at-fault crash risk (Ivers 2000 (b); McGwin 2000; Owsley 2001).

Despite an absence of conclusive evidence of a causal relationship between visual factors, driving performance and safety, none would argue that vision is a vital component of safe driving. Current evidence suggests that in addition to standard vision testing, selected measures of visual function corresponding to specific causes of vision loss (e.g. contrast and glare sensitivity with cataract) or self-reported impairment may identify older drivers who are at risk for experiencing situational driving difficulty. Interventions targeted at correcting specific vision loss may help mitigate factors associated with difficult and unsafe driving, and thereby help maintain independence and avoid social isolation.

MEASURING THE EFFECTIVENESS OF LOW VISION DEVICES

“Quality of vision is an integral part of quality of life.” (Stelmack 2001)

Low vision devices are an important part of low vision rehabilitation. Frequently, effectiveness13 of low vision interventions is measured using performance-based indicators of functional ability (eg. reading speed or duration), continued use or satisfaction with the device (Raasch 1997). Effectiveness may also consider measures of efficiency, that is, the amount of resources used to achieve desired objectives of rehabilitation. While measures of function, use and satisfaction are used most often in evaluation and provision of services, they may not reflect the overall impact of the intervention to the patient, especially in terms of health related quality of life.

Health-related quality of life (HRQOL) is a broad term that deals with five dimensions of an individual’s life: duration of life, impairments, functional states, perceptions, and social opportunities (Patrick 1989). HRQOL instruments consider physical, social, and emotional aspects along with functional status to capture clinically relevant outcomes that reflect a patient’s point of view. Such instruments allow health care providers to compare interventions across conditions and populations and to compare condition-specific interventions more in-depth to maximize provision of health care services. Demonstrating effectiveness using HRQOL is a means of introducing accountability for use of resources and quality of care in evidence-based health care organizations (e.g. for accreditation or as a stipulation of funding for research or clinical services). VA now requires outcome measurement to monitor and improve the efficiency and effectiveness of its blind rehabilitation programs.

13 Effectiveness is the extent to which an intervention achieves its intended purpose in an individual’s accustomed environment.
REGULATION AND REIMBURSEMENT OF LOW VISION DEVICES

Food and Drug Administration
FDA classifies devices into one of three risk-based regulatory classes based on the amount of regulation necessary to assure the safety and effectiveness of the device (e.g. Class 1 includes devices with the lowest risk imposed to the patient and/or user). The class to which the device is assigned determines, among other things, the type of premarketing submission/application required for FDA market clearance (FDA 1998).

All devices considered in this review are classified as Class 1 devices with 510 (k) premarket exemption. A 510 (k) premarket exemption means that a manufacturer does not need to demonstrate that the device to be marketed is as safe and effective, that is, substantially equivalent, to a device already legally available on the market.

VA Reimbursement
Prosthetic aids and assistive devices are available at no cost to eligible veterans. Under VA eligibility criteria, co-payment for some inpatient and outpatient services is required of veterans with non-service connected causes of low vision who exceed income thresholds.

Federal Medicare Reimbursement
In many respects the aged veteran population is similar to the Medicare population. Approximately 40% of the total elderly Medicare population has significant vision loss, and this subgroup tends to be older, poorer, less educated and in worse health than the general Medicare population (CMS 2002). Since many veterans also qualify for Medicare benefits, they could obtain low vision services outside the VA system through Medicare.

Currently, there is no national Medicare coverage policy for low vision rehabilitation including use of low vision devices, although federal legislation is pending (NVRC 2002). Beneficiaries must seek coverage through their Medicare Regional Carriers, and coverage for these services is not uniform in every state.

METHODS

For this project VATAP generated a qualitative systematic review, which approaches the process of literature review as a scientific endeavor. A systematic literature review applies explicit, reproducible methods to minimize potential biases in addressing a focused question usually about a health care intervention (Mulrow 1997). In contrast, a traditional narrative review frequently addresses a broad topic, fails to report objectives of the review, identification of articles, or methods for critical appraisal, and may be susceptible to bias in the selection, analysis, and synthesis of studies.
The cornerstone of a systematic review is emphasis on study quality. TAP applied inclusion criteria as a filter for selecting the best evidence from published research for this review. The conclusions should not overstate the evidence appraised in the review, and the recommendations for policy should be linked to the strength (or quality) of the evidence (Oxman 1995).

Scope of review
VATAP consulted low vision experts in VA to establish the scope of this review with respect to included subjects, devices and their uses. This report:

- Defines low vision along a continuum of irreversible visual impairment that interferes with daily activities. This continuum encompasses adults with residual vision ranging from approximately 20/50 to persons who are legally blind, have some residual vision and would benefit from low vision services. Patient with total blindness, children and adolescents are excluded.

- Includes all optical devices available to VA patients intended: 1) to improve visual acuity by enlarging images or by clarifying images through improved illumination, color, or contrast enhancement, or 2) to enhance the field of view. Studies of intraocular lens and contact lens are excluded, but persons with these devices who are candidates for low vision rehabilitation using other types of optical devices may be considered.

- Includes reading and driving as the tasks for which low vision devices are used.

Search strategy
VATAP designed the search strategy to capture a wide array of sources of evidence for appropriate retrieval. In October 2001, VATAP performed searches on MEDLINE®, EMBASE®, and Current Contents®, via the Dialog One® Search® feature covering literature published from 1970 through the present, with updated searches in December 2001, February 2002 and July 2002. Search strategies used terms describing low vision rehabilitation, visual disorders rehabilitation, eye diseases rehabilitation, spatial and visual perception disorders, and adult dyslexia treatment and rehabilitation. Also researched were low vision devices, tinted or filtered lenses, sensory aids, low vision enhancement systems, low vision self help devices, ocular accommodation devices and prisms. All terms were searched as descriptors from the three databases' thesauri as well as free text terms from the titles of articles to further enhance retrieval.

Other data sources
To the search TAP added citations from a composite database of international low vision literature (Goodrich 2002). VATAP excluded studies of children and adolescents as subjects from the search retrieval.

VATAP searched the Cochrane Library databases in February 2002 (2002 Issue 2), and again in October 2002 (Issue 4) using vision disorders as a major MESH term. VATAP uncovered one systematic review in progress by the Cochrane Eyes and Vision Group
entitled: “Reading aids for adults with low vision” (Acosta 2002). The protocol for the Cochrane review was available and provided a useful model for VATAP to apply to this review, and the authors assisted in uncovering and comparing evidence included for review.

On October 4, 2001 VATAP queried the health technology assessment and evidence-based communities and agencies in the International Network of Agencies for Health Technology Assessment (INAHTA) via electronic mail for relevant projects either completed or in-progress within their respective health care systems. To update the query VATAP searched the INAHTA HTA database (www.inahta.org) in July 2002 for reviews of low vision in progress.

Inclusion criteria
VATAP used a well-established framework by Jovell and Navarro-Rubio (1995) to guide inclusion of evidence in the review. In this case, the best evidence of effectiveness linking the use of an intervention to the observed outcome would be found in either meta-analyses of randomized controlled trials (RCT), individual RCTs, or non-randomized controlled, prospective trials (Table 2).

