Clinical observations of vitreoretinal surgery for four different phenotypes of X-linked congenital retinoschisis

Chen Zhao, Qi Zhang, Hai-Ying Jin, Pei-Quan Zhao

Department of Ophthalmology, Xinhua Hospital, School of Medicine, Shanghai Jiao Tong University, Shanghai 200092, China

Co-first authors: Chen Zhao and Qi Zhang

Correspondence to: Pei-Quan Zhao. Department of Ophthalmology, Xinhua Hospital, School of Medicine, Shanghai Jiao Tong University, No. 1665 Kongjiang Road, Shanghai 200092, China. zhaopeiquan@xinhuaomed.com.cn

Received: 2017-04-19        Accepted: 2017-09-25

Abstract

• AIM: To evaluate the outcomes of vitreoretinal surgery for four different phenotypes of X-linked retinoschisis (XLRS).

• METHODS: This study included thirty-one eyes of 25 patients who developed XLRS with severe complications. Among the 31 eyes, there were 7 eyes with vitreous hemorrhage, 8 eyes with retinal detachment and vitreous hemorrhage, and 16 eyes with rhegmatogenous retinal detachment. All of the patients underwent standard three-port pars plana vitrectomy. All of the eyes were divided into 4 groups before surgery according to a modified classification scheme, with the first three groups as follows: group A, with foveal cystic schisis; group B with macular lamellar schisis; and group C with foveolamellar changes. Peripheral schisis was not present in these 3 groups; however, group D was a complex group with both macular and peripheral changes. One year after surgery, we analyzed the best corrected visual acuity and postoperative anatomical and functional outcomes of these 4 groups.

• RESULTS: There were 7 eyes in group A (22.6%), 1 eye in group B (3.2%), 15 eyes in group C (48.4%) and 8 eyes in group D (25.8%). Postoperative anatomical and functional outcomes were satisfactory at the last visit, while the mean visual acuity was increased to 0.27±0.11, with a significant difference (t=-1.049, P=0.000) compared with the results before surgery (0.1±0.08). Visual acuity was improved in 23 eyes (74.2%). Complications were found in three eyes: two eyes with proliferative vitreoretinopathy and traction retinal detachment 10 and 12mo after surgery, respectively; and one eye with vitreous hemorrhage 15mo after surgery. These eyes were in groups C and D. The retinas remained attached in these 3 eyes after reoperation.

• CONCLUSION: We should pay greater attention to XLRS with foveolamellar changes because it is the most changeable phenotype. Once complications occur, vitreoretinal surgery can significantly improve visual acuity and restore the anatomic structure of the retina.

