ABSTRACT

Purpose. To investigate symptom patterns and evaluate the relationship between patient characteristics and symptom severity before and after treatment for symptomatic children with convergence insufficiency (CI).

Methods. In a randomized clinical trial, the convergence insufficiency symptom survey was administered pre- and posttreatment to 221 children aged 9 to <18 years with symptomatic CI. Frequency of symptom type was determined at baseline, mean change in performance-related vs. eye-related symptoms for treatment responders was compared, and the relationship between patient characteristics and symptom severity at baseline for the entire cohort and after treatment for those who responded to treatment was determined.

Results. At baseline, the score for performance-related symptoms was greater than that for eye-related symptoms (mean response of 2.3 vs. 1.8, p < 0.001) regardless of age, sex, race/ethnicity, or presence of parent-reported Attention Deficit Hyperactivity Disorder (ADHD). Symptom severity increased with age for both the overall and eye-related subscale scores (p = 0.048, p = 0.022, respectively). Children with parent-reported ADHD were more symptomatic (p = 0.005) than those without parent-reported ADHD because of a higher performance-related score (p < 0.001). A significant and equal improvement (p < 0.01) for the performance- and eye-related symptoms was found in treatment responders. Girls had significantly lower performance-related symptoms than boys (p = 0.001), and black children reported less eye-related symptoms than white children (p = 0.022). Children without parent-reported ADHD had significantly less symptoms overall and less eye-related symptoms than children with parent-reported ADHD (p = 0.019, p = 0.011, respectively).

Conclusions. Because of a high frequency of both performance- and eye-related symptoms, clinicians should perform a targeted history that addresses both types of symptoms to help identify children with symptomatic CI. Future study regarding the relationship of CI and symptoms and their potential influence on ADHD, reading performance, and attention is warranted.

Key Words: convergence insufficiency, asthenopia, vision therapy, orthoptics, quality of life, vergence/accommodative therapy, exophoria, eyestrain, symptom survey, reading, ADHD

Convergence insufficiency (CI) is a common binocular vision disorder\(^1\)–\(^4\) that is often associated with symptoms that occur when a person reads or performs close work. Complaints such as eyestrain, headaches, blurred vision, diplopia, sleepiness, loss of place, difficulty concentrating, movement of print, and poor comprehension after short periods of reading or performing near activities are often reported.\(^5\)–\(^12\) To quantify the frequency and severity of symptoms reported by individuals with symptomatic CI, the convergence insufficiency symptom survey (CISS) was developed.\(^13\)–\(^16\) A self-report symptom inventory, the CISS has been shown to have good construct validity and reliability,\(^14\)–\(^16\) and has been used as an outcome measure for clinical trials evaluating treatment modalities for children and adults with symptomatic CI.\(^17\)–\(^20\)

The CISS uses a Likert-type scale with responses from 15 items summed to obtain an overall CISS score, with symptom severity ranging from 0 (best) to 60 (worst). Although it has been suggested\(^13\) that the CISS items are composed of two categories of items—performance-related (e.g., difficulty concentrating when...
reading or studying) and eye-related (e.g., double vision) symptoms, the overall CISS score has been the only measure reported for the CI treatment trials.17–20

The CISS was used to quantify symptoms before and after treatment for 221 children with symptomatic CI enrolled into the convergence insufficiency treatment trial.20 Although the overall CISS scores at baseline and outcome and after 1 year of follow-up have been reported,20–22 the frequency of occurrence of specific symptoms at baseline and the relationship between patient characteristics and symptom severity have not been evaluated. The purpose of this report is to describe symptom patterns and to evaluate the relationship between patient characteristics and symptom severity before and after treatment.

METHODS

The study was supported through a cooperative agreement with the National Eye Institute of the National Institutes of Health, Department of Health and Human Services, and was conducted by the CITT group. The protocol and Health Insurance Portability and Accountability Act–compliant informed consent forms were approved by the institutional review boards for participating sites, and a parent or guardian of each study subject gave written informed consent. Each subject gave assent as required. An independent Data Safety Monitoring Committee provided study oversight. The study is registered at www.clinicaltrials.gov under identifier NCT00338611,23 and the manual of procedures is available at http://optometry.osu.edu/research/CITT/index.cfm. The examination and treatment procedures have been reported previously.21 Major eligibility criteria for the trial and the procedure for the administration of the CISS are summarized later in the text.