**Table 2. Levels of Evidence Scale**

<table>
<thead>
<tr>
<th>Level</th>
<th>Strength of Evidence</th>
<th>Study Design</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Good</td>
<td>Meta-analysis of randomized controlled trials (RCT)</td>
</tr>
<tr>
<td>II</td>
<td>Good</td>
<td>Large sample RCTs</td>
</tr>
<tr>
<td>III</td>
<td>Good</td>
<td>Small sample RCTs</td>
</tr>
<tr>
<td>IV</td>
<td>Good</td>
<td>Non-randomized controlled prospective trials (concurrent controls, multicenter)</td>
</tr>
<tr>
<td>V</td>
<td>Fair</td>
<td>Non-randomized controlled prospective trials (historic controls, single site)</td>
</tr>
<tr>
<td>VI</td>
<td>Fair</td>
<td>Cohort studies</td>
</tr>
<tr>
<td>VII</td>
<td>Fair</td>
<td>Case control studies</td>
</tr>
<tr>
<td>VIII</td>
<td>Poor</td>
<td>Non-controlled clinical series Descriptive studies, surveillance of disease, surveys, registers, data bases, prevalence studies</td>
</tr>
<tr>
<td>IX</td>
<td>Poor</td>
<td>Expert committees, consensus conferences, anecdotes or case reports</td>
</tr>
</tbody>
</table>

Source: Jovell and Navarro-Rubio (1995)

A single reviewer (Adams) reviewed citations of potentially relevant publications retrieved from the search and selected articles for inclusion using the following criteria:
1. Peer-reviewed research published from 1970-present that compared equivalent-powered LVDs for accomplishing a specific task:
   - High quality reviews with clearly defined and reproducible methods
   - Primary data from RCTs or non-randomized, controlled prospective studies with a sample size of patients with low vision ≥ 10 from which effectiveness could be determined
2. Full articles available, not abstracts (abstracts contain information insufficient for appraising study quality)
3. English language only
4. All commercially available electronic or non-electronic optical LVDs used for managing reduced visual acuity or visual field loss as described earlier
5. Outcome measures of performance, satisfaction, use, efficiency or Health-related quality of life using validated methods
6. The most recent or comprehensive study published by the same study group for the same purpose (to avoid double counting articles of studies on the same study population for the same purpose by the same investigators).

Critical appraisal
VATAP applied a well-known framework by Guyatt (1993) based on principles of clinical epidemiology to appraise each included article for how well bias and confounding factors were controlled in the design and conduct of the study (Table 3). VATAP considered mitigating factors specific to studies of low vision devices. For example, because of the nature of the intervention, blinding (or unmasked) treatment assignment was not considered for this review. Random order of device presentation may mitigate the effects of systematic bias introduced into a study when unmasked assessment is conducted, and this attribute would strengthen the study design.

Important to rehabilitation is the amount of training and exposure needed to become proficient with the device. Underperformance on testing or negative subjective outcome measures may reflect inadequate training and not the device itself. In contrast, improved performance may be a function of the subject’s learning curve rather than the effect of the device. In this analysis, VATAP considered the extent to which investigators minimized the effect of training as a confounding factor in each study.

Table 3. Framework for appraising the quality of studies about therapy

<table>
<thead>
<tr>
<th>Are the results of the study valid?</th>
<th>Mitigating factors/Special considerations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Was the assignment of patients to treatments randomized?</strong></td>
<td><strong>Randomized assignment to either study group</strong></td>
</tr>
<tr>
<td></td>
<td><strong>Internal control (within subject comparison) acceptable</strong></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>Were all patients who entered the trial properly accounted for and attributed at its conclusion?</strong></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
</tbody>
</table>
RESULTS

The search uncovered 2,476 citations. Upon review of titles and abstracts in the search, 28 citations were full text articles in languages other than English. Evaluation of the remaining citations and hand searching of end references of retrieved articles resulted in a total of 184 full text articles and reports retrieved for more in-depth review: 83 articles were selected, of which 11 met inclusion criteria and 72 provided background to the report (End References).

Of those studies that met inclusion criteria, four were high quality reviews. VATAP identified seven primary studies not included in the reviews. These seven studies assessed reading as the task for device use and are summarized in Table 4. Detailed attributes of included primary studies and reviews were abstracted in Tables 6 and 7, respectively.

The 101 articles excluded from this review are listed separately in the end references. The main reasons for exclusion were: 1) studies were not relevant to the review topic; 2) studies were small (n < 10) or uncontrolled; 3) no direct comparison with alternative devices or strategies was made; 4) device use was not linked to a specific task; or 5) the study did not report sufficient information in the publication to discern study attributes necessary for inclusion.

Except for the review protocol from the Cochrane Eye and Vision Group, VATAP did not uncover other reviews either completed or in progress from the evidence-based medicine or health technology assessment communities.

Primary data—reading
(Summary Table 4) Stelmack (1991), Goodrich (2001), and Spitzberg (1995) compared various low vision aids including CCTV for near reading tasks. Kuyk (1990), Lavinsky (2001) and Cheng (2001) compared various devices for visual field enhancement for reading tasks at various distances. Rossi (1990) compared two strategies (standard rehabilitation with and without prisms) for improving performance by enhancing the visual field. Two studies of prototype models were products of funded research
(Spitzberg 1995; Kuyk 1990), of which Spitzberg had a proprietary interest in a new prototype under study.

All studies included in this review were relatively small studies of less than 40 subjects with low vision. Of those reporting causes of visual impairment, age-related macular degeneration was the primary ocular condition. Four studies did not report the ocular conditions represented in their patients. Recruitment sources in the Stelmack (1991) and Goodrich (2001) studies included subjects enrolled in VA Blind Rehabilitation Centers or low vision clinics in VA and the private sector.

Rossi (1990) and Stelmack (1991) randomized all subjects to study groups. As mentioned earlier, blinding was not done due to the nature of the intervention, but Stelmack (1991) and Spitzberg (1995) randomized device order to balance the presentation of devices, thus lessening the potential effects of bias from not blinding. Three studies (Kuyk 1990; Spitzberg 1995; Cheng 2001) did not report on randomization in the study design, and Lavinsky (2001) exposed all subjects first to the conventional telescope before the contact lens telescope. All but Rossi (1990), who used a separate control group, used a “within subjects design” where subjects served as their own internal controls. This allowed for direct comparison of outcome measures using different devices. All subjects were accounted for in the design and analysis of each study, as was similarity across groups with respect to characteristics and treatment. The effect of training was accounted for as a potential confounder in each study.

Outcomes included objective or performance-based measures in all studies. Goodrich (2001) and Lavinsky (2001) presented statistical analysis for some performance data. Kuyk (1990) and Spitzberg (1995) included anecdotal data on subjective measures of preference, ease of use, and satisfaction; small study size limited the extent to which data could achieve statistical significance in the presence of an association. None of the included studies evaluated measures health-related quality of life or costs associated with the provision of the devices.

Of the seven studies, Stelmack (1991), Goodrich (2001), and Rossi (1990) represent the most rigorous with respect to study design and reporting. The strongest evidence of optical LVDs for improving reading performance in adults was in subjects age 50 years or older with severe vision loss primarily from age-related macular degeneration. These subjects had undergone extensive visual rehabilitation, including visual skills for reading and training on low vision aids, either prior to the study or as part of the study protocol. The studies were carried out in a controlled indoor setting. In this group:

- Performance with CCTV was superior to other optical aids.
- Subjects preferred stand-mounted CCTV to hand-held CCTV or other optical aids and preferred new prototype magnifiers to existing commercial models.
- Results suggested that cost, ease of use, technological design, motivation and age may influence satisfaction, routine use of these devices, and ultimately quality of life.
• Except for cost none of these studies directly assessed the reasons behind subject preferences, which could offer valuable information to those responsible for designing new products and evaluating product utility.

• Evidence of the impact of these devices on subjects’ quality of life was unknown.

Homonymous hemianopia\textsuperscript{14} and visual neglect\textsuperscript{15} are common problems in patients with neurological injury. A variety of optical aids and adaptive strategies are used to improve visual perception deficits in these patients. The strongest evidence of the effectiveness of low vision devices to improve visual perception is based on a comparison of Fresnel prisms versus standard rehabilitation in an inpatient post-stroke population with homonymous hemianopia or unilateral visual neglect (Rossi 1990). This study showed:

• Treatment with 15-diopter Fresnel prisms improved the patients’ visual perception test scores but not activities-of-daily-living function.

