Congenital cataracts and other disorders of visual deprivation can damage the developing visual system of a child, and permanently reduce central and peripheral vision. Therefore early diagnosis and surgery, appropriate refractive error correction, amblyopia therapy and long-term follow-up are essential. It is well known that recovery of normal visual function after cataract surgery is more probable in adults as compared to children due to impairment of the developing visual system.[2,3]

The prevalence of congenital cataract has been reported from 1 to 15 per 10,000 children worldwide, whereas it ranges from 1 to 3 per 10,000 births in developing countries.[4] The number of blind children due to congenital cataracts globally and in developing countries are 200,000 and 133,000, respectively.[5]

Red reflex examination at birth is an easy method to screen for congenital cataracts leading to early diagnosis and timely surgery.[6] The sensitivity and specificity of red reflex examination with no pupil dilation immediately after birth have been reported as 85 and 38.50%, respectively for detection of all types of cataracts.[6,7]

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of congenital ocular diseases,[7] thus pediatricians should be aware of the possibility of false positive results.[8]

Accompanying abnormalities such as ocular hypoplasia and early or late surgical complications such as inflammation and glaucoma may adversely affect visual outcomes in children undergoing congenital cataract surgery.[1]

Birch et al showed that late surgery may be considered as one of the leading causes of decreased best-corrected visual acuity (BCVA), and only 53% of cases with history of late congenital cataract surgery achieved BCVA of 0.60 LogMAR or better.[9]

According to the literature, the best visual outcomes may be achieved when surgery is performed during the first 6 weeks of age in unilateral cases and within 5 to 8 weeks of age in bilateral cases.[10] Furthermore, age at surgery and time interval between affected fellow eyes in developed countries have been reported less than one year and one week, respectively.[11‑14] However, in a recent study by the authors of the current perspective, these figures were 3 years and 3 months,[1] respectively. This difference may be explained by lack of routine early screening for media opacities in our neonates resulting in late diagnosis and late surgery, poor economic status, or a combination of the above-mentioned causes.

Based on the literature, mean BCVA is 0.9 (range, 0.2‑1.4) LogMAR in unilateral and 0.57 (range, 0.2‑1.0) LogMAR in bilateral congenital cataract subjects.[10,15,16] Our results were consistent with these findings (0.90 LogMAR in unilateral and 0.35 LogMAR in bilateral cases).[1] Since the prognosis for good VA in unilateral cases is disappointing as compared to bilateral cases, earlier diagnosis, surgery, and regular follow-up are mandatory.

Amblyopia is the major cause of visual impairment following congenital cataract surgery.[10,11] It was the reason for decreased vision in 76% of cases in the study by Ledoux et al[17] and was also observed in 56% of our cases[1] which can be due to unilateral cataract, late surgery, longer time interval between fellow eyes and lack of compliance with amblyopia therapy.

Strabismus following congenital cataract has been reported from 13% to 86% in the literature,[16,17] and was observed in 47.6% of our cases.[1] Ocular deviation in these children represents unequal reduced VA in their both eyes due to unilateral or asymmetric bilateral lens opacities.

Non-desirable surgical outcomes have been reported even in subjects with early cataract surgery, wearing appropriate glasses, contact lenses or implantation of intraocular lenses (IOL), which has been due to the lack of long-term follow-up (at least up to the age of 10 years) stressing the fundamental role of follow-up visits for management of possible complications in children with congenital cataracts.[9]

In the study by Rajavi et al,[1] risk factors for lower postoperative visual acuity included unilateral cataract, nystagmus, female gender (due to discrimination), strabismus and all types of congenital cataract except the zonular type. Furthermore, lack of parental compliance for treatment of amblyopia, longer surgical interval, and delayed surgery were other risk factors for not gaining an acceptable VA after the operation.[9,15,18,19]

Female gender is still considered as the second-grade sibling in the traditional families with the excuse of sons’ family responsibility in the future[1] therefore, parents are recommended to concern over the health of their children regardless of their gender.

Posterior capsular opacity (PCO) and glaucoma are major complications of cataract surgery in children.[10,17] Although the possibility of PCO is reduced by posterior capsulotomy and anterior vitrectomy during cataract surgery, glaucoma is still considered as a significant late complication in these cases requiring specific attention.

All practitioners, caregivers and parents should be aware of the following points:

1. Screening for red reflex opacities in all neonates is highly recommended, and suspected cases should be referred to a pediatric ophthalmologist for further examination. The importance and simplicity of performing this test makes it a feasible screening method.

2. Parents should be informed about the long therapeutic process and aware that cataract surgery is a starting and not the ending point of treatment for a child with congenital cataract.

3. Early cataract surgery, particularly in unilateral cases, is recommended (<6 weeks, based on the general neonatal health).[10] Contact lens fitting rapidly after surgery and IOL implantation at the appropriate age is also recommended.

4. Bilateral congenital cataracts should be operated before strabismus or nystagmus appears (<10 weeks)[10] with no longer than one-week interval between fellow eyes. Refractive error correction by aphakic glasses or contact lenses is recommended. IOL implantation at the appropriate age (>1 year) is suggested.

5. Due to the possibility of ocular deviation secondary to congenital lens opacity, all strabismic children should have slit lamp examination prior to strabismus surgery.

6. After congenital cataract surgery, amblyopia treatment should be started as soon as possible together with periodic follow-up examinations in order to achieve a satisfactory visual outcome.

7. The practitioners should consider the possibility of PCO, elevated IOP and amblyopia at each follow-up visit especially in cases with concurrent microphthalmia and/or associated congenital anomalies.
It is expected that equal and fair medical care should be provided to all children regardless of gender.

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There are no conflicts of interest.

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