Accuracy of Noncycloplegic Retinoscopy, Retinomax Autorefractor, and SureSight Vision Screener for Detecting Significant Refractive Errors

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PURPOSE. To evaluate, by receiver operating characteristic (ROC) analysis, the ability of noncycloplegic retinoscopy (NCR), Retinomax Autorefractor (Retinomax), and SureSight Vision Screener (SureSight) to detect significant refractive errors (RE) among preschoolers.

METHODS. Refraction results of eye care professionals using NCR, Retinomax, and SureSight (n = 2588) and of nurse and lay screeners using Retinomax and SureSight (n = 1452) were compared with masked cycloplegic retinoscopy results. Significant RE was defined as hyperopia greater than +3.25 diopters (D), myopia greater than 2.00 D, astigmatism greater than 1.50 D, and anisometropia greater than 1.00 D interocular difference in hyperopia, greater than 3.00 D interocular difference in myopia, or greater than 1.50 D interocular difference in astigmatism. The ability of each screening test to identify presence, type, and/or severity of significant RE was summarized by the area under the ROC curve (AUC) and calculated from weighted logistic regression models.

RESULTS. For detection of each type of significant RE, AUC of each test was high; AUC was better for detecting the most severe levels of RE than for all REs considered important to detect (AUC 0.97–1.00 vs. 0.92–0.95). The area under the curve of each screening test was high for myopia (AUC 0.97–0.99). Noncycloplegic retinoscopy and Retinomax performed better than SureSight for hyperopia (AUC 0.92–0.99 and 0.90–0.98 vs. 0.85–0.94, P < 0.02), Retinomax performed better than NCR for astigmatism greater than 1.50 D (AUC 0.95 vs. 0.90, P = 0.01), and SureSight performed better than Retinomax for anisometropia (AUC: 0.85–1.00 vs. 0.76–0.96, P ≤ 0.07). Performance was similar for nurse and lay screeners in detecting any significant RE (AUC 0.92–1.00 vs. 0.92–0.99).

CONCLUSIONS. Each test had a very high discriminatory power for detecting children with any significant RE.

Keywords: vision screening, refractive error, children’s vision

Significant refractive errors are the most prevalent and treatable vision problems in preschool children.1 A high prevalence of refractive error in young children has been established.2–10 Previous literature and clinical practice guidelines have identified an association between significant refractive error and amblyopia and strabismus.11–18 Research has also revealed educational and cognitive implications of uncorrected hyperopia.19–22 Therefore, significant refractive error is important to detect with vision screening.23–26

The Vision In Preschoolers (VIP) Study Group showed that screening tests of refraction (noncycloplegic refraction [NCR], Retinomax autorefractor [Retinomax], and SureSight Vision Screener [SureSight]) had excellent testability and performed best in identifying preschool children with VIP-targeted vision disorders, including significant refractive error, strabismus, and amblyopia.23 Furthermore, the VIP Study Group has shown that the Retinomax and SureSight can be used by trained eye care professionals, nurse screeners, or lay screeners to detect significant refractive errors.23–25

Although previous studies have evaluated detection of refractive error in preschool children using the Retinomax,23–25,27–38 or SureSight,23–25,36–43 age range often varies amongst studies and the majority of studies focus on agreement between autorefraction and a gold standard measure of refraction. Few studies have performed receiver operating characteristic (ROC) curve analysis to examine the ability of NCR, Retinomax, or SureSight to detect or screen for specific types of refractive error. Because there is not complete agreement regarding the best specificity level for use in screening, ROC curve analysis is helpful to compare the

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relevant methods are described briefly below.