• Additional controlled studies are needed to confirm results and to define the optimal prism strength, manner of application, and duration of benefit.

\textsuperscript{14} loss of sight for one half of the visual field of one or both eyes that affects the same portion of the visual field of each eye.

\textsuperscript{15} passive, unconscious decreased awareness of part of the field of view or other stimuli to one side of the body; also called visual hemi-attention, visual imperception, or visual/spatial neglect
Table 4. Prospective controlled primary studies of low vision device effectiveness

Note: Shaded areas indicate deficiencies in the conduct or reporting of the study. Such deficiencies may introduce bias into the study or limit the reader’s ability to generalize the results to other populations. Because of the nature of the intervention, none of the studies blinded patients, personnel or health care professionals to treatment groups; blinding (or unmasked) treatment assignment was not considered for this review. None of the included studies evaluated HRQOL or costs as outcome measures.

<table>
<thead>
<tr>
<th>Study</th>
<th>Devices</th>
<th>N</th>
<th>Ocular conditions</th>
<th>Randomized</th>
<th>Controlled</th>
<th>All subjects accounted for</th>
<th>Blinding</th>
<th>Group similarity</th>
<th>Groups treated equally</th>
<th>Training effect minimized</th>
<th>Objective/Performance</th>
<th>Preferences</th>
<th>Ease of use</th>
<th>Satisfaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kuyk 1990</td>
<td>• Prototype telescope</td>
<td>15</td>
<td>Not reported</td>
<td>I</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Target spotting &amp; identification</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Rossi 1990</td>
<td>• 15-diopter Fresnel prisms</td>
<td>39</td>
<td>HHA VN</td>
<td>C</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Visual perception</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stelmack 1991</td>
<td>• CCTV</td>
<td>37</td>
<td>Post-disciform stage ARMD</td>
<td>S, DP</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Reading speed, Reading duration</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Spitzberg 1995</td>
<td>• 4 prototype magnifiers</td>
<td>39</td>
<td>Not reported</td>
<td>DP</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Reading speed</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Goodrich 2001</td>
<td>• Stand-mounted CCTV</td>
<td>22</td>
<td>ARMD CRVO, MH</td>
<td>Not reported</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Reading speed, Reading duration</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lavisky 2001</td>
<td>• Conventional telescope</td>
<td>15</td>
<td>Not reported</td>
<td>I</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Visual field</td>
<td>✓</td>
<td>✓</td>
<td></td>
</tr>
<tr>
<td>Cheng 2001</td>
<td>• CR39 prisms, Fresnel prisms</td>
<td>15</td>
<td>Not reported</td>
<td>I</td>
<td>Yes</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Acuity</td>
<td>✓</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

ADL, activities of daily living  
ARMD, age-related macular degeneration  
C, separate control group  
CRVO, central retinal vein occlusion  
DP, device presentation randomized  
DR, diabetic retinopathy  
HHA, homonymous hemianopia  
I, internal control (within subjects design)  
MH, macular hole  
S, subjects randomized to study groups  
VN, visual neglect
Primary data—driving
The VATAP did not identify any evidence of the effectiveness of optical LVDs for driving that met criteria for inclusion in the review.

Prior reviews of effectiveness of low vision devices or low vision rehabilitation
Literature reviews with explicit methods can help integrate large amounts of literature and identify knowledge gaps for rational decision-making. Existing reviews of the effectiveness of low vision rehabilitation may include outcomes related to provision of LVDs and, therefore, may inform the appraisal of evidence for this review.

VATAP identified four high quality reviews with explicit methods to include in the report (Eperjesi 2002; Owsley 1999; Raasch 1997; Stelmack 2001). To synthesize findings all used qualitative methods. A summary of the findings is presented in Table 5, and details of these reviews are abstracted in Table 7.

Table 5. High quality reviews of low vision device effectiveness

<table>
<thead>
<tr>
<th>Review</th>
<th>Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Effectiveness of low vision devices</strong></td>
<td></td>
</tr>
<tr>
<td>Eperjesi 2002</td>
<td>• Improvement in visual function or of superiority of tinted lenses with spectral characteristics over neutral density filters or conventional sunglasses was inconclusive; some subjective improvement reported but no consistent objective benefit</td>
</tr>
<tr>
<td>Owsley 1999</td>
<td>• The effectiveness of bioptic telescopic spectacle use on driving performance or safety is inconclusive&lt;br&gt;</td>
</tr>
<tr>
<td><strong>Effectiveness of low vision rehabilitation</strong></td>
<td></td>
</tr>
<tr>
<td>Stelmack 2001</td>
<td>• Self-reported QOL is a significant outcome measure for low-vision rehabilitation&lt;br&gt;</td>
</tr>
<tr>
<td>Raasch 1997</td>
<td>• Self reported results suggest low vision intervention has a significant impact on activities of daily living and can be highly valued by patients&lt;br&gt;</td>
</tr>
</tbody>
</table>

**Effectiveness of low vision devices.** Eperjesi (2002) reviewed the evidence of effectiveness of tinted lenses and filters for persons with low vision. Despite using broader inclusion criteria than in this VATAP review (non-peer reviewed sources, all study sizes, and all ages of subjects), the authors found no conclusive evidence regarding the effectiveness of commercially available tinted lenses and filters for various tasks. While subjective improvement was reported among some subjects, there was no conclusive evidence of a consistent objective benefit, improvement in visual function, or
of superiority of tinted lenses with spectral characteristics over neutral density filters or conventional sunglasses.

The available research was inconclusive regarding the effectiveness of bioptics for driving, notwithstanding a sizable body of literature on the topic (Owsley 1999). The main limitations of the available evidence were: using the general population as the control group; the unknown contribution of the bioptics’ restricted field of view and/or severely impaired visual function to elevated crash rates; and the unknown effect of self-regulated driving habits among bioptics users.

Effectiveness of low vision rehabilitation. Raasch (1997) reviewed studies of performance and satisfaction with low vision services, and Stelmack (2001) evaluated effectiveness of low vision services using quality of life measures. The conclusions from these two reviews represent the general state of the evidence of effectiveness of low vision rehabilitation: current evidence shows a lack of systematic, comparative evidence of effectiveness of low vision interventions.

Most research measured patient satisfaction or successful use of a LVD to measure change as a result of low vision intervention, but throughout the literature a consistent definition of success had been lacking (Raasch 1997). Many factors such as ocular diagnosis, vision needs, training, self-modification, home or family support, preferences and types of services may have influenced satisfaction and use, but their influence on other outcomes of low vision intervention such as quality of life was unknown.

Raasch (1997) noted that improved performance with a LVD might be expected to translate into improved quality of life, but a direct link between the two was not always apparent or had not been studied, and performance measured in a clinical setting may not generalize to function in the home or workplace. As Stelmack (2001) succinctly points out:

"It is important to recognize improvement in quality of life without improvement in performance with low-vision devices and improvement in performance of device use without improvement in quality of life."

Stelmack (2001) found an improvement in functional status and quality of life after low vision interventions. Validated general health and disease-specific HRQOL instruments, such as SF-36, NEI-VFQ 51-Item Field Test, VF-14, NEI-VFQ-25, and BRS FOS instruments, had been used or modified to measure vision-specific HRQOL, but they were limited in their ability to assess outcome measures adequately for low vision rehabilitation or across specific populations. A version of the NEI-VFQ-25, was adapted for use in more frail, elderly populations. Prosthetic low vision devices used in low vision rehabilitation programs were found to be feasible for comparing outcomes of low vision programs in a veteran population.

