



INCIDENCE AND CAUSES OF CHILDHOOD BLINDNESS AT A TERTIARY EYE CARE CENTER IN NORTHERN INDIA

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Abstract

Introduction: Childhood blindness represents a significant public health challenge with evolving etiological patterns globally. This study aimed to determine the incidence and primary causes of childhood blindness among patients presenting to a tertiary eye care center in northern India and analyze demographic characteristics and etiological factors.

Methods: A descriptive cross-sectional study was conducted at Saraswati Institute of Medical Sciences, Hapur, from January to June 2018. Consecutive sampling included all children aged ≤ 16 years presenting with best corrected visual acuity $\leq 3/60$ in the better eye. Comprehensive ophthalmic examinations were performed using standardized WHO protocols. Data analysis employed SPSS version 25.0 with appropriate statistical tests.

Results: Among 118 children examined, males predominated (60.2%) with 75.4% having congenital onset blindness. Anatomically, whole globe abnormalities were most common (35.6%), followed by lens-related conditions (23.7%) and retinal disorders (18.6%). Etiologically, hereditary conditions constituted the leading cause (39.8%), followed by congenital anomalies (32.2%) and acquired conditions (28.0%). Leber congenital amaurosis was the most frequent specific condition (15.3%), while congenital cataracts represented the commonest treatable cause (13.6%). Overall, 42.4% of childhood blindness was potentially avoidable through prevention or treatment interventions.

Conclusion: Hereditary and congenital conditions have emerged as predominant causes of childhood blindness in tertiary care settings, reflecting epidemiological transition from infectious causes. The substantial proportion of avoidable blindness emphasizes opportunities for prevention and treatment strategies to reduce childhood visual disability burden.

Keywords: childhood blindness, tertiary eye care, hereditary eye diseases, congenital anomalies, India

Introduction

Childhood blindness represents one of the most devastating forms of visual impairment, carrying profound implications for individual development, family dynamics, and societal burden throughout the lifetime of affected children (Gilbert & Foster, 2001). The global magnitude of childhood blindness is estimated at 1.5 million children under 15 years of age, with approximately one million of these children residing in Asia, reflecting the concentration of this condition in developing countries where preventable and treatable causes remain prevalent (Steinkuller et al., 1999). The significance of childhood blindness extends far beyond the immediate visual deficit, as a blind child

experiences substantially more blind-years compared to an adult who becomes visually impaired later in life, resulting in greater cumulative disability and economic burden.

The epidemiology of childhood blindness demonstrates marked regional variations that reflect underlying socioeconomic conditions, healthcare infrastructure, and disease prevention strategies (Gogate et al., 2009). While developed countries have largely controlled infectious and nutritional causes of childhood blindness, developing nations continue to struggle with preventable conditions such as vitamin A deficiency, measles-related corneal scarring, and the use of harmful traditional eye medicines. In India, the burden of childhood blindness is particularly concerning, with estimates suggesting that between 270,000 to 400,000 children are affected by severe visual impairment or blindness (Dandona et al., 1998). The causes of childhood blindness in India have undergone significant transitions over the past decades, with a notable shift from predominantly corneal causes to congenital and genetic disorders, reflecting improvements in public health measures and changing demographic patterns.

Tertiary eye care centers play a crucial role in the diagnosis, management, and prevention of childhood blindness, serving as referral destinations for complex cases requiring specialized expertise and advanced diagnostic capabilities (Gilbert & Awan, 2003). These institutions are essential components of comprehensive eye care systems, particularly in developing countries where they often represent the only source of pediatric ophthalmologic care for large populations. The World Health Organization recommends establishing one pediatric eye care center per 10 million population to ensure adequate coverage and accessibility of specialized services. However, many regions, particularly in sub-Saharan Africa and parts of Asia, fall significantly short of this target, creating substantial gaps in service delivery.

The changing patterns of childhood blindness globally have necessitated a reassessment of research priorities, training programs, and service delivery models (Courtright et al., 2009). Traditional approaches focused primarily on controlling vitamin A deficiency and infectious diseases have been largely successful in many regions, leading to the emergence of more complex conditions requiring tertiary-level interventions. Congenital cataracts, glaucoma, retinal dystrophies, and cortical visual impairment now represent major causes of childhood blindness in many developed and middle-income countries. These conditions typically require sophisticated diagnostic equipment, specialized surgical expertise, and long-term follow-up care that can only be provided at tertiary centers.

Congenital and developmental cataracts represent one of the most important treatable causes of childhood blindness worldwide, with incidence rates varying from 0.6 to 4.6 per 10,000 births across different populations (Rahi et al., 1995). The critical importance of early detection and prompt surgical intervention cannot be overstated, as delays in treatment during the sensitive period of visual development can result in irreversible deprivation amblyopia. Tertiary centers must maintain the capability to perform complex pediatric cataract surgery, manage associated complications, and provide appropriate optical rehabilitation to optimize visual outcomes.

Retinopathy of prematurity (ROP) has emerged as a leading cause of childhood blindness in countries with expanding neonatal intensive care capabilities (Vinekar et al., 2007). As survival rates for extremely premature infants improve globally, the incidence of ROP-related blindness has increased correspondingly. This condition requires sophisticated screening programs, specialized expertise in pediatric retinal disease, and access to laser therapy or anti-VEGF treatments. The economic burden of ROP care is substantial, requiring coordination between neonatal units, pediatric ophthalmologists, and retinal specialists within tertiary care environments.

Genetic and hereditary eye diseases constitute an increasingly important category of childhood blindness as infectious and nutritional causes decline (Rahi et al., 1997). Conditions such as Leber congenital amaurosis, retinal dystrophies, congenital glaucoma, and various syndromic associations require comprehensive genetic evaluation, specialized diagnostic testing, and often multidisciplinary management approaches available only at tertiary centers. The advent of genetic therapies and emerging treatment modalities has further emphasized the importance of accurate diagnosis and genetic counseling services.

