ABSTRACT

Objectives: To examine the experience of infants, children and their parents, the role of ophthalmologists and other health, social care and education professionals in the certification and registration processes and examine the relationship between certification and referrals and pathways to support.

Design: Qualitative study.

Setting: Telephone interviews with health and, social care professionals, qualified teachers of children and young people with vision impairment (QTVIs) and parents of infants/children in England.

Participants: 52 health, social care and education professionals who are part of the certification or registration process. 26 parents of infants and children with vision impairment.

Results: Referrals to education do not require a Certificate of Vision Impairment (CVI); however, the majority of parents received support from education and social services only after an offer of the CVI, which was often dependent on having a formal diagnosis. Parents stated they wanted support sooner, particularly parents of children with additional complex needs who experienced longer delays. Areas with multidisciplinary, well-supported teams and support roles such as eye clinic liaison officers (ECLOs) appeared to have more reliable referral pathways.

Conclusions: For infants and children with vision impairment, there should be a consistent mechanism for triggering education and social care support even with uncertainty about diagnosis and/or prognosis. All professionals involved in the certification and registration processes (ophthalmologists, optometrists, ECLOs, orthoptists, social workers, QTVIs) can better communicate the value and benefits of certification and registration.

INTRODUCTION

The purpose of the Certificate of Vision Impairment (CVI) for both children and adults is to provide a reliable route for someone with sight loss to be brought to the attention of social care. Certification and registration is transformative for adults as the referral to social services and the support triggered substantially improves their lives. While for infants and children, social services often have an important role to play, and registration also brings financial and practical benefits to families, the main provider of support is the local authority specialist vision impairment (VI) education service. Early assessment and interventions (eg, low-vision aids) can decrease risks of delayed development in motor, cognitive, language and social domains. In the UK, qualified teachers of children and young people with
vision impairment (QTVIs) most commonly provide training and support to enable parents to assist their child’s early development.

Information is parents’ greatest need at the period of establishing a diagnosis of their child’s ophthalmic disorder.8 During this critical period when an infant/child’s VI is being diagnosed, parents state they want support to both accept their infant/child’s sight loss and to learn how to maximise the remaining vision and develop adaptive skills.9–11 However, research consistently finds parents do not receive this information from health professionals at an early stage, leaving parents frustrated12 and children with delayed development.13 14

Through interviews with health and social care and education professionals and parents, this research explores what triggers referral for support and whether clinicians depend on a firm diagnosis or a CVI to act as a trigger, or needs of/requests from patients and parents. In addition, it examines the impact of waiting for referrals to education and provides examples of good practice and identifies ways of ensuring prompt referrals. The research follows on from previous research on adults and the CVI and registration process1 (see box 1), whereas the aim of this research was to examine issues related to the certification and registration processes in infants and young people.

### Box 1 The certification and registration processes

Certification and registration are two separate processes and involve a number of stages. First, the Certificate of Vision Impairment (CVI) is completed by a consultant ophthalmologist who establishes a child’s eligibility for certification as either sight impaired (SI) or severely sight impaired (SSI). The completed CVI is then sent to the local Social Services Department who ‘offer’ registration, as it is a voluntary choice. Registration offers practical and financial benefits and concessions.15 The CVI should act as a referral for a social care assessment, and where eligible, the offer of support.

**Support offered by education follows a separate referral pathway and is not dependant on certification or registration** but based on a child’s need.

The UK Department of Health recommends that infants and young children who have congenital ocular abnormalities leading to visual defects should be certified as SI unless they are obviously SSI.16 Children aged 4 and over should be certified as SSI or SI according to their binocular corrected vision. No other UK-specific guidelines exist for children. Between 1982 and 2011, there has been more than twofold increase in new blind and partial-sight registration in children in England.17 However, the rise in registration rates is inconsistent across England.18 The cause of this irregularity is estimated to be due to both the ‘combination of a genuinely higher prevalence of visual impairment locally and under-reporting of visual impairment nationally’.19 The criteria for certification as blind or partially sighted are defined by the Department of Health. Generally, to be certified (and subsequently registered) as SSI (blind) a person’s sight needs to fall into one of the following categories, while wearing glasses or contact lenses that they need:

- Visual acuity of less than 3/60 with a full visual field
- Visual acuity of between 3/60 and 6/60 with a moderate reduction of field of vision, such as tunnel vision
- Visual acuity of 6/60 or above, but with a very reduced field of vision especially if a lot of sight is missing especially in the lower part of the field

To be certified (and subsequently registered) as SI (partially sighted) a person’s sight has to fall into one of the following categories, while wearing glasses or contact lenses that they need:

- Visual acuity of 3/60 to 6/60 with a full field of vision
- Visual acuity of up to 6/24 with a moderate reduction of field of vision or with a central part of vision that is cloudy or blurry
- Visual acuity of up to 6/18 or better if a large part of their field of vision, for example, a whole half of their vision, is missing or a lot of their peripheral vision is missing.

