

BMJ Open

BMJ Open is committed to open peer review. As part of this commitment we make the peer review history of every article we publish publicly available.

When an article is published we post the peer reviewers' comments and the authors' responses online. We also post the versions of the paper that were used during peer review. These are the versions that the peer review comments apply to.

The versions of the paper that follow are the versions that were submitted during the peer review process. They are not the versions of record or the final published versions. They should not be cited or distributed as the published version of this manuscript.

BMJ Open is an open access journal and the full, final, typeset and author-corrected version of record of the manuscript is available on our site with no access controls, subscription charges or pay-per-view fees (<http://bmjopen.bmj.com>).

If you have any questions on BMJ Open's open peer review process please email info.bmjopen@bmj.com

BMJ Open

A phenomenological approach to childhood cataract treatment using semi-structured interviews: How might we improve provision of care?

Journal:	<i>BMJ Open</i>
Manuscript ID	bmjopen-2018-024869
Article Type:	Research
Date Submitted by the Author:	29-Jun-2018
Complete List of Authors:	Hamm, Lisa; University of Auckland, School of Optometry and Vision Science Boluk, Karla; University of Waterloo, Recreation and Leisure Studies Black, Joanna; University of Auckland, Optometry and Vision Science Dai, Shuan; University of Auckland School of Medicine, Ophthalmology; Auckland District Health Board, Ophthalmology Thompson, Benjamin; University of Waterloo, Optometry and Vision Science
Keywords:	Paediatric cataract, parental stress, semi-structured interviews, adherence

SCHOLARONE™
Manuscripts

Peer Review Only

Title: A phenomenological approach to childhood cataract treatment using semi-structured interviews: How might we improve provision of care?

Running head: Family experience of childhood cataract treatment

Word count: 4328

Authors:

Lisa Hamm¹, Karla Boluk², Joanna Black¹, Shuan Dai³, Benjamin Thompson^{1,4}

Affiliations:

¹ University of Auckland, School of Optometry and Vision Science

² University of Waterloo, Recreation and Leisure Studies

³ University of Auckland, Dept. Of Ophthalmology

⁴ University of Waterloo, School of Optometry and Vision Science

Corresponding author: Lisa Hamm
E: l.hamm@auckland.ac.nz, P: +64 (0) 9 923 9774
A: 89 Grafton Road, Grafton, Auckland, 1010, New Zealand

Author Contributions

LH, JB, SD and BT designed the study. LH and SD recruited the participants. LH conducted the interviews and transcribed the data. KB and LH independently coded and analysed the transcribed data, and then interpreted the data in an iterative manner. LH wrote the first draft the manuscript, and all authors assisted with revisions. All authors were involved in the final proofing process and agreed to be accountable for the content of the manuscript.

Acknowledgements

We would like to thank all the families who took the time to talk to us about thier experiences, as well as David Welch for his early contributions to the project.

Financial support: Lisa Hamm was supported by Education New Zealand. Education New Zealand had no role in data collection, interpretation or reporting.

1
2
3 31 **Conflict of interest:** None of the above authors have any proprietary interests or
4
5 32 conflicts of interest related to this submission.
6
7 33

8
9
10 34 **Keywords**

11 35 Paediatric cataract, parental stress, semi-structured interviews, adherence
12
13 36

14
15
16 37 **Data Statement**

17 38 Full transcripts of interviews are not available to protect participants' anonymity
18
19
20
21 39

40 Abstract

41 **Purpose:** To understand how we might improve the provision of medical care for children
42 with cataracts.

43 **Design:** A phenomenological design was employed. Semi-structured interviews were
44 conducted to capture rich descriptions of the phenomena. Our goal in the interview and the
45 analysis was to understand the sources of distress associated with treatment for cataract
46 and deprivation amblyopia which 1) could be addressed by the medical community and 2)
47 related to treatment adherence.

48 **Setting:** Interviews were conducted by a non-clinician researcher in New Zealand (NZ) in a
49 location chosen by informants. In NZ the red eye reflex screening test is performed shortly
50 after birth, and surgery to remove paediatric cataracts is publicly funded.

51 **Participants:** Families of children who had a history of cataract in Auckland, NZ were posted
52 an invitation to participate. Twenty families were interviewed.

53 **Results:** Our analysis illustrated that informants described a wide range of experiences,
54 from declined cataract surgery to full adherence to medical advice including years of
55 patching for more than four hours a day. Across these experiences, we identified three
56 relevant themes; delays in diagnosis, communication between the parent and clinician, and
57 parental social support networks.

58 **Conclusion:** The medical community may be better placed to support families dealing with
59 childhood cataract by improving detection of childhood cataract, building appropriate
60 communication pathways, and promoting social support, with an emphasis on empathetic,
61 individualised care.

62

1
2
3 63 **Article Summary**

4
5 64 **Strengths and limitations of this study**

6
7 65 *Strengths:*

- 8
9
10 66 • Informants had a broad range of experiences, from very positive to very negative,
11 likely to reflect the diversity of the cohort of interest.
12
13 68 • Data from interviews were rich and themes converged across diverse experiences.

14
15
16 69 *Limitations:*

- 17
18 70 • The impact of culture was not a specific focus of the interviews. Our iterative
19 analysis indicated that cultural factors may enhance understanding of families'
20 experiences. This is recommended as an area of further research.
21
22 72
23 73 • The authors are generally invested in adherence with ophthalmological
24 recommendations. This bias is clearly stated in our purpose.
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

76 Introduction

77 Elimination of visual impairment from childhood cataract is one of the key objectives of the
78 VISION2020 initiatives¹. The diagnosis and removal of a childhood cataract requires
79 specialised expertise and surgical facilities, and significant funding has been allocated
80 internationally to meet these needs. However, the vision loss associated with bilateral or
81 unilateral paediatric cataract is not fully resolved with surgical removal of the opaque lens
82 (for overview see^{2,3}). A replacement lens needs to be implanted, or compensatory contact
83 lens or spectacle correction prescribed. Children often require follow up surgeries and
84 ongoing assessment of appropriate refraction. Furthermore, the risk of deprivation
85 amblyopia increases sharply if the cataract is not removed shortly after onset^{4,5}. Amblyopia
86 is caused by abnormal visual cortex development due to disrupted visual experience and
87 impairs vision in the affected eye(s)⁶. Amblyopia due to congenital cataract is a rare, but its
88 effects on vision are severe due to pronounced visual deprivation at an early age^{4,5}. The
89 treatment for amblyopia involves occlusion (patching) or penalization (atropine cycloplegia)
90 of the stronger eye to promote use of the weaker eye. These are home-based therapies
91 which can be difficult for families to implement⁷. Therefore, the efficacy of childhood
92 cataract treatment is affected by a range of factors including early detection, prompt uptake
93 of surgical intervention, and dedicated commitment to follow up care by the family and
94 medical community (for discussion see⁸).

95 Each step in the treatment pathway for childhood cataract has been investigated, including
96 screening⁹⁻¹³, factors contributing to delayed utilization of surgical services¹⁴, parental stress
97 associated with childhood cataract treatment^{15,16} and compliance with amblyopia
98 treatment¹⁷⁻²¹. Across this predominantly quantitative international body of work, the
99 services that are available, affordable, and accepted by the community being studied vary
100 considerably. In this study, we were interested in the experience of childhood cataract in a
101 setting where advanced ophthalmological services are well established and publicly funded.
102 A qualitative study in the USA found that 'treatment' was one of 6 key themes decreasing
103 quality of life for children with a history of cataracts²². Our project compliments this by
104 focusing on families' lived experience (phenomenology) through the treatment pathway.
105 Our aim was to identify sources of distress which 1) could be addressed by the medical
106 community and 2) related to adherence with recommendations from their ophthalmologist.

107 **Methodology**

108 **Study context**

109 The study was conducted in Auckland, New Zealand (NZ), where comprehensive childhood
110 vision screening systems are in place (red eye reflex exam at birth and at the 6 week check,
111 pre and in school acuity checks [www.wellchild.org.nz/health-info-resources/health-](http://www.wellchild.org.nz/health-info-resources/health-topic/vision)
112 [topic/vision](http://www.wellchild.org.nz/health-info-resources/health-topic/vision)), specialised paediatric cataract surgery is publically funded and prioritised, and
113 subsidies are available for refractive correction and occlusion therapy for families who
114 require financial assistance or have particularly high visual needs. The study cohort was
115 culturally diverse and approximately a third had lived and received medical advice outside of
116 NZ. The first author carried out the semi-structured interviews. This author is a non-clinical
117 researcher (not involved in any of the participant's care), and it was made clear to all
118 informants that participation in the study would not impact the care they received.

119 **Patient and Public Involvement**

120 This project was a first step towards understanding how the medical community could
121 improve provision of treatment for childhood cataract; as such the project was to
122 understand their experiences and needs. The project was part of a larger study about the
123 consequences of²³, and potential for rehabilitation after²⁴ visual deprivation due to
124 childhood cataract, with this step focused on understanding patient experience of the
125 treatment pathway. A lay summary of results are available to all participants, and
126 information will also be disseminated at a community follow up meeting or 'hui'.
127

128 **Participant recruitment**

129 In phenomenological studies a purposive sampling strategy is supported in order to recruit
130 participants who have experienced the phenomenon under study²⁵. Accordingly, research
131 participants were intentionally contacted following a medical records search. Inclusion
132 criteria included history of a visually significant paediatric cataract (congenital,
133 developmental and traumatic cases) and a current age of at least four. Exclusion criteria
134 included severe developmental disorders or severe ocular disease unrelated to paediatric
135 cataract, noted on ophthalmological records. Invitation letters were posted to thirty-nine
136 parents resulting in seventeen positive responses, each of whom were invited to participate

137 in the study, including one parent of a non-verbal child with autism (autism was not noted
138 on the ophthalmological record). One additional family made contact independently; their
139 son had not had surgery to remove his cataract. Two additional families were made aware of
140 the project through their specialist. A total of twenty semi-structured interviews were
141 carried out. The study complied with the tenets of the declaration of Helsinki and was
142 approved by ethics committees from the University of Auckland and the Auckland District
143 Health Board. Participants provided informed consent, and children who were able provided
144 assent.

145 **Semi-Structured interviews**

146 Interviewing is the primary data collection strategy in phenomenological studies²⁵. As such,
147 semi-structured interviews were employed focussing on the lived experiences and personal
148 perceptions of the treatment process for childhood cataract and secondary deprivation
149 amblyopia. Interviews were conducted in a private location of the participant's choice and
150 audio recorded (except in one case due to excessive background noise). We requested that
151 both caregivers and the affected child were present, and we encouraged all attendees to
152 contribute to the conversation. Interviews were approximately one hour in length for each
153 family, but varied in an attempt to obtain rich data. Semi-structured interviews were
154 transcribed by the first author. Potentially identifiable information was removed or
155 anonymized, and pseudonyms were assigned (pseudonyms reflect most common NZ names
156 for birth year). The approximate timing of key treatment events were derived from each
157 interview (narrative summary) and used to provide context for understanding
158 phenomenology (lived experience).

159 **Phenomenology**

160 In line with a descriptive phenomenological approach²⁶⁻²⁸ our primary goal was to develop
161 an in-depth appreciation of each family's experience of childhood cataract treatment. Our
162 specific research interest was to understand sources of distress which 1) could be addressed
163 by the medical community and 2) related to adherence to medical recommendations.

164 Husserl's²⁶ approach to phenomenology, as made explicit by Giorgi²⁷, and summarised by
165 Wertz²⁸ includes four steps, which we carried out as follows. 1) *Open reading*; we set aside

166 theoretical and personal biases and attempted to understand each experience as a whole.
167 2) *Meaning units*; we divided each transcript into units of meaning or 'themes'. 3)
168 *Psychological reflection*; we reflected on the relevance of the themes to our research
169 interest, and how they related to each other. This was an iterative process including re-
170 reading, summarising, grouping together, and redefining meaning units to better reflect
171 informant experience. 4) *Structural understanding and description*; we consolidated relevant
172 emergent ideas, which we present as three themes and six sub themes. The first and second
173 author mutually carried out an analysis of the transcripts and then compared notes which
174 resulted in the final presentation of themes. We used SRQR reporting guidelines for
175 qualitative research²⁹.

176 Research questions are approached from the frame of our own experiences. Our framing
177 included that 1) a critique of family's responses to a challenging life event was outside our
178 scope and 2) we trusted the evidence suggesting that prompt removal of cataracts and
179 adherence with secondary treatments (such as patching) are in the best interest of the child.
180 We structured our research question to highlight our perspective rather than bracketing our
181 experience from our interpretation; we focused on what the *medical community* could do to
182 alleviate distress, particularly in ways that could promote *adherence*. In the process of
183 seeking to find meaning across the experiences of our informants, we tended towards a
184 constructivist epistemology; fundamentally believing that meaning is constructed rather
185 than discovered. Through iteration, we understood our constructed meaning to be best
186 exemplified in terms of negative and positive experiences of the emergent themes.
187 Therefore the themes presented reflect encouragement to families at their best, and
188 exasperated distress at their worst.