Because (1) prevalence and type of refractive error vary across different pediatric licensed eye care professionals in mobile medical units in Phase I and by nurse and lay screeners in a school screening setting in Phase II. The VIP Study personnel were trained and certified in each procedure. In Phase 1, year 01, NCR and Retinomax (in automatic mode) (Right Manufacturing, Virginia Beach, VA) were performed. The eye care professional performed NCR using a streak retinoscope and retinoscopy lens rack or handheld trial lenses while the child watched an animated target on a video screen. The child wore retinoscopy spectacles, corresponding to the screener’s working distance, to control accommodation. In Phase 1, year 02 and in Phase 2, the Retinomax and SureSight Vision Screener (in child mode) (Welch Allyn, School Health Corp., Hanover Park, IL) were used. As part of a comprehensive eye examination, cycloplegic retinoscopy was performed on all children on a later day by a different pediatric licensed eye care professional who was trained and certified in the procedures and who was masked to the screening results. Cycloplegic retinoscopy was performed 30 to 40 minutes after instillation of drops including 1% cyclopentolate.

**Measurement of Refractive Error**

Each child’s vision was screened using two tests of refraction. Testing was performed by pediatric licensed eye care professionals in mobile medical units in Phase I and by nurse and lay screeners in a school screening setting in Phase II. The VIP Study personnel were trained and certified in each procedure. In Phase 1, year 01, NCR and Retinomax (in automatic mode) (Right Manufacturing, Virginia Beach, VA) were performed. The eye care professional performed NCR using a streak retinoscope and retinoscopy lens rack or handheld trial lenses while the child watched an animated target on a video screen. The child wore retinoscopy spectacles, corresponding to the screener’s working distance, to control accommodation. In Phase 1, year 02 and in Phase 2, the Retinomax and SureSight Vision Screener (in child mode) (Welch Allyn, School Health Corp., Hanover Park, IL) were used. As part of a comprehensive eye examination, cycloplegic retinoscopy was performed on all children on a later day by a different pediatric licensed eye care professional who was trained and certified in the procedures and who was masked to the screening results. Cycloplegic retinoscopy was performed 30 to 40 minutes after instillation of drops including 1% cyclopentolate.

**Definitions of Vision Disorders and Classification of Children**

Results from cycloplegic retinoscopy were used to classify children with respect to the presence or absence of each type of significant refractive error. If either or both eyes had significant refractive error, the child was considered to have significant refractive error. Hyperopia and myopia were defined as greater than 3.25 D or 2.00 D, respectively, in any meridian in either eye. Children with greater than 1.50 D difference between principal meridians were classified as having astigmatism. Anisometropia was defined as greater than 1.00 D interocular difference in hyperopia (most hyperopic meridian), or greater than 3.00 D interocular difference in myopia (most myopic meridian), or greater than 1.50 D interocular difference in astigmatism. Each type of significant refractive error was further classified into a hierarchy of groups or levels of severity (Table 1).

Children were classified as a screening pass or fail based upon the child’s worse eye and using the following results for each screening test of refraction: most positive meridian for hyperopia, most negative meridian for myopia, cylinder for astigmatism, and maximum interocular difference for anisometropia.

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**Table 1. Definitions of Significant Refractive Errors in the VIP Study**

<table>
<thead>
<tr>
<th>Refractive Error</th>
<th>Group 1</th>
<th>Groups 1 and 2</th>
<th>Groups 1, 2, and 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hyperopia</td>
<td>≥+5.0 D</td>
<td>&gt;+3.25 D with IOD in SE of ≥+0.5 D</td>
<td>&gt;+3.25 D with IOD in SE of ≤+0.5 D</td>
</tr>
<tr>
<td>Myopia</td>
<td>≥6.0 D</td>
<td>≥4.0 D</td>
<td>&gt;2.0 D</td>
</tr>
<tr>
<td>Astigmatism</td>
<td>≥2.5 D</td>
<td>&gt;1.5 D</td>
<td>N/A</td>
</tr>
<tr>
<td>Anisometropia (IOD)</td>
<td>&gt;2.0 D hyperopia, &gt;3.0 D astigmatism, or &gt;6.0 D myopia</td>
<td>1.0 D hyperopia, &gt;1.5 D astigmatism, or &gt;3.0 D myopia</td>
<td>N/A</td>
</tr>
</tbody>
</table>

IOD, intraocular difference; N/A, not applicable.

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**Methods**

This is a secondary data analysis of the VIP data. The VIP Study methods have been published in detail previously. The relevant methods are described briefly below.