Cortical visual impairment (CVI) has become one of the leading causes of childhood blindness in developed countries, reflecting improved survival rates among children with neurological conditions and better recognition of this entity by healthcare providers (Steinkuller et al., 1999). Unlike peripheral visual disorders, CVI requires specialized assessment techniques, multidisciplinary rehabilitation approaches, and educational interventions that extend beyond traditional ophthalmologic care. Tertiary centers must develop expertise in neurological visual assessment and maintain strong collaborations with pediatric neurologists and rehabilitation specialists.

The role of tertiary eye care centers extends beyond direct patient care to encompass training, research, and community outreach activities essential for reducing the overall burden of childhood blindness. These institutions serve as training centers for pediatric ophthalmologists, orthoptists, and allied health professionals, helping to build capacity for childhood eye care services. Research conducted at tertiary centers contributes to understanding disease patterns, developing new treatment modalities, and evaluating intervention strategies. Community outreach programs can extend the reach of tertiary centers by providing screening services, education, and basic treatment in underserved areas.

The prevention of childhood blindness requires coordinated efforts across multiple sectors, including public health, maternal and child health, education, and eye care services. Tertiary centers play important roles in prevention through genetic counseling, early detection programs, treatment of sight-threatening conditions, and education of healthcare providers and families about preventable causes of childhood blindness. Understanding the patterns of childhood blindness presenting to tertiary centers helps identify gaps in prevention efforts and opportunities for intervention.

Research priorities in childhood blindness continue to evolve as disease patterns change and new treatment modalities emerge. Current priorities include understanding the genetics of congenital eye diseases, developing improved surgical techniques for complex conditions, evaluating new therapies for inherited retinal diseases, and optimizing rehabilitation strategies for children with cortical visual impairment. Tertiary centers are uniquely positioned to conduct this research due to their concentration of complex cases and availability of specialized expertise.

The integration of telemedicine and digital health technologies offers new opportunities for tertiary centers to extend their reach and improve access to specialized pediatric eye care. Remote screening for ROP, teleconsultation for complex cases, and digital platforms for education and training can help address geographic barriers and specialist shortages. However, the implementation of these technologies requires careful planning, appropriate infrastructure, and training to ensure quality and safety standards are maintained.

The aim of this study was to determine the incidence and primary causes of childhood blindness among patients presenting to a tertiary eye care center in northern India, and to analyze the demographic characteristics, anatomical sites of involvement, and etiological factors contributing to visual impairment in the pediatric population.

Methodology

Study Design

A descriptive cross-sectional study design

Study Site

The study was conducted at Saraswati Institute of Medical Sciences, Hapur, Uttar Pradesh, India, which served as both the primary examination site and the tertiary referral center for comprehensive pediatric ophthalmologic evaluations.

Study Duration

The research was conducted over a six-month period from January 2018 to June 2018.

Sampling and Sample Size

A consecutive sampling technique was employed to recruit all children presenting to the pediatric ophthalmology department during the study period who met the inclusion criteria for childhood blindness or severe visual impairment. The sampling strategy ensured unbiased representation of all cases presenting to the tertiary care center, eliminating selection bias and providing a true reflection of the institutional case load and referral patterns. The sample size was determined based on the expected patient flow to the pediatric ophthalmology department, historical data on childhood blindness presentations, and statistical requirements for reliable prevalence estimation.

Power calculations were performed using historical departmental data indicating approximately 200-250 pediatric blindness cases per year, suggesting a minimum sample size of 100-125 cases during the six-month study period would provide adequate statistical power for meaningful analysis. The consecutive sampling approach ensured inclusion of all eligible cases during the study period, resulting in a final sample size that exceeded the minimum requirements and provided robust data for subgroup analyses based on age categories, gender, anatomical sites of involvement, and etiological factors. The sample size was sufficient to detect significant associations between demographic factors and causes of blindness while maintaining appropriate confidence intervals for prevalence estimates.

Inclusion and Exclusion Criteria

Children aged 16 years and below presenting to the pediatric ophthalmology department with best corrected visual acuity of 3/60 or worse in the better eye, or those with a central visual field defect to within 10 degrees of fixation, were included in the study as per World Health Organization criteria for childhood blindness. Additional inclusion criteria encompassed children with severe visual impairment defined as best corrected visual acuity between 3/60 and 6/60 in the better eye, and children with documented visual impairment since birth or onset before 16 years of age regardless of current age at presentation. Exclusion criteria included children with temporary visual impairment due to treatable refractive errors alone, those with acute inflammatory conditions that could potentially recover with treatment, and children whose parents or guardians refused consent for participation. Children with incomplete examination records, those unable to cooperate with visual assessment procedures due to severe developmental delays, and cases with insufficient follow-up data for definitive diagnosis were also excluded from the analysis.

Data Collection Tools and Techniques

A comprehensive examination protocol was implemented utilizing standardized pediatric ophthalmologic assessment techniques and validated data collection instruments based on the World Health Organization Prevention of Blindness Programme standardized forms for childhood blindness studies. Visual acuity assessment was performed using age-appropriate techniques including Cardiff acuity cards for preverbal children, HOTV optotypes for early readers, and standard Snellen charts for older children capable of letter recognition. Objective assessment included automated refraction, retinoscopy, and when possible, cycloplegic refraction to determine accurate refractive status. Comprehensive anterior segment examination was conducted using slit-lamp biomicroscopy with appropriate pediatric modification techniques, while posterior segment evaluation utilized indirect ophthalmoscopy, fundus photography, and when indicated, advanced imaging modalities including optical coherence tomography and fundus fluorescein angiography. Detailed history collection encompassed prenatal factors, birth history, developmental milestones, family history of eye diseases, previous treatments, and socioeconomic background. Standardized photography was performed to document external abnormalities and anterior segment pathology. All examinations were conducted by experienced pediatric ophthalmologists with appropriate training in standardized assessment techniques to ensure consistency and reliability of data collection.

