What do parents of children with dysphagia think about their MDT? A qualitative study

Emma Cowpe (Jebson),1 Ben Hanson,2 Christina H Smith3

ABSTRACT

Objectives: To seek the experiences and perspectives of parents caring for children with dysphagia, with emphasis on their experiences of working within their child’s multidisciplinary team (MDT).

Setting: This research was completed in community settings, within families’ homes across the UK.

Participants: 14 families self-selected to participate in the study. Criteria specified that participants must care for a child under the age of 18 and to decrease ambiguity the term ‘diagnosis of dysphagia’ was defined as the need for modified (thickened) fluids. Exclusion criteria: caring for an adult over the age of 18; diet and fluid modifications for reasons other than dysphagia (eg, for symptomatic treatment of gastro-oesophageal reflux disease. Participants were interviewed within their homes using a semistructured questionnaire and data was analysed using a descriptive phenomenological approach through use of thematic coding and constant comparison. Themes and relationships were inductively generated from the data.

Results: Participants universally expressed a desire to be involved with their child’s MDT; this study identified the following facilitators and barriers to collaboration: accessing services, professional knowledge and professional skillset. Participants described three means of responding to these barriers: reacting emotionally, seeking solutions and making decisions.

Conclusions: This study recorded in-depth reports of participants’ experiences of working with healthcare providers. Despite government-driven efforts towards person-centred healthcare and social care, participants shared accounts of times when this has not occurred, describing a negative impact on the well-being and quality of life of their child and family.

BACKGROUND

Swallowing difficulties (dysphagia) can vary in aetiology, symptomology and severity and affect children with a variety of medical diagnoses. Up to 90% of children with neurodevelopmental disorders such as cerebral palsy and Down syndrome experience signs of dysphagia including coughing, choking, chest infections, poor weight gain and breathlessness,1 often from birth or early infancy.

Children with gastro-oesophageal reflux disease (GORD) and those born prematurely are also at increased risk of childhood dysphagia.2–4 Thus dysphagia can be a symptom of different underlying medical conditions and is diagnosed and treated by a variety of medical specialists.

Children with complex and pervasive needs require support in a range of inpatient and outpatient specialist settings including hospitals, clinics and schools. Children may be in contact with a range of health and care professionals including paediatricians, general practitioners (GPs), speech and language therapists, physiotherapists, dieticians and social workers; for these children and their families well-coordinated input is essential for provision of individualised care.5–7 Multidisciplinary teams (MDTs) are commissioned to provide joined up, patient-centred support and seek to place children and their families at the centre of decision-making processes.1 8 9 Government initiatives such as the Early Support keyworking scheme6 and the Common Assessment Framework10 have been set up throughout the UK to improve care quality by providing a framework for regular MDT discussion; avoiding repetition of information and reducing carer burden; and providing a single point of contact for better coordination of care and support.6
A number of studies have explored parent experience of caring for children with dysphagia although typically research has focused on the needs of carers of children with complex needs which may or may not involve dysphagia as a symptom, or has been confined to specific settings such as schools or Child Development Centres. Mahant et al, Craig et al and Sullivan et al have studied the experiences of parents of children requiring gastrostomy feeding; these studies highlighted the importance and value placed by parents on oral feeding, and acknowledged the complexities involved in decision-making when considering how to balance risks of oral intake with maximising quality of life (QoL) for children with dysphagia. Recommendations for practice include the need for good-quality information and support for families and the need for professionals to understand caregiver’s perspectives and priorities and to adopt a model of partnership working with the families under their care. Studies looking at specialist feeding clinics within the USA and the UK describe the intricate and multifaceted nature of feeding and swallowing disorders and highlight the importance of multidisciplinary working to ensure holistic, child-centred and family-centred assessment, treatment and support. This paper seeks to build on this research and explore the issues pertinent to families caring for children with dysphagia in the UK, and to gain their perspectives on the care they have received.