189

190 Results

191 Narrative summary

192 Informants had a wide range of experiences. Key treatment events are summarised as
193 cataract diagnosis, cataract removal, refractive correction and amblyopia treatment.
194 Timelines for each participant are presented in Figure 1, based on estimates from interview
195 data. Some participants had congenital cataracts which were present at birth, others had

1
2
3 196 cataracts which were detected or developed during early childhood (developmental), and
4 197 others had traumatic cataracts resulting from an eye injury. It is of note that some parents
5 198 were unsure whether their child had congenital or developmental cataracts.
6
7

8
9 199 There were no cases in which access to surgery after diagnosis limited care, however
10 200 perception of delayed detection of cataract was an important issue raised. We have
11 201 expanded on this in the first theme. Unlike most families who prioritized prompt cataract
12 202 removal, Emily and Benjamin's families both declined surgery initially. Both later took up the
13 203 services, expressing a mixture of frustration and regret. We explore these cases more within
14 204 the second and third themes.
15
16
17
18

19
20 205 The experience with home implementation of follow up treatment was also diverse.
21 206 Although no families noted cost of glasses, contacts or patches to be prohibitive, parents
22 207 expressed concern that acquiring these items and navigating subsidies was complicated and
23 208 could be stressful. Only children with very early surgery used contact lenses instead of an
24 209 intraocular lens (IOL). Of the six surgeries done before six months of age, half (Olivia, James
25 210 and Jack) remained aphakic and used a contact lenses rather than IOLs. Each of these
26 211 families described the use of the contact lenses as the most stressful part of their
27 212 experience, with two of the three families describing traumatic events related to inserting,
28 213 removing and/or losing contact lenses. These two families advocated for further support or
29 214 re-structuring of contact lens use.
30
31
32
33
34
35
36
37

38 215 Patching experience ranged from none (including children with bilateral cataracts who had
39 216 equal visual acuity loss in both eyes, and children with unilateral cataract who were given a
40 217 very poor prognosis), to four hours per day over several years (Olivia, Oliver, Grace and
41 218 Hannah). The most challenging aspect of care was described as patching, only superseded by
42 219 use of contact lenses in infancy in the small subset of children for whom it was required. For
43 220 those who used glasses, feedback was very positive, except for the one participant
44 221 prescribed glasses in her teenage years (Georgia), who felt uneasy with the aesthetic aspects
45 222 of wear.
46
47
48
49
50

51
52 223
53
54

55 224
56
57
58
59
60

225 **Figure 1** *Timeline of treatment pathway*

226

227 **Phenomenological summary**

228 Three themes and several sub-themes emerged from the phenomenological analysis related
229 to how the medical system might better mitigate parental distress and encourage
230 adherence. These are summarized in Table 1.

231

232 **Table 1. Summary of themes and sub-themes**

1. Improved detection and diagnosis
a) Red eye reflex exam at birth
b) Referrals and process
2. Communicating with medical community
a) Building rapport
b) Provision of information about cataract and deprivation amblyopia
c) Foresight about treatment pathway
3. Social support
a) Advice from social networks
b) Availability of emotional and practical support

233

234 **Improved detection and diagnosis**

235 The detection and diagnosis of a cataract was a complex experience for families. It elicited
236 predictable emotions such as disbelief, fear and uncertainty, but also varied experiences
237 related to the family's understanding of timing. If parents understood the timing of
238 diagnosis to be linked to their child's visual outcome, emotions related to perceived
239 efficiency of the screening test and processes surrounding referral were heightened. For
240 those parents who were informed about the cataract promptly, the negative emotions

241 related to an unexpected diagnosis were balanced by a sentiment of more solemn
 242 appreciation. By contrast, anger arose for parents if the medical community was perceived
 243 to be responsible for a delay. In Table 2 we highlight 2 subcategories which were the targets
 244 of the frustration or appreciation. Table 2A highlights cases in which a person perceived to
 245 be responsible for the red eye reflex screening test (most commonly the midwife) was
 246 directly targeted for appreciation or anger. Table 2B highlights the importance of the
 247 process whereby the concern was communicated between professionals, culminating in a
 248 diagnosis.

249 **Table 2. Improved detection and diagnosis.** Examples of positive (left column) and negative
 250 (right column) experiences relating to timing of diagnosis.
 251

A – Red eye reflex screening test

She's been a midwife for ages, and because of that experience, she picked up it. (Lily's mother)

We were pretty angry at our midwife because we felt that she should have picked that up. (Jack's mother)

B – Referrals and process

We got sent to a follow up...to test the red eye reflex in the left eye and basically look in detail because they were expecting the possibility of a cataract. So it was diagnosed pretty much right from the start, she would have been about two days old when it was found and diagnosed. (Olivia's mother)

She [Allied medical worker] put him on a wait list and it took until about 15 months, it was a long time, so I was thinking I guess it's not a big deal. ...when we saw [ophthalmologist] he basically outlined right from the beginning that there was no hope. I found it ... sorry ... there is still a bit of anger there. (Riley's mother)

252
 253 In the left column of Table 2 we can see examples of effective screening, referrals and
 254 following up on red flags. By contrast, the right column highlights cases where an
 255 opportunity existed for a cataract to be detected earlier, and was missed. These experiences
 256 suggest that any strategy to improve medical provision for childhood cataracts needs to
 257 start with improved training for screeners, both in terms of administering the test and

258 procedures for prompt referral. Anger with a representative of the medical community,
259 exemplified by Jack and Riley's mother's comments, but shared by many informants, had a
260 lasting impact. A parent's perception of a delay in diagnosis compromised trust in the
261 medical community, which hindered ongoing communication. This was particularly the case
262 if the specialist managing the child's ongoing care was seen to be associated with this initial
263 perceived error. Although experience with detection and diagnosis was an important part of
264 ongoing communication with the medical community, many other factors also contributed
265 to communication, these expanded on within the next theme.

266

267 ***Communicating with the medical community***

268 Once diagnosed, the relationship a family had with the specialist, and to a lesser degree, the
269 allied medical staff, was an important part of their experience, permeating across treatment
270 events. While some families lit up while describing the kindness and competence displayed
271 by the medical professionals they had come to know well, for others anger and frustration
272 expanded when reflecting on their interactions with the medical community.

273 Rapport between the parents and the specialist appeared to be particularly important, as it
274 formed the foundation of the ongoing relationship with the medical team. Example quotes
275 are presented in Table 3A, with the left panel containing an example of a positive
276 impression, and the right showing an example of a negative impression.

277 Parental knowledge about childhood cataract ranged widely between participant families.
278 Some families had a scientific background, and could understand and appreciate a detailed
279 explanation. Others had limited background knowledge, but effective communication
280 allowed sufficient understanding for the family to feel comfortable with recommendations.
281 Other parents did not feel as though their questions were addressed sufficiently. In Table
282 3B, we provide examples of two informants for whom understanding of the condition had
283 an influence on the uptake of surgery. Additionally, some families experienced significant
284 distress over misunderstandings, or gaps of communication. For example, Sophie and Lily's
285 mothers were distraught at the time of the interview about perceived connections between
286 behaviours during pregnancy and the cause of the cataract. Accordingly, it appeared

1
2
3 287 important for the specialist to understand what the relevant information was for each
4 288 family, and to communicate it at an appropriate level.
5
6

7 289 This was particularly poignant for communication about potential future events. Childhood
8 290 cataract has a long treatment pathway, much of which is dependent on parents to
9 291 implement. When families became exasperated with home based aspects of treatment,
10 292 those who were prepared in advance were not as discouraged by these setbacks, whereas
11 293 families who had not anticipated the challenges appeared to be quite distressed by them.
12 294 Furthermore, the lack of foresight fostered a distrust of the specialist if parents perceived
13 295 that information was withheld. Table 3C provides one example from a family who was
14 296 warned of the difficulties from the outset (and appreciated the foresight), compared to a
15 297 family who felt their specialist had not communicated information which would have been
16 298 useful.
17
18
19
20
21
22
23
24

25 299 Whether the communication between the medical community and the parent was
26 300 perceived as positive was individualized and nuanced, but across participants it included
27 301 aspects of rapport, clear communication about the condition and of upcoming potential
28 302 challenges.
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

304 **Table 3. Communicating with medical community.** Examples of positive (left column) and
 305 negative (right column) experiences relating to communication.
 306

A - Building rapport	
<i>We went in and met the paediatric ophthalmologist, and he was just a lovely, lovely man. Yeah, it was just great (Aiden's mother)</i>	<i>I don't want to go to [...] anymore. He's quite 'nothing can be done, nothing can be done' (Riley's mother)</i>
B - Provision of information about cataract and deprivation amblyopia	
<i>Asked whether specialist's description of the condition made sense to her: Yes, it did. Otherwise we could never have agreed to the surgery (Lily's mother)</i>	<i>I was avoiding the surgery from when she was three months onwards [...] they were not answering questions that I was asking to my satisfaction (Emily's mother)</i>
C - Foresight about the treatment pathway	
<i>When she was first diagnosed, the specialist sat us down with this one week old baby and said this is going to be a long, hard road (Olivia's mother)</i>	<i>Why hadn't (paediatric ophthalmologist) told us about this? We'd been going for three months! [...] It's not smooth, it's not transparent, and that's what's hard (Oliver's mother)</i>

307

308 **Social support**

309 Throughout the interviews, it became apparent that the provision of care was improved if
 310 there was an awareness of how recommendations fit within the family's social context. This
 311 includes perspectives or values held by an individual family and the resources available to
 312 them. It was common for families to include anecdotes about how friends or family
 313 members reinforced ideas, provided advice, helped pay for travel, supervised a child who
 314 was patching, looked after other children, or helped remove a contact out of an infant's eye.
 315 In many cases, such a support person happened to have a related occupation (for example,
 316 an optometrist, nurse, or physician) which provided extra comfort as parents navigated the

1
2
3 317 treatment pathway. When this was not available, it appeared to impact on parental
4 318 experience and capacity to comply with treatment.

5
6
7 319 Advice that parents received from friends or family was an important part of their
8
9 320 experience. When families were confronted by something new or unfamiliar, for example,
10 321 when the vision problem was first detected or surgery was offered, families who had others
11 322 encouraging them to engage with the medical system were likely to move forward with
12 323 recommendations (example in Table 4A - left). On the other hand, some social networks
13 324 were sceptical of the medical community, making it difficult or distressing for parents to
14 325 seek, or act on, medical advice. Families who felt tension between the advice from the
15 326 medical community and the advice from friends and family experienced an increased burden
16 327 during an already challenging time (Table 4A - right). Additionally, if this tension existed, and
17 328 the family member who was given the information did not have the decision making power
18 329 within the family (as was the case for Benjamin's family – further expanded on in Table 4B
19 330 right), communication from the medical community was less effective.

20
21
22 331 Beyond social networks overtly advising with or against medical advice, there was an
23 332 important emotional and practical role for this extended group. It was not specifically stated
24 333 that a phone call from a friend, or a ride from a family member directly mitigated distress or
25 334 delays along the treatment pathway, but the absence of this support appeared to be
26 335 associated with both. Emily's mother provided a particularly impactful example of this; she
27 336 was overwhelmed when surgery was recommended, and she declined it, but four years later
28 337 when she joined a community group allowing social support, she decided to comply with
29 338 recommendations (Table 4B - left). Like Emily's mother, several families simply felt
30 339 overwhelmed at diagnosis or during amblyopia treatment, and did not have the emotional
31 340 or practical support they needed to overcome this feeling. The example we provide in Table
32 341 4B – right, is from Benjamin's family, and it also highlights the inter-relationship with
33 342 practical support, advice, and communication. It is of particular note that contact lens use
34 343 required additional practical support; Olivia and Jack's families described requiring 2 or more
35 344 adults to extract the contact from their infant's eye.

345 As discussed within 'communication', the family's social context also varied widely across
 346 informants. It was further complicated with changes over time; a school or job change, for
 347 example, impacted the balance of needs and resources a family had.

348 **Table 4. Social Support.** Examples of positive (left column) and negative (right column)
 349 experiences relating to social support
 350

A – Advice from social networks

Her grandfather, who is a GP, saw that she was looking at things close and her eye was turning in. We took her to an optometrist and they picked up that there was something there, and they referred us to the ophthalmologist who said 'yes that looks like a cataract to me, we'll send you to [city]'. And that all happened really quickly (Grace's mother)

The doctor is trying to give us good advice [...] the sooner we get it done the better. And my mum and dad are pretty much saying 'No'. So that was another kinda difficulty [...] Yeah, an emotional, mental thing as well with the family's input – what they think about it (Lily's mother)

B – Availability of emotional and practical support

When describing decision to go through with the surgery four years after it was recommended: It was just talking about it I guess with the [Community worker] that I was having the get-togethers with [...] she was lovely. I would talk with her and we would talk with [child] and she would even offer to drive us out there and stuff like that. Because that was another issue, just the transport (Emily's mother)

We took him to the hospital and they were saying right then and there to do a little surgery [...] I was carrying [another child], and my husband didn't want him to have the surgery (Benjamin's mother)

351

352

353 The parent's perspective on the role they had within their child's care was an important
354 influencer of adherence. This perceived role was related to parenting strategies and belief
355 systems; factors less likely to be directly addressed by the medical community, and therefore
356 not a focus of our analysis. We note simply that parents who perceived the condition as
357 severe from the outset, and had the resources to both advocate for their child within the
358 medical community and be creative in their implementation of home based treatments,
359 tended to experience less distress and be more compliant with recommendations. Several
360 families expressed interest in some form of network which would allow families to share
361 these experiences and strategies directly.

362

363 Discussion

364 The goal of our descriptive phenomenological approach²⁶⁻²⁸ was to develop an in-depth
365 appreciation of the individual experience of childhood cataract treatment; with a specific
366 interest in understanding sources of distress which 1) could be addressed by the medical
367 community and 2) related to adherence to medical recommendations.