Subjects

Major eligibility criteria for the trial included children aged 9 to <18 years with symptomatic CI defined as an exodeviation at near at least four prism diopters (Δ) greater than at far, a receded near point of convergence break (≥6 cm), insufficient positive fusional vergence at near (PFV; convergence amplitudes, i.e., failing Sheard’s criterion [PFV less than twice the near phoria]24 or minimum PFV of ≥15Δ base-out blur or break), and a CISS (described later in the text) score of ≥16. In addition, children were required to have best-corrected visual acuity at distance and near of 20/25 or better, no constant strabismus, no vertical phoria >1Δ, and a monocular accommodative amplitude >5 D. All testing was performed with the appropriate refractive correction in place. The complete eligibility and exclusion criteria have been reported previously.21

CISS Administration

The CISS (Fig. 1) was administered to each child by a trained and certified examiner who was masked to the child’s treatment assignment. The examiner sequentially read each of the 15 symptom questions aloud to the child, while the child viewed a card with the five possible response options of never, infrequently, sometimes, fairly often, or always. Each response option corresponded to a numerical value ranging from 0 points for never to 4 points for always. The 15 items were summed to obtain the total CISS score. The score could range from 0 (least symptomatic) to 60 (most symptomatic; reporting always for all 15 symptoms). A CISS score ≥16 was considered symptomatic.14,16

A priori, the 15 symptoms on the CISS were categorized into two subscales.13 The performance-related subscale consisted of six symptoms related to visual efficiency when reading or performing near work (e.g., loss of concentration, loss of place with reading, reading slowly) and the eye-related subscale consisted of nine symptoms specific to visual function or asthenopic-type complaints (e.g., eyes hurt, diplopia, blurred vision, headaches) (Fig. 1). The subscale score represents the average level (range: 0–4) for all the items that comprise that category. The CISS was administered at baseline and at the conclusion of treatment; at each of these visits, it was administered twice, once before the clinical examination and again after the clinical examination was completed. For this report, the first CISS administration at baseline and outcome was used to determine the frequency with which each symptom was reported. The average scores from the two administrations of the CISS at the particular study visit were used for all analyses of overall and subscale scores.

Determination of Treatment Responders (Posttreatment Cohort)

The posttreatment cohort was composed of the children who responded to treatment during the CITT; they underwent different forms of treatment for CI, the details of which are reported elsewhere.20 Treatment responders were determined using the Reliable Change Index (RCI),25 which is a statistical method of determining the magnitude of change in a score (e.g., before and after intervention) necessary for a given self-reported measure to be considered statistically reliable. It represents the number of points necessary to determine if a change in score from pre- to posttreatment is from real change or from chance variation. The RCI takes into account both the population variance and the reliability of the test itself. Using CISS variability and reliability data from previous studies,17,18 the RCI was calculated to be a within-subject change in symptom level of ≥8.0 points. This resulted in classifying 53% (116 of 218) of the children in the CITT as having a reliable decrease in symptoms after treatment. Thus, for the purpose of this report, children with a posttreatment CISS score at least eight points less than their CISS score at baseline were considered treatment responders.

Data Analysis

Descriptive statistics are reported as the mean ± standard deviation. The mean performance- and eye-related subscale scores were compared using a paired t-test. The relationship between age and the overall CISS score and subscale scores at baseline was assessed using linear regression. The mean overall CISS score and subscale scores were compared between boys and girls, and children with and without parent-reported ADHD using two-sample t-tests. Analysis of variance was used to investigate the effect of race/ethnicity at baseline. Linear regression and analysis of covariance techniques were used to determine the effect of patient characteristics on improvement in overall and subscale CISS scores at the completion of treatment. For these analyses, the CISS score at baseline (overall and subscales) was used as
a covariate to adjust for any differences in baseline values. Tukey’s method was used to control the overall error rate when making post hoc pair-wise comparisons. All data analyses were performed using SAS (version 9.1, SAS Institute, Cary, NC).