The only additional advice with regard to infants and young people are:

- Infants and young children who have congenital ocular abnormalities leading to visual defects should be certified as SI unless they are obviously severely sight impaired.
- Children aged 4 and over should be certified as severely sight impaired or sight impaired according to the binocular corrected vision.20

**MATERIALS AND METHODS**

**Sample**

The purposive sample seeks to provide an overview of the certification and registration processes in five National Health Service (NHS) areas. The study of health, social care and education professionals was conducted in five areas in England identified as having differences in sight loss registration rates (lower, average or higher registration rates than might be expected for the size of the child population in that area)21 and whether or not they employed specialist paediatric ophthalmologists. We purposely selected areas that would provide us with, as Morse22 states, ‘excellent examples’ as well as more ordinary or common practices. Purposive sampling is based on knowledge of a population.23 In this research, those professionals involved in certifying and supporting infants and children with VI and parents of children who are certified as severely sight impaired or sight impaired were deliberately selected for interview.

Hospital consultants were identified by Royal National Institute of Blind People (RNIB) or the advisory group; subsequent healthcare staff were identified by the consultants first contacted. The research advisory group included health, education and social care professionals. Education (QTVIs) and social care interviewees were identified by healthcare professionals or one of the...
authors. Parents were recruited via invitation calls sent from RNIB and Action for Blind People (posted on their websites) and letters sent by interviewed QTVIs. The request for interviews was also placed on social media websites of parent support groups—this was beyond our control but was very welcome as it led to a number of parents being interviewed. Parents were recruited from across the whole of England.

After examination of the National Research Ethics Framework the research was deemed a service evaluation and ethics approval was not necessary. None of the parental participants were identified or approached via NHS; instead, we used membership of charities and patient support organisations. Informed consent was obtained by all participants at the beginning of each interview. Personal identifying information was not recorded.

Interviews and data analysis
The interviews consisted of semistructured questions covering the following themes:

- Description and/or experience of certification and registration processes;
- Attitudes to and meaning of certification and registration;
- Role and relationships with relevant stakeholders (health, education, social services);
- Improving experiences and systems.

The nature of the interviews encouraged participants to feel comfortable to raise issues they felt relevant to their own experiences. The advisory group provided guidance on the interview questions’ consistency, validity and ambiguity.

Each transcript was read and analysed multiple times. Interview data were analysed thematically, similar to previous research with adults. A list of deductive codes was initially created; inductive codes emerged during the second level of the thematic analysis.

All interviews were digitally recorded with the participant’s consent, lasted between 10 and 50 min and were transcribed verbatim. The interviews were completed between March and July 2014.

Extracts are referenced with the type of interviewee and interview number—parent (Par); ophthalmologist (Ophth); secretary/administrator (Adm); nurse (Nur); optometrist (Optom); eye clinic liaison officer (ECLO); orthoptist (orth); social services staff including managers, rehabilitation officers, administrators (SS) and QTVIs and a manager (QTVIs). The terms ‘certification’ and ‘registration’ were used inconsistently by most interviewees; hence these terms are amended in the text for clarity.

RESULTS
Interviewees
Seventy-eight participants were interviewed. Hospital interviews were with a range of staff involved in the certification process (see table 1). Of the 12 consultants interviewed, 10 were qualified for over 10 years, the remaining 2 consultants were qualified for over 5 years.

Twenty-seven per cent (n=7) of parents stated they had an income below £15 000/annum. Two parents (8%) classified themselves as Asian, slightly below England’s population of mixed, Asian, African-American and Chinese ethnicities which make up 12% of England’s population.

Specialist education services in seven local authorities associated with the hospitals and representatives from seven social care departments were also interviewed. In one area, representatives from different social services and education departments were interviewed resulting in professionals from eight local authority areas being interviewed.

Referral processes
Across England, the typical patient pathway when an infant or child has a VI is a direct referral from health to education, usually via letter. Once a referral from the hospital is received by education, a QTVI makes initial contact, usually by telephone. All but one of the areas under analysis referred directly from hospital to
education. In one area undergoing a lengthy reorganisation, QTVIs were not alerted when children were certified and stated ‘it’s a little bit hit and miss when we find out’ (QTVI3).