368 Strengths and Limitations

369 We were able to recruit a wide variety of participants, with a diverse range of experiences
370 despite childhood cataract being relatively rare. We were able to conduct rich interviews in
371 which we found some informants were very appreciative of the medical community's role in
372 their experience of medical treatment for childhood cataract, while others felt as though
373 they were let down by the medical community. This diversity, and our theoretical framework
374 allowed us to identify themes and construct meaningful summaries which could be
375 presented in positive and negative terms. In other words, for each potential factor we could
376 consider whether the lack impaired a family's experience, and the presence enhanced it. In
377 some cases this directly linked to whether a family adhered to medical recommendations. In
378 this way, the diversity of informant experience is likely to aid in quality and the
379 generalisability of our analysis.

1
2
3 380 However, there were some limitations in our design and analysis. During the interviews we
4 381 did not focus specifically on cultural frameworks for health. However, upon iterative
5 382 reflection it was our impression that a deep understanding of culture may have facilitated
6 383 our understanding of informant's experiences. Further expanding on this, perhaps a
7 384 symbolic interactionism theoretical perspective, would be a valuable direction for future
8 385 research in order to understand underlying cultural reasons for choices, particularly related
9 386 to declining available surgery. Furthermore, we started with the belief that adherence with
10 387 ophthalmological recommendations is the best for children. Therefore discussion around
11 388 whether a parent's decision to decline surgery or cease patching was in fact best for the
12 389 child was outside the scope of this project. Similarly, parental motivation was not a focus of
13 390 our analysis, but is part of the theoretical framework related to adherence³⁰. Following up
14 391 with participants on these points as part of our dissemination step will be a good
15 392 opportunity to enhance future research.

26 393 **Summary and Implications of findings**

27
28 394 The varied experiences of the diverse informants we interviewed revealed opportunities for
29 395 systemic improvement in childhood cataract treatment. Early detection and prompt medical
30 396 treatment are critical for good visual outcomes^{4 5}. Delayed detection was a perceived issue
31 397 for many of our informants. More effective implementation of the red eye reflex exam may
32 398 improve this situation^{12 31}. Survey data from practitioners in New Zealand reflected this
33 399 opportunity for improvement, specifically highlighting the value of better training for those
34 400 administering early eye exams¹³. Our work supports the value of such an initiative.

35
36
37 401 Addressing issues that are social rather than organizational is perhaps more complex. The
38 402 experiences expressed by our informants suggests that improved communication between
39 403 the patient and the health care community, and promoting strong social support networks
40 404 are particularly important.

41
42
43 405 These are not new ideas. For example, Shudy et al.³² thoughtfully use the term 'sense of
44 406 partnership with the staff' in their review of the impact of paediatric illness on a family,
45 407 which mirrors the first emergent theme in our analysis. Jackson et al.³³ discuss how close
46 408 social support networks improve coping in families of children with brain tumours, with
47 409 ideas closely related to our second theme of social support. Jackson et al. provide a

1
2
3 410 framework for conceptualizing social support, breaking it down into formal (medical) and
4 411 informal (friends/family), and use informational, emotional, and practical support as
5 412 qualifiers. Information is typically the domain of formal support, whereas the emotional
6 413 support most effectively comes from informal connections³³.

9
10 414 In our cohort, the impact of these factors was most dramatic in relation to uptake of surgical
11 415 services. Both families who declined surgery struggled to establish good communication with
12 416 a specialist, and did not have strong informal social support networks in place. The specific
13 417 concerns these families raised are also not new. Insufficient funds for travel or other indirect
14 418 costs, lack of information, and/or the parent targeted for education not having decisional
15 419 power are reasons presented in similar work in lower resource areas for delayed
16 420 presentation for available surgery¹⁴ or follow up³⁴.

17 421 In addition to these poignant cases, additional families in our cohort felt overwhelmed at
18 422 various stages of the treatment pathway, particularly when attempting patching and contact
19 423 lens wear. Contact lens use in infants (and to a lesser degree patching) has been associated
20 424 with higher levels of parental stress¹⁶. Parental stress is known to contribute to decreased
21 425 adherence with amblyopia treatment in cases of strabismic and anisometropic amblyopia²⁰.
22 426 These ideas appeared to be consistent with accounts from our informants. Again, good
23 427 communication and strong social support appeared to mitigate these factors. Although care
24 428 must be taken when generalizing, we suggest that perhaps communication and practical
25 429 support may have been factors which account for the difference between the results of
26 430 Drews et al¹⁶ which revealed an association between the use of contact lenses following
27 431 paediatric cataract surgery and parental stress and those of Celano et al¹⁵ which did not.
28 432 Notably, the data reported by Celano et al¹⁵ were collected in the context of a clinical trial, in
29 433 which extensive information and practical support were provided.

30 434 The participant families in this study had diverse backgrounds and needs, highlighting the
31 435 importance of empathetic and individualized care. Although there were many professionals
32 436 involved in patient care (for example nurses, orthoptists and low vision organizations),
33 437 paediatric ophthalmologists were generally seen as the primary providers of information.
34 438 The demand on a single specialist of providing sensitive, individualised support to a diverse
35 439 community may be significant.

1
2
3 440 In some cases print materials can help supplement communication of important information
4 441 and encourage treatment compliance¹⁹, however information alone is often insufficient to
5 442 promote action³⁵. Patient care may benefit from the provision of accurate information and
6
7 443 practical support from different professionals. Some research has described the role of a
8
9 444 'childhood blindness coordinator' to be particularly beneficial to fill this gap^{34 36}. This role can
10 445 involve helping parents understand the condition, providing foresight about the treatment
11 446 pathway, as well as providing practical support such as text message reminders and travel
12 447 planning³⁴. In terms of emotional support, formal networks tend to be less effective than
13 448 informal support networks³³. However, Ireys et al. showed the benefit of peer support
14 449 groups for mothers of children with chronic illness³⁷, and Emily's family provides a good
15 450 example of how a caring community volunteer can provide effective support. These are all
16 451 avenues which could be further explored to help families dealing with childhood cataract.

452 **Conclusion**

453 The message from the literature about childhood cataract is that without a team of
454 supportive professionals working with affected families, the pathway from detection to
455 rehabilitation will be difficult, and outcomes will be compromised^{2 3 8}. Our work supports
456 this, and emphasises that to achieve the best visual outcomes for these children, we need to
457 improve screening practices, communicate effectively and consider creative ways to support
458 families with surgical uptake and post-surgical follow up.

459

460 **References**

- 461 1. Gilbert C, Foster A. Childhood blindness in the context of VISION 2020 - The right to sight.
462 *B World Health Organ* 2001;79(3):227-32.
- 463 2. Medsinghe A, Nischal KK. Pediatric cataract: Challenges and future directions. *Clin*
464 *Ophthalmol* 2015;9:77-90. doi: 10.2147/OPTH.S59009
- 465 3. Oscar A, Veleva N, Chernodrinska V, et al. Childhood cataract. *Pediatrics* 2014;54(1):58-
466 61.
- 467 4. Birch EE, Cheng C, Stager Jr DR, et al. The critical period for surgical treatment of dense
468 congenital bilateral cataracts. *J AAPOS* 2009;13(1):67-71.
- 469 5. Birch EE, Stager DR. The critical period for surgical treatment of dense congenital
470 unilateral cataract. *Invest Ophthalmol Vis Sci* 1996;37(8):1532-38.
- 471 6. Holmes JM, Clarke MP. Amblyopia. *Lancet* 2006;367(9519):1343-51. doi: 10.1016/S0140-
472 6736(06)68581-4
- 473 7. Wang J. Compliance and patching and atropine amblyopia treatments. *Vision Res*
474 2015;114:31-40. doi: <http://dx.doi.org/10.1016/j.visres.2015.02.012>
- 475 8. Repka MX. Monocular infantile cataract: Treatment is worth the effort. *Arch Ophthalmol-*
476 *Chic* 2010;128(7):931-33. doi: 10.1001/archophthalmol.2010.130
- 477 9. Saiju R, Yun S, Yoon PD, et al. Bruckner red light reflex test in a hospital setting. *KUMJ*
478 2012;10(38):23-26.
- 479 10. Gräf M. Early detection of ocular disturbances in children. Brückner's transillumination
480 test (red reflex) - A must in childhood screening. *Deutsches Arz* 2007;104(11):724-29.
- 481 11. Shija F, Shirima S, Lewallen S, et al. Comparing key informants to health workers in
482 identifying children in need of surgical eye care services. *International Health*
483 2012;4(1):1-3.
- 484 12. Fry M, Wilson GA. Scope for improving congenital cataract blindness prevention by
485 screening of infants (red reflex screening) in a New Zealand setting. *J Paediatr Child*
486 *H* 2005;41(7):344-46. doi: 10.1111/j.1440-1754.2005.00628.x
- 487 13. Raoof N, Dai S. Red reflex screening in New Zealand: a large survey of practices and
488 attitudes in the Auckland region. *N Z Med J* 2016;129(1438):38-43. [published Online
489 First: 2016/07/23]
- 490 14. Bronsard A, Geneau R, Shirima S, et al. Why are children brought late for cataract
491 surgery? Qualitative findings from Tanzania. *Ophthalmic Epidemiol* 2008;15(6):383-
492 88.
- 493 15. Celano M, Hartmann EE, Drews-Botsch CD. Parenting stress in the infant aphakia
494 treatment study. *J Pediatr Psychol* 2013;38(5):484-93. doi: 10.1093/jpepsy/jst009
- 495 16. Drews C, Celano M, Plager DA, et al. Parenting stress among caregivers of children with
496 congenital cataracts. *J AAPOS* 2003;7(4):244-50. doi: 10.1016/S1091-8531(03)00118-
497 6
- 498 17. Drews-Botsch CD, Celano M, Kruger S, et al. Adherence to occlusion therapy in the first
499 six months of follow-up and visual acuity among participants in the infant Aphakia
500 treatment study (IATS). *Invest Ophthalmol Vis Sci* 2012;53(7):3368-75.
- 501 18. Drews-Botsch CD, Hartmann EE, Celano M. Predictors of adherence to occlusion therapy
502 3 months after cataract extraction in the Infant Aphakia Treatment Study. *J AAPOS*
503 2012;16(2):150-55.

- 1
2
3 504 19. Loudon SE, Fronius M, Looman CWN, et al. Predictors and a remedy for noncompliance
4 505 with amblyopia therapy in children measured with the occlusion dose monitor.
5 506 *Invest Ophthalmol Vis Sci* 2006;47(10):4393-400.
6 507 20. Loudon SE, Passchier J, Chaker L, et al. Psychological causes of non-compliance with
7 508 electronically monitored occlusion therapy for amblyopia. *Brit J Ophthalmol*
8 509 2009;93(11):1499-503. doi: 10.1136/bjo.2008.149815
9 510 21. Tjiam AM, Akcan H, Ziylan F, et al. Sociocultural and psychological determinants in
10 511 migrants for noncompliance with occlusion therapy for amblyopia. *Graef Arch Clin*
11 512 *Exp* 2011;249(12):1893-99.
12 513 22. Castañeda YS, Cheng-Patel CS, Leske DA, et al. Quality of life and functional vision
13 514 concerns of children with cataracts and their parents. *Eye (Basingstoke)*
14 515 2016;30(9):1251-59. doi: 10.1038/eye.2016.134
15 516 23. Hamm L, Chen Z, Li J, et al. Interocular suppression in children with deprivation
16 517 amblyopia. *Vision Res* 2017;133:112-20. doi: 10.1016/j.visres.2017.01.004
17 518 24. Hamm LM, Chen Z, Li J, et al. Contrast-balanced binocular treatment in children with
18 519 deprivation amblyopia. *Clin Exp Optom* 2018;101(4):541-52. doi: 10.1111/cxo.12630
19 520 25. Starks H, Brown Trinidad S. Choose Your Method: A Comparison of Phenomenology,
20 521 Discourse Analysis, and Grounded Theory. *Qualitative Health Research*
21 522 2007;17(10):1372-80. doi: 10.1177/1049732307307031
22 523 26. Husserl E. Ideas: General introduction to pure phenomenology. New York: Collier Books
23 524 1962.
24 525 27. Giorgi A. Psychology as a human science: A phenomenologically based approach. New
25 526 York: Harper & Row 1970.
26 527 28. Wertz FJ. A phenomenological psychological approach to trauma and resilience. Five
27 528 Ways of Doing Qualitative Analysis. New York: The Guildford Press 2011.
28 529 29. O'Brien BC, Harris IB, Beckman TJ, et al. Standards for reporting qualitative research: a
29 530 synthesis of recommendations. *Academic medicine : journal of the Association of*
30 531 *American Medical Colleges* 2014;89(9):1245-51. doi:
31 532 10.1097/acm.0000000000000388 [published Online First: 2014/07/01]
32 533 30. McGrady ME, Ryan JL, Brown GA, et al. Topical review: Theoretical frameworks in
33 534 pediatric adherence-promotion interventions: Research findings and methodological
34 535 implications. *Journal of Pediatric Psychology* 2015;40(8):721-26. doi:
35 536 10.1093/jpepsy/jsv025
36 537 31. Litmanovitz I, Dolfin T. Red reflex examination in neonates: The need for early screening.
37 538 *Israel Med Assoc J* 2010;12(5):301-02.
38 539 32. Shudy M, De Almeida ML, Ly S, et al. Impact of pediatric critical illness and injury on
39 540 families: A systematic literature review. *Pediatrics* 2006;118(SUPPL. 3):S203-S18. doi:
40 541 10.1542/peds.2006-0951B
41 542 33. Jackson AC, Enderby K, O'Toole M, et al. The role of social support in families coping with
42 543 childhood brain tumor. *J Psychosoc Oncol* 2009;27(1):1-24. doi:
43 544 10.1080/07347330802614634
44 545 34. Kishiki E, Van Dijk K, Courtright P. Strategies to improve follow-up of children after
45 546 surgery for cataract: Findings from Child Eye Health Tertiary Facilities in sub-Saharan
46 547 Africa and South Asia. *Eye (Basingstoke)* 2016;30(9):1234-41. doi:
47 548 10.1038/eye.2016.169
48 549 35. Blödt S, Kaiser M, Adam Y, et al. Understanding the role of health information in
49 550 patients' experiences: secondary analysis of qualitative narrative interviews with