• KEYWORDS: X-linked retinoschisis; complications; vitreoretinal surgery

DOI:10.18240/ijo.2018.06.15

INTRODUCTION

X-linked retinoschisis (XLRS; OMIM: 312700) is a relatively common early onset retinal degenerative disease that affects males early in life[1]. XLRS may spontaneously regress or may be stationary or progressive. Roughly 20%[2] of all patients with XLRS may progress to retinal detachment (RD), and in up to one third of such patients[3], vitreous hemorrhage (VH) develops, which causes severe vision loss. Until now, no satisfying treatment has been available to halt the formation and development of schisis in patients with XLRS[4]. Surgical interventions are required for XLRS patients with severe complications such as RD and VH. In the fundus, different phenotypes are described: macular foveal schisis is regarded as the most typical change, radially oriented intraretinal foveomacular cysts are seen in a spoke-wheel configuration. In addition, approximately 50% of all cases have peripheral retinoschisis[1,3,5], some of which threatening the macula area around fovea. The optical coherence tomography (OCT) shows the splitting of retina usually located at the inner nuclear layer (INL), outer plexiform layer and outer nuclear layer (ONL), with small cystic spaces between the split retinal layers in the perifoveal region[6]. Do different phenotypes lead to different incidences of complications and prognosis? Our study aimed to examine a group of XLRS patients who underwent vitreoretinal surgery because of the severe complications with progressive XLRS. In the present study, we analyze the relationship of severe complications incidence rate and different phenotypes based on a modified classification scheme[7,8] according to its fundus and OCT image of XLRS. Our indications for surgery are discussed, and the results of our treatment are reported.
SUBJECTS AND METHODS
This was a retrospective study of 31 eyes of 25 patients treated with vitrectomy who were diagnosed with XLRS and suffered from severe complications. We retrospectively reviewed all of the case notes of children with XLRS who underwent vitreoretinal surgery in our department from March 2009 to March 2015, identifying 31 consecutive eyes. Patients were excluded if the follow-up was less than 1 y. A total of 31 eyes were enrolled in this study from Shanghai Jiao Tong University Affiliated with XinHua Hospital (Shanghai). We confirmed XLRS based on clinical evidence, including patient symptoms, fundus changes, OCT, fluorescein fundus angiography (FFA), and electroretinography (ERG). Diagnostic criteria including: 1) age of onset is under 15 years old; 2) severe impact in best corrected visual acuity (BCVA) (<20/60); 3) OCT images show the foveal cystic schisis, macular lamellar schisis with or without peripheral schisis; 4) ERG shows selectively reduced b wave. FFA results amplitude confirming the diagnosis. Baseline clinical characteristics and demographic data were collected from the medical records and included sex, age, eye examinations, diagnosis, and treatment. Preoperative data were obtained at the visit prior to surgical treatment and included BCVA, slit-lamp and both direct and indirect ophthalmoscopy, ERG and OCT. According to the presurgery results on ophthalmoscope and OCT, all of the eyes were divided into 4 groups based on a modified classification scheme[7-8], in which there were typical changes in fundus and/or in OCT. In the fundus, radially oriented intraretinal foveomacular cysts are seen in a spoke-wheel configuration in most cases (Figure 1), whereas OCT shows the most extended foveal schisis, with large, low, reflective cysts which are surrounded perifoveally by many smaller cysts, located in the INL and another deeper schisis appearing in the photoreceptolayer (Figure 2). The vascular changes were remarkable, appearing as extensive vascular sheathing in some cases (Figure 3). The incidence of preoperative complications was also analyzed preoperatively in the four different groups. Besides the cases when the complication occurred affect the result of OCT, we chose the data when it was first diagnosed. The surgical procedures were all performed by one surgeon (Zhao PQ). All of the patients underwent pars plana vitrectomy without lensectomy associated with internal limiting membrane peeling. Photocoagulation was applied to the retinal holes and degeneration areas. Gas or silicone oil was used to fill the eyes in different situations. After surgery, the patients were reviewed at 1, 3 and 6 mo, and further surgical interventions were performed when required.

We then obtained data from the postoperative clinical assessments and at each follow-up visit. BCVA, slit-lamp and direct and indirect ophthalmoscopy and OCT were performed in all of the participants at every visit after surgery (6 and 12 mo follow-ups), at which we obtained information about visual function and anatomical replacement. At the last visit (12 mo), we separated the patients into 4 groups. We calculated BCVA with a Snellen chart transferred the light perception (LP), hand move, and finger count to 1/800, 1/400, and 1/200, respectively, using statistics[9]. We compared the preoperative results and the data from 12-month follow-up visit by mean±standard deviation. “Favorable” outcomes were reported when BCVA increased by one line or more. Anatomically, the area of schisis was calculated by OCT, and we compared the preoperative and postoperative results; “favorable” outcomes were reported when the retina was flat when examined by
Clinical observation of XLRS

Table 1 Classification scheme of XLRS

<table>
<thead>
<tr>
<th>Groups</th>
<th>Foveal cystic schisis</th>
<th>Macular lamellar schisis</th>
<th>Peripheral schisis</th>
<th>BCVA mean±SD</th>
<th>Incidence (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>0.16±0.05</td>
<td>22.6</td>
</tr>
<tr>
<td>B</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>0.1</td>
<td>3.2</td>
</tr>
<tr>
<td>C</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td>0.08±0.13</td>
<td>48.4</td>
</tr>
<tr>
<td>D</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>0.1±0.17</td>
<td>25.8</td>
</tr>
<tr>
<td>Total</td>
<td></td>
<td></td>
<td></td>
<td>0.1±0.08</td>
<td></td>
</tr>
</tbody>
</table>

BCVA: Best corrected visual acuity.

ophthalmoscope. The paired sample t-test, in SPSS statistical software, version 13.0 (Chicago, IL, USA), was used to compare the preoperative and postoperative data. P values of 0.05 or less were considered statistically significant.