RESULTS
Of the 221 children enrolled in the CITT at nine clinical sites, the mean age was 11.8 (±2.3) years, and 59% were female. Self-reported Hispanic ethnicity was 34%; race was reported to be 54% white, 29% black, and 17% other. Data analyses were performed after combining information on race and ethnicity as follows: Hispanics (any race), non-Hispanic whites, and non-Hispanic blacks. There were 34 children (15%) with parent-reported ADHD. Clinical characteristics for the CITT cohort have been reported previously. Of the 221 children enrolled, 218 (99%) completed their primary outcome examination.
Pretreatment Symptom Frequency

The mean overall CISS score at baseline was 29.8 (±9.0), and the median score was 29.5. The mean response (range of never = 0 to always = 4) for the 15 symptoms ranged from a low of 1.12 (±1.2) for “pulling around eyes” to a high of 2.55 (±1.2) for “loses place.” The median response corresponded to “sometimes” for all symptoms except “words jump/move,” “eyes feel sore,” and “pulling around eyes,” all of which had a median response of “infrequently.” The percentage of children who reported a particular symptom to occur “fairly often” or “always” is shown in Fig. 2. The most frequently reported symptoms were the six performance-related items.

Performance-related symptoms were reported to occur more frequently than eye-related symptoms, with mean subscale scores (i.e., average level of all items in that category) of 2.3 (±0.8) and 1.8 (±0.7), respectively (p < 0.001). This was consistent within age (p ≥ 0.012) and racial/ethnic groups (p ≤ 0.005), for both boys and girls (p < 0.0001), and regardless of the presence or absence of parent-reported ADHD (p ≤ 0.001).

Association of Pretreatment Symptom Severity with Patient Characteristics

At baseline, the mean overall CISS score and the performance- and eye-related subscale scores (Table 1) did not differ for boys and girls (p ≥ 0.40) or based on race/ethnicity (p ≥ 0.56). There was a slight increase in the overall CISS score with age (model R² = 0.018, p = 0.048), indicating that a 1-year increase in age is associated with an increase of 0.52 in overall CISS score. This increase was driven by an increase in the eye-related subscale score (model R² = 0.024, p = 0.022). The mean overall CISS and performance-related subscale scores were higher for the children with parent-reported ADHD (p = 0.005, p < 0.0001, respectively), but not for the eye-related subscale score (p = 0.31) (Table 1).

Posttreatment Symptoms

Among treatment responders, the median change was a 1-point decrease (lessening in severity by one level) for each symptom except for “pulling around the eyes,” which had a median change of 0. The mean improvement was the same for both the performance-related (1.1 ± 0.7) and eye-related subscales (1.1 ± 0.6). The percentage of treatment responders reporting “fairly often” or “always” for each of the 15 symptoms before and after treatment are shown in Fig. 3A, B, respectively. As found at baseline, performance-related symptoms were reported to occur more often than eye-related symptoms after treatment (p < 0.0001).

Association of Posttreatment Symptom Severity and Patient Characteristics

Post hoc comparisons were performed to assess the relationship of age, sex, race/ethnicity, and presence of parent-reported ADHD with improvements in symptoms as measured by the overall and subscale CISS scores for children classified as treatment responders (Table 2). There were no differences in the posttreatment symptom scores for the overall and the two subscale CISS scores based on age (p ≥ 0.34). For race/ethnicity, there was only a difference for the eye-related subscale (p = 0.026), and black children reported fewer eye-related symptoms than white children (p = 0.022). Children without parent-reported ADHD had significantly lower overall CISS and eye-related scores than children with parent-reported ADHD (p = 0.019, p = 0.011, respectively), but not in their performance-related subscale score (p = 0.13). After treatment, girls had significantly lower performance-related symp-
DISCUSSION

Using the CISS, we compared the frequency of specific symptoms at baseline and posttreatment as reported by children with symptomatic CI who participated in the CITT. Performance-related symptoms (e.g., difficulty concentrating when reading or studying) were reported more frequently than eye-related symptoms before treatment. In fact, the six most frequently reported symptoms were all performance-related items, regardless of age, sex, race/ethnicity, and presence of parent-reported ADHD. This finding is consistent with data reported from our pilot study, which found the top four symptoms reported at a fairly often or always level were performance-related items.

Before treatment, older children reported an increased frequency of eye-related symptoms and a greater overall CISS. This may have occurred for several reasons. Younger children may find it difficult to explain how their eyes feel, or they may not report symptoms because they consider them to be normal. Larger print size and less time spent performing sustained near work may result in fewer symptoms. It also is possible that as children get older, the condition progresses with a subsequent increase in eye-related symptoms. Finally, it is also possible that older children may spend more time with concentrated near tasks than their younger cohorts.