**Subjects**

During the VIP Study, 3- to 5-year-old Head Start preschool children (n = 4040) were enrolled at one of five VIP clinical centers (New England College of Optometry, Boston, MA; Northeastern State University Oklahoma College of Optometry, Tahlequah, OK; The Ohio State University College of Optometry, Columbus, OH; Pennsylvania College of Optometry at Salus University, Philadelphia, PA; and University of California Berkeley School of Optometry, Berkeley, CA). The proportion of children with vision problems was enriched in the VIP Study by recruiting all children who failed the local Head Start vision screening and a random sample of children who passed the screening. The VIP Study adhered to the tenets of the Declaration of Helsinki and was approved by the appropriate local institutional review boards associated with each VIP center. Parents or legal guardians of participating children provided written informed consent/parental permission prior to testing.
Table 2. Frequency Distribution of Significant Refractive Error by Hierarchy

<table>
<thead>
<tr>
<th></th>
<th>Group 1 Only n (%)</th>
<th>Groups 1 and 2 n (%)</th>
<th>Groups 1, 2, and 3 n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>VIP Phase 1, year 1, N = 1142</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any significant refractive error</td>
<td>123 (10.8)</td>
<td>222 (19.4)</td>
<td>240 (21.0)</td>
</tr>
<tr>
<td>Myopia</td>
<td>8 (0.7)</td>
<td>13 (1.1)</td>
<td>28 (2.5)</td>
</tr>
<tr>
<td>Hyperopia</td>
<td>56 (4.9)</td>
<td>96 (8.4)</td>
<td>122 (10.7)</td>
</tr>
<tr>
<td>Astigmatism</td>
<td>64 (5.6)</td>
<td>131 (11.5)</td>
<td>131 (11.5)</td>
</tr>
<tr>
<td>Anisometropia</td>
<td>10 (0.9)</td>
<td>49 (4.3)</td>
<td>49 (4.3)</td>
</tr>
<tr>
<td>VIP Phase 1, year 2, N = 1446</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any significant refractive error</td>
<td>146 (10.1)</td>
<td>262 (18.1)</td>
<td>299 (20.7)</td>
</tr>
<tr>
<td>Myopia</td>
<td>7 (0.5)</td>
<td>9 (0.6)</td>
<td>25 (1.7)</td>
</tr>
<tr>
<td>Hyperopia</td>
<td>67 (4.6)</td>
<td>116 (8.0)</td>
<td>168 (11.6)</td>
</tr>
<tr>
<td>Astigmatism</td>
<td>82 (5.7)</td>
<td>156 (10.8)</td>
<td>N/A</td>
</tr>
<tr>
<td>Anisometropia</td>
<td>10 (0.7)</td>
<td>47 (3.3)</td>
<td>N/A</td>
</tr>
<tr>
<td>VIP Phase II, N = 1452</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any significant refractive error</td>
<td>198 (13.6)</td>
<td>337 (23.2)</td>
<td>380 (26.2)</td>
</tr>
<tr>
<td>Myopia</td>
<td>9 (0.6)</td>
<td>11 (0.8)</td>
<td>39 (2.7)</td>
</tr>
<tr>
<td>Hyperopia</td>
<td>80 (5.5)</td>
<td>124 (8.5)</td>
<td>182 (12.5)</td>
</tr>
<tr>
<td>Astigmatism</td>
<td>114 (7.9)</td>
<td>218 (15.0)</td>
<td>N/A</td>
</tr>
<tr>
<td>Anisometropia</td>
<td>19 (1.3)</td>
<td>56 (3.9)</td>
<td>N/A</td>
</tr>
</tbody>
</table>