- 1
2
3 551 people diagnosed with cancer in Germany. *BMJ Open* 2018;8(3) doi:
4 552 10.1136/bmjopen-2017-019576
5 553 36. Courtright P, Bowman R, Gilbert C, et al. Childhood cataract in Africa. 2008
6 554 37. Ireys HT, Chernoff R, DeVet KA, et al. Maternal outcomes of a randomized controlled
7 555 trial of a community-based support program for families of children with chronic
8 556 illnesses. *Arch Pediat Adol Med* 2001;155(7):771-77.
9
10 557
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

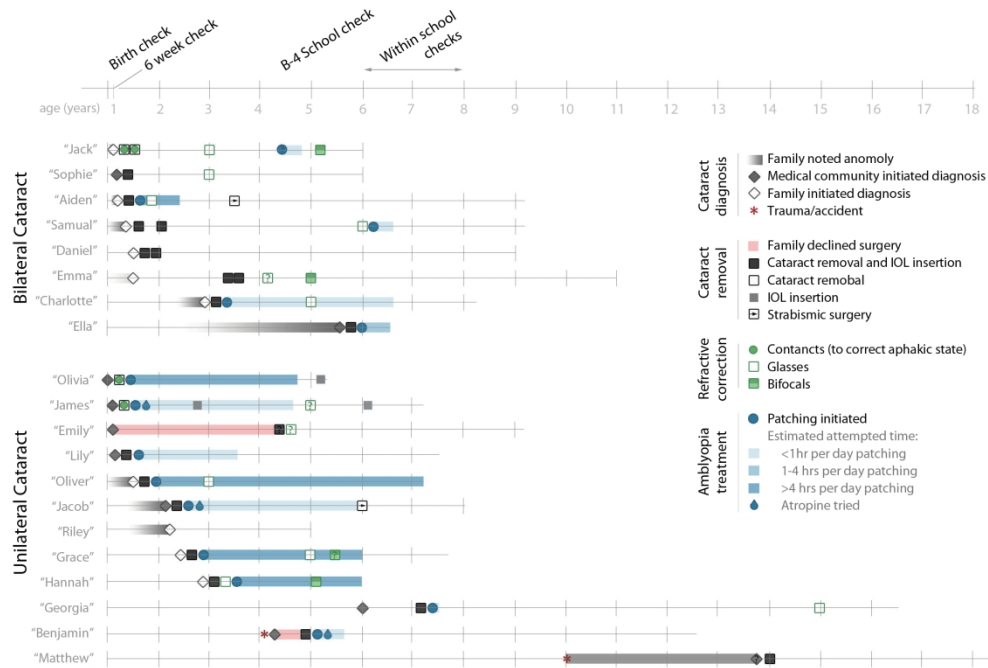


Figure 1 Timeline of treatment pathway

Participants are categorised into cases of unilateral or bilateral cataract. An asterisk by a patient pseudonym indicates a family history of paediatric cataract. Cataract diagnosis is depicted with a diamond. If a medical professional, or a community screen initiated the appointment in which a diagnosis occurred, the diamond is grey, whereas the diamond is white if the family initiated the appointment at which the diagnosis was made. A grey bar preceding a diamond indicates that abnormal behaviour or ocular appearance was noticed by the family prior to diagnosis. Treatment is represented by a square; black represents a surgical intervention, and green optical. Red bars between diagnosis and treatment indicate treatment was recommended during this time, but the family declined. IOL = intraocular lens

939x647mm (72 x 72 DPI)

Reporting checklist for qualitative study.

Based on the SRQR guidelines.

Instructions to authors

Complete this checklist by entering the page numbers from your manuscript where readers will find each of the items listed below.

Your article may not currently address all the items on the checklist. Please modify your text to include the missing information. If you are certain that an item does not apply, please write "n/a" and provide a short explanation.

Upload your completed checklist as an extra file when you submit to a journal.

In your methods section, say that you used the SRQR reporting guidelines, and cite them as:

O'Brien BC, Harris IB, Beckman TJ, Reed DA, Cook DA. Standards for reporting qualitative research: a synthesis of recommendations. *Acad Med.* 2014;89(9):1245-1251.

		Reporting Item	Page Number
Title	#1	Concise description of the nature and topic of the study identifying the study as qualitative or indicating the approach (e.g. ethnography, grounded theory) or data collection methods (e.g. interview, focus group) is recommended	1
	#2	Summary of the key elements of the study using the abstract format of the intended publication; typically includes background, purpose, methods, results and conclusions	3
Problem formulation	#3	Description and significance of the problem / phenomenon studied: review of relevant theory and empirical work; problem statement	5
Purpose or research question	#4	Purpose of the study and specific objectives or questions	5
Qualitative approach and research paradigm	#5	Qualitative approach (e.g. ethnography, grounded theory, case study, phenomenology, narrative research) and	7-8

guiding theory if appropriate; identifying the research paradigm (e.g. postpositivist, constructivist / interpretivist) is also recommended; rationale. The rationale should briefly discuss the justification for choosing that theory, approach, method or technique rather than other options available; the assumptions and limitations implicit in those choices and how those choices influence study conclusions and transferability. As appropriate the rationale for several items might be discussed together.

1			
2			
3			
4			
5			
6			
7			
8			
9			
10			
11			
12			
13			
14	Researcher	#6	8
15	characteristics and		
16	reflexivity	Researchers' characteristics that may influence the research, including personal attributes, qualifications / experience, relationship with participants, assumptions and / or presuppositions; potential or actual interaction between researchers' characteristics and the research questions, approach, methods, results and / or transferability	
17			
18			
19			
20			
21			
22			
23			
24			
25	Context	#7	6
26		Setting / site and salient contextual factors; rationale	
27			
28	Sampling strategy	#8	6
29		How and why research participants, documents, or events were selected; criteria for deciding when no further sampling was necessary (e.g. sampling saturation); rationale	
30			
31			
32			
33			
34			
35	Ethical issues pertaining	#9	7
36	to human subjects	Documentation of approval by an appropriate ethics review board and participant consent, or explanation for lack thereof; other confidentiality and data security issues	
37			
38			
39			
40	Data collection methods	#10	7
41		Types of data collected; details of data collection procedures including (as appropriate) start and stop dates of data collection and analysis, iterative process, triangulation of sources / methods, and modification of procedures in response to evolving study findings; rationale	
42			
43			
44			
45			
46			
47			
48			
49			
50	Data collection	#11	7
51	instruments and	Description of instruments (e.g. interview guides, questionnaires) and devices (e.g. audio recorders) used for data collection; if / how the instruments(s) changed over the course of the study	
52	technologies		
53			
54			
55			
56			
57	Units of study	#12	6-7 and
58		Number and relevant characteristics of participants, documents, or events included in the study; level of	8-9
59			
60			

participation (could be reported in results)

(also
Figure 1)

1			
2			
3			
4	Data processing	#13 Methods for processing data prior to and during analysis, including transcription, data entry, data management and security, verification of data integrity, data coding, and anonymisation / deidentification of excerpts	6-7
5			
6			
7			
8			
9			
10			
11	Data analysis	#14 Process by which inferences, themes, etc. were identified and developed, including the researchers involved in data analysis; usually references a specific paradigm or approach; rationale	7
12			
13			
14			
15			
16			
17			
18	Techniques to enhance trustworthiness	#15 Techniques to enhance trustworthiness and credibility of data analysis (e.g. member checking, audit trail, triangulation); rationale	8
19			
20			
21			
22			
23	Syntheses and interpretation	#16 Main findings (e.g. interpretations, inferences, and themes); might include development of a theory or model, or integration with prior research or theory	Table 1
24			
25			
26			
27			
28	Links to empirical data	#17 Evidence (e.g. quotes, field notes, text excerpts, photographs) to substantiate analytic findings	Tables 2-4
29			
30			
31			
32	Intergration with prior work, implications, transferability and contribution(s) to the field	#18 Short summary of main findings; explanation of how findings and conclusions connect to, support, elaborate on, or challenge conclusions of earlier scholarship; discussion of scope of application / generalizability; identification of unique contributions(s) to scholarship in a discipline or field	17-20
33			
34			
35			
36			
37			
38			
39			
40			
41			
42	Limitations	#19 Trustworthiness and limitations of findings	17-18
43			
44	Conflicts of interest	#20 Potential sources of influence of perceived influence on study conduct and conclusions; how these were managed	8 and 18
45			
46			
47			
48			
49			
50	Funding	#21 Sources of funding and other support; role of funders in data collection, interpretation and reporting	1
51			
52			
53			

The SRQR checklist is distributed with permission of Wolters Kluwer © 2014 by the Association of American Medical Colleges. This checklist can be completed online using <https://www.goodreports.org/>, a tool made by the [EQUATOR Network](#) in collaboration with [Penelope.ai](#)

BMJ Open

A phenomenological approach to childhood cataract treatment in New Zealand using semi-structured interviews: How might we improve provision of care?

Journal:	<i>BMJ Open</i>
Manuscript ID	bmjopen-2018-024869.R1
Article Type:	Research
Date Submitted by the Author:	24-Oct-2018
Complete List of Authors:	Hamm, Lisa; University of Auckland, School of Optometry and Vision Science Boluk, Karla; University of Waterloo, Recreation and Leisure Studies Black, Joanna; University of Auckland, Optometry and Vision Science Dai, Shuan; University of Auckland School of Medicine, Ophthalmology; Auckland District Health Board, Ophthalmology Thompson, Benjamin; University of Waterloo, Optometry and Vision Science
Primary Subject Heading:	Ophthalmology
Secondary Subject Heading:	Paediatrics, Patient-centred medicine, Qualitative research
Keywords:	Paediatric cataract, parental stress, semi-structured interviews, adherence

SCHOLARONE™
Manuscripts

Title: A phenomenological approach to childhood cataract treatment in New Zealand using semi-structured interviews: How might we improve provision of care?

Running head: Family experience of childhood cataract treatment

Word count: 4328

Authors:

Lisa Hamm¹, Karla Boluk², Joanna Black¹, Shuan Dai³, Benjamin Thompson^{1,4}

Affiliations:

¹ University of Auckland, School of Optometry and Vision Science

² University of Waterloo, Recreation and Leisure Studies

³ University of Auckland, Dept. Of Ophthalmology

⁴ University of Waterloo, School of Optometry and Vision Science

Corresponding author: Lisa Hamm
E: l.hamm@auckland.ac.nz, P: +64 (0) 9 923 9774
A: 89 Grafton Road, Grafton, Auckland, 1010, New Zealand

Author Contributions

LH, JB, SD and BT designed the study. LH and SD recruited the participants. LH conducted the interviews and transcribed the data. KB and LH independently coded and analysed the transcribed data, and then interpreted the data in an iterative manner. LH wrote the first draft the manuscript, and all authors assisted with revisions. All authors were involved in the final proofing process and agreed to be accountable for the content of the manuscript.

Acknowledgements

We would like to thank all the families who took the time to talk to us about their experiences, as well as David Welch for his early contributions to the project.

1
2
3 28 **Financial support:** Lisa Hamm was supported by Education New Zealand. Education
4
5 29 New Zealand had no role in data collection, interpretation or
6
7 30 reporting.

8
9 31 **Conflict of interest:** None of the above authors have any proprietary interests or
10
11 32 conflicts of interest related to this submission.

13
14 33
15
16 34 **Keywords**

17
18
19 35 Paediatric cataract, parental stress, semi-structured interviews, adherence
20
21 36

22
23 37 **Data Statement**

24
25
26 38 Full transcripts of interviews are not available to protect participants' anonymity
27
28
29 39

40 Abstract

41 **Purpose:** To understand how we might improve the provision of medical care for children with
42 cataracts.

43 **Design:** A phenomenological design was employed. Semi-structured interviews were conducted
44 to capture rich descriptions of the phenomena. Our goal in the interview and the analysis was
45 to understand the sources of distress associated with treatment for cataract and deprivation
46 amblyopia which 1) could be addressed by the medical community and 2) related to treatment
47 adherence.

48 **Setting:** Interviews were conducted by a non-clinician researcher in New Zealand (NZ) in a
49 location chosen by informants. In NZ the red reflex screening test is performed shortly after
50 birth, and surgery to remove paediatric cataracts is publicly funded.

51 **Participants:** Families of children who had a history of cataract in Auckland, NZ were posted an
52 invitation to participate. Twenty families were interviewed.