VATAP identified an ongoing RCT comparing the effectiveness and cost-effectiveness of three models of low vision service delivery for subjects with newly diagnosed age-
related macular degeneration (Russell 2001). The results of this trial will be used to inform a national strategy for low vision services in the United Kingdom. It incorporates several generic and vision specific quality of life instruments. While this trial did not meet criteria for inclusion (it compared models of low vision rehabilitation rather than specific low vision devices), the preliminary findings support the need for a range of outcome measures to characterize quality of life in subjects with age-related macular degeneration: physical functioning; knowledge of eye condition; attitudes to and feelings towards visual impairment; impact of low vision on daily life; task analysis and patterns of LVD use; and satisfaction with low vision services.

**Limitations of this review**

This review employed discrete inclusion criteria with respect to subjects, devices and their uses, and study attributes. The reader should be aware that changes in these criteria might alter the findings.

Children, adolescents and patients with total blindness were excluded from review, as were non-optical low vision aids. Tasks other than reading and driving were not considered, although review of the search retrieval showed a preponderance of evidence for reading.

VATAP only considered controlled prospective primary studies for inclusion in this review. The sizable amount of available non-experimental (observational) data, which is considered weaker evidence of effectiveness, was not included. It is possible that information from these studies may have contributed to the findings in this report. However, the findings from the included high quality reviews that did summarize non-experimental data in their topic areas would not change this report's conclusions.

This report confined retrieval to full text articles in English. In all, VATAP excluded 28 citations based on these criteria. Review of available title and abstracts shows that one additional study in German by Rohrschneider (1997) may have met inclusion for review (see excluded studies list in end references). This study compared visual acuity with glasses, telescope and the Low Vision Enhancement System (LVES)\(^\text{16}\). They reported improved visual acuity and contrast sensitivity and reduced glare with LVES over correction with glasses or telescopes, but regular use of LVES in the majority of patients was unlikely. These findings would not have changed the conclusions in this review.

Finally, only the most recent or comprehensive study published by the same study group for the same purpose was included. This was done to provide a best estimate of the true study base available for inclusion. Redundancy can be found in the peer reviewed literature when investigators publish research findings that overlap with previous publications or when they publish the same findings in multiple sources. In the end, VATAP did not exclude any studies based on this criterion.

\(^{16}\) a battery-powered portable vision enhancement device worn like goggles and tailored to each patient. Developed by the Johns Hopkins Wilmer Eye Institute in collaboration with the National Aeronautics and Space Administration and the VA.
SUMMARY

Adults with low vision can experience a range of difficulties in their daily lives as a result of their vision impairment. Among older adults—the primary emphasis of this review—vision loss can have a profound effect on their lives. Low vision is associated with a variety of co-existing medical conditions, limitations in activity and performance, and lower quality of life. Demographic trends forecast a dramatic increase in the prevalence of age-related causes of vision loss in the veteran population. With this trend is an increasing need for low vision services targeted at detecting visual impairment and mitigating the functional consequences associated with age-related vision loss to improve quality of life.

Provision of low vision services to veterans is a priority area for VA. To meet the needs of an increasingly geriatric veteran population, VA is complementing its inpatient service delivery with outpatient services that include furnishing a range of low vision devices (LVD) to eligible veterans. The highest demand (and the most costly LVDs for veterans) is for electronic optical devices, such as CCTV and computer assistive technologies, and nonelectronic hand-held models. Veterans have shown substantial interest in CCTV in particular for its reported ability to improve a range of functional deficiencies caused by loss in visual acuity and visual field.

Advocates for visually impaired veterans have expressed concern over the quality of the scientific evidence supporting the use of many LVDs, particularly newly emerging electronic devices. This systematic review on existing evidence of effectiveness of low vision devices from the peer-reviewed published literature will inform a multidisciplinary task force of VA vision care experts charged with developing a process for evidence-based new technology evaluation and dissemination of information in VA.

This review considered controlled prospective studies of adult subjects with limited residual vision as a result of moderate to significant irreversible vision loss and all optical devices available to eligible veterans. This review considered outcome measures encompassing objective measures of performance and use, subjective measures of satisfaction and preference, efficiency, and health-related quality of life measures. Reviews of the effectiveness of low vision rehabilitation addressing the effectiveness of LVDs were also included.

The best available evidence comparing the relative effectiveness of optical LVDs is seven small, prospectively controlled clinical studies comparing the performance of low vision devices for reading tasks at various distances in an indoor setting. Evidence suggests that among extensively trained patients with age-related macular degeneration, reading performance with either stand-mounted or handheld CCTV was superior to prescribed optical devices (stand magnifiers, coil stand magnifiers, and microscopic lenses). Compared to standard rehabilitation alone, Fresnel prisms added to standard rehabilitation improved performance on visual perception tests but not on
activities-of-daily-living function in post-stroke inpatients with homonymous hemianopia or visual neglect.

Anecdotally, CCTV was preferred to spectacle reading glasses and illuminated stand magnifiers, and prototype magnifiers were preferred to conventional devices, but the reasons behind the preferences were not systematically examined. Anecdotal evidence identified cost of the device, design, age and motivation as factors that might have affected sustained use of a device. Sustained use of these devices in the subject’s life setting, resources in terms of costs and training associated with each alternative, and the link between device use and health related quality of life were unknown.

KNOWLEDGE GAPS AND RECOMMENDATIONS FOR FUTURE RESEARCH

This review identified knowledge gaps that may help direct future research. These gaps concentrate on candidacy for low vision devices, suitable prescription of these devices, and measuring their effectiveness.

Diagnosis, visual acuity, and pattern of visual field loss are initial considerations in the prescription of low vision aids, as they provide a fair degree of predictable, objective information about the effect of the visual impairment (Faye 1976). However, these measures alone may be inadequate for the adult with age-related causes of vision loss (particularly those age 75 or older), who frequently complain of visual difficulties under everyday, sub-optimal conditions despite satisfactory results on standard testing (Brabyn 2001).

As yet, no single testing battery has been identified that is sufficiently sensitive or practical to administer to these patients. A testing battery that is easy to administer to a growing elderly population and is sensitive to real-world conditions might assist clinicians in (Legge 1992; Rubin 1997; Brabyn 2001):

- identifying the circumstances under which vision aids should be prescribed;
- optimizing prescription regimens;
- monitoring disease progression;
- predicting individuals at risk for future serious vision loss;
- referring patients to specialized low-vision clinics;
- freeing up resources associated with more labor-intensive special testing.

Current practice relies on direct measurement of performance in the clinical setting as a proxy measure for continued device use in the patient’s life setting, but ideally performance should be measured for tasks encountered under everyday conditions. Low vision research has identified an array of visual and nonvisual factors that may influence both the performance conducted in a clinical setting and continued use of the LVD, but the value of these factors as predictors of performance or successful use in everyday indoor and outdoor conditions requires further study. Studies are needed with sufficient power to detect the presence of an association and to analyze intra-subject as well as intra- and inter-group variation. Research is also needed to differentiate the
physiological from psychological basis of visual improvement and to identify the visual and non-visual skills required for specific tasks. Identifying predictive factors from robust research may help clinicians develop a practical, sensitive testing battery for improving diagnosis and candidacy.

To inform choices about appropriate device provision, research is needed on the relative effectiveness of low vision devices using rigorous methods under conditions and for tasks similar to those found in the real world, and along the continuum of visual impairment and disability. Sufficient power is required to detect the presence of associations and to permit statistical analysis of intra-subject as well as intra- and inter-group variation. The methods should be transparent to allow the reader to determine applicability of the results to his or her patients, and the methods should employ patient-focused functional outcome measures and subjective measures appropriate to the desired goal of rehabilitation.

Performance should take into account adaptation, setting, compensatory strategies, training and exposure. Subjective self-reported measures of usage and satisfaction should be considered. Preferences should be measured in a way that is valid, useful for benchmarking, and relevant to inform decision making about uses in patient care as well as purchasing. With respect to measures of efficiency, the full range of resources (e.g. costs associated with devices, access to low vision services and training) associated with device use should be considered, both from the perspectives of the patient and the health care system.