Statistical Analysis

We assessed the accuracy of each screening test for detecting any significant refractive error and each type of refractive error (hyperopia, myopia, astigmatism, and anisometropia) using ROC curve analysis. The details of ROC curve analysis applied to the VIP Study data have been described previously by Ying et al.\textsuperscript{45} with respect to the performance of NCR, Retinomax, and SureSight in detecting preschoolers with one or more targeted vision disorders (amblyopia, strabismus, and significant refractive error).\textsuperscript{45} An ROC curve plots the sensitivity against the false positive rate (i.e., 1-specificity), in which each point reflects values obtained at a different cutpoint value from a continuous or ordinal measure. By calculating the area under the ROC curve (AUC), the ROC analysis provides a summary of the discriminative ability for a screening test, and allows a quick comparison of discriminative ability among different screening tests. The AUC has a value from 0.0 to 1.0. An AUC greater than 0.82–0.92 (SureSight: 0.85–1.00; Retinomax: 0.90–0.98) indicates good discriminative ability for a screening test, and allows a quick comparison of discriminative ability among different screening tests.

The Retinomax performed better than NCR for the detection of astigmatism greater than 1.50 D (AUC 0.95 vs. 0.88–1.00) as compared with any significant anisometropia (0.88–1.00) as compared with any significant anisometropia (0.95–1.00) as compared with any significant anisometropia (0.88–1.00) as compared with any significant anisometropia (0.88–1.00). The AUCs were next highest for hyperopia (NCR: 0.95–1.00; Retinomax: 0.95–1.00; SureSight: 0.95–1.00; Fig. 4, Supplementary Table S2), and astigmatism (NCR: 0.90–0.96; SureSight: 0.97–0.98; Retinomax: 0.90–0.96; Fig. 2, Supplementary Table S4), with higher AUCs for the most severe (highest) levels of hyperopia and from 0.85 to 0.94 for any hyperopia greater than 3.25 D (Fig. 3, Supplementary Table S3). The AUCs were somewhat lower for anisometropia (NCR: 0.82–0.92; SureSight: 0.85–1.00; Retinomax: 0.76–0.96), with higher AUCs for the most severe (highest) level of anisometropia (0.88–1.00) as compared with any significant anisometropia (0.76–0.91; Fig. 5, Supplementary Table S5).

The Retinomax performed better than NCR for the detection of myopia, hyperopia, astigmatism, and anisometropia are shown in Figures 1 to 5, respectively. Supplementary Tables S1 through S5 show the AUC for any significant refractive error, myopia, hyperopia, astigmatism, and anisometropia, respectively, by hierarchy (groups of levels of severity) and for each type of test and screener. For the detection of any significant refractive error overall, the AUC of each screening test was high. The AUC ranged from 0.97 to 1.00 for detecting the most severe (highest) levels of RE, and ranged from 0.92 to 0.93 for detecting any significant refractive error (Fig. 1, Supplementary Table S1). The AUCs were highest for myopia (NCR: 0.99; SureSight: 0.97–0.98; Retinomax: 0.97–0.98; Fig. 2, Supplementary Table S2), and astigmatism (NCR: 0.90–0.96; SureSight: 0.95–0.99; Retinomax: 0.95–0.99; Fig. 4, Supplementary Table S4). The AUCs were next highest for hyperopia (NCR: 0.92–0.99; SureSight: 0.85–0.94; Retinomax: 0.90–0.98), with the AUC ranging from 0.92 to 0.99 for the most severe (highest) levels of RE, and ranged from 0.92 to 0.93 for detecting any significant refractive error (Fig. 1, Supplementary Table S1). The AUCs were highest for myopia (NCR: 0.99; SureSight: 0.97–0.98; Retinomax: 0.97–0.98; Fig. 2, Supplementary Table S2), and astigmatism (NCR: 0.90–0.96; SureSight: 0.95–0.99; Retinomax: 0.95–0.99; Fig. 4, Supplementary Table S4). The AUCs were next highest for hyperopia (NCR: 0.92–0.99; SureSight: 0.85–0.94; Retinomax: 0.90–0.98), with the AUC ranging from 0.92 to 0.99 for the most severe (highest) levels of hyperopia and from 0.85 to 0.94 for any hyperopia greater than 3.25 D (Fig. 3, Supplementary Table S3). The AUCs were somewhat lower for anisometropia (NCR: 0.82–0.92; SureSight: 0.85–1.00; Retinomax: 0.76–0.96), with higher AUCs for the most severe (highest) level of anisometropia (0.88–1.00) as compared with any significant anisometropia (0.76–0.91; Fig. 5, Supplementary Table S5).
The detection of hyperopia was better for NCR and Retinomax than for SureSight (AUC NCR: 0.92–0.99 and Retinomax: 0.90–0.98 versus SureSight: 0.85–0.94, \( P \leq 0.02 \) for Retinomax versus SureSight). The differences between Retinomax and SureSight reached statistical significance for all three types of screeners for one or more severity levels of hyperopia (Fig. 3, Supplementary Table S3). The detection of anisometropia was better for SureSight than for Retinomax (AUC 0.85–1.00 vs. 0.76–0.96, \( P \leq 0.07 \)), with differences reaching statistical significance for nurse and lay screeners at both severity levels of anisometropia and for licensed eye care professionals for any significant anisometropia (Fig. 5, Supplementary Table S5).