53 **Results:** Our analysis illustrated that informants described a wide range of experiences, from
54 declined cataract surgery to full adherence to medical advice including years of patching for
55 more than four hours a day. Across these experiences, we identified three relevant themes;
56 timing of diagnosis, communication between the parent and clinician, and parental social
57 support networks.

58 **Conclusion:** The medical community may be better placed to support families dealing with
59 childhood cataract by improving detection of childhood cataract, building appropriate
60 communication pathways, and promoting social support, with an emphasis on empathetic,
61 individualised care.

62

63 Article Summary

64 Strengths and limitations of this study

65 *Strengths:*

- 66 • Informants had a broad range of experiences, from very positive to very negative, likely
67 to reflect the diversity of the cohort of interest.
- 68 • Data from interviews were rich and themes converged across diverse experiences.

69 *Limitations:*

- 70 • The impact of culture was not a specific focus of the interviews. Our iterative analysis
71 indicated that cultural factors may enhance understanding of families' experiences. This
72 is recommended as an area of further research.
- 73 • The authors are generally invested in adherence with ophthalmological
74 recommendations. This bias is clearly stated in our purpose.

75

76 Introduction

77 Elimination of visual impairment from childhood cataract is one of the key objectives of the
78 VISION2020 initiatives^{1,2}. The removal of the opaque lenses preventing a child from seeing
79 requires specialised expertise, surgical facilities and various consumables, at an estimated base
80 cost of approximately \$300 USD³⁻⁵ in the developing countries to an estimated \$5000 USD in
81 developed countries such as New Zealand. Although childhood cataracts are rare (impacting
82 approximately 2.5 in 10 000 children⁶), the investment in treatment is advocated for globally
83 due to the improvements in quality of life⁷ and long term economic benefits⁴. Given the impact,
84 removal of childhood cataracts is prioritized, with many non-governmental organizations
85 (NGOs) supporting communities unable to make this investment⁸⁻¹⁰. Despite NGO support,
86 disparities exist, with childhood cataract accounting for approximately 20-30% of childhood
87 blindness in lower income areas^{11,12}, and closer to 5%¹³ in higher income countries¹⁴.

88 However, the vision loss associated with bilateral or unilateral paediatric cataract is not fully
89 resolved with merely surgical removal of the opaque lens (for overviews see¹⁵⁻¹⁷). The cataract
90 needs to be identified and appropriate referrals made^{18,19}. A replacement lens needs to be
91 implanted, or compensatory contact lens or spectacle correction prescribed¹⁵. Children often
92 require follow up surgeries and ongoing assessment of appropriate refraction¹⁵. Furthermore,
93 there is a risk of secondary conditions, such as glaucoma²⁰ and the risk of deprivation amblyopia
94 increases sharply if the cataract is not removed shortly after onset^{21,22}. Deprivation amblyopia is
95 abnormal visual cortex development caused by visual deprivation, typically by a cataract, in one
96 or both eyes early in life²³. Deprivation amblyopia leads to long-lasting visual impairment that
97 persists after cataract removal²³. The impact of deprivation amblyopia on the more basic
98 aspects of vision (contrast sensitivity and visual acuity) of the affected eye(s) is more
99 pronounced following unilateral²² than bilateral²¹ cataract, although the symptoms are varied
100 and dependent on the duration of visual deprivation²⁴. The treatment for amblyopia involves
101 occlusion (patching) or penalization (atropine cycloplegia) of the stronger eye to promote use of
102 the weaker eye²³. These are home-based therapies which can be difficult for families to
103 implement²⁵. Taken together, the efficacy of childhood cataract surgery is affected by a range of

104 factors including early detection, prompt uptake of surgical intervention, and dedicated
105 commitment to follow up care by the family and medical community (for discussion see^{15 26}).

106 Each step in the treatment pathway for childhood cataract has been investigated, including
107 screening^{19 27-30}, factors contributing to delayed utilization of surgical services³¹, parental stress
108 associated with childhood cataract treatment^{32 33}, compliance with amblyopia treatment³⁴⁻³⁸
109 and the impact of cataract removal on quality of life^{7 39}. Across this international body of work,
110 the services that are available, affordable, and accepted by the community being studied vary
111 considerably. Studies of social barriers to accessing care, delayed cataract removal and poor
112 follow up tend to be done in lower income countries^{7 19 31 40}, whereas research into issues such
113 as increasing adherence with amblyopia treatment, tend to cluster in higher income countries²⁵
114 ³⁴⁻³⁷. The research generally reflects the predominant hierarchical needs within communities⁴¹.
115 However, socio-economic disparities exist within even high-income countries and there is
116 growing recognition that wider research is needed to allow the best visual outcomes for all
117 children^{38 41}.

118 In this study, we were interested in the experience of childhood cataract in Auckland, New
119 Zealand (NZ), where ethnicity is diverse (67% European, 15% Maori, 11% other, 9% Asian, 7%
120 pacific peoples, 1% Middle Eastern/Latin American/Africa - from 2006 census data), and
121 advanced ophthalmological services are now well established and publicly funded. In the early
122 1970s cataract accounted for 22.5% of the children registered as blind in NZ ⁴², whereas a more
123 recent report indicates substantial improvements in the treatment of childhood cataract, with
124 only approximately 4% of children experiencing blindness or low vision attributed to childhood
125 cataract⁴³. There is likely further room for improvement, with a recent report suggesting
126 detection of childhood cataract is suboptimal³⁰. There is no published research in NZ about
127 adherence with follow up or the extent of secondary amblyopia following childhood cataract
128 surgery. In the current study we aimed to understand how childhood cataract and its treatment
129 impacted NZ families, as a starting point to address potential gaps in the system. A qualitative
130 study in the USA found that 'treatment' itself was one of 6 key themes decreasing quality of life
131 for children with a history of cataracts³⁹. Our project probes this idea by focusing on families'

132 lived experience (phenomenology) throughout the treatment pathway. Our aim was to identify
133 sources of distress which 1) could be addressed by the medical community and 2) related to
134 adherence with recommendations from their ophthalmologist.

135

136 Methodology

137 Study context

138 The study was conducted in Auckland, New Zealand (NZ), where comprehensive childhood
139 vision screening systems are in place (red reflex exam at birth and at the 6 week check, pre and
140 in school acuity checks) specialised paediatric cataract surgery is publically funded and
141 prioritised, and subsidies are available for refractive correction, occlusion therapy and
142 transportation costs for families who require financial assistance and/or have particularly high
143 visual needs. The study cohort was culturally diverse (reflecting national averages) and
144 approximately a third had lived and received medical advice outside of NZ. The first author
145 carried out the semi-structured interviews. This author is a non-clinical researcher (not involved
146 in any of the participant's care), and it was made clear to all informants that participation in the
147 study would not impact the care they received.

148 Patient and Public Involvement

149 This project was a first step towards understanding how the medical community could improve
150 provision of treatment for childhood cataract; as such the project was to understand their
151 experiences and needs. The project was part of a larger study about the consequences of⁴⁴, and
152 potential for rehabilitation after⁴⁵ visual deprivation due to childhood cataract, with this step
153 focused on understanding patient experience of the treatment pathway. A lay summary of
154 results will be available to all participants, and information will also be disseminated at a
155 community follow up meeting or 'hui'.

156

157 Participant recruitment

1
2
3 158 In phenomenological studies a purposive sampling strategy is supported in order to recruit
4
5 159 participants who have experienced the phenomenon under study⁴⁶. Accordingly, research
6
7 160 participants were intentionally contacted following a medical records search. Inclusion criteria
8
9 161 included history of a visually significant paediatric cataract (congenital, developmental and
10
11 162 traumatic cases) and a current age of at least four. Exclusion criteria included severe
12
13 163 developmental disorders or severe ocular disease unrelated to paediatric cataract, noted on
14
15 164 ophthalmological records. Invitation letters were posted to thirty-nine parents resulting in
16
17 165 seventeen positive responses, each of whom were invited to participate in the study, including
18
19 166 one parent of a non-verbal child with autism (autism was not diagnosed around the time of
20
21 167 cataract surgery). One additional family made contact independently; their son had not had
22
23 168 surgery to remove his cataract. Two additional families were made aware of the project
24
25 169 through their specialist. A total of twenty semi-structured interviews were carried out. The
26
27 170 study complied with the tenets of the declaration of Helsinki and was approved by ethics
28
29 171 committees from the University of Auckland and the Auckland District Health Board.
30
31 172 Participants provided informed consent, and children who were able provided assent.

31 173 **Semi-Structured interviews**

32
33
34 174 Interviewing is the primary data collection strategy in phenomenological studies⁴⁶. As such,
35
36 175 semi-structured interviews were employed focussing on the lived experiences and personal
37
38 176 perceptions of the treatment process for childhood cataract and secondary deprivation
39
40 177 amblyopia. Interviews were conducted in a private location of the participant's choice and audio
41
42 178 recorded (except in one case due to excessive background noise). We requested that both
43
44 179 caregivers and the affected child were present, and we encouraged all attendees to contribute
45
46 180 to the conversation. Interviews were approximately one hour in length for each family, but
47
48 181 varied in an attempt to obtain rich data. Semi-structured interviews were transcribed by the
49
50 182 first author. Potentially identifiable information was removed or anonymized, and pseudonyms
51
52 183 were assigned (pseudonyms reflect most common NZ names for birth year). The approximate
53
54 184 timing of key treatment events were derived from each interview (narrative summary) and used
55
56 185 to provide context for understanding phenomenology (lived experience).

186 Phenomenology

187 Phenomenology is a qualitative data analysis strategy used in various fields as a tool to delve
188 into personal lived experience⁴⁷⁻⁴⁹. This approach is particularly important for childhood
189 medical research, in which parental choices impact childhood health outcomes and these
190 choices exist within a complex context. Our primary goal was to develop an in-depth
191 appreciation of each family's experience of childhood cataract treatment. We felt this goal was
192 best addressed by a descriptive phenomenological approach.

193 Husserl's⁴⁷ approach to phenomenology, as made explicit by Giorgi⁴⁸, and summarised by
194 Wertz⁴⁹ includes four steps, which we carried out as follows. 1) *Open reading*; we attempted to
195 understand each experience as a whole. 2) *Meaning units*; we divided each transcript into units
196 of meaning or 'themes'. 3) *Psychological reflection*; we reflected on the relevance of the
197 themes to our research interest, and how they related to each other. This was an iterative
198 process including re-reading, summarising, grouping together, and redefining meaning units to
199 better reflect informant experience. 4) *Structural understanding and description*; we
200 consolidated relevant emergent ideas, which we present as three themes and seven sub
201 themes. The first and second author mutually carried out an analysis of the transcripts and then
202 compared notes which resulted in the final presentation of themes. We used SRQR reporting
203 guidelines for qualitative research⁵⁰.

204 Research questions are approached from the frame of our own experiences. Our framing
205 included that 1) we trusted the evidence suggesting that prompt removal of cataracts and
206 adherence with amblyopia treatments (such as patching) are in the best interest of the child,
207 and 2) a critique of family's responses to a challenging life event was outside our scope. We
208 structured our research question to highlight our perspective rather than bracketing our
209 experience from our interpretation; we focused on what the *medical community* could do to
210 alleviate distress, particularly in ways that could promote *adherence*. In the process of seeking
211 to find meaning across the experiences of our informants, we tended towards a constructivist
212 epistemology; fundamentally believing that meaning is constructed rather than discovered.
213 Through iteration, we understood our constructed meaning to be best exemplified in terms of

1
2
3 214 negative and positive experiences of the emergent themes. Therefore the themes presented
4
5 215 reflect encouragement to families at their best, and exasperated distress at their worst.
6
7
8 216
9

10 217
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

218 Results

219 Narrative summary

220 Informants had a wide range of experiences. Key treatment events are summarised as cataract
221 diagnosis, cataract removal, refractive correction and amblyopia treatment. Timelines for each
222 participant are presented in Figure 1, based on estimates from interview data. Some
223 participants had congenital cataracts which were present at birth, others had cataracts which
224 were detected or developed during early childhood (developmental), and others had traumatic
225 cataracts resulting from an eye injury. It is of note that some parents were unsure whether their
226 child had congenital or developmental cataracts.

227 There were no cases in which access to surgery after diagnosis limited care, however perception
228 of delayed detection of cataract was an important issue raised. For one child (Riley), parents
229 were told cataract removal would have too poor a prognosis to justify surgery, given the delay
230 in detection. We have expanded on delays in diagnosis in the first theme. Unlike most families
231 who prioritized prompt cataract removal, Emily and Benjamin's families both declined surgery
232 initially. Both later took up the services, phenomena we further explore within the second and
233 third themes.

234 The experience with home implementation of follow up treatment was diverse. Although no
235 families noted cost of glasses, contacts or patches to be prohibitive, parents expressed concern
236 that acquiring these items and navigating subsidies was complicated and could be stressful. For
237 those who used glasses, feedback was very positive, except for the one participant prescribed
238 glasses in her teenage years (Georgia), who felt uneasy with the aesthetic aspects of wear. Only
239 children with very early surgery used contact lenses instead of an intraocular lens (IOL). Of the
240 six surgeries done before six months of age, half (Olivia, James and Jack) remained aphakic and
241 used a contact lenses rather than IOLs. Each of these families described the use of the contact
242 lenses as the most distressing part of their experience, with two of the three families describing
243 traumatic events related to inserting, removing and/or losing contact lenses. These two families
244 advocated for further support or re-structuring of contact lens use. Patching experience ranged
245 from none (including children with bilateral cataracts who had equal visual acuity loss in both

246 eyes, and children with unilateral cataract who were given a very poor prognosis), to four hours
 247 per day over several years (Olivia, Oliver, Grace and Hannah). The most challenging aspect of
 248 care was described as patching, only superseded by use of contact lenses in infancy in the small
 249 subset of children for whom it was required.