RESULTS

All 25 of the patients were male (100%), and the mean age at presentation was 7.47 years old (range 5 to 15y). Among the 31 eyes, there were 7 eyes with VH, 8 eyes with RD and VH, and 16 eyes with rhegmatogenous retinal detachment (RRD). The presurgical BCVA was from LP to 0.25, and the mean was 0.1±0.08. According to the results of ophthalmoscopy and OCT, peripheral retinoschisis occurred in 8 eyes (25.8%), whereas foveal involvement was present in all of the affected patients (Table 1). In these 8 eyes, peripheral retinoschisis presented as a sharply delineated detachment of the inner retinal sheath, limited to the periphery or mid-periphery; 2 of these 8 eyes presented several internal splits in the retina, which generated to vitreous veils. The incidence of complications preoperatively, respectively, in the four different groups is shown in Table 2. The most impressive outcome was that VH occurred in 5 of 8 eyes in group D (62.5%).

The mean follow-up period in this study was 37.2mo (range 12 to 58mo) and we recorded the data when 12-month after surgery as the last visit. After surgery, the patients achieved clear refractive media and good retinal replacement. At the last visit, 23 eyes (74.2%) had a “favorable” outcome with better visual acuity. Among these 23 eyes, 5 eyes were treated with photocoagulation before surgery because of the progressive extent of schisis; and dam-type endolaser was done for peripheral retinal holes and retinoschisis areas on 3 eyes in group D during the surgery (Figure 4). In this study, postoperative BCVA, anatomical replacement and the incidence of secondary complications were also reported (Table 3). The postoperative BCVA of our 31 eyes was significantly better in all of the four groups, with a mean final BCVA of 0.27±0.11. The difference was significant (t=1.04, P=0.000), compared with the BCVA before surgery (0.1±0.08). Among the 8 eyes rated “unfavorable”, 3 eyes in group C and D were treated with a second vitreoretinal surgery because of tractional RD recurrence after the first surgery. The second surgeries in theses 3 eyes were performed successively at the 10, 12 and 15mo follow-ups, respectively. However, according to the postoperative BCVA, they were considered “unfavorable”.

DISCUSSION

XLRS is one of the most common X-linked, recessively inherited, bilateral, blindness-inducing diseases. It is limited almost exclusively to male subject[3]. The differential diagnosis of XLRS essentially depends on the stage of the disease. XLRS can be progressive or stationary, or it can spontaneously regress. In most cases, it remains static. However, it will lead to blindness if severe complications occur. In this study, we focused on XLRS stages suffering severe complications. The incidence of VH was approximately 50% (including RD+VH). VH can be seen in retinopathy of prematurity (ROP), intracranial hemorrhage, thrombophilic diseases, or retinal dysplasia[10-11]. The most important differential diagnosis is ROP. Patients with ROP will have a history of premature
delivery and perinatal complications, and they will develop vascularization of the retinal periphery in the majority of eyes after involution of the disease\(^{[12-13]}\). In comparison, intrachisis or VH in XLRS results from the rupture of unsupported retinal vessels or rarely from neovascularization. Therefore, we do not use anti-VEGF drugs as we do in ROP before surgery. It is not difficult to distinguish these diseases, especially by using gene detection. Because the genotypes of XLRS have no significant connection with phenotype, we believe that the latter is more practical than the former. Furthermore, spectral domain OCT allowed high-resolution imaging and characterization of the features in XLRS, and it highlighted the variability of the same genetic disease\(^ {[14]}\). Therefore, in this study, we chose to divide the eyes into 4 groups by phenotype, according to typical characteristics on fundus and OCT images.