For significant refractive error overall, performance was similar for nurse and lay screeners (AUC 0.92–1.00 vs. 0.92 to 0.99), with generally small, nonsignificant differences. NCR, Retinomax, and SureSight all had very high discriminatory power for detecting children with any significant refractive error when administered by pediatric licensed eye care professionals, pediatric nurse screeners, or lay screeners (Fig. 1, Supplementary Table S1). Differences in the AUC for the detection of each type of refractive error for pediatric nurse and lay screeners also were small and generally nonsignificant.

To provide the failure criteria that maximize the sensitivity for detecting presence of any significant refractive error by each screening test, we pooled data from all VIP phases because of the similarity in the ROC across all phases and determined the sensitivities at several specificity levels ranging from 50% to 90%. The sensitivities for detecting any significant refractive error, the most severe (group 1) level of refractive error, and each specific type of refractive error are provided in Supplementary Table S6. The corresponding failure criteria for detection of any significant refractive error are also provided in Supplementary Table S6. The cutoffs for failure criteria for detecting any significant refractive error differed among screening tests, depending on the type of refractive error. For example, for detection of significant refractive error overall at 90% specificity, the referral criteria varied from \((-1.00 \text{ D (SureSight)}\) to \(-1.75 \text{ D (NCR)}\) to \(-2.75 \text{ D (Retinomax)}\) for myopia, from 1.50 D (Retinomax) to 2.5 D
Second, an analysis of the VIP Study data by Ying et al. showed that both SureSight and Retinomax were useful instruments for detection of one or more targeted vision disorders for each test including the most severe conditions. Referral criteria for plegic SureSight had similar and high accuracy in detecting levels and types of refractive error. However, the SureSight showed better detection of anisometropia than that of the Retinomax, indicating there was apparently more fixation targets. Identification of astigmatism was better with the Retinomax than with NCR, indicating there was apparently more relaxation of accommodation. Although the test distance for the SureSight is further from the child than that of the Retinomax, we found significantly better detection of hyperopia with Retinomax. Therefore, the Retinomax target may provide better relaxation of accommodation. However, the SureSight showed better detection of anisometropia than the Retinomax, indicating there was apparently more consistent accommodation between the two eyes perhaps due to either the more remote test distance or differences in the fixation targets. Identification of astigmatism was better with the Retinomax than with NCR. An association has previously been shown among hyperopia and anisometropia and astigmatism, which may in part explain the excellent performance for all three tests for detection of overall significant refractive error difference despite differences among the tests for detection of specific levels and types of refractive error.

These results support the conclusions of two previous studies. Specifically, Cordonnier and De Maertelaer showed that both SureSight and Retinomax were useful instruments for vision screening, with the use of unique referral criteria. Second, an analysis of the VIP Study data by Ying et al. showed that NCR, noncycloplegic Retinomax, and noncycloplegic SureSight had similar and high accuracy in detecting preschoolers with one or more targeted vision disorders including the most severe conditions. Referral criteria for detection of one or more targeted vision disorders for each test of refraction have been reported by Ying et al. This study shows similar performance of each test of refraction for the detection of specific types of refractive error and overall refractive error and adds to the findings of Ying et al., which showed comparable performance between tests for detection of significant vision disorders overall.