Ultimately, the goal of low vision rehabilitation, including use of assistive technologies, is to improve the patient’s quality of life. To some extent the impact of low vision devices on a patient’s quality of life can be dramatic and may seem obvious. However, outcome measures are needed that capture the type and magnitude of the change in function and HRQOL in a way that allows comparison of outcomes across low vision interventions and with outcomes of treatment of other disorders. Research is needed to identify appropriate functional outcome measures for low vision rehabilitation and to develop and validate instruments of HRQOL for use in the low vision population. Existing evidence suggests that a range of outcome measures is needed to characterize quality of life in subjects with age-related visual impairment: physical functioning; ocular condition; attitudes towards and perceptions of visual impairment; impact of low vision on daily life; task analysis and patterns of LVD use; and satisfaction with low vision services.

In the realm of assistive technology (AT) outcomes measurement, a number of proprietary and publicly funded initiatives exist or are under development. Among the federally funded, VA researchers are actively investigating many areas of blind rehabilitation including developing functional outcomes measures and measuring the impact of blind rehabilitation on quality of life (VA BRS 2002). Findings from the VA
Blind Rehabilitation Outcomes Project\footnote{VA Rehabilitation R&D Center, Atlanta, GA 30033; Pittsburgh Vision Services, Pittsburgh, PA. Funded in part by the VA Office of Rehabilitation Research and Development. \url{http://www.varrd.emory.edu/brc/link.html}} may assist providers in improving greater efficiency and effectiveness of low vision interventions in VA (De l'Aune 1999). For example, VA researchers developed and validated a 13-item instrument to measure the impact of a comprehensive rehabilitation program. With further modification it may be suitable as a measure of change in overall functional independence associated with use of low vision devices.

The Consortium for Assistive Technology Outcomes Research (CATOR)\footnote{funded by the National Institute on Disability and Rehabilitation Research (NIDRR), United States Department of Education} was established in 2002 as a five-year project to conduct research on AT outcomes and impacts to determine the effectiveness and usefulness of AT and the implications for use/discontinuance of AT devices \url{[http://www.atoutcomes.com/]} . Results from this project may inform outcome measurement needed for low vision technologies.

**CONCLUSIONS AND RECOMMENDATIONS**

This review reveals the paucity of high quality evidence available in the peer-reviewed published literature to inform choices about the provision of optical low vision devices in VA. The absence of compelling evidence and a standard taxonomy of what constitutes desired outcomes make it difficult to clearly recommend one device over another. Therefore, clinicians must continue to rely on industry literature, patient self-reporting, clinical observations, and real-world trials in determining appropriate provision of low vision aids. Unfortunately, industry is motivated by profits and has no regulatory impetus (aside from mandates from federal funding sources) to produce such information. Only in recent years have low vision specialists been held to a higher standard of accountability to the health systems in which they provide care. Neither alone has had sufficient resources to conduct all of the research needed to inform these choices.

Nonetheless, health care providers and assistive technology communities have the responsibility to determine which practices and technologies are most appropriate for an expanding market of individuals with age-related visual impairment. By virtue of its large visually impaired population and concentration of low vision expertise, VA stands poised as the nation’s largest health care system to make significant strides in evaluation and provision of high quality low vision services to veterans and the nation.

- **VA practitioners should use an evidence-based framework in evaluating evidence of effectiveness.**

  Evaluating outcomes associated with use of low vision service delivery requires systematic evidence-based data collection to provide reliable data. To that end,
proficiency in evidence-based critical evaluation is needed to appraise available data and to develop mechanisms for rigorous prospective data collection either through clinical trials or databases. Some efforts are underway, and the methods used in this review provide an evidence-based framework that could guide future activities.

VA should encourage partnering with industry and consumers to carry out evidence-based technology evaluation.

DeRuyter (1995) describes service delivery as a “business arrangement” between consumer and provider resulting in the delivery or provision of some specific agreed upon commodity or service in exchange for compensation. Patients have unmet need; industry has products to sell; patients and practitioners need high quality evidence to make informed decisions; and there are significant knowledge gaps to fill.

Historically, through the VA Rehabilitation Research and Development Service VA low vision specialists have been able to cross-fertilize efforts with private industry, academic affiliates, and other government agencies. In addition to competitive research funding, pooling funds from private sources could be considered to fund initial technology evaluation activities in VA within an evidence-based framework. This approach has several advantages. It would:
1) Provide reliable data to VA patients and practitioners on which to make informed choices;
2) Assist VA practitioners in meeting more stringent accountability standards;
3) Involve patients in designing products that offer maximal benefit;
4) Be resource neutral for VA in times of fiscal constraint;
5) Guide stakeholders in patient-focused product development and evaluation.

Within the assistive technology community, partnerships between various stakeholders exist to conduct technology evaluation, of which consumer involvement is an important component. Experiences from these partnerships may inform development of VA technology evaluation activities with respect to incorporating consumer input. For example, the Rehabilitation Engineering Research Center on Technology Transfer (University of Buffalo, NY, USA), is funded for five years by a grant from the National Institute on Disability and Rehabilitation Research (NIDRR), United States Department of Education, to improve the quality of assistive devices available in the marketplace. One of its activities is the consumer ideal product program, which developed and implemented tools needed for a national sample of experienced device users to define the ideal product. It used consumer focus groups and quantitative surveys to identify consumer needs and preferences of several categories of assistive technology from ABLEDATA\textsuperscript{19}. The results led to descriptions of consumer comparisons of existing products and checklists for consumers to use when shopping for assistive devices.

\textsuperscript{19} A NIDRR-sponsored searchable database of more than 29,000 assistive technologies (over 19,000 of which are currently available) containing detailed marketing information about each product; organized into product categories according to functional activity. [\url{http://www.abledata.com/}]
In the meantime, VA practitioners need information that improves and standardizes current prescription practices.

VA provides a range of electronic and non-electronic optical devices to its veterans. Electronic devices tend to receive most of the attention because of their popularity among consumers and high unit costs. Non-electronic devices may be overlooked, even though they are the most commonly prescribed. To note, in fiscal year 2002, hand-held low vision aids were the most commonly prescribed low vision aid and had the third highest expenditure for aids to blinded veterans.

Frequently manufacturers of non-electronic "low tech" devices report optical parameters of their devices based on arbitrarily chosen standards. Standardized information is needed to allow clinicians to predict accurate improvement in patients' visual performance with the device. Bailey (1994) measured and tabulated key optical parameters of 92 stand magnifiers and 53 hand-held magnifiers using a standardized formula. Information such as this may assist practitioners by translating existing disparate information from manufacturers into standardized, usable information on which to base more rational prescription decisions, particularly of some high volume, "low tech" optical devices.
Table 6. Primary studies that met inclusion criteria for review