250

251

252 **Figure 1** *Timeline of treatment pathway*

253

254 **Phenomenological summary**

255 Three themes and several sub-themes emerged from the phenomenological analysis related to
 256 how the medical system might better mitigate parental distress and encourage adherence.

257 These are summarized in Table 1.

258

259 **Table 1. Summary of themes and sub-themes**

1. Detection and diagnosis
a) Red reflex exam at birth
b) Referrals and process
2. Communication with medical community
a) Rapport
b) Provision of information about cataract and deprivation amblyopia
c) Foresight about treatment pathway
3. Social support
a) Advice from social networks
b) Availability of emotional and practical

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

260

261

support

For peer review only

262 ***Improved detection and diagnosis***

263 The detection and diagnosis of a cataract was a complex experience for families. If parents
264 understood the timing of diagnosis to be linked to their child's visual outcome, emotions
265 related to perceived efficiency of the screening and processes surrounding referral were
266 heightened. Specifically, for those parents who were informed about the cataract promptly, the
267 negative emotions related to an unexpected diagnosis were balanced by deep appreciation. By
268 contrast, delays associated with perceived error elicited anger and distrust in the medical
269 community.

270 Table 2A highlights the important role of midwives in early screening with the red reflex test;
271 while Lily's mother expresses appreciation from prompt detection (left), Jack's mother express
272 anger about a perceived missed detection (right). Table 2B highlights the importance of the
273 process from detection to diagnosis. Olivia's mother was impressed with the prompt referral
274 pathway and communication between medical staff (left), whereas Riley's mother expressed
275 anger that the medical community had not appropriately referred her son to a specialist.

276 These experiences suggest that any strategy to improve medical provision for childhood
277 cataracts in NZ needs to start with improved training for midwives, general practitioners and
278 pediatricians, both in terms of administering the red reflex test and procedures for prompt
279 referral. Anger with a representative of the medical community, exemplified by Jack and Riley's
280 mother's comments, but shared by many informants, had a lasting impact. A parent's
281 perception of a delay in diagnosis compromised trust in the medical community, which
282 hindered ongoing communication.

283

284

Table 2. Improved detection and diagnosis. Examples of positive (left column) and negative (right column) experiences relating to timing of diagnosis.

A – Red reflex screening test	
<i>She's been a midwife for ages, and because of that experience, she picked up it. (Lily's mother)</i>	<i>We were pretty angry at our midwife because we felt that she should have picked that up. (Jack's mother)</i>
B – Referrals and process	
<i>We got sent to a follow up...to test the red eye reflex in the left eye and basically look in detail because they were expecting the possibility of a cataract. So it was diagnosed pretty much right from the start, she would have been about two days old when it was found and diagnosed. (Olivia's mother)</i>	<i>She [Allied medical worker] put him on a wait list and it took until about 15 months, it was a long time, so I was thinking I guess it's not a big deal. ...when we saw [ophthalmologist] he basically outlined right from the beginning that there was no hope. I found it ... sorry ... there is still a bit of anger there. (Riley's mother)</i>

288

289

290 **Communication with the medical community**

291 Once diagnosed, the relationship a family had with the specialist, and to a lesser degree, the
 292 allied medical staff, was an important part of their experience, permeating across treatment
 293 events. While some families lit up while describing the kindness and competence displayed by
 294 the medical professionals they had come to know well, others relived the anger and frustration
 295 when reflecting on their interactions with the medical community. Rapport between the
 296 parents and the specialist appeared to be particularly important, as it formed the foundation of
 297 the ongoing relationship with the medical team. If there was a connection made and trust

298 established (Table 3A) discourse was positive, whereas if a family was put off by the specialist
299 (Table 3B) the treatment pathway was impaired if not halted.

300 Beyond rapport, provision of understandable information about the condition was critical.

301 What level of information was understandable varied between informants. Some families had a
302 scientific background, and could appreciate a detailed technical explanation while others had
303 limited background knowledge. Across this spectrum of varied backgrounds, effective
304 communication allowed sufficient understanding for the family to feel comfortable with
305 recommendations. In Table 3B, we provide examples of two informants for whom
306 understanding of the condition directly influenced uptake of surgery.

307 Appropriate, understandable communication was also important for parents to prepare for
308 potential future events. When families became exasperated with home based aspects of
309 treatment, those who were prepared in advance (Table 3C - left) were more resilient to the
310 setbacks. On the other hand, families who had not anticipated the challenges, or were not
311 provided useful information to face these challenges in advance (Table 3C – right) became
312 increasingly distressed by them, and lost trust in the system.

313 Whether the communication between the medical community and the parent was perceived as
314 positive was individualized and nuanced, but across participants it included aspects of rapport,
315 clear communication about the condition and of upcoming potential challenges.

316

317

Table 3. Communicating with medical community. Examples of positive (left column) and negative (right column) experiences relating to communication.

A - Building rapport	
<i>We went in and met the paediatric ophthalmologist, and he was just a lovely, lovely man. Yeah, it was just great (Aiden's mother)</i>	<i>I don't want to go to [...] anymore. He's quite 'nothing can be done, nothing can be done' (Riley's mother)</i>
B - Provision of information about cataract and deprivation amblyopia	
<i>Asked whether specialist's description of the condition made sense to her: Yes, it did. Otherwise we could never have agreed to the surgery (Lily's mother)</i>	<i>I was avoiding the surgery from when she was three months onwards [...] they were not answering questions that I was asking to my satisfaction (Emily's mother)</i>
C - Foresight about the treatment pathway	
<i>When she was first diagnosed, the specialist sat us down with this one week old baby and said this is going to be a long, hard road (Olivia's mother)</i>	<i>Why hadn't (paediatric ophthalmologist) told us about this? We'd been going for three months! [...] It's not smooth, it's not transparent, and that's what's hard (Oliver's mother)</i>

321

322 **Social support**

323 Throughout the interviews, it became apparent that provision of care improved if the medical
 324 community understood the family's social context. This includes perspectives or values held by
 325 an individual family and the resources available to them. Advice that parents received from
 326 friends or family was an important part of their social context. Families whose social networks
 327 encouraged them to engage with the medical system were likely to move forward with

328 recommendations (example in Table 4A - left). On the other hand, some social networks were
329 sceptical of the medical community, making it difficult or distressing for parents to seek, or act
330 on, medical advice. Families who felt tension between the advice from the medical community
331 and the advice from friends and family experienced an increased burden during an already
332 challenging time (Table 4A - right).

333 Beyond social networks overtly advising with or against medical advice, there was an important
334 emotional and practical role for this extended group. It was not specifically stated that a phone
335 call from a friend, or an offer to baby-sit other children directly mitigated distress or delays
336 along the treatment pathway, but the absence of this type of support appeared to be associated
337 with both. For example, Benjamin's mother wanted to go through with cataract removal surgery
338 initially, however, she was pregnant and did not have support from her family network (Table
339 4B – right). Like Benjamin's mother, Emily's mother initially declined surgery. However she was
340 able to revisit this decision when she had more emotional and practical support (Table 4B - left).
341 Families were almost ubiquitously overwhelmed at some point during diagnosis or during
342 amblyopia treatment, and only some had the emotional and practical support they needed to
343 follow through with recommendations. It is of particular note that contact lens use required
344 additional practical support; Olivia and Jack's families described requiring 2 or more adults to
345 extract the contact from their infant's eye. Both families had excellent adherence at all stages of
346 the treatment pathway, and substantial support networks.

347 As discussed within 'communication', the family's social context also varied widely across
348 informants. It was further complicated with changes over time; a school or job change, for
349 example, impacted the balance of needs and resources a family had. To fully comply with
350 recommendations, parents needed to have the resources to cope with new challenges,
351 advocate for their child, and be creative in their implementation of home based treatments.
352 Supportive social networks appeared critical to meet this high bar.

353

354

355 **Table 4. Social Support.** Examples of positive (left column) and negative (right column)
 356 experiences relating to social support

357

A – Advice from social networks

Her grandfather, who is a GP, saw that she was looking at things close and her eye was turning in. We took her to an optometrist and they picked up that there was something there, and they referred us to the ophthalmologist who said ‘yes that looks like a cataract to me, we’ll send you to [city]’. And that all happened really quickly
 (Grace’s mother)

The doctor is trying to give us good advice [...] the sooner we get it done the better. And my mum and dad are pretty much saying ‘No’. So that was another kinda difficulty [...] Yeah, an emotional, mental thing as well with the family’s input – what they think about it (Lily’s mother)

B – Availability of emotional and practical support

When describing decision to go through with the surgery four years after it was recommended: It was just talking about it I guess with the [Community worker] that I was having the get-togethers with [...] she was lovely. I would talk with her and we would talk with [child] and she would even offer to drive us out there and stuff like that. Because that was another issue, just the transport
 (Emily’s mother)

We took him to the hospital and they were saying right then and there to do a little surgery [...] I was carrying [another child], and my husband didn’t want him to have the surgery (Benjamin’s mother)

358

359

360 Discussion

361 The goal of our descriptive phenomenological approach⁴⁷⁻⁴⁹ was to develop an in-depth
362 appreciation of the individual experience of childhood cataract treatment; with a specific
363 interest in understanding sources of distress which 1) could be addressed by the medical
364 community and 2) related to adherence to medical recommendations.

365 Strengths and Limitations

366 We were able to recruit a wide variety of participants, with a diverse range of experiences
367 despite childhood cataract being relatively rare. We were able to conduct rich interviews in
368 which we found some informants were very appreciative of the medical community's role in
369 their experience of medical treatment for childhood cataract, while others felt as though they
370 were let down by the medical community. This diversity, and our theoretical framework allowed
371 us to identify themes and construct meaningful summaries which could be presented in positive
372 and negative terms. In other words, for each potential factor we could consider whether the
373 lack impaired a family's experience, and the presence enhanced it. In some cases this directly
374 linked to whether a family adhered to medical recommendations. In this way, the diversity of
375 informant experience is likely to aid in quality and the generalisability of our analysis.

376 However, there were some limitations in our design and analysis. The phenomenological
377 approach does not prioritise counting the frequency with which certain experiences occur.
378 However, it is of interest to know for example, how often parents decline available cataract
379 surgery, or how many families were able to comply with occlusion therapy. We addressed this
380 briefly by including such details in our 'narrative summary'. However, such quantitative
381 questions are best answered with a different methodology. Conversely, a limitation could be
382 that we did not go deep enough into respondents' experiences. For example, during the
383 interviews we did not focus specifically on cultural frameworks for health. Upon iterative
384 reflection it was our impression that a deep understanding of culture may have facilitated our
385 understanding of informant's experiences. Further expanding on this, perhaps a symbolic
386 interactionism theoretical perspective would be a valuable direction for future research in order
387 to understand underlying cultural reasons for choices. Furthermore, we started with the belief

388 that adherence with ophthalmological recommendations is the best for children. Therefore
389 discussion around whether a parent's decision to decline surgery or cease patching was in fact
390 best for the child was outside the scope of this project. Similarly, parental motivation was not a
391 focus of our analysis, but is part of the theoretical framework related to adherence⁵¹. Following
392 up with participants on these points as part of our dissemination step will be a good opportunity
393 to enhance future research.

394 **Summary and Implications of findings**

395 The varied experiences of the diverse informants we interviewed revealed opportunities for
396 systemic improvement in childhood cataract treatment. Early detection and prompt medical
397 treatment are critical for good visual outcomes^{21 22}. Delayed detection was a perceived issue for
398 many of our informants. More effective implementation of the red reflex exam may improve
399 this situation^{18 29}. Survey data from practitioners in New Zealand reflected this opportunity for
400 improvement, specifically highlighting the value of better training for those administering early
401 eye exams³⁰. Our work supports the value of such an initiative.

402 Addressing issues that are social rather than organizational is perhaps more complex. The
403 experiences expressed by our informants suggest that improved communication between the
404 patient and the health care community, and promoting strong social support networks are
405 particularly important. These are not new ideas. Literature concerned with chronic pediatric
406 conditions have described similar phenomena. For example, Shudy et al.⁵² highlighted the
407 importance of a 'sense of partnership with the staff' mirroring the second emergent theme in
408 our analysis. Similarly, Jackson et al.⁵³ discuss how close social support networks improve coping
409 in families of children with brain tumours, with ideas closely related to our third theme of social
410 support. Within our cohort, the impact of these factors was most dramatic in relation to uptake
411 of surgical services. Within the themes of communication and social support, the specific
412 concerns these families raised reflect barriers experienced in lower resources areas. These
413 include indirect costs such as travel^{31 41}, lack of appropriate information^{11 31}, beliefs about
414 health³¹ and/or the parent targeted for education not having decisional power³¹.