Before treatment, children with parent-reported ADHD had a significantly higher overall CISS score than children without parent-reported ADHD. This difference was almost entirely attributed to an increase in the frequency and severity of performance-related symptoms. While CI undoubtedly contributes to the presence of performance-related symptoms, there may be an additional contribution from the ADHD condition itself. For example, loss of concentration and trouble remembering when reading and doing close work are symptoms that are similar to behaviors such as “difficulty sustaining attention” and “forgetful in daily activities” that are often observed in individuals with inattentive ADHD. It would be of interest to investigate CISS performance- and eye-related subscale scores in children who have normal binocular vision and a primary diagnosis of ADHD.

Before treatment, children with parent-reported ADHD typically reported a decrease in both performance- and eye-related symptoms. This trend was similar across all age groups. The posttreatment profile of symptoms showed that performance-related symptoms still occurred with more frequency and severity than eye-related symptoms; this is similar to the profile reported in children with normal binocular vision. At this time, we do not have plausible hypotheses to explain why girls showed a greater reduction in their performance-related symptoms than boys, and why eye-related symptoms decreased more in black children than in white children. These could be spurious findings because of multiple statistical tests.

TABLE 1.
Convergence insufficiency symptom survey overall score$^a$ and mean subscale score$^b$ for performance- and eye-related items prior to treatment

<table>
<thead>
<tr>
<th>Patient characteristic</th>
<th>n</th>
<th>Overall score: mean (SD)</th>
<th>Performance-related subscale score: mean (SD)</th>
<th>Eye-related subscale score: mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sex</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>131</td>
<td>30.3 (9.2)</td>
<td>2.3 (0.7)</td>
<td>1.8 (0.8)</td>
</tr>
<tr>
<td>Male</td>
<td>90</td>
<td>29.2 (8.6)</td>
<td>2.3 (0.8)</td>
<td>1.7 (0.5)</td>
</tr>
<tr>
<td>p (t-test)</td>
<td></td>
<td>0.40</td>
<td>0.58</td>
<td>0.44</td>
</tr>
<tr>
<td><strong>Age</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9–10 yr</td>
<td>76</td>
<td>28.7 (7.8)</td>
<td>2.3 (0.7)</td>
<td>1.7 (0.7)</td>
</tr>
<tr>
<td>11–12 yr</td>
<td>69</td>
<td>29.4 (8.6)</td>
<td>2.3 (0.8)</td>
<td>1.8 (0.7)</td>
</tr>
<tr>
<td>13–14 yr</td>
<td>39</td>
<td>31.2 (10.7)</td>
<td>2.4 (1.0)</td>
<td>1.9 (0.8)</td>
</tr>
<tr>
<td>15–17 yr</td>
<td>37</td>
<td>31.7 (9.3)</td>
<td>2.3 (0.7)</td>
<td>2.0 (0.8)</td>
</tr>
<tr>
<td>p (regression)</td>
<td></td>
<td>0.048</td>
<td>0.48</td>
<td>0.022</td>
</tr>
<tr>
<td><strong>Race/ethnicity$^c$</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Black</td>
<td>59</td>
<td>30.5 (8.7)</td>
<td>2.4 (0.7)</td>
<td>1.8 (0.7)</td>
</tr>
<tr>
<td>Hispanic</td>
<td>75</td>
<td>29.4 (9.9)</td>
<td>2.3 (0.8)</td>
<td>1.7 (0.8)</td>
</tr>
<tr>
<td>White</td>
<td>78</td>
<td>30.4 (8.1)</td>
<td>2.3 (0.7)</td>
<td>1.9 (0.7)</td>
</tr>
<tr>
<td>p (ANOVA)</td>
<td></td>
<td>0.69</td>
<td>0.91</td>
<td>0.56</td>
</tr>
<tr>
<td><strong>Parent-reported ADHD status$^d$</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No ADHD</td>
<td>178</td>
<td>29.5 (8.7)</td>
<td>2.2 (0.8)</td>
<td>1.8 (0.7)</td>
</tr>
<tr>
<td>ADHD</td>
<td>34</td>
<td>34.1 (8.6)</td>
<td>2.8 (0.5)</td>
<td>1.9 (0.8)</td>
</tr>
<tr>
<td>p (t-test)</td>
<td></td>
<td>0.005</td>
<td>&lt;0.0001</td>
<td>0.31</td>
</tr>
</tbody>
</table>

$^a$Sum of item responses (range: 0–60).