Under Special Educational Needs and Disability legislation, an infant or child’s entitlement to specialist educational support is entirely independent of whether or not they have a CVI. All QTVIs interviewed confirmed children do not need to be certified or registered in order to receive educational support.

(Certification and registration) doesn’t open doors [to QTVIs]...we can offer all the resources and things whether they are registered or not. (QTVI2)

All parents interviewed stated the lead professional supporting their infants and children came from education rather than social care (where adults receive their support).

Parents and ophthalmologists report different experiences
The interviews found differences in opinions between the experiences of parents seeking support for their VI infants and children and ophthalmologists’ stated practice. Parents stated they wanted referrals to education prior to diagnosis, but doctors appeared to wait until they had a firm diagnosis before referring. The offer of a CVI appeared to trigger ophthalmologists to refer to education services.

Parents wait for diagnosis and offer of CVI before being referred
Despite processes outlining that a CVI is not needed for an education referral, two-thirds of parents (n=18) stated they were not referred to support until after the CVI was issued to their infants/children. These parents stated ophthalmologists delayed or did not mention either certification or referral to education until they had reached a firm diagnosis.

(Ophthalmologists) really didn’t do a lot before (certification), that’s what I’m angry about, those years before school, more wasn’t done. Now when I look online and stuff you see all of these nursery schools for VI children and afternoon sessions and play sessions. There was none of that, I didn’t know about any of that. (Par14)

One parent of a child diagnosed with idiopathic nystagmus at age 9 months waited until he was 3 to be certified, by this stage she had referred her child to a QTVI through her own networks and received no referrals to support or information about her child’s VI from the hospital. In this case, certification was prompted only when the child participated in a research project and the research staff recommended it to provide additional support (Par6).

For some parents, the lack of referral or access to information had a lasting and significant impact on their lives. One parent was told by a paediatric ophthalmologist to treat their child eventually certified as sight impaired simply as ‘a normal baby’.

The doctor at *** just said to me ‘just treat him like a normal baby. Just carry on exactly like you are’...if they’d certified him and I had had proper advice they might’ve said to do as much as you can to stimulate him. She said just treat him like a normal baby...If he had been certified and I had spoken to a QTVI earlier and been given more advice. (Par14)

Education and social care professionals confirmed what parents had said—they often encountered children who were not referred to support until the CVI was issued or a firm diagnosis reached. One QTVI confirmed that some children go without support because they are not referred as ophthalmologists wait to see if vision improves. QTVIs stated infants and children with complex needs often presented late to their services.

it’s often the more complex needs children that...we haven’t had a referral to the service for, the education service and that is often because they are very young. (QTVI8)

Ophthalmologists state referral is offered before certification
While the majority of parents wanted support before certification, all 12 ophthalmologists, 3 ECLOs and 6 orthoptists stated they referred infants and children to QTVIs and education before they reached a stage where they offered certification.

You don’t have to be certified to get access to the VI teaching service. So we’re not depriving them of something by them not being certified...I don’t leap in and certify until I absolutely know where they are going to end up. And I’m still going to do all the other things that I would do for them, referral for low vision aids, for teaching service, specialist optometry and so on. It’s on my list but I wouldn’t say I, I personally don’t leap in and certify early on. (Oph3)

Paediatric ophthalmologists with close relationships with QTVIs and education stated they frequently discussed cases with QTVIs.

The first thing we tend to do is refer to VI team, even before certification, so they have a little bit of input from this team before they actually get information about registration, education team, enhanced service...We rely on them heavily as well. (Oph2)

Other areas had internal systems, working closely with orthoptists to ensure children were offered support when they needed it.

We have a system where the orthoptist would see all those children as well and would normally make a referral to learning support services but as I say, we wouldn’t certify them until their eye-sight deteriorates to appropriate standards...We’re also going to be referring people who
aren’t certified, from that point of view they are equally
going to have access to learning support. (Oph12)

Reasons for differences between parents’ and clinicians’

experiences

Numerous issues arose suggesting reasons for the disson-
ance between parents and ophthalmologists’ experi-
ences of referrals and certification. First, vision typically
improves with age and it is standard practice for ophthal-
mologists to wait months, in some cases years, to identify
a diagnosis.