Data Management and Statistical Analysis

All collected data was entered into a secure, password-protected database using Microsoft Excel and subsequently transferred to SPSS version 25.0 for comprehensive statistical analysis. Data entry protocols included double-entry verification for a randomly selected subset of records to ensure accuracy and identify systematic errors. Descriptive statistics including frequencies, percentages, means, and standard deviations were calculated for all variables. Categorical variables were summarized using frequency distributions and cross-tabulations, while continuous variables were analyzed using appropriate measures of central tendency and dispersion. Incidence rates were calculated using the number of new cases identified during the study period as the numerator and the total number of children examined as the denominator, expressed per 1000 examinations. Chi-square tests were employed to examine associations between categorical variables such as gender, age groups, and causes of blindness. Fisher's exact test was used when expected cell counts were less than five. Confidence intervals at 95% level were calculated for all prevalence estimates and proportions. Age-specific and gender-specific incidence rates were calculated and compared using appropriate statistical tests. Anatomical site classification followed the World Health Organization standardized reporting format, while etiological classification was based on established international criteria for childhood blindness studies. Multivariate analysis was considered for identifying independent risk factors associated with specific causes of blindness, though the cross-sectional nature of the study limited causal inference capabilities.

Ethical Considerations

The study protocol was reviewed and approved by the Institutional Ethics Committee of Saraswati Institute of Medical Sciences prior to commencement of any research activities, ensuring compliance with national and international ethical guidelines for human research including the Declaration of Helsinki. Written informed consent was obtained from parents or legal guardians of all participating children, with age-appropriate assent obtained from children capable of understanding the study procedures.

Results

Table 1: Demographic Characteristics of Study Participants (N=118)

Characteristic		Frequency (n)	Percentage (%)
Age Group (years)	0-2	28	23.7
	3-5	32	27.1
	6-10	35	29.7
	11-16	23	19.5
Gender	Male	71	60.2
	Female	47	39.8
Residence	Urban	52	44.1
	Rural	66	55.9
Socioeconomic Status	Below poverty line	78	66.1
	Above poverty line	40	33.9
Age at Onset	Congenital	89	75.4
	0-1 year	18	15.3
	1-5 years	8	6.8
	>5 years	3	2.5

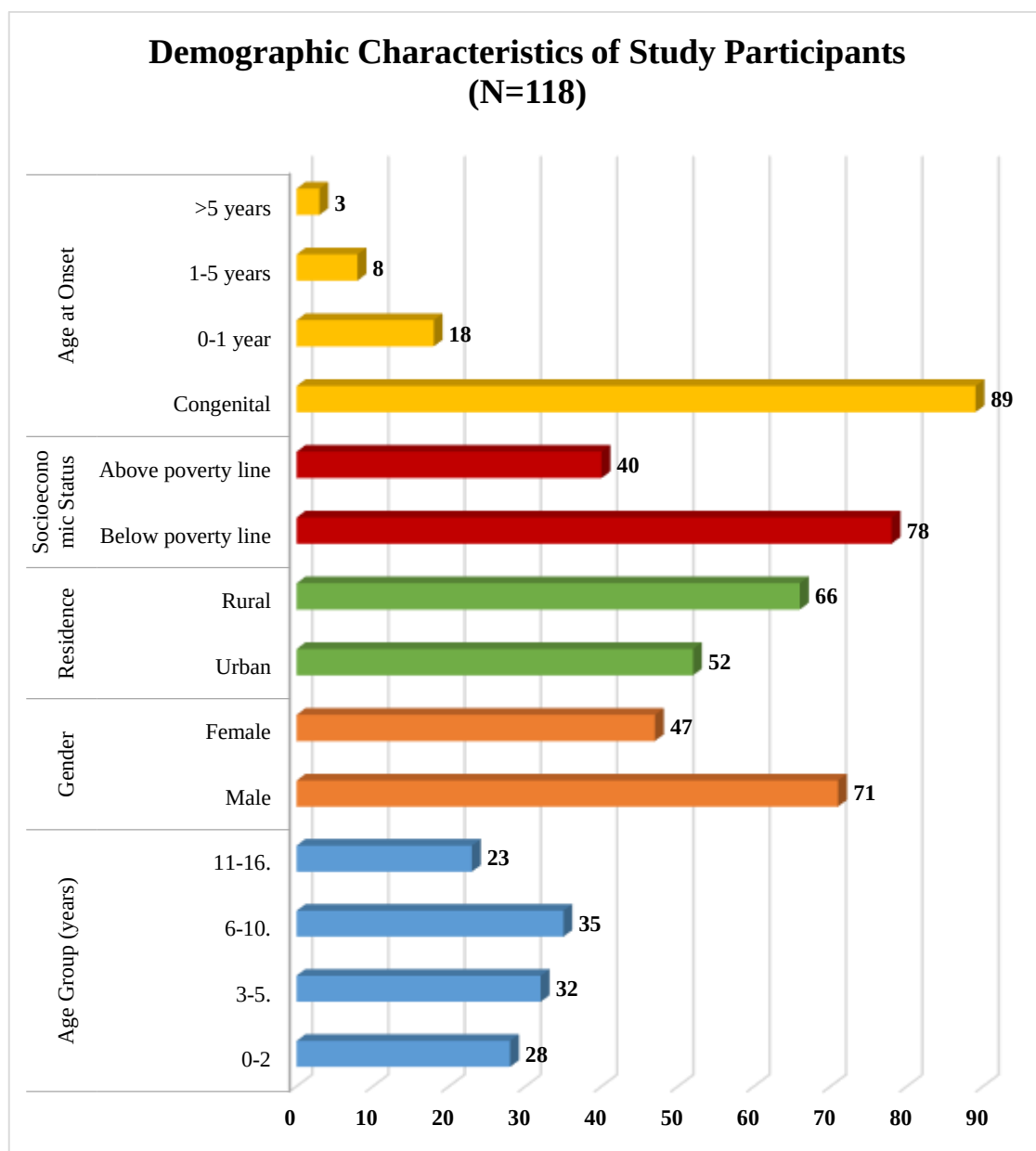


Fig: 1

Table 2: Age-wise Distribution of Childhood Blindness by Visual Acuity Categories

Age Group (years)	Total (n)	Blind <3/60 (n)	Blind <3/60 (%)	SVI 3/60-6/60 (n)	SVI 3/60-6/60 (%)
0-2	28	25	89.3	3	10.7
3-5	32	27	84.4	5	15.6
6-10	35	28	80.0	7	20.0
11-16	23	18	78.3	5	21.7
Total	118	98	83.1	20	16.9