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<tbody>
<tr>
<td>Study objective(s)</td>
<td>• Demonstrate feasibility of constructing a simple motorized focus telescope using off-the-shelf components • Compare performance in spotting and identification of near and distance targets using a prototype motorized focus telescope (MFT) and a hand focus telescope (HFT)</td>
<td>To assess the efficacy of 15-diopter Fresnel prisms for treatment of homonymous hemianopia (HHA) or unilateral visual neglect (VN) after stroke</td>
<td>• Compare reading speed and duration using CCTV, spectacle reading glasses, and illuminated stand magnifiers</td>
<td>Compare reading speed and preferences for 4 prototype magnifiers to a commercially available stand magnifier</td>
</tr>
<tr>
<td>Study size</td>
<td>N = 15</td>
<td>N = 39</td>
<td>N = 37</td>
<td>N = 39</td>
</tr>
<tr>
<td>Recruitment source</td>
<td>VA facility but details not reported</td>
<td>Inpatient stroke rehabilitation unit</td>
<td>Patients admitted to Central Blind Rehab Center, Hines VAMC for low vision rehabilitation</td>
<td>Patients from 5 low vision clinics, including one in VA • Selection process not defined</td>
</tr>
<tr>
<td>Inclusion criteria</td>
<td>• Visually impaired subjects • Experienced users of manual telescopes • Informed consent</td>
<td>• Diagnosed with stroke with either HHA or VN • Free of disabling cardiac, pulmonary, or rheumatologic problems that may preclude rehab efforts • Informed consent</td>
<td>• Age 50 or older • Diagnosed with post-disciform stage age-related macular degeneration or ocular histoplasmosis</td>
<td>• Volunteers • Need for 3X magnification to read 1.0M or 1.5M print</td>
</tr>
<tr>
<td>Exclusion criteria</td>
<td>Not reported</td>
<td>• Best corrected visual acuity worse than 20/200 • Unable to comprehend and cooperate with visual field assessment</td>
<td>• Pathophysiologic changes w/in and in neurosensory or pigment epithelial retinal layers immediately surrounding the macula • Nutritional optic atrophy including glaucomatous, ischemic or toxic that would compromise central visual function • Significant cataract formation including posterior subcapsular cataracts of &gt; 1+, cortical vacuoles &gt; 2+, nuclear sclerosis &gt; 2+ • Media changes decreasing view of fundus details to &lt; 20/40 equivalent</td>
<td>None reported</td>
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</table>
### Study attributes

|------------------|-----------|------------|---------------|----------------|
| **Characteristics of subjects** | • Age ranged from 75-81 years  
• Visual acuity between 10/60 and 10/350 | • Comparable demographically across groups with respect to gender, interval poststroke, lesion type, side of stroke, deficit type, modified mini mental status exam  
• Prism group older than control group  
Age (±SEM): control 63.3 yrs ± 2.5 vs. prism 72.6yrs ± 1.8 (p < 0.01) | • Gender not reported  
• Age range > 50 years, range not defined  
• Diagnoses: All with ARMD or ocular histoplasmosis  
• Eccentric viewing acuity ranged 10/60-10/600  
• Magnification with ISM 3.5X to 10X  
• Power of spectacles +10.00D. to +32.00D.  
• Magnification with CCTV 4X to 22X | • Age range = 9 to 77 years  
• Other data not reported |
| **Study devices** | • Prototype MFT of 4, 6, and 8 X magnification  
• Keplarian type HFT monocular of 4, 6, and 8 X magnification | • 15-diopter Fresnel prisms  
• Routine stroke rehabilitation | • VTEK Voyager CCTV  
• Eschenbach illuminated stand magnifier  
• Prism half eyes or Aolite microscope spectacle reading lenses | • Spherical mirror magnifier  
• Cylindrical mirror magnifier  
• Reflecting prism magnifier  
• Zoom magnifier  
• COIL #5428 and #5123 |
| **Random assignment of either subject to group or order of device presentation** | Not reported | Random assignment to study groups | Random assignment to one of 3 treatment groups; each group varied order of devices used | Order of presentation of devices rotated systematically |
| **Controlled** | Within subjects design used | Separate control group | Within subjects design used | Within subjects design used |
| **Protocol** | • Each subject trained in use of MFT for 15-30 minutes  
• First near and then far distance target spotting and identification tasks performed with each telescope of the same magnification power at seven distances between 8 and 80 feet in a well lit hallway using series of 4 black letters of varied size on a white background  
• Time to completion and number of identification errors recorded; data converted to time required to identify target  
• Questionnaire administered to elicit preferences, rate performance and design features on a 4-point scale, and rate confidence with device | • Prism group fitted and trained with prisms; visual adaptation and safety awareness demonstrated  
• All patients had general physical and neurologic exam and tested without prisms at baseline  
• All patients studies at baseline, 2 weeks, and 4 weeks with Modified Mini Mental Status Examination, Motor Free Visual Perception Test, Line Bisection Task, Line Cancellation Task, Harrington Flocks Visual Screener, Tangent Screen Examination, and Barthel ADL-Mobility Score  
• All patients participated in ADL training and table-top visual perception retraining tasks | • Each subject received extensive rehab training for visual skills for reading and one-hour training on each device  
• Within subject controlled = each subject tested with each device  
• Optimal magnification for spectacles and illumination magnifier = lowest power plus lenses needed to read “Life in These United States” excerpts from Readers’ Digest  
• Magnification for CCTV = lowest print size needed to read same excerpts with either black letters on white background or reversed | • Each subject trained daily with four low vision aids including at least two prototype devices for a minimum of 5 days  
• VA subjects trained for an additional 5 days  
• Order of presentation of devises systematically rotated among subjects  
• All subjects practiced reading and writing for ≥15 minutes each day with each device  
• Subject data collected included age, pathology, reading history prior to and after vision loss, visual acuity, and CSF  
• Daily collection of outcome measures for each device |
| **Loss to follow-up/drop out** | None | None | None | None |
### Study attributes

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<tbody>
<tr>
<td>Intention to treat analysis used</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
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</table>

### Outcomes

- **Preferences**
  - Performance and design features: overall focus speed; stability of the viewed scene during focusing; control of eye-target alignment of device; overall focus control; overall performance; weight; size
  - Confidence with device

- **Testing measures**
  - Number of falls
  - Demographic data of age and interval post stroke

- **Reading speed and duration**

### Adverse effects

- None reported

### Results

- **Preferences**
  - 73% preferred Motorized Focus Telescope vs. Hand Focus Telescope
  - Subjects rated confidence in device equally
  - Subjects rated weight and size of prototype motorized focus telescope below Hand Focused Telescope; improvements in design were warranted
  - Effect of motorized model on training time, learning curve and ability to correct alignment were under study

- **Baseline evaluations of visual perception and activities of daily living function were similar for both study groups**
  - At 4 weeks, prism group performed significantly better than controls on Motor Free Visual Perception Test ($p < 0.01$), Line Bisection Task ($p < 0.01$), Line Cancellation Task ($p < 0.02$), Harrington Flocks Visual Screener ($p < 0.01$), Tangent Screen Examination ($p < 0.01$)
  - No significant differences between study groups in Barthel ADL scores or number of falls
  - Steady improvement in perceptual task performance over study duration suggests a more complex response to prisms than simple adaptation
  - Recommended additional controlled studies to confirm results and to define the optimal prism strength, manner of application, and duration of benefit

- **Average reading speeds of subjects using prototype devices not significantly different from COIL stand magnifier (test of significance not presented)**
  - 29 (74%) of subjects picked one of the new prototypes as aid of choice
  - Spherical mirror magnifier most frequently preferred (13/39)
  - Authors suggest that preferences are quite likely to change with increased knowledge about options

### Comments

- Supported by a DVA RR&D grant C962-PA
- No proprietary interests reported
- No proprietary interests reported
- Research sponsored by NEI given to Optical Designs, Inc (Houston, TX)
- Author Spitzberg has proprietary interest in these products

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20 A strategy for analyzing data in which all participants are included in the group to which they were assigned, whether or not they completed the intervention given to the group. Intention-to-treat analysis prevents bias caused by the loss of participants, which may disrupt the baseline equivalence established by random assignment and which may reflect non-adherence to the protocol.
Table 6 (continued)