The etiology of XLRS is unclear\(^{[15]}\), but the vitreous dystrophy theory, the Müller cell theory of defects and the retinal vascular dystrophy theory are often mentioned. Due to inherited dystrophy in the interior retina, especially in the Müller cells or in the vitreous cortex, the interior retina is detached from the retinal nerve fiber layer (RNFL); thus, the incidences of secondary detachment and VH are 5% to 20% and 33%, respectively. Once these complications occurs, XLRS becomes a vision-threatening disease that requires early surgery in the majority of cases\(^{[2,16-17]}\). We found in this study that the eyes with the complication of VH were all in group D (combined peripheral schisis). With the presence of peripheral retinoschisis in the RNFL, vascular changes were prominent in these cases. The retinal vessels could remain running freely through the vitreous cavity if the inner sheet of the schisis degenerated. The extensive vascular sheathing with dendritic opacified retinal vessels; but no clear retinal neovascularization was found in our cases. The investigators believed that in some instances, VH was caused by traction on the retinal blood vessels\(^ {[18]}\). Other less common signs of vitreous traction were tenting of the inner retinal layer (12%) or blood vessels (4%) and geographic areas of white without pressure (35%\(^{[19]}\)). Because the sample size of this study was small, we should perform more studies to confirm the presumptions about the relevance of peripheral schisis and VH. However, it would help to perform FFA of the peripheral schisis to provide the correct diagnosis and treatment.

During the course of XLRS, secondary complications, including RD and VH, can occur and may lead to poor outcomes. Thus, which phenotype of XLRS is the most dangerous and most easily causes severe complications? Our study aimed to study cases with severe complications, with foveal involvement occurring in 96.8% of eyes, whereas Wille\(^ {[20]}\) reported that macular retinoschisis occurred in all affected patients. In our report, foveal retinoschisis was present in 98% of involved individuals. Peripheral retinoschisis, which is characterized by intraretinal splitting involving the nerve fiber layer\(^ {[12-21]}\), occurred in approximately 25.8% of eyes (group D), which is less than the 50% that Pachydiaki and Young\(^ {[22]}\) reported. In our study, the inner layer of the peripheral schisis was immobile and elevated in a stable configuration. In only one case, the original area of detachment was caused by a combination of inner and outer layer holes in areas of retinoschisis. Using primarily scleral buckling, external drainage of subretinal fluid was popular for treating RRD in XLRS. However, currently, this technique is usually performed only for more peripheral breaks. Considering that scleral buckling cannot relieve the traction on the retina thoroughly as well as the limitations of globe growth, we prefer vitrectomy\(^ {[4]}\), especially if the outer-layer holes are large and posterior. At the same time, inner layer retinotomy, internal drainage of schisis fluid and subretinal fluid, internal limiting membrane peeling, endolaser, gas-fluid exchange with long-acting gases and postoperative positioning of the patient, and use of heavy perfluorocarbon liquids to tamponade the schisis and detachment were used during surgery. It is clear that a variety of surgical techniques have yielded successful results, although it is also obvious that the best method for treating RRD associated with XLRS has not yet been determined. As we know, a perfect internal limiting membrane peel plays an important role in relieving the traction on the retina\(^ {[23-25]}\). As we showed in this study, successful vitreoretinal surgery in these affected eyes led to good outcomes. To achieve better BCVA in children, lens-sparing vitrectomy, combined with internal limiting membrane peeling, was performed in every case. Luna et al\(^ {[26]}\) reported that peripheral retinal laser photoagulation before lens-sparing vitrectomy for ROP eyes affected surgical outcomes. We also believe that it would work in XLRS. In our study, 5 eyes underwent photoagulation before surgery. During the surgery, the surgeon considered it difficult to manipulate the vitreous in the eyes that underwent pre-surgical laser photoagulation, although all of them had “favorable” outcomes. We also had 8 eyes in which BCVA did not improve after surgery. Among these 8 eyes, 6 (75%) had foveal cystic schisis pre-surgically. We believe the bad outcomes were also related to repeated surgeries and to long-term detachment, which leads to photoreceptor apoptosis\(^ {[27]}\).

We also had 3 cases that underwent a second surgery, all in groups C and D, in patients with foveomacular changes. Because it is the most changeable phenotype, we should pay greater attention to it and undertake longer follow-ups. Although a perfect internal limiting membrane peeling was important in these two groups, dam-type endolaser was necessary for peripheral retinal holes and retinoschisis areas (mostly in group D), which were effectively prevented from extending to the macula. Furthermore, we will continue to investigate whether photocoagulation can form a barrier against
Clinical observation of XLRS


14 Leng T. Two cases of X-linked retinoschisis with different spectral domain optical coherence tomography findings. *Clin Ophthalmol* 2012;6:1563-1565.