1
2
3 415 In addition to these poignant cases in which surgery was declined, additional families in our
4
5 416 cohort felt overwhelmed at various stages of the treatment pathway, particularly when
6
7 417 attempting patching and contact lens wear (struggles more consistent with the literature from
8
9 418 higher resourced countries). Contact lens use in infants (and to a lesser degree patching) has
10
11 419 been associated with higher levels of parental stress³³. Parental stress is known to contribute to
12
13 420 decreased adherence with amblyopia treatment in cases of strabismic and anisometric
14
15 421 amblyopia³⁷. These ideas appeared to be consistent with accounts from our informants. Again,
16
17 422 good communication and strong social support appeared to mitigate these factors. Although
18
19 423 care must be taken when generalizing, we suggest that perhaps communication and practical
20
21 424 support may have been factors which account for the difference between the results of Drews
22
23 425 et al³³ which revealed an association between the use of contact lenses following paediatric
24
25 426 cataract surgery and parental stress and those of Celano et al³² which did not. Notably, the data
26
27 427 reported by Celano et al³² were collected in the context of a clinical trial, in which extensive
28
29 428 information and practical support were provided.

30 429 Although all families found parts of the treatment pathway challenging at times, cultural socio
31
32 430 economic factors appeared to exasperate the challenges. Some work suggests that if a family
33
34 431 does not feel connected with the dominant society (or the person representing the society as
35
36 432 the medical professional) their child is less likely to adhere to treatment for amblyopia³⁸. In our
37
38 433 diverse cohort the concept appeared to extend beyond patching to cataract removal, which has
39
40 434 more substantial consequences. Although there were many professionals involved in patient
41
42 435 care, paediatric ophthalmologists were generally seen as the primary providers of information.
43
44 436 This means the demand on a single specialist of providing sensitive, individualised support to a
45
46 437 diverse community may be significant.

47 438 In some cases print materials can help supplement communication of important information
48
49 439 and encourage treatment compliance³⁶, however information alone is often insufficient to
50
51 440 promote action⁵⁴. Patient care may benefit from the provision of accurate information and
52
53 441 practical support from different professionals. Some research has described the role of a
54
55 442 'childhood blindness coordinator' to be particularly beneficial to fill this gap^{40 55}. This role can

1
2
3 443 involve helping parents understand the condition, providing foresight about the treatment
4
5 444 pathway, as well as providing practical support such as text message reminders and travel
6
7 445 planning⁵⁵. In terms of emotional support, formal networks tend to be less effective than
8
9 446 informal support networks⁵³. However, Ireys et al. showed the benefit of peer support groups
10
11 447 for mothers of children with chronic illness⁵⁶, and Emily's family provides a good example of
12
13 448 how a caring community volunteer can provide effective support. Research within the childhood
14
15 449 disability service in New Zealand has suggested the role of a 'cultural case worker' may facilitate
16
17 450 both improved communication and supplement social support for families with culturally or
18
19 451 linguistically diverse backgrounds⁵⁷, working as 'patient advocates' rather than expert medical
20
21 452 staff. These are all avenues that could be further explored to help families dealing with
22
23 453 childhood cataract.

24 454 Although the system for childhood cataract treatment in NZ is technically robust, it does not
25
26 455 appear to have sufficient support for families who are unable to effectively advocate for
27
28 456 themselves. The current study suggests strategies from research in lower income countries may
29
30 457 help some NZ families. Examples are the community roles of 'key informants' and 'childhood
31
32 458 blindness coordinators', which facilitate education, detection process, and communication. New
33
34 459 Zealand has organisations that could fill these roles, such as the Blind and Low Vision Network
35
36 460 New Zealand (BLENNZ, which provides comprehensive support for children, from infancy to 17
37
38 461 years of age, with moderate to severe visual impairment) and social workers or caseworkers.
39
40 462 However, their services are not well utilized by families due to the lack of awareness among
41
42 463 general population and medical practitioners. Better training for midwives, pediatricians and
43
44 464 general practitioners for early detection and referral as well as better awareness of existing
45
46 465 support services are likely to be important steps towards improving provision of care for
47
48 466 childhood cataract.

49 467 **Conclusion**

50
51 468 Childhood cataract is a rare and challenging condition^{15 16 26}, its treatment requires much from
52
53 469 families during what is already an overwhelming phase of life. Our work suggests that
54
55 470 availability of surgery and funding for secondary costs for those who need it is insufficient to

1
2
3 471 achieve the best visual outcomes for NZ children with cataracts. We need to improve screening
4
5 472 practices, including streamlining referral pathways to specialised paediatric ophthalmologists.
6
7 473 We need to find ways to communicate effectively with diverse families, ensuring the condition
8
9 474 and its treatment is understood and family's questions are answered. Finally, we need to
10
11 475 consider creative ways to support families with surgical uptake and post-surgical follow up. This
12
13 476 requires awareness family context, including available emotional and practical support.
14

15 477 References

-
- 17 478 1. Gilbert C, Foster A. Childhood blindness in the context of VISION 2020 - The right to sight.
18 479 *Bulletin of the World Health Organization* 2001;79(3):227-32.
- 20 480 2. Courtright P, Hutchinson AK, Lewallen S. Visual impairment in children in middle- and lower-
21 481 income countries. *Archives of Disease in Childhood* 2011;96(12):1129-34. doi:
22 482 10.1136/archdischild-2011-300093
- 24 483 3. Gogate P, Dole K, Ranade S, et al. Cost of pediatric cataract surgery in Maharashtra, India.
25 484 *International Journal of Ophthalmology* 2010;10(7):1248-52. doi: 10.3969/j.issn.1672-
26 485 5123.2010.07.004
- 27 486 4. Shamanna BR, Muralikrishnan R. Childhood cataract: Magnitude, management, economics
28 487 and impact. *Community Eye Health* 2004;17(50):17-18.
- 30 488 5. Evans CT, Lenhart PD, Lin D, et al. A cost analysis of pediatric cataract surgery at two child eye
31 489 health tertiary facilities in Africa. *J AAPOS* 2014;18(6):559-62. doi:
32 490 10.1016/j.jaapos.2014.08.005
- 33 491 6. Rahi JS, Dezateux C. Measuring and interpreting the incidence of congenital ocular
34 492 anomalies: Lessons from a national study of congenital cataract in the UK. *Investigative*
35 493 *Ophthalmology and Visual Science* 2001;42(7):1444-48.
- 37 494 7. Kalia A, Gandhi T, Chatterjee G, et al. Assessing the impact of a program for late surgical
38 495 intervention in early-blind children. *Public health* 2017;146:15-23. doi:
39 496 10.1016/j.puhe.2016.12.036 [published Online First: 2017/04/14]
- 41 497 8. Sinha P, Chatterjee G, Gandhi T, et al. Restoring vision through "Project Prakash": the
42 498 opportunities for merging science and service. *PLoS Biol* 2013;11(12):e1001741. doi:
43 499 10.1371/journal.pbio.1001741
- 44 500 9. Levine S, Øverland L, Ramson P. The miracle workers: obstacles and opportunities for
45 501 restoring sight to children in KwaZulu-Natal. *Anthropology Southern Africa* 2014;37(1-
46 502 2):81-93. doi: 10.1080/23323256.2014.940193
- 48 503 10. Lewallen S, Schmidt E, Jolley E, et al. Factors affecting cataract surgical coverage and
49 504 outcomes: a retrospective cross-sectional study of eye health systems in sub-Saharan
50 505 Africa. *BMC Ophthalmol* 2015;15:67. doi: 10.1186/s12886-015-0063-6
- 51 506 11. Courtright P. Childhood cataract in sub-Saharan Africa. *Saudi J Ophthalmol* 2012;26(1):3-6.
52 507 doi: 10.1016/j.sjopt.2011.10.006
- 54 508 12. Kemmanu V, Hegde K, Giliyar SK, et al. Prevalence of Childhood Blindness and Ocular
55 509 Morbidity in a Rural Pediatric Population in Southern India: The Pavagada Pediatric Eye
56
57
58
59
60

- 1
2
3 510 Disease Study-1. *Ophthalmic Epidemiology* 2016;23(3):185-92. doi:
4 511 10.3109/09286586.2015.1090003
5
6 512 13. Rahi JS, Cable N. Severe visual impairment and blindness in children in the UK. *Lancet*
7 513 2003;362(9393):1359-65. doi: 10.1016/S0140-6736(03)14631-4
8 514 14. Koay CL, Patel DK, Tajunisah I, et al. A comparative analysis of avoidable causes of childhood
9 515 blindness in Malaysia with low income, middle income and high income countries.
10 516 *International Ophthalmology* 2015;35(2):201-07. doi: 10.1007/s10792-014-9932-x
11
12 517 15. Medsinghe A, Nischal KK. Pediatric cataract: Challenges and future directions. *Clinical*
13 518 *Ophthalmology* 2015;9:77-90. doi: 10.2147/OPTH.S59009
14 519 16. Oscar A, Veleva N, Chernodrinska V, et al. Childhood cataract. *Pediatrriya* 2014;54(1):58-61.
15
16 520 17. Lewallen S, Roberts H, Hall A, et al. Increasing cataract surgery to meet Vision 2020 targets;
17 521 experience from two rural programmes in east Africa. *Br J Ophthalmol*
18 522 2005;89(10):1237-40. doi: 10.1136/bjo.2005.068791
19 523 18. Litmanovitz I, Dolfin T. Red reflex examination in neonates: The need for early screening.
20 524 *Israel Medical Association Journal* 2010;12(5):301-02.
21
22 525 19. Shija F, Shirima S, Lewallen S, et al. Comparing key informants to health workers in
23 526 identifying children in need of surgical eye care services. *International Health*
24 527 2012;4(1):1-3.
25 528 20. Baden C, Shija F, Lewallen S, et al. Glaucoma after pediatric cataract surgery in a population
26 529 with limited access to care. *J AAPOS* 2013;17(2):158-62. doi:
27 530 10.1016/j.jaapos.2012.11.009
28
29 531 21. Birch EE, Cheng C, Stager Jr DR, et al. The critical period for surgical treatment of dense
30 532 congenital bilateral cataracts. *Journal of AAPOS* 2009;13(1):67-71.
31 533 22. Birch EE, Stager DR. The critical period for surgical treatment of dense congenital unilateral
32 534 cataract. *Investigative Ophthalmology and Visual Science* 1996;37(8):1532-38.
33
34 535 23. Holmes JM, Clarke MP. Amblyopia. *Lancet* 2006;367(9519):1343-51. doi: 10.1016/S0140-
35 536 6736(06)68581-4
36 537 24. Hamm LM, Black J, Dai S, et al. Global processing in amblyopia: a review. *Front Psychol*
37 538 2014;5:583. doi: 10.3389/fpsyg.2014.00583
38
39 539 25. Wang J. Compliance and patching and atropine amblyopia treatments. *Vision Research*
40 540 2015;114:31-40. doi: <http://dx.doi.org/10.1016/j.visres.2015.02.012>
41 541 26. Repka MX. Monocular infantile cataract: Treatment is worth the effort. *Archives of*
42 542 *Ophthalmology* 2010;128(7):931-33. doi: 10.1001/archophthalmol.2010.130
43
44 543 27. Saiju R, Yun S, Yoon PD, et al. Bruckner red light reflex test in a hospital setting. *Kathmandu*
45 544 *University Medical Journal* 2012;10(38):23-26.
46 545 28. Gräf M. Early detection of ocular disturbances in children. Brückner's transillumination test
47 546 (red reflex) - A must in childhood screening. *Deutsches Arzteblatt* 2007;104(11):724-29.
48 547 29. Fry M, Wilson GA. Scope for improving congenital cataract blindness prevention by
49 548 screening of infants (red reflex screening) in a New Zealand setting. *Journal of*
50 549 *Paediatrics and Child Health* 2005;41(7):344-46. doi: 10.1111/j.1440-1754.2005.00628.x
51
52 550 30. Raoof N, Dai S. Red reflex screening in New Zealand: a large survey of practices and
53 551 attitudes in the Auckland region. *The New Zealand medical journal* 2016;129(1438):38-
54 552 43. [published Online First: 2016/07/23]