Each of the three tests of refraction evaluated in this study are among the best performing screening tests and each may be used as a stand-alone vision screening procedure. Because many strabismic children have significant refractive error, these screening tests of refraction identify many children with strabismus. However, a test of refraction also may be combined with a test of eye alignment/stereopsis in order to attain some improvement in sensitivity for detection of strabismus. These findings suggest that other differences such as personnel needed, ease of use/interpretation, cost, and testing time may be considered in selecting which of these tests to use for screening. Both Retinomax and SureSight may be performed by eye care professionals, trained nurses, or lay screeners, while NCR requires more extensive training. Differences in the detection of significant refractive error between pediatric nurse and lay screeners were small. However, it is important to note that these results apply to the use of these tests of refraction by comparably trained personnel. Software is available for the SureSight (when minus cylinder format) which incorporates the VIP referral criteria (School Health Corp.) and displays an asterisk on the printout for each child who meets the VIP referral criteria, thus facilitating interpretation of the results. Manual interpretation of Retinomax results is needed as similar software is not available.

Noncycloplegic retinoscopy, Retinomax, and SureSight have been shown to perform somewhat better than screening tests of monocular acuity for detection of one or more visual disorders, the most severe visual disorders and significant refractive error. Because the recommended vision screening for children is frequently a screening test of monocular acuity alone, future research should compare the sensitivities of screening tests of monocular acuity alone with that of screening using combinations of tests (i.e., one of the best tests of monocular acuity and one of the best tests of refraction) in order to determine whether adding a screening test of noncycloplegic refraction to a test of monocular visual acuity improves detection of visual disorders including significant refractive error.

The strengths of this study include the standard training of screeners and application of screening protocols and standardized cycloplegic refractive error measurements performed by study-certified optometrists and ophthalmologists on all participating children. In addition, VIP Study participants were Head Start preschool children who were geographically, racially, and ethnically diverse. The enriched sample over representing preschool children with vision disorders provided a large number of preschool children with significant refractive error. However, a limitation of the study is the small number of children with myopia. Although children recruited to participate in the VIP Study include a higher percentage of children who failed an initial Head Start screening, and were thus more likely to have vision disorders, the analysis of detection of refractive error overall and for specific types of refractive error by NCR, Retinomax, and SureSight is generalizable to other preschool children.

In conclusion, AUC was excellent for the most severe (highest) levels of each type of refractive error and very good to excellent for the detection of any significant refractive error. Each test had a very high power for detecting preschool children with any significant refractive error.
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APPENDIX

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Philadelphia, PA: Pennsylvania College of Optometry

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Menacker, MD (GSE/LPS); Graham Quinn, MD, MSCE (GSE/LPS); Janet Schwartz, OD (GSE/LPS); Brandy Scombordi-Raghu, OD (GSE/LPS); Janet Swiatocha, OD (GSE/LPS); Edward Zikoski, OD (GSE/LPS); Jennifer Lin, MD (GSE); Leslie Kennedy (LS/PL); Rosemary Little (LS/PL); Geneva Moss (LS/PL); Latricia Rorie (LS); Shirley Stokes (LS/PL); Jose Figueroa (LS/VD); Eric Nesmith (LS); Gwen Gold (BPC/NHC/PL); Ashanti Carter (PL); David Harvey (LS/VD); Sandra Hall, RN (NS); Lisa Hildebrand, RN (NS); Margaret Lapsley, RN (NS); Cecilia Quenzer, RN (NS); Lynn Rosenbach, RN (NHC/NS).

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Study Center, Columbus, OH: The Ohio State University College of Optometry

Paulette Schmidt, OD, MS (PI); Beth Haas (Study Coordinator).

Coordinating Center, Philadelphia, PA: University of Pennsylvania, Department of Ophthalmology

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Maryann Redford, DDS, MPH.

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