Characteristics of Advanced Stages of Retinopathy of Prematurity

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Abstract

Purpose: To evaluate the incidence and risk factors of severe visual impairment and retinopathy of prematurity (ROP)-related blindness in a tertiary eye hospital in Iran

Methods: In a cross-sectional study, clinical data of premature infants screened for ROP in Farabi Eye Hospital during 2003-2007 were reviewed. Characteristics of advanced stages of ROP (stage 4 or 5) were determined and compared to other stages of ROP.

Results: Among 1053 infants, 380 (36.1%) had ROP, 91 infants (8.6%) had advanced stages of ROP in at least one eye (none of them had a history of previous ROP screening); 40 of them had bilateral stage 5. The mean gestational age (GA) and birth weight (BW) in infants with advanced ROP were 28.3±2.3 (24-35) weeks and 1267±398 (450-2600) g, respectively. 74.7% of infants with advanced ROP had been examined after 9 weeks of infantile age. Only 23.1% (21/91) had recommendation for eye examination by their neonatologists. Lower GA was the independent risk factor of developing advanced ROP (P< 0.001).

Conclusion: We observed a high incidence of advanced ROP (stage 4 and or 5) in our settings; most of them were not referred for screening examination during the first few weeks and had delayed eye examination. Among the possible risk factors, GA was the only factor related with development of advanced ROP.

Keywords: Blindness, Visual Impairment, Retinopathy of Prematurity


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Introduction

Childhood blindness can cause significant loss of productivity for the child and society; therefore, control of childhood blindness is a priority of WHO’s vision 2020 program.\(^1\) It has been estimated that about 75% of the total 1.4 million blind children live in developing countries.\(^1\) Retinopathy of prematurity (ROP) is one of the potentially preventable causes of childhood blindness\(^2\) which afflicts 50,000 infants each year.\(^3\) Based on the infant mortality rate in our country, it is predicted that Iran is one of the countries with high risk of severe visual impairment due to ROP and more data about it is needed.\(^4,5\)

Many studies have addressed the natural course and risk factors of ROP\(^6-10\); some have focused on risk factors of severe ROP\(^11-15\) and some have assessed risk factors of progression into advanced ROP and retinal detachment after treatment.\(^16-18\) However, it is not yet clear what causes the infant to have regressed ROP, and what causes the infant to progress into retinal detachment; thus, the question remains that if there is any specific risk factor for late stages of ROP. We undertook this study to report characteristics of advanced ROP in a tertiary eye hospital in Iran to provide preliminary evidence about visual impairment and blindness due to ROP in our country, and to assess risk factors of late stages of ROP.

Methods

The study was conducted in pediatric retina and vitreous division of Farabi Eye Hospital, affiliated with Tehran University of Medical Sciences, during October 2003 to November 2007. Data of premature infants screened for ROP were retrieved from infants’ medical records. The retrieved data were: the result of complete eye exam (according to the international classification of ROP\(^19\)), infantile age at first eye exam, neonatologist’s recommendation for eye examination, birth weight (BW), gestational age (GA), neonatal age, gender, singleton or multiple gestations, oxygen therapy, mechanical ventilation, respiratory distress syndrome, phototherapy, intraventricular hemorrhage, necrotizing entrocolitis, sepsis (blood culture positive) and blood transfusion. Data of the last 9 factors were available as yes/no and were analyzed as dichotomous variables. The statistical program of SPSS 15 was used for database analysis. The chi square test and independent sample t-test were employed to evaluate potential dichotomous and continuous risk factors respectively.

Definition In this study:

- **Mild ROP** was defined as ROP that needed no treatment.
- **Severe ROP** was defined as ROP that needed treatment (according to the criteria of early treatment for retinopathy of prematurity).\(^20\)
- **Advanced ROP** was defined as stage 4 or 5 of ROP in at least one eye.
- **Visual impairment** was defined as advanced ROP less than bilateral stage 5.

Bilateral stage 5 of disease was considered as blindness.

Results

During the period of the study, 1060 premature infants were examined, 7 were excluded because their complete eye examination result was not available. From 1053 infants, 380 (36.1%) had ROP, 91 of them (8.6%) were in advanced stages of ROP and 3.8% (\(40/1053\)) had bilateral stage 5 of ROP. From 91 infants with advanced ROP, 53 (58.2%) were males; 43 (47.3%) were born in Tehran and the rest were born in other cities and none of them had a history of previous ROP screening. The mean GA and BW of infants with advanced ROP are compared with other stages of ROP in table 1. The mean infantile age at first ophthalmic examination was 27.5±31.7 weeks in advanced ROP while it was 6.7±2.9 weeks in infants with other stages of ROP, the difference was statistically significant (P<0.001). The first eye exam of 74.7% (\(68/91\)) of infants with advanced ROP was done after 9 weeks of infantile age. Only 21 of 91 infants (23.1%) had recommendation for eye examination during the first few weeks by their neonatologists, and 33 (36.32%) did not have it (the reason of their attendance for retinal exam is summarized in table 2). The data was unavailable in the rest of 37 infants.