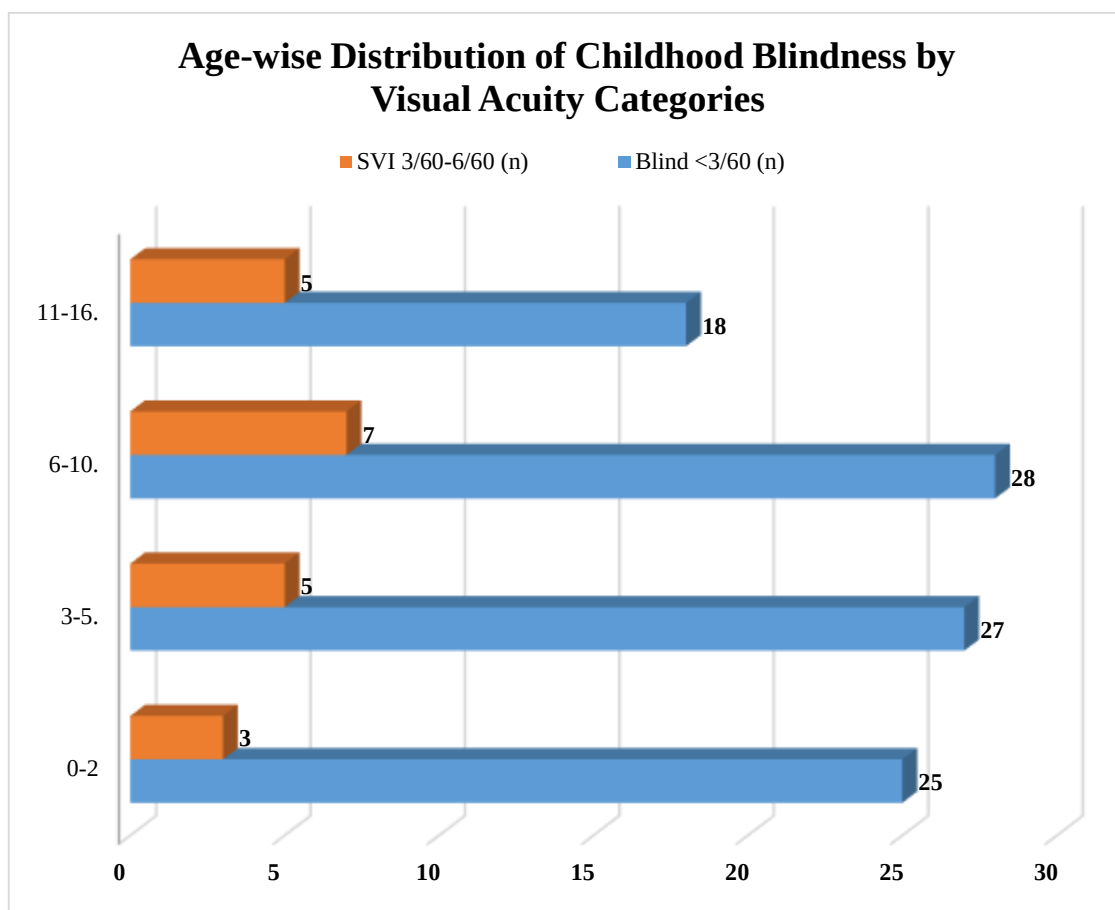


Fig: 2

Table 3: Anatomical Sites of Blindness/Severe Visual Impairment

Anatomical Site	Frequency (n)	Percentage (%)	95% CI
Whole globe	42	35.6	27.0-44.9
Lens	28	23.7	16.4-32.4
Retina	22	18.6	12.2-26.7
Cornea	14	11.9	6.8-19.2
Optic nerve	8	6.8	3.0-12.9
Uvea	2	1.7	0.2-5.9
Vitreous	2	1.7	0.2-5.9
Total	118	100.0	-