It’s not clear cut. Certain conditions where it is, there are
certain where it isn’t. Some children with delayed visual
maturation, where you don’t really know how much their
vision will improve or to what level, so some of those it’s
quite a waiting game or you’re not able to assess the
child’s vision very accurately. And so you’re waiting for
more accurate clinical information. (Orth4)

The difficulty in measuring visual acuity (VA) and VI
in infants was mentioned by most ophthalmologists and
many stated they waited to offer certification until they
(or orthoptists) could measure VA or until test results
were received.

It can be extremely difficult to measure VA in
very young children, small babies. So what may appear,
for example, to be a non-seeing baby at 12 weeks with
something like delayed visual maturation may actually
turn out to be a baby with perfectly normal sight in six
weeks, or it may be much longer. I think it’s a delayed
visual maturation if I think it’s a baby with much more
severe visual diagnosis then it’s based on that. (Oph11).

What appears to be happening is that some ophthal-
mologists associate both the offer of certification and
referral to support with determining a firm diagnosis.

This issue may be a particular problem in hospitals
without paediatric ophthalmology specialist consultants.
While this is a small sample, consultants working in dif-
f erent tertiary centres stated parents often arrived at
their hospitals looking for support, having not been
referred during their visit to the first hospital.

If you have (ophthalmologists) who do not have specialist
training in paediatric ophthalmology they don’t under-
stand the pathways and the needs of the children and
they don’t think to refer them to the VI services and they
don’t think that a child can use a Low Vision Assessment
and they don’t think that actually the parents do actually
need help filling in the Disability Living Allowance form.
(Oph8)

Ophthalmologists who certified numerous children in
a year had more flexible attitudes to certification and
the Department of Health (DH) guidance.

I tend to be rather flexible with interpreting the guide-
lines. I don’t think visual acuity is the best way of doing
it in children. For instance, children with nystagmus might
have better vision than 6/18 but they obviously have
visual problems in terms of tracking and visual percep-
tion. Same with some of the milder forms of cerebral VI
where their visual acuity can be good but they are con-
fronted with a real world where they are overloaded with
visual information and they really struggle. I tend to
adopt a more functional approach to it. Based on fact
that acuity are guidelines rather than strict legal defini-
tions. (Oph7)

While interpreting the DH guidance flexibly can
benefit those on the borderlines or waiting for a diagno-
sis, it can also mean certification is offered inconsistently.

Another issue that arose was related to the offer of cer-
tification for children with complex needs. Some health
professionals stated they delayed the offer of certification
for these children because they felt their parents already
had so much to deal with and the benefits of certification
would not be worthwhile. Some ophthalmologists
stated they believed parents of infants/children with
complex needs might not want to discuss certification
but made this assumption without actually discussing the
offer with parents.

They’ve got multiple agency involvement it probably isn’t
very important to register that child if they are going
blind...many of our children who need that are multiply
handicapped are already getting a lot of support and I
think they feel they just don’t want another label.
(Oph6).

How to ensure early and consistent support

Multidisciplinary teams

Ophthalmologists stated working in multidisciplinary
teams reduced possible delays in certification or referrals
for support.

Many of our children come from multi-disciplinary teams
where we supply staff to check vision for these children.
(Oph6)

Across the eight local authority areas studied, four
areas held joint meetings between health, education and
SS, meetings were usually held once a term or a few
times a year. These joint meetings facilitated relation-
ships, making it easier for professionals to pick up the
phone or send an email when there is a problem—or
before an issue becomes a problem.

I don’t think we could offer a good service if we didn’t
work together as a team. (Orth3)

Direct referral pathways

To encourage prompt and consistent referrals, two areas
created referral forms to directly refer from health to
education as children were getting lost in systems and
referrals were not happening fast enough.
We get parents to sign a consent form to say they are happy to share information so we can liaise quite easily. (Orth3)

The DH has a recommended pathway for referral prior to certification, using the Referral of Visual Impairment (RVI). Although designed primarily for use with adults, it can be used with children as well. This referral does not require the patient (adult or child) to be certified and registered. In interviews, the RVI was not mentioned once by ophthalmologists (although they were not directly asked if they used the RVI). Ophthalmologists referred to QTVIs and social care without the need for a RVI.

**Eye clinic liaison officers**

Intermediaries such as specialist nurses, ECLOs and Family Support Unit provided a reliable referral route for parents to education and other services for specialist support. Parents who attended children’s hospitals, where most have roles such as ECLOs or Family Support Units, very much valued the information and support they offered.