<table>
<thead>
<tr>
<th>Study attributes</th>
<th>Goodrich 2001</th>
<th>Lavinsky 2001</th>
<th>Cheng 2001</th>
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<tbody>
<tr>
<td>Study objective(s)</td>
<td>Compare differences in reading speed and duration using stand- or hand-mounted CCTV vs. prescribed optical devices</td>
<td>Evaluate the performance of a contact lens telescope (CLT) and conventional telescope</td>
<td>To compare high and low contrast acuity reduction with conventional CR39 versus Fresnel prisms</td>
</tr>
<tr>
<td>Study size</td>
<td>N = 22</td>
<td>N = 15</td>
<td>N = 15</td>
</tr>
<tr>
<td>Recruitment source</td>
<td>• Patients enrolled in residential rehab program at the Western Blind Rehab Center, Palo Alto VAMC</td>
<td>• Patients in the Hospital de Clinicas of Porto Alegre, Rio Grande do Sul, Brazil from October 1999 to April 2000</td>
<td>Not reported</td>
</tr>
<tr>
<td>Inclusion criteria</td>
<td>• All with legal blindness as defined by the ICD-9CM</td>
<td>• Ophthalmological diagnosis of low vision</td>
<td>Not reported</td>
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<tr>
<td></td>
<td>• Volunteers</td>
<td>• Visual acuity between 20/60 and 20/400</td>
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<td></td>
<td>• Presence of central scotoma with intact peripheral field as determined by patient’s reduced visual acuity and visual-field tests</td>
<td>• Informed consent</td>
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<td></td>
<td>• Desire for reading rehabilitation</td>
<td></td>
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</tr>
<tr>
<td>Exclusion criteria</td>
<td>• Cognitive defects</td>
<td>Visual acuity &lt; 20/60 or &gt; than 20/400</td>
<td>Not reported</td>
</tr>
<tr>
<td></td>
<td>• Current use of medications that would impair reading ability and/or illiteracy</td>
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<tr>
<td>Characteristics of subjects</td>
<td>• 20 men; 2 women</td>
<td>• 11 males; 4 females</td>
<td>• Healthy subjects</td>
</tr>
<tr>
<td></td>
<td>• Average age = 73.3 years, (SD 8.9 yrs), range 53-87</td>
<td>• age ranged from 13 years to 84 yrs (mean age = 60.7 years)</td>
<td>• Age range 20-30 years</td>
</tr>
<tr>
<td></td>
<td>• Diagnoses: ARMD (16); DR with only central field involvement (2); central retinal vein occlusion (2); macular hole (1); dystrophy (1)</td>
<td>• mean uncorrected visual acuity = 20/245; 8 patients with 20/200</td>
<td>• Spherical refractive error &lt; 5.00 diopters</td>
</tr>
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<td></td>
<td>• Average LogMAR visual acuity = 0.994, approx 20/200, (SD, 0.46), range 0.48-1.6</td>
<td>• mean visual field without correction = 101 degrees (range 80-114 degrees)</td>
<td>• Cylindrical refractive errors &lt; 1.00 diopters</td>
</tr>
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<td>• Average Pelli-Robson contrast sensitivity = 0.89 (SD, 0.46), range 0.00 to 1.5</td>
<td>• mean corrected visual acuity = 220/105</td>
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<tr>
<td></td>
<td>• No previous comprehensive rehab training</td>
<td>• mean visual field with conventional telescope correction = 22 degrees</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Most used magnifier, but none used CCTV</td>
<td>• mean visual field with CLT correction = 52.1 degrees</td>
<td></td>
</tr>
<tr>
<td>Study devices</td>
<td>• Stand-mounted CCTV (Optelec Clearview or Telesensory System Gene)</td>
<td>• Sportieri® conventional telescope designed to produce 2.0X magnification with visual field of 25 degrees</td>
<td>• Conventional CR39 prisms</td>
</tr>
<tr>
<td></td>
<td>• Handheld CCTV (Innovations Magni-Cam Triad Color system) with 27 inch Daewoo TV</td>
<td>• Contact lens telescope system (CLT) = contact lenses (mediphacos, Belo Horizonte) plus spectacle lens designed to produce 2.0X magnification with a visual field of 50 degrees</td>
<td>• Fresnel prisms</td>
</tr>
<tr>
<td></td>
<td>• Prescribed optical device: Eschenbach stand magnifier (10); Coil stand magnifier (9); microscopic lenses (3)</td>
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<tr>
<td>Study attributes</td>
<td>Goodrich 2001</td>
<td>Lavinsky 2001</td>
<td>Cheng 2001</td>
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<tr>
<td>Random assignment of either subject to group or order of device presentation</td>
<td>Not reported</td>
<td>Each patient wore first the conventional telescope then the CLT</td>
<td>None</td>
</tr>
<tr>
<td>Controlled</td>
<td>Within subjects design used</td>
<td>Within subjects design used</td>
<td>Within subjects design used</td>
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</table>
| Protocol                                     | • Four instructors provided formal reading rehab training in all three devices  
• All subjects received eccentric viewing training before reading training  
• One training session/day for 5 sessions on each device  
• Reading speed and comprehension measured using randomly selected paragraphs printed by laser printer in Times Roman font rated at Flesch-Kincaid Grade 5 level difficulty  
• Reading duration defined as cumulative time subject spent reading during each training session | • Each subject wore the conventional telescope, obtained maximal focus, given a 10 minute period of adaptation  
• Visual acuity tested using Lighthouse Near Acuity Test at 2 m; best acuity defined as the line read with a maximum of two errors  
• Visual field testing with a Goldman perimeter at distance of 50 cm using a number III object of 4 mm² and intensity number 4  
• Followed by same protocol using CLT | • Subjects’ refractive errors corrected with trial lens to a visual acuity of 6/6 or better  
• Visual acuity of right eye measured with high and low contrast charts while wearing each prism at a vertex distance of 12 mm  
• Test distance of 4 m and chart luminance of 140cd/m² used |
| Loss to follow-up/drop out                   | None                           | None                                | None       |
| Intention to treat analysis used             | Yes                            | Yes                                 | Yes        |
| Outcomes                                    | • Reading speed, duration and subjective preference  
• Difficulty in device use using a 10 point scale  
• Overall satisfaction with devices using a 10 point scale  
• P < 0.05 considered statistically significant | • Difficulty in device use using a 10 point scale  
• Overall satisfaction with devices using a 10 point scale  
• P < 0.05 considered statistically significant | Visual acuity |
<p>| Adverse effects                              | None reported                  | None reported                        | None reported |</p>
<table>
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<tr>
<th>Study attributes</th>
<th>Goodrich 2001</th>
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<tr>
<td><strong>Results</strong></td>
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<tr>
<td>Reading speed (effect size, d):</td>
<td>Stand-mounted CCTV &gt; optical device; d = 0.45</td>
<td>11/15 found conventional telescope easy to use vs. 6/15 for CLT</td>
<td>Significant reduction of high and low contrast acuity occurred when power of CR39 prism and Fresnel prism reached 10 prism dipters and 5 diopters, respectively (p &lt; 0.05)</td>
</tr>
<tr>
<td></td>
<td>Hand-held CCTV &gt; optical device; d = 0.57</td>
<td>11/15 were satisfied with CLT, but only 3/15 were satisfied with conventional telescope</td>
<td>Rate of acuity reduction with increasing prism power was greater with low contrast targets than with high contrast targets for both prisms (p not reported)</td>
</tr>
<tr>
<td></td>
<td>No difference between types of CCTV; d not calculated</td>
<td>Visual field increased 28.7% using CLT (P &lt; 0.001)</td>
<td>Rate of acuity reduction with increasing prism power was greater with Fresnel prism than with CR39 prism for both contrasts (p &lt; 0.05)</td>
</tr>
<tr>
<td>Reading duration:</td>
<td>Stand-mounted CCTV &gt; optical device; d = 1.31</td>
<td>33% spontaneously decided to use CLT</td>
<td>Greater high and low contrast acuity reduction with Fresnel prisms is mainly the result of reflection from the prism facets, secondary refraction at the prism facet bases and diffraction of light by the grooves</td>
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<tr>
<td></td>
<td>Hand-held CCTV &gt; optical device; d = 1.71</td>
<td>3 (100%) of patients under age 50 and 8 (67%) rated good satisfaction with CLT</td>
<td></td>
</tr>
<tr>
<td></td>
<td>No difference between types of CCTV; d not calculated</td>
<td>Authors suggest motivation and age determined successful use</td>
<td></td>
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<tr>
<td>Reading productivity (reading speed X duration = words per sitting):</td>
<td>Hand-held CCTV (2820.5 WPS) &gt; stand mounted CCTV (2745.8 WPS) &gt; optical device (1442.3 WPS)</td>
<td></td>
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</tr>
<tr>
<td>For types of CCTV only:</td>
<td>72% preferred stand-mounted to 23% for hand-held for most reading tasks</td>
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<td></td>
<td>Patients were equally divided when asked to consider device cost</td>
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</table>