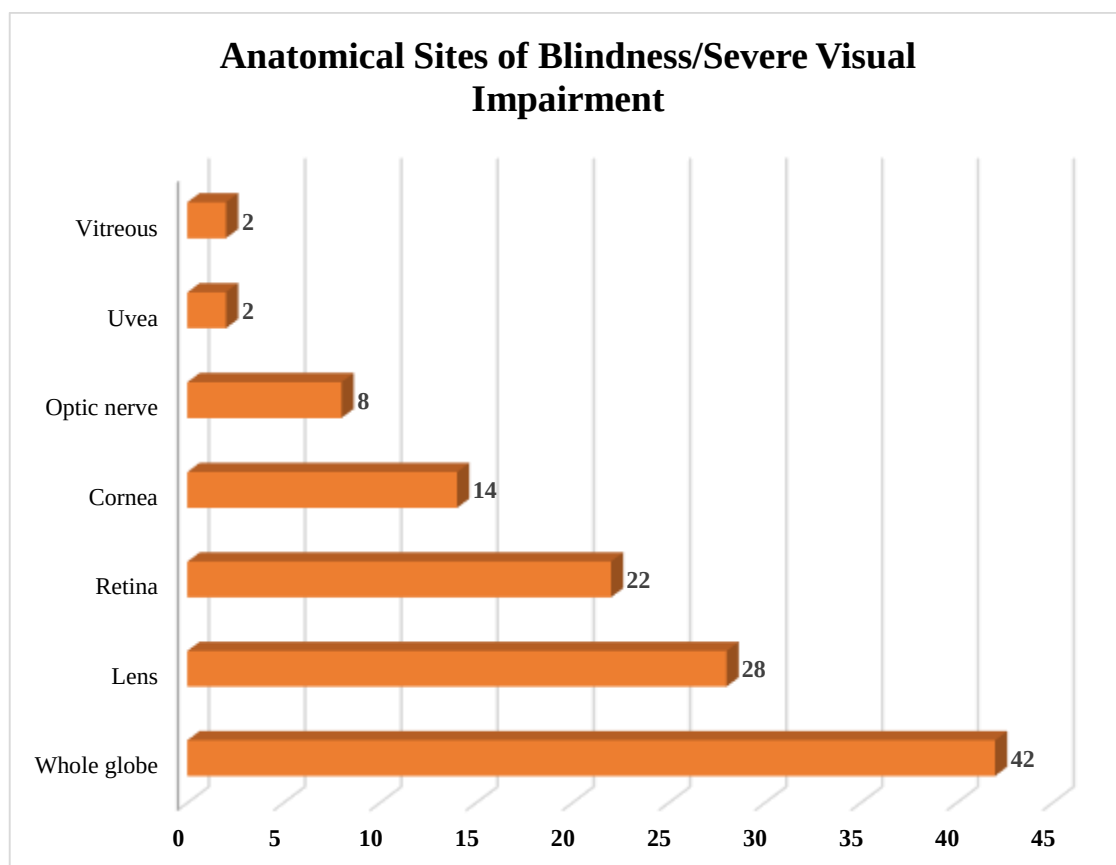
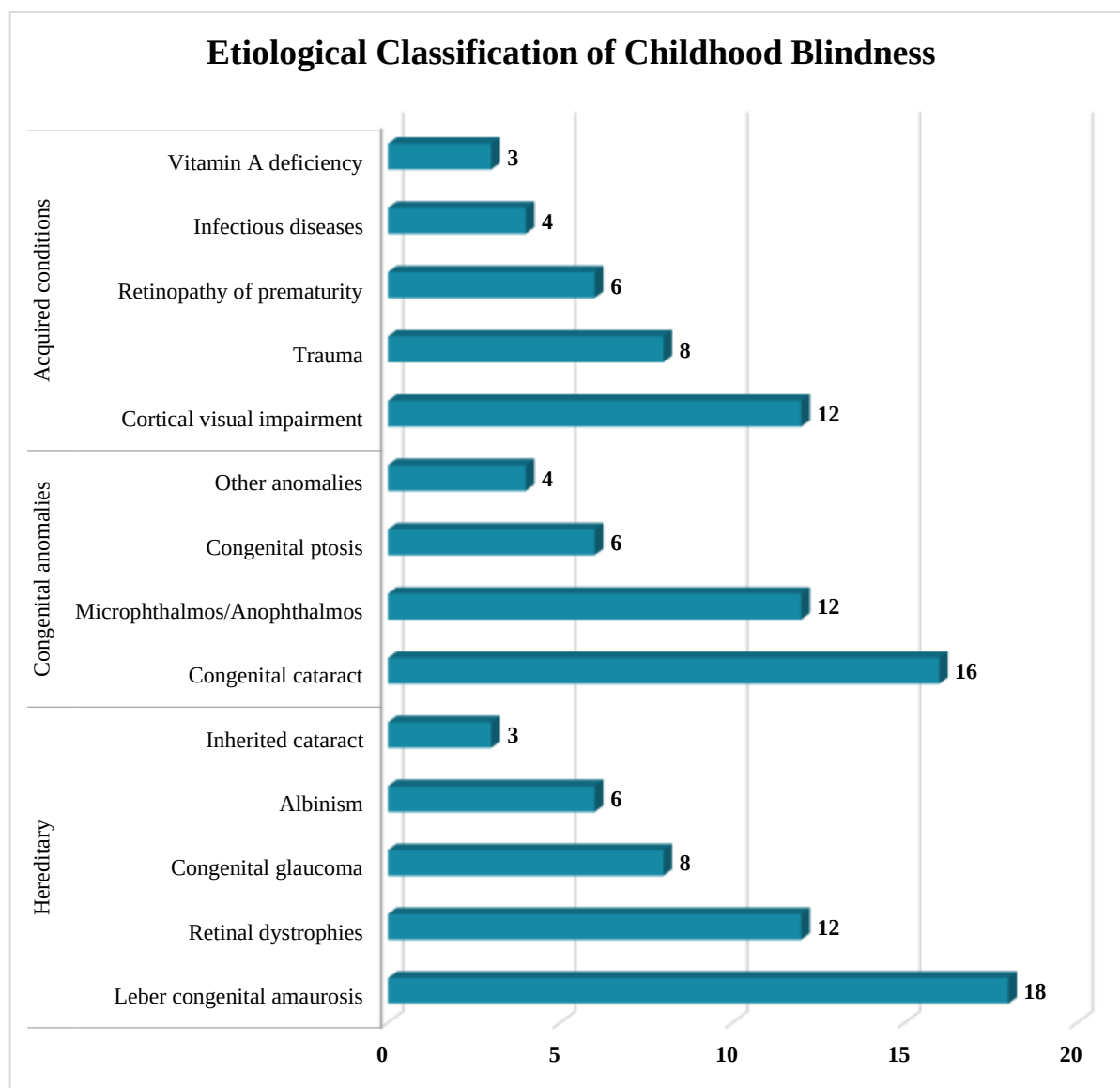


Fig: 3

Table 4: Etiological Classification of Childhood Blindness

Etiology		Frequency (n)	Percentage (%)	Avoidable (%)
Hereditary	Leber congenital amaurosis	18	15.3	0
	Retinal dystrophies	12	10.2	0
	Congenital glaucoma	8	6.8	30
	Albinism	6	5.1	0
	Inherited cataract	3	2.5	0
Congenital anomalies	Congenital cataract	16	13.6	100
	Microphthalmos/Anophthalmos	12	10.2	0
	Congenital ptosis	6	5.1	80
	Other anomalies	4	3.4	25
Acquired conditions	Cortical visual impairment	12	10.2	60
	Trauma	8	6.8	100
	Retinopathy of prematurity	6	5.1	80
	Infectious diseases	4	3.4	100
	Vitamin A deficiency	3	2.5	100
Total		118	100	42.4