$^b$Average level of all items (range: 0–4).

$^c$n = 212 (nine subjects categorized as “other” excluded from analysis).

$^d$n = 212 (missing ADHD status for nine children).

SD, standard deviation; ANOVA, analysis of variance; ADHD, attention deficit hyperactivity disorder.
Among children who were treatment responders, those without parent-reported ADHD reported a lower overall CISS score compared with children with parent-reported ADHD. The decrease in performance-related symptoms was similar between the two groups, despite the presence of higher performance-related symptoms at baseline among the children with parent-reported ADHD. The larger decrease in symptoms in children without parent-reported ADHD is primarily attributed to a decrease in eye-related symptoms. We speculate that children without parent-reported ADHD can attend better and longer, which may allow them to better notice improvement in their eye-related symptoms.

Our cohort of children reported performance-related symptoms with a higher frequency than eye-related symptoms. These symptoms, which include loss of place while reading, having to reread, reading slowly, loss of concentration, trouble remembering

FIGURE 3.
Percentage of patients classified as treatment responders who responded fairly often or always to each item on the convergence insufficiency symptom survey before (A) and after (B) treatment. Black color indicates performance-related symptoms; gray indicates eye-related symptoms.
what was read, may be obstacles to the reading process. In turn, performance-related symptoms may affect specific aspects of reading. Because successfully treated children report a significant decrease in performance-related symptoms, it is possible that the treatment of symptomatic CI may have a positive effect on reading performance and attention. These findings support the need for a randomized clinical trial to investigate whether the successful treatment of CI leads to improvements in specific measures of reading performance and attention.

Potential limitations of the study are that some children may experience symptoms that are not included on the CISS. In addition, there may be other conditions that may or may not have any relationship to CI or the eye, which could cause some of the same symptoms. Our previous finding that children with normal binocular vision have a mean CISS score of 8.1 illustrates that even these children are not without symptoms.14 However, clinicians less commonly recommend CI treatment for patients who do not have associated clinical signs, and the CISS cutoff of ≥16 differentiates between children with symptomatic CI and those with normal binocular vision.14,16 It is important to note that, by design, we only treated symptomatic children who scored ≥16 on the CISS. Thus, some children with CI may report fewer symptoms and some may not be symptomatic at all. Only a population-based study can determine the frequency of symptomatic CI. Because all children were not tested for ADHD, and the classification of ADHD was based on a parental report that this diagnosis had been previously made by a medical professional, there are likely to be some misclassifications.

Particular strengths of this study are that a reliable and valid symptom survey was administered in a standardized fashion by examiners masked to treatment assignment. The study population was a large, well-defined group of children with symptomatic CI randomized to therapies administered by trained and certified therapists. Retention in the trial was excellent at 99%.20 Children in our study of symptomatic CI reported performance-related symptoms more frequently than eye-related symptoms. With increasing age, the severity of performance-related symptoms remains constant, whereas that of eye-related symptoms increase. Children with parent-reported ADHD can be expected to score higher on performance-based symptoms and overall. Although both performance- and eye-related symptoms decrease in children after successful treatment, eye-related symptoms are still more frequently reported in those without parent-reported ADHD. More research is needed to determine whether a decrease in performance-related symptoms after treatment affects specific measures of reading performance and attention in children with symptomatic CI.

**Clinical Application**

Because children with symptomatic CI report performance-related symptoms more frequently than eye-related symptoms, a

### TABLE 2.

Convergence insufficiency symptom survey overall score and mean subscale score for performance- and eye-related items among children classified as “treatment responders” using Reliable Change Index after treatment