Seven of forty infants (17.5%) with stage 5 in both eyes had neonatal recommendation for eye examination before 9 weeks of
neonatal age and 17 (42.5%) did not have this recommendation. Two of them (5%) were recommended by another physician to have eye examination when they were older than 9 weeks.

In statistical analyses, only lower GA was identified as risk factor of having advanced ROP compared to other stages of ROP (P<0.001), while there was no significant association between advanced ROP and other factors. Using ROP grouping of: none, mild, severe and advanced; a linear correlation with GA was seen as is shown in figure 1. The difference of GA among these 4 group was statistically significant (P<0.001). Using the same ROP grouping, BW showed a linear correlation with ROP except for advanced ROP (Figure 2). The difference of BW among the first three groups was statistically significant (P<0.02); however, the difference between severe and advanced ROP was not significant (P=0.9).

<table>
<thead>
<tr>
<th>Stages of ROP</th>
<th>Mean gestational age (Minimum-Maximum)</th>
<th>Mean birth weight (Minimum-Maximum)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Advanced retinopathy of prematurity</td>
<td>28.3±2.3 (24-35) weeks</td>
<td>1267±398 (450-2600) g</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Other stages of retinopathy of prematurity</td>
<td>29.8±2.3 (24-36) weeks</td>
<td>1333±379 (560-2800) g</td>
<td>&gt;0.05</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Reason</th>
<th>No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parental notice to their infant’s low vision</td>
<td>9 (27.3%)</td>
</tr>
<tr>
<td>Parental notice to other ophthalmic problems</td>
<td>4 (12.1%)</td>
</tr>
<tr>
<td>Being recommended by another physician when they were older than 9 weeks</td>
<td>7 (21.2%)</td>
</tr>
<tr>
<td>Being recommended by neonatologist to seek care at 3rd month</td>
<td>1 (3.0%)</td>
</tr>
<tr>
<td>Unavailable data</td>
<td>12 (36.4%)</td>
</tr>
<tr>
<td>Total</td>
<td>33</td>
</tr>
</tbody>
</table>

**Figure 1.** Correlation between retinopathy of prematurity and gestational age

**Table 1.** Statistical analysis of the role of gestational age and birth weight on advanced retinopathy of prematurity
Discussion

In our study, the incidence of advanced ROP (stage 4 or 5) was 8.6% (N=91). This number consists of 3.8% of bilateral stage 5 (N=40) and 2.3% (N=24) of unilateral stage 5; these infants were considered as visually impaired or blind. Although high incidence of stage 4 or 5 of ROP has recently been reported from a tertiary eye hospital in China, most of the other recent studies have not reported the same findings. For example, one study has reported the incidence of advanced ROP to be 1.8% in infants born below 27 weeks of GA, and many other studies have not reported any advanced ROP in their settings. Specially in more mature infants. While better control of risk factors is a reason, performing regular screening programs which results in identification of ROP before formation of the advanced disease is another reason. However, the results of this study showed that we have a high percent of infants who come too late for retinal exam. The study from China also reported the same results about the time of screening exams, whereas a report from New Zealand showed that only one of 85 infant (1.2%) was examined after 9 weeks (at 11 weeks) and had stage 4 ROP.

In a study from our center, during 1999-2002, the infants with severe visual impairment due to ROP had a first eye examination at 25.1±13.8 weeks, and only 36% of them had neonatologists’ recommendation for eye examination. Comparison of these data with the results of the present study (27.5±31.7 and 23.1% respectively), may show that there has been no major improvement in the referral situation during the past decade in our country. This emphasizes the role of implementing a national ROP screening protocol, and to educate ophthalmologists, neonatologists and parents about this sight-threatening disease.

Presentation of ROP in infants with higher GA or BW in this study, suggests that there should be set a wider screening criteria for detecting premature infants with ROP in Iran, since UK and AAPPOS screening guidelines may not be applicable in our society.

In our study, lower GA was related with developing ROP, in addition, it was related with having severe or advanced ROP. Many studies have shown the role of lower GA in developing ROP; also there are studies which have shown the role of lower GA on progression of ROP to its severe form. BW also had significant correlation with development of ROP, the difference between mean BW in mild and severe ROP was also significant. However, BW had no significant difference between severe and advanced
ROP. Maybe because the infants usually gain a little weight in lower GAs, and they add more weight only in later GAs; thus, we can not see statistical difference in BW of infants with different stages of ROP who were born in lower GAs.

This was a unit-based retrospective study and there were some limitations in it. First, the incidence of advanced ROP in our study may not be representative of all of our country, as not all the infants eligible for ROP screening are screened in Iran, due to the lack of a uniformed guideline for their screening. Another reason for non-representation of our data is that some patients are referred to Farabi Eye Hospital just because they have advanced ROP; thus, the observed percent of advanced ROP is increased. Therefore, larger population-based studies about ROP are needed in Iran.

**Conclusion**

In summary, we observed a high incidence of ROP related visual impairment and blindness. Many of these infants did not have recommendation for eye examination and were examined too late. These findings indicate that we need a national screening program for retinopathy of prematurity in our country.

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**References**