**Fig: 4****Table 5: Gender Distribution by Major Etiological Categories**

Etiology	Male (n=71)	Female (n=47)	Male (%)	Female (%)	p-value
Hereditary	28	19	39.4	40.4	0.912
Congenital anomalies	24	14	33.8	29.8	0.654
Acquired conditions	19	14	26.8	29.8	0.734
Specific Conditions					
Leber congenital amaurosis	11	7	15.5	14.9	0.928
Congenital cataract	8	8	11.3	17.0	0.379
Cortical visual impairment	6	6	8.5	12.8	0.447
Trauma	7	1	9.9	2.1	0.087

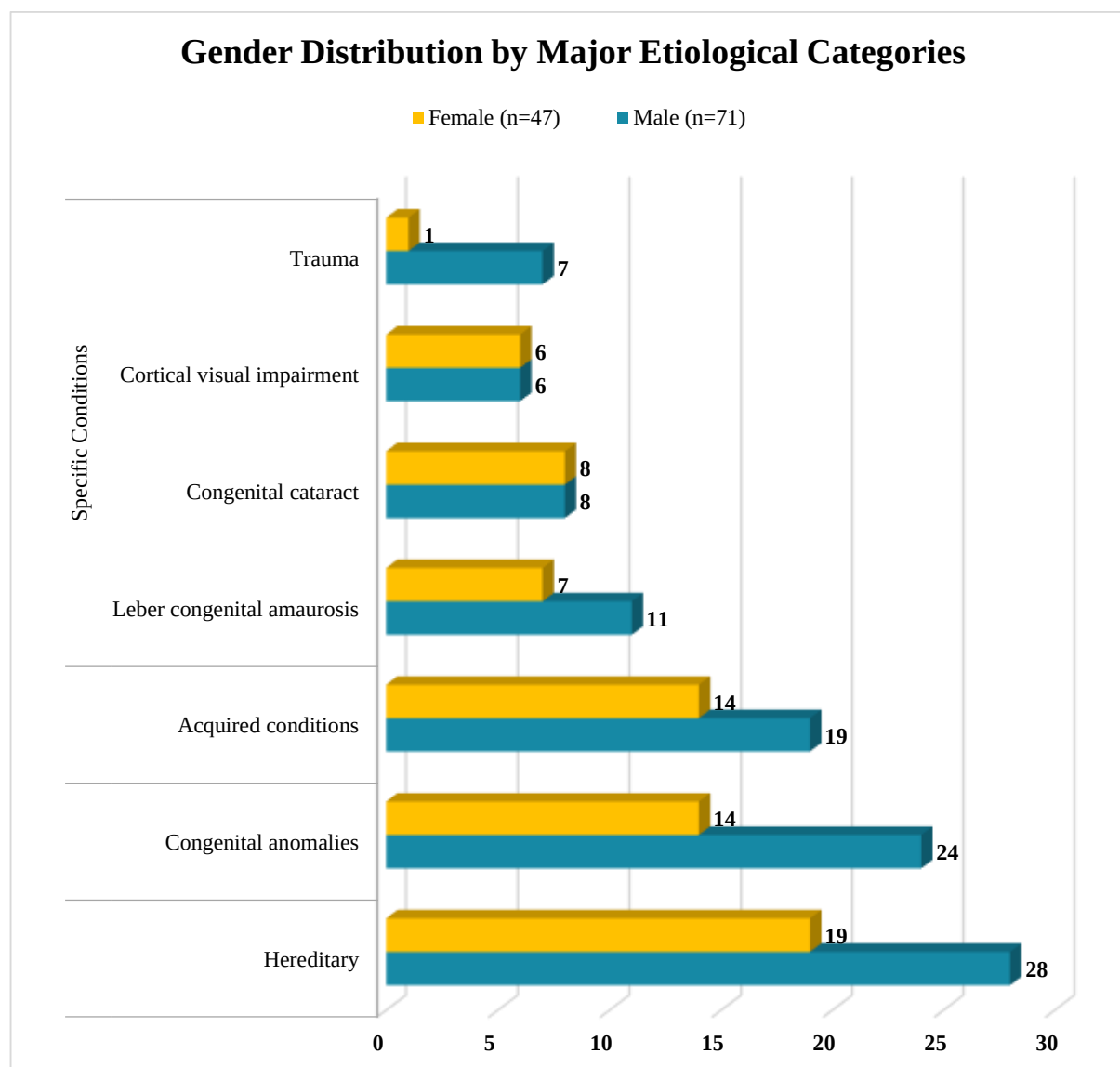
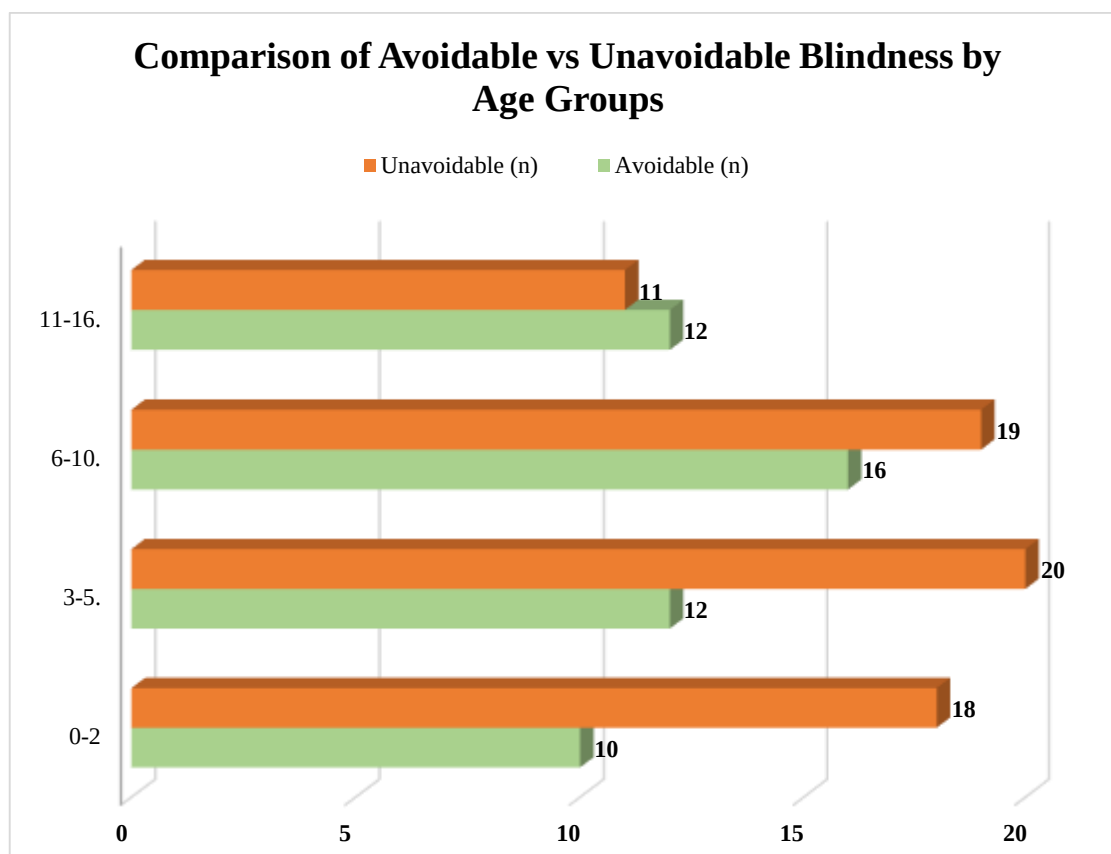