- 1
2
3 553 31. Bronsard A, Geneau R, Shirima S, et al. Why are children brought late for cataract surgery?
4 554 Qualitative findings from Tanzania. *Ophthalmic Epidemiol* 2008;15(6):383-88.
5 555 32. Celano M, Hartmann EE, Drews-Botsch CD. Parenting stress in the infant aphakia treatment
6 556 study. *Journal of Pediatric Psychology* 2013;38(5):484-93. doi: 10.1093/jpepsy/jst009
7 557 33. Drews C, Celano M, Plager DA, et al. Parenting stress among caregivers of children with
8 558 congenital cataracts. *Journal of AAPOS* 2003;7(4):244-50. doi: 10.1016/S1091-
9 559 8531(03)00118-6
10 560 34. Drews-Botsch CD, Celano M, Kruger S, et al. Adherence to occlusion therapy in the first six
11 561 months of follow-up and visual acuity among participants in the infant Aphakia
12 562 treatment study (IATS). *Investigative Ophthalmology and Visual Science*
13 563 2012;53(7):3368-75.
14 564 35. Drews-Botsch CD, Hartmann EE, Celano M. Predictors of adherence to occlusion therapy 3
15 565 months after cataract extraction in the Infant Aphakia Treatment Study. *Journal of*
16 566 *AAPOS* 2012;16(2):150-55.
17 567 36. Loudon SE, Fronius M, Looman CWN, et al. Predictors and a remedy for noncompliance with
18 568 amblyopia therapy in children measured with the occlusion dose monitor. *Investigative*
19 569 *Ophthalmology and Visual Science* 2006;47(10):4393-400.
20 570 37. Loudon SE, Passchier J, Chaker L, et al. Psychological causes of non-compliance with
21 571 electronically monitored occlusion therapy for amblyopia. *British Journal of*
22 572 *Ophthalmology* 2009;93(11):1499-503. doi: 10.1136/bjo.2008.149815
23 573 38. Tjiam AM, Akcan H, Ziylan F, et al. Sociocultural and psychological determinants in migrants
24 574 for noncompliance with occlusion therapy for amblyopia. *Graefes Archive for Clinical*
25 575 *and Experimental Ophthalmology* 2011;249(12):1893-99.
26 576 39. Castañeda YS, Cheng-Patel CS, Leske DA, et al. Quality of life and functional vision concerns
27 577 of children with cataracts and their parents. *Eye (Basingstoke)* 2016;30(9):1251-59. doi:
28 578 10.1038/eye.2016.134
29 579 40. Courtright P, Bowman R, Gilbert C, et al. Childhood cataract in Africa. 2008
30 580 41. Lenhart PD, Courtright P, Wilson ME, et al. Global challenges in the management of
31 581 congenital cataract: proceedings of the 4th International Congenital Cataract
32 582 Symposium held on March 7, 2014, New York, New York. *J AAPOS* 2015;19(2):e1-8. doi:
33 583 10.1016/j.jaapos.2015.01.013
34 584 42. Sturman D. Blindness in childhood in New Zealand. *Transactions of the Ophthalmological*
35 585 *Society of New Zealand* 1975;Vol. 27:45-52.
36 586 43. Chong C, Dai S. Cross-sectional study on prevalence, causes and avoidable causes of visual
37 587 impairment in Māori children. *New Zealand Medical Journal* 2013;126(1379)
38 588 44. Hamm L, Chen Z, Li J, et al. Interocular suppression in children with deprivation amblyopia.
39 589 *Vision Res* 2017;133:112-20. doi: 10.1016/j.visres.2017.01.004
40 590 45. Hamm LM, Chen Z, Li J, et al. Contrast-balanced binocular treatment in children with
41 591 deprivation amblyopia. *Clin Exp Optom* 2018;101(4):541-52. doi: 10.1111/cxo.12630
42 592 46. Starks H, Brown Trinidad S. Choose Your Method: A Comparison of Phenomenology,
43 593 Discourse Analysis, and Grounded Theory. *Qualitative Health Research*
44 594 2007;17(10):1372-80. doi: 10.1177/1049732307307031
45 595 47. Husserl E. Ideas: General introduction to pure phenomenology. New York: Collier Books
46 596 1962.

- 1
2
3 597 48. Giorgi A. Psychology as a human science: A phenomenologically based approach. New York:
4 598 Harper & Row 1970.
- 5 599 49. Wertz FJ. A phenomenological psychological approach to trauma and resilience. Five Ways
6 600 of Doing Qualitative Analysis. New York: The Guildford Press 2011.
- 7 601 50. O'Brien BC, Harris IB, Beckman TJ, et al. Standards for reporting qualitative research: a
8 602 synthesis of recommendations. *Academic medicine : journal of the Association of*
9 603 *American Medical Colleges* 2014;89(9):1245-51. doi: 10.1097/acm.0000000000000388
10 604 [published Online First: 2014/07/01]
- 11 605 51. McGrady ME, Ryan JL, Brown GA, et al. Topical review: Theoretical frameworks in pediatric
12 606 adherence-promotion interventions: Research findings and methodological implications.
13 607 *Journal of Pediatric Psychology* 2015;40(8):721-26. doi: 10.1093/jpepsy/jsv025
- 14 608 52. Shudy M, De Almeida ML, Ly S, et al. Impact of pediatric critical illness and injury on
15 609 families: A systematic literature review. *Pediatrics* 2006;118(SUPPL. 3):S203-S18. doi:
16 610 10.1542/peds.2006-0951B
- 17 611 53. Jackson AC, Enderby K, O'Toole M, et al. The role of social support in families coping with
18 612 childhood brain tumor. *Journal of Psychosocial Oncology* 2009;27(1):1-24. doi:
19 613 10.1080/07347330802614634
- 20 614 54. Blödt S, Kaiser M, Adam Y, et al. Understanding the role of health information in patients'
21 615 experiences: secondary analysis of qualitative narrative interviews with people
22 616 diagnosed with cancer in Germany. *BMJ Open* 2018;8(3) doi: 10.1136/bmjopen-2017-
23 617 019576
- 24 618 55. Kishiki E, Van Dijk K, Courtright P. Strategies to improve follow-up of children after surgery
25 619 for cataract: Findings from Child Eye Health Tertiary Facilities in sub-Saharan Africa and
26 620 South Asia. *Eye (Basingstoke)* 2016;30(9):1234-41. doi: 10.1038/eye.2016.169
- 27 621 56. Ireys HT, Chernoff R, DeVet KA, et al. Maternal outcomes of a randomized controlled trial of
28 622 a community-based support program for families of children with chronic illnesses.
29 623 *Archives of Pediatrics and Adolescent Medicine* 2001;155(7):771-77.
- 30 624 57. Mortensen A, Latimer S, Yusuf I. Cultural case workers in child disability services: an
31 625 evidence-based model of cultural responsiveness for refugee families. *Kotuitui*
32 626 2014;9(2):50-59. doi: 10.1080/1177083X.2014.911752

627

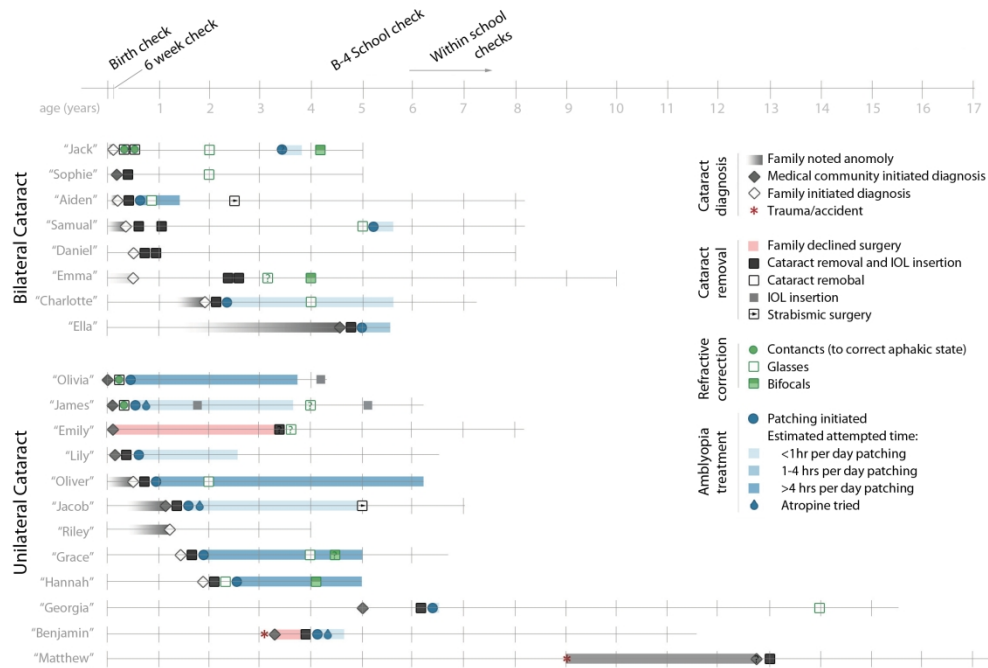


Figure 1. Timeline of treatment pathway. Participants are categorized into cases of unilateral or bilateral cataract. An asterisk by a patient pseudonym indicates a family history of paediatric cataract. Cataract diagnosis is depicted with a diamond. If a medical professional, or a community screen initiated the appointment in which a diagnosis occurred, the diamond is grey, whereas the diamond is white if the family initiated the appointment at which the diagnosis was made. A grey bar preceding a diamond indicates that abnormal behaviour or ocular appearance was noticed by the family prior to diagnosis. Treatment is represented by a square; black represents a surgical intervention, and green optical. Red bars between diagnosis and treatment indicate treatment was recommended during this time, but the family declined. IOL = intraocular lens

Reporting checklist for qualitative study.

Based on the SRQR guidelines.

Instructions to authors

Complete this checklist by entering the page numbers from your manuscript where readers will find each of the items listed below.

Your article may not currently address all the items on the checklist. Please modify your text to include the missing information. If you are certain that an item does not apply, please write "n/a" and provide a short explanation.

Upload your completed checklist as an extra file when you submit to a journal.

In your methods section, say that you used the SRQR reporting guidelines, and cite them as:

O'Brien BC, Harris IB, Beckman TJ, Reed DA, Cook DA. Standards for reporting qualitative research: a synthesis of recommendations. *Acad Med.* 2014;89(9):1245-1251.

		Reporting Item	Page Number
Title	#1	Concise description of the nature and topic of the study identifying the study as qualitative or indicating the approach (e.g. ethnography, grounded theory) or data collection methods (e.g. interview, focus group) is recommended	1
	#2	Summary of the key elements of the study using the abstract format of the intended publication; typically includes background, purpose, methods, results and conclusions	3
Problem formulation	#3	Description and significance of the problem / phenomenon studied: review of relevant theory and empirical work; problem statement	5
Purpose or research question	#4	Purpose of the study and specific objectives or questions	5
Qualitative approach and research paradigm	#5	Qualitative approach (e.g. ethnography, grounded theory, case study, phenomenology, narrative research) and	7-8

guiding theory if appropriate; identifying the research paradigm (e.g. postpositivist, constructivist / interpretivist) is also recommended; rationale. The rationale should briefly discuss the justification for choosing that theory, approach, method or technique rather than other options available; the assumptions and limitations implicit in those choices and how those choices influence study conclusions and transferability. As appropriate the rationale for several items might be discussed together.

1			
2			
3			
4			
5			
6			
7			
8			
9			
10			
11			
12			
13			
14	Researcher	#6	8
15	characteristics and		
16	reflexivity	Researchers' characteristics that may influence the research, including personal attributes, qualifications / experience, relationship with participants, assumptions and / or presuppositions; potential or actual interaction between researchers' characteristics and the research questions, approach, methods, results and / or transferability	
17			
18			
19			
20			
21			
22			
23			
24			
25	Context	#7	6
26		Setting / site and salient contextual factors; rationale	
27			
28	Sampling strategy	#8	6
29		How and why research participants, documents, or events were selected; criteria for deciding when no further sampling was necessary (e.g. sampling saturation); rationale	
30			
31			
32			
33			
34			
35	Ethical issues pertaining	#9	7
36	to human subjects	Documentation of approval by an appropriate ethics review board and participant consent, or explanation for lack thereof; other confidentiality and data security issues	
37			
38			
39			
40	Data collection methods	#10	7
41		Types of data collected; details of data collection procedures including (as appropriate) start and stop dates of data collection and analysis, iterative process, triangulation of sources / methods, and modification of procedures in response to evolving study findings; rationale	
42			
43			
44			
45			
46			
47			
48			
49			
50	Data collection	#11	7
51	instruments and	Description of instruments (e.g. interview guides, questionnaires) and devices (e.g. audio recorders) used for data collection; if / how the instruments(s) changed over the course of the study	
52	technologies		
53			
54			
55			
56			
57	Units of study	#12	6-7 and
58		Number and relevant characteristics of participants, documents, or events included in the study; level of	8-9
59			
60			

participation (could be reported in results)

(also
Figure 1)

1			
2			
3			
4	Data processing	#13 Methods for processing data prior to and during analysis, including transcription, data entry, data management and security, verification of data integrity, data coding, and anonymisation / deidentification of excerpts	6-7
5			
6			
7			
8			
9			
10			
11	Data analysis	#14 Process by which inferences, themes, etc. were identified and developed, including the researchers involved in data analysis; usually references a specific paradigm or approach; rationale	7
12			
13			
14			
15			
16			
17			
18	Techniques to enhance trustworthiness	#15 Techniques to enhance trustworthiness and credibility of data analysis (e.g. member checking, audit trail, triangulation); rationale	8
19			
20			
21			
22			
23	Syntheses and interpretation	#16 Main findings (e.g. interpretations, inferences, and themes); might include development of a theory or model, or integration with prior research or theory	Table 1
24			
25			
26			
27			
28	Links to empirical data	#17 Evidence (e.g. quotes, field notes, text excerpts, photographs) to substantiate analytic findings	Tables 2-4
29			
30			
31			
32	Intergration with prior work, implications, transferability and contribution(s) to the field	#18 Short summary of main findings; explanation of how findings and conclusions connect to, support, elaborate on, or challenge conclusions of earlier scholarship; discussion of scope of application / generalizability; identification of unique contributions(s) to scholarship in a discipline or field	17-20
33			
34			
35			
36			
37			
38			
39			
40			
41			
42	Limitations	#19 Trustworthiness and limitations of findings	17-18
43			
44	Conflicts of interest	#20 Potential sources of influence of perceived influence on study conduct and conclusions; how these were managed	8 and 18
45			
46			
47			
48			
49			
50	Funding	#21 Sources of funding and other support; role of funders in data collection, interpretation and reporting	1
51			
52			
53			

The SRQR checklist is distributed with permission of Wolters Kluwer © 2014 by the Association of American Medical Colleges. This checklist can be completed online using <https://www.goodreports.org/>, a tool made by the [EQUATOR Network](#) in collaboration with [Penelope.ai](#)