When we had hospital appointments, we were up there quite a bit in the first few months...(ECLOs) would sit in on the appointments that we had...the information they first gave us, it was really helpful and it was nice to know someone was...We would’ve felt really isolated and not had a clue basically. I don’t really know what we would’ve done. (Par22)

Ophthalmologists stated these intermediary roles could potentially solve many of the problems faced by parents of infants and children with VI in providing timely information and support.

I think ECLOs are one of the things that would really help transform care for children with VI over the years. (Oph5)

However, even when these personnel were in hospitals, referring to these intermediary roles was not consistent, instead it was a subjective decision made by ophthalmologists. Parents were very disappointed when children’s hospitals/Centres of Excellence failed to provide support or information through these intermediary roles (as they were not referred). One parent, despite seeing orthoptists, consultants and optometrists at a children’s hospital, did not receive the support she expected.

I was supposed to get a journal— from birth up to about five, about certain things you can do with blind children to help them. I never received it and they promised me it. (Par18).

**DISCUSSION**

**Delayed referrals are unnecessary**

The referral to specialist support was often unnecessarily delayed in the process of establishing a diagnosis. Even though most childhood visual disorders are present from infancy, our research confirmed there is often a ‘prolonged period of uncertainty before the final diagnosis is achieved’. Previous research found 21% of parents waited less than a month to have their child’s VI diagnosed while 25% waited over a year. They also found 40% of children with multiple disabilities had to wait more than a year to have their VI diagnosed.

The purpose of certification is to formally refer a person to social services; however, it should not be the only prompt to a referral. While the ophthalmologists who took part in this study were aware of the importance of speedy referral of infants and children to the specialist education service, evidence from parents indicates that this is not consistent across the country. Delays in referring to education may be due to doctors waiting to obtain a firm diagnosis; health staff not understanding the role of the QTVI (particularly ophthalmologists without a paediatric specialism and, therefore, an understanding that ‘education’ is not just about the school years); and a lack of clear referral processes in some areas.

For parents of infants and children with complex needs, the delays could be longer as children are not always referred to ophthalmology departments or offered eye examinations and vision assessment by other health professionals. VI in children with complex needs is often underidentified and can take some time to diagnose. The number of children with neurodevelopmental disorders is contributing to the increase in the prevalence of VI, yet diagnosing VI in infants and children with complex needs is more difficult too, and therefore this cohort is more likely to be overlooked.

This suggests not all eligible infants and children are being certified and that they and their families may be missing out on important financial and practical support. Early support is crucial for infants and children with VI and their families, to support children’s cognitive development, communication, social and independence skills. Referrals to the specialist teacher (QTVI) from the local authority education advisory service is not dependent on certification.

**Refer when support is needed, not when certification is offered**

The compassion shown by ophthalmologists, orthoptists, optometrists and ECLOs was clearly apparent, but this is not enough—these sentiments need to translate into actions, so that infants and children with VI are promptly and consistently referred to the support they need to secure the best start to their lives. There should be a formal mechanism for triggering QTVI and social care support even if there is uncertainty about diagnosis and prognosis, such as formal referral pathways.
Previous studies found value in infants and children managed by multidisciplinary teams ‘to ensure comprehensive and integrated intervention.’ This research demonstrates the value of multidisciplinary teams as well as intermediaries to ensure referrals and support are offered to infants and children with VI.

A consistent mechanism for triggering educational and social support for children with VI and their families, even when diagnosis and eventual level of visual function are uncertain, would improve child development and families’ experience.

**LIMITATIONS OF RESEARCH**

**Bias**

Overall, the number of participants interviewed in each category was small; as such, the findings should be considered indicative; however, saturation/replication levels were reached in all interview groups, suggesting confidence in the findings.9 We sought to reduce the potential for selection bias by selecting from a broad group of parents; however, all children needed to be certified. We aimed to minimise reporting bias by defining the research questions at the beginning of the project and the researcher (TB) was experienced in the topic and with the interview population.5 In addition, the advisory group provided objective guidance in forming the research questions and in reviewing the findings.

**Sample**

The original aim of the research was to interview parents from the same areas as the professionals (similar to the sample researched for the adult research).1 As there are fewer infants and children certified compared with adults, it was decided to widen the sample to include all parents of infants and children certified in all parts of England. As such, it is difficult to provide a snapshot of each area. Despite this limitation, the research is able to identify key themes arising from each interview cohort.

In addition, other professionals involved in the certification and registration process were occasionally mentioned by interviewees (eg, community paediatricians, support workers delivering portage (home-visiting educational service for preschool children with additional support needs and their families)); however, they were not interviewed as the research focused on the key workers providing support and information to parents and carers of infants/young people with sight loss.

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