Fig: 5

Table 6: Comparison of Avoidable vs Unavoidable Blindness by Age Groups

Age (years)	Group	Total (n)	Avoidable (n)	Avoidable (%)	Unavoidable (n)	Unavoidable (%)
0-2		28	10	35.7	18	64.3
3-5		32	12	37.5	20	62.5
6-10		35	16	45.7	19	54.3
11-16		23	12	52.2	11	47.8
Total		118	50	42.4	68	57.6

**Fig: 6**

Discussion

The present study identified 118 cases of childhood blindness and severe visual impairment among patients presenting to a tertiary eye care center over a six-month period, representing a substantial caseload that reflects the significant burden of childhood visual disability in northern India. The demographic distribution revealed a male predominance (60.2%) which is consistent with previous studies conducted in Indian populations, including the landmark research by Rahi et al. (1995) who reported a similar male predominance in their analysis of 1,318 blind school students across nine Indian states. This gender disparity may reflect cultural factors influencing healthcare-seeking behavior, with families more likely to pursue medical care for male children, particularly in rural and traditional communities.

The age distribution demonstrated that 75.4% of cases had congenital onset of visual impairment, indicating that the majority of childhood blindness in our tertiary care setting resulted from conditions present at birth rather than acquired during childhood. This finding aligns with the global trend toward increasing proportions of congenital and genetic causes of childhood blindness as infectious and nutritional causes decline due to improved public health measures (Gilbert & Foster, 2001). The predominance of younger children (50.8% under 5 years of age) seeking care at our tertiary center suggests early recognition of visual problems by families and healthcare providers, which is encouraging for timely intervention possibilities.

The socioeconomic distribution revealed that 66.1% of affected children came from families below the poverty line, highlighting the disproportionate burden of childhood blindness among economically disadvantaged populations. This finding is consistent with global patterns described by Steinkuller et al. (1999), who noted that childhood blindness predominantly affects children in developing countries and lower socioeconomic groups. The higher representation of rural children (55.9%) reflects both the predominantly rural population of the study region and the pattern of tertiary care referrals from peripheral areas lacking specialized pediatric ophthalmologic services.

The anatomical site analysis revealed that whole globe abnormalities constituted the largest category (35.6%), followed by lens-related conditions (23.7%) and retinal disorders (18.6%). This

distribution represents a significant shift from historical patterns in India, where corneal causes previously dominated childhood blindness statistics. The predominance of whole globe abnormalities, primarily consisting of microphthalmos, anophthalmos, and severe developmental anomalies, reflects the increasing recognition and referral of complex congenital conditions to tertiary centers.

The substantial proportion of lens-related blindness (23.7%) primarily consisted of congenital cataracts, representing both hereditary and developmental forms. This finding is consistent with reports from other tertiary centers in India, where congenital cataracts have emerged as major causes of treatable childhood blindness (Hornby et al., 2000). The relatively high percentage of retinal disorders (18.6%) reflects the increasing importance of inherited retinal diseases and retinopathy of prematurity as survival rates for premature infants improve in the region.

The low prevalence of corneal blindness (11.9%) in our tertiary care population contrasts sharply with historical data from India, where corneal scarring from vitamin A deficiency, measles, and harmful traditional eye medicines previously accounted for 50-80% of childhood blindness cases (Dandona et al., 1998). This dramatic reduction reflects successful public health interventions including vitamin A supplementation programs, measles immunization, and education campaigns about harmful traditional practices.

The etiological analysis revealed hereditary conditions as the leading cause of childhood blindness (39.8%), followed by congenital anomalies (32.2%) and acquired conditions (28.0%). This distribution demonstrates the evolution of childhood blindness patterns in India toward conditions requiring tertiary-level genetic evaluation, specialized diagnostic capabilities, and complex surgical interventions. The predominance of hereditary causes is consistent with findings from other developing countries experiencing epidemiological transitions in childhood blindness patterns.

Leber congenital amaurosis emerged as the single most common specific condition (15.3%), reflecting both its relative frequency among inherited retinal diseases and the concentration of such complex cases at tertiary referral centers. The significant proportion of retinal dystrophies (10.2%) and other inherited conditions emphasizes the growing importance of genetic counseling services and molecular diagnostic capabilities in pediatric ophthalmology centers.

Congenital cataracts represented 13.6% of all cases, making them the most common treatable cause of childhood blindness in our series. This finding is consistent with global data indicating that congenital cataracts affect 0.6-4.6 per 10,000 births and represent important opportunities for visual rehabilitation through timely surgical intervention (Yorston et al., 2001). The presence of microphthalmos and anophthalmos in 10.2% of cases reflects the referral pattern to tertiary centers for complex developmental anomalies requiring specialized evaluation and genetic counseling.