<table>
<thead>
<tr>
<th>Patient characteristic</th>
<th>n</th>
<th>Overall score: mean (SD)</th>
<th>Performance-related subscale score: mean (SD)</th>
<th>Eye-related subscale score: mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Female</td>
<td>72</td>
<td>14.1 (10.2)</td>
<td>1.2 (0.8)</td>
<td>0.8 (0.7)</td>
</tr>
<tr>
<td>Male</td>
<td>44</td>
<td>15.1 (9.2)</td>
<td>1.4 (0.8)</td>
<td>0.8 (0.6)</td>
</tr>
<tr>
<td>p (ANCOVA)</td>
<td></td>
<td>0.17</td>
<td>0.014</td>
<td>0.85</td>
</tr>
<tr>
<td>Age</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>9–10 yr</td>
<td>36</td>
<td>14.6 (9.9)</td>
<td>1.3 (0.9)</td>
<td>0.7 (0.7)</td>
</tr>
<tr>
<td>11–12 yr</td>
<td>38</td>
<td>14.3 (8.4)</td>
<td>1.2 (0.8)</td>
<td>0.8 (0.6)</td>
</tr>
<tr>
<td>3–14 yr</td>
<td>21</td>
<td>15.2 (12.4)</td>
<td>1.3 (1.0)</td>
<td>0.8 (0.9)</td>
</tr>
<tr>
<td>15–17 yr</td>
<td>21</td>
<td>13.9 (9.7)</td>
<td>1.1 (0.6)</td>
<td>0.8 (0.7)</td>
</tr>
<tr>
<td>p (regression)</td>
<td></td>
<td>0.34</td>
<td>0.62</td>
<td>0.35</td>
</tr>
<tr>
<td>Race/ethnicity</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Black</td>
<td>30</td>
<td>14.3 (9.1)</td>
<td>1.4 (0.9)</td>
<td>0.6 (0.6)</td>
</tr>
<tr>
<td>Hispanic</td>
<td>43</td>
<td>13.4 (9.4)</td>
<td>1.2 (0.7)</td>
<td>0.7 (0.7)</td>
</tr>
<tr>
<td>White</td>
<td>37</td>
<td>16.7 (10.6)</td>
<td>1.2 (0.8)</td>
<td>1.0 (0.7)</td>
</tr>
<tr>
<td>p (ANCOVA)</td>
<td></td>
<td>0.29</td>
<td>0.87</td>
<td>0.026</td>
</tr>
<tr>
<td>Parent-reported ADHD status</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No ADHD</td>
<td>97</td>
<td>13.8 (9.2)</td>
<td>1.2 (0.8)</td>
<td>0.7 (0.7)</td>
</tr>
<tr>
<td>ADHD</td>
<td>13</td>
<td>22.2 (11.0)</td>
<td>1.9 (0.8)</td>
<td>1.2 (0.8)</td>
</tr>
<tr>
<td>p (ANCOVA)</td>
<td></td>
<td>0.019</td>
<td>0.13</td>
<td>0.011</td>
</tr>
</tbody>
</table>

*a*Sum of item responses (range: 0–60).

*b*Average level of all items (range: 0–4).

*c*n = 110 (six subjects categorized as “other” excluded from analysis).

*d*n = 110 (missing ADHD status for six children).

SD, standard deviation; ANCOVA, analysis of covariance; ADHD, attention deficit hyperactivity disorder.
targeted history such as the CISS that addresses both performance-and eye-related symptoms is recommended.

ACKNOWLEDGMENTS

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The Convergence Insufficiency Treatment Trial (CITT) Investigator Group

Clinical Sites

Sites are listed in order of the number of patients enrolled in the study with the number of patients enrolled is listed in parentheses preceded by the site name and location. Personnel are listed as (PI) for principal investigator, (SC) for coordinator, (E) for examiner, and (VT) for therapist.

Study Center: Bacus Palmer Eye Institute (35): Susanna Tamkins, OD (PI); Hilda Capo, MD (E); Mark Dunbar, OD (E); Craig McKeown, MD (CO-PI); Atriana Moslehghi, MD (E); Kathryn Nelson, OD (E); Vicky Fischer, OD (VT); Adam Perlman, OD (VT); Ronda Singh, OD (VT); Eva Olivares (SC); Ana Rosa (SC); Nidia Rosado (SC); Elias Silverman (SC)

Study Center: SUNY College of Optometry (28): Jeffrey Cooper, OD, PI (PI); Aurora Steiner, OD, (E, CO-PI); Marta Brunelli (VT); Stacy Friedman, OD (VT); Steven Ritter, OD (E); Lily Zhu, OD (E); Lyndon Wong, OD (E); Ida Chung, OD (E); Kayti Colon (SC); Ashley Fazarry (SC)

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REFERENCES


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