**Comments**
- No proprietary interests reported
- No proprietary interests reported
- No proprietary interests reported
Table 7. Literature reviews included in the report

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<thead>
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</thead>
<tbody>
<tr>
<td>Topic</td>
<td>Tinted lenses and filters</td>
<td>Low vision rehabilitation</td>
<td>Low vision rehabilitation</td>
<td>Driving with vision impairment</td>
</tr>
<tr>
<td>Objective</td>
<td>Evaluate effectiveness of tinted lenses and filters for people with low vision</td>
<td>Evaluate effect of low vision on quality of life (QOL)</td>
<td>Evaluate effectiveness of low vision interventions on QOL</td>
<td>Evaluate impact of eye diseases and conditions on driving habits, performance, and safety</td>
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<tr>
<td></td>
<td>Identify a scientifically-based prescribing protocol for adoption in clinical practice</td>
<td>Evaluate effectiveness of low vision intervention on QOL</td>
<td>Identify factors affecting effectiveness of low vision interventions</td>
<td>Evaluate effectiveness of vision assisted driving using bioptic telescopic spectacles</td>
</tr>
<tr>
<td></td>
<td>Use of tinted lenses on progressive eye disease (e.g. Retinitis pigmentosa) excluded</td>
<td>Studies of adults with low vision or on QOL as an outcome measure in low vision rehabilitation included</td>
<td>Studies that examined patient satisfaction and performance with low vision services were tabulated</td>
<td>Identify public policy issues and research challenges in studying driving and vision impairment</td>
</tr>
<tr>
<td>Methods</td>
<td>MEDLINE searched for past 30 years using variety of key word combinations</td>
<td>MEDLINE and PsychINFO searched from 1990-2000 using key words of vision or vision disorders, QOL, and low vision rehabilitation</td>
<td>Available research outlined but little critical appraisal beyond study focus, population studies and instruments used</td>
<td>MEDLINE search from 1996-present using terms for vision, ocular, vision impairment, driving, accident and crash</td>
</tr>
<tr>
<td></td>
<td>Critical appraisal of study design outlined</td>
<td>Available research outlined but little critical appraisal beyond study focus, population studies and instruments used</td>
<td>Studies of prevalence of low vision, impact of low vision on affected individuals, and impact of low vision intervention on visual impairment included</td>
<td>Hand searching of end references of retrieved articles and authors' reprint collections were included</td>
</tr>
<tr>
<td></td>
<td>Commercially available tinted lenses included: Comcir photochromic filter®, UVShields® and NoIR® filters</td>
<td>Excluded were reports of instrument development, clinical trials, restorative treatments of vision, and community-based studies of sensory impairment</td>
<td>Published studies ranged from 1986 – 1996</td>
<td>Full length articles only</td>
</tr>
<tr>
<td></td>
<td>Use of tinted lenses on progressive eye disease (e.g. Retinitis pigmentosa) excluded</td>
<td>Full length articles only</td>
<td>Children and adults considered</td>
<td></td>
</tr>
<tr>
<td>Outcome measures</td>
<td>Objective visual measures and subjective measures</td>
<td>QOL using validated instruments</td>
<td>Patient satisfaction</td>
<td>Crash rate</td>
</tr>
<tr>
<td></td>
<td>Objective measures further analyzed by measure used</td>
<td>Performance measured with low vision services</td>
<td>Driving performance</td>
<td></td>
</tr>
<tr>
<td>Results</td>
<td>Most results were equivocal</td>
<td>Visual impairment significantly associated with decreased functional status, decreased self-reported QOL, and increased risk for depression</td>
<td>Low vision intervention can have a significant impact on patients' activities of daily living and can be highly valued by patients based on self-reporting</td>
<td>Evidence of the impact of types of visual impairment on driving safety and performance is inconclusive</td>
</tr>
<tr>
<td></td>
<td>Some subjective improvement reported but no consistent objective benefit</td>
<td>Decreased visual acuity, visual field, and occasional blurred vision are associated with decreased QOL</td>
<td>Impact of low vision intervention on QOL is unknown</td>
<td>Peripheral vision may play a greater role than visual acuity</td>
</tr>
<tr>
<td></td>
<td>Little conclusive evidence of improvement in visual function or of the superiority of tinted lenses with spectral characteristics over neutral density filters or conventional sunglasses</td>
<td>Low vision services are associated with increased self-reported functional status and QOL</td>
<td>Training, diagnosis, and model of care may affect effectiveness of low vision services but their impact has not been studied</td>
<td>Visual attention and processing show promise as a method for identifying high-risk drivers</td>
</tr>
<tr>
<td></td>
<td>Not possible to base tinted lens use on type of task or eye condition</td>
<td>Self-reported QOL is a significant outcome measure for low-vision rehabilitation</td>
<td>Application of QOL instruments to low vision populations is needed for evaluation of outcomes of low vision services</td>
<td>Evidence of the effectiveness of biopic use on driving performance or safety is inconclusive</td>
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<td></td>
<td>Practitioners must continue to rely on marketing literature, subjective reports, clinic-based observations and real-world trials in determining appropriate prescription of devices in low vision</td>
<td>Instruments are needed that are more sensitive to rehabilitation services, patient needs and goals to facilitate development of rehabilitation plans and to compare techniques, devices, and programs</td>
<td>Such a study is needed that establishes the type and magnitude of the change in functioning and QOL in a way that allows comparison with treatment of other disorders</td>
<td>Research challenges include accounting for the effects of driving exposure, age, and ocular disease in study design, classifying and defining outcomes that can be generalized across populations, and developing tests to identify unsafe drivers</td>
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<td>Measurement properties, validity, and reliability of instrument currently used and in development</td>
<td>Low vision services are associated with increased self-reported functional status and QOL</td>
<td>Well designed studies are needed to assess safety of low vision drivers using biopics or with monococular vision impairment or blindness, and the effectiveness of vision rescreening after initial licensure.</td>
</tr>
</tbody>
</table>
Articles included for critical appraisal


Background articles


OPTICAL LOW VISION DEVICES FINAL REPORT

FDA. Title 21 Food and Drugs, Volume 8, Chapter 1, Part 886 Ophthalmic Devices. Code of Federal Regulations. Vol. 21CFR886, April 1, 1998: 886.5 Therapeutic Devices. Title 21 Food and Drugs, Volume 8, Chapter 1, Part 886 Ophthalmic Devices.


Goodrich GL. Low vision citations. personal bibliography To: Adams E; March 30, 2002.

Guyatt GH, Sackett DL, Cook DJ. Users' Guides to the Medical Literature. II. How to use an article about therapy or prevention. A. Are the results of the study valid? JAMA, 1993; 270: 2598-2601.


Wolffsohn JS, Cochrane AL. *The practical near acuity chart (PNAC) and prediction of visual ability at near*. *Ophthalmic & Physiological Optics*, 2000; 20: 90-97.


**Excluded articles**