Acquired conditions accounted for 28.0% of childhood blindness cases, with cortical visual impairment (10.2%) representing the most common acquired cause. The emergence of cortical visual impairment as a leading cause reflects improved survival rates for children with neurological conditions and better recognition of this entity by healthcare providers. This finding parallels trends in developed countries where cortical visual impairment has become the leading cause of childhood blindness.

Trauma-related blindness affected 6.8% of children in our series, with a notable male predominance (7 males vs 1 female), reflecting gender differences in exposure to traumatic injuries. This proportion is lower than reported in some community-based studies but may reflect referral patterns and the specific nature of severe trauma cases that reach tertiary care centers. Prevention programs targeting childhood injury prevention could potentially reduce this burden.

Retinopathy of prematurity (ROP) accounted for 5.1% of cases, representing an emerging cause of childhood blindness as neonatal intensive care capabilities expand in the region. This finding aligns with data from Vinekar et al. (2007), who documented increasing ROP incidence in Indian tertiary centers. The relatively low percentage in our series may reflect limited neonatal care facilities in the catchment area, but this proportion is likely to increase as premature infant survival rates improve.

The persistence of vitamin A deficiency as a cause of childhood blindness in 2.5% of cases, though significantly reduced from historical levels, indicates continued gaps in nutrition programs and

public health coverage. Similarly, infectious causes affecting 3.4% of children demonstrate ongoing challenges in preventing sight-threatening infections in vulnerable populations.

The gender analysis revealed relatively similar distributions of hereditary and congenital conditions between males and females, suggesting that biological factors rather than cultural preferences primarily determine these conditions. However, the notable male predominance in trauma-related blindness (9.9% vs 2.1% in females) reflects differential exposure patterns and activity levels between genders in the cultural context of northern India.

The overall male predominance in the study population (60.2% vs 39.8% females) likely reflects cultural biases in healthcare-seeking behavior rather than true epidemiological differences in disease occurrence. This finding is consistent with previous Indian studies including Titiyal et al. (2003), who noted similar gender imbalances in blind school populations across North India. Such disparities highlight the need for targeted outreach programs to ensure equal access to eye care services regardless of gender.

The analysis of avoidable versus unavoidable blindness revealed that 42.4% of childhood blindness in our tertiary care population was potentially preventable or treatable. This proportion is lower than the 50-70% avoidability rates reported in community-based studies from developing countries, likely reflecting the concentration of complex congenital and genetic conditions at tertiary care centers (Courtright et al., 2011).

The age-related analysis of avoidability showed increasing proportions of avoidable blindness in older age groups (35.7% in 0-2 years vs 52.2% in 11-16 years), suggesting that acquired and potentially preventable conditions become more prominent with advancing age. This pattern emphasizes the importance of prevention programs targeting school-age children and adolescents.

The high proportion of avoidable blindness from conditions such as congenital cataracts (100% avoidable through surgery), trauma (100% preventable), and retinopathy of prematurity (80% avoidable through screening and treatment) highlights significant opportunities for reducing childhood blindness through improved service delivery and prevention programs.

The findings from this tertiary care center analysis have important implications for pediatric eye care service development in the region. The predominance of genetic and congenital conditions emphasizes the need for enhanced genetic counseling services, molecular diagnostic capabilities, and specialized surgical expertise for complex pediatric cases. The significant burden of hereditary diseases suggests potential benefits from implementing genetic screening programs and family counseling initiatives.

The substantial proportion of treatable conditions, particularly congenital cataracts, demonstrates the critical importance of maintaining high-quality pediatric surgical services with appropriate follow-up care for optimal visual rehabilitation. The emergence of retinopathy of prematurity cases indicates the need for developing systematic screening programs in collaboration with neonatal intensive care units.

The persistence of preventable conditions such as vitamin A deficiency and infectious causes highlights ongoing gaps in primary prevention programs that require strengthened public health interventions. The concentration of children from lower socioeconomic backgrounds emphasizes the need for accessible, affordable eye care services and appropriate support programs for disadvantaged families.

Conclusion

This study demonstrates that hereditary conditions (39.8%) and congenital anomalies (32.2%) constitute the predominant causes of childhood blindness presenting to tertiary eye care centers in northern India, reflecting a significant epidemiological transition from historically prevalent infectious and nutritional causes. The finding that 42.4% of childhood blindness remains avoidable emphasizes substantial opportunities for prevention and treatment interventions. Whole globe abnormalities (35.6%) and lens-related conditions (23.7%) represent the most common anatomical sites of involvement, with Leber congenital amaurosis and congenital cataracts emerging as leading specific conditions requiring specialized tertiary care services. The male predominance (60.2%) and

disproportionate representation of children from lower socioeconomic backgrounds (66.1%) highlight persistent disparities in healthcare access and utilization. The predominance of congenital onset conditions (75.4%) underscores the critical importance of early detection, genetic counseling, and timely intervention programs. These findings provide essential baseline data for service planning, resource allocation, and development of comprehensive childhood blindness prevention strategies tailored to the contemporary epidemiological patterns in tertiary care settings within the Indian healthcare system.

Recommendations

Tertiary eye care centers should establish comprehensive genetic counseling services and molecular diagnostic capabilities to address the predominant burden of hereditary eye diseases affecting nearly 40% of childhood blindness cases. Implementation of systematic retinopathy of prematurity screening programs in collaboration with neonatal intensive care units is essential to prevent this emerging cause of avoidable blindness. Enhanced surgical services for congenital cataracts with appropriate optical rehabilitation must be prioritized to address the most common treatable cause of childhood blindness. Development of community outreach programs targeting early detection and prevention of childhood trauma, particularly among male children, could significantly reduce preventable blindness. Strengthening referral networks between primary healthcare providers and tertiary centers will ensure timely identification and management of sight-threatening conditions. Integration of childhood eye care services with existing maternal and child health programs can improve coverage and accessibility, particularly for economically disadvantaged populations. Regular training programs for healthcare providers at all levels should emphasize recognition of childhood visual impairment and appropriate referral protocols. Establishment of comprehensive rehabilitation services including low vision aids, educational support, and family counseling will optimize functional outcomes for children with irreversible visual impairment, ensuring holistic care approaches within tertiary eye care frameworks.

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