METHOD
Participants
This cross-sectional study involved parents and carers of children with a diagnosis of dysphagia, aged between 2 and 11 years. Information about the study was distributed through special schools, charities and local and national support groups in the form of letters, online forums and newsletters. Parents self-selected by contacting the researcher based on the information provided, and gave written consent to participate in the study. Fifteen parents were initially recruited; one interview was discounted from analysis as the individual with dysphagia was 24 years old. Analysis was completed with interviews from 14 parents; the children’s characteristics are summarised in table 1.

Procedure
Audiorecorded, in-depth interviews were considered the most appropriate form of data collection; providing participants with dedicated time and space to share their individual experiences and perceptions.

Participants were contacted and interviews were scheduled at their convenience. Thirteen parents were interviewed at the family home and one parent was interviewed at her place of work. Although all participants consented to audiorecording, each parent was reminded of their right to withdraw from the study at any time. Given the emotive nature of the issues being discussed, breaks were offered throughout the interview process to enable a period of time ‘off recording’ if required. Informal discussion prior to and following each interview allowed for debriefing and reflection without participants feeling under pressure of being recorded.

Semistructured questionnaires formed the basis of interviews; this provided a degree of structure and standardisation in questioning while allowing for deviation from the ‘script’. Adopting a semistructured approach allowed for more natural conversation during interviews, avoiding an overtly formal interview style and maximising the richness of each participant’s account. Given the sensitivity of the topics discussed, freedom to deviate from structured, formal questioning was essential. Parents were asked questions, including open/closed and content-mapping/dimension-mapping questions, to gain general and more specific information relating to their child’s swallowing history, initial support received, ongoing needs and current support. Interviews took approximately 1 h and each audiorecording was transcribed prior to analysis.

The researcher kept a journal for additional notes throughout the data collection process. Although journal entries were not formally used in data analysis, they were used to document preliminary codes and themes that began to emerge at the time of each interview, as well as notes on follow-up questions that had been asked, and general attitudes of interviewees at the time (eg, positivity/negativity).

Analysis
Interviews were transcribed and returned to participants to verify accuracy. Two parents made minor clarifications to comments within their manuscript; one parent providing further detail of the chronology of her daughter’s medical interventions and another providing the full name of their child’s thickener. The parent’s amendments were recorded prior to coding.

Qualitative analysis methods were identified as most appropriate for this data set, given the richness of participants’ accounts and the complexity of issues discussed. Descriptive phenomenology was selected in order to best describe participants’ experiences and to uncover common themes within and between participants’ accounts. Methods of thematic coding and constant comparison were employed to generate codes inductively from the data, rather than impose preconceived themes constructed by the researcher. In turn, the codes that emerged led to the construction of themes. During coding a transcript was selected at random and manually coded line by line with short phrases summarising each ‘unit’; examples included negative emotions, creative problem solving and service flexibility. Units varied in length; comprising of phrases, sentences or a number of lines, with one code applied to each unit.
Additional transcripts were then added to the set and manually coded, with codes being continually compared within and between scripts and amended where required. As analysis progressed codes were grouped into broader themes. For reliability of the coding, 25% of the coded transcripts were validated by an independent researcher with experience in qualitative data analysis.

RESULTS

Fourteen participants completed the study; participants cared for a child between the ages of 2 and 11 years; mean age 5.8 years. Of the 14 participants involved, 12 were under the care of a paediatrician when their child’s swallowing difficulties were first investigated; 13 had contact from a speech and language therapist; 8 from physiotherapy; 8 from dietetics and 7 were seen by an occupational therapist. Eight families had contact from a health visitor; eight from the GP; four had involvement from school or nursery staff; one family had input from a school nurse and one from respite staff.

One overriding theme emerged from the data, present in all 14 interviews: all parents and carers expressed their desire to be involved in their child’s care through close involvement in the MDT. Three subthemes emerged relating to factors particularly facilitating or hindering their ability to be involved within the MDT. A further three subthemes emerged as parents described their responses to the barriers they have experienced (figure 1). Participants’ quotes are provided verbatim alongside their identifying number as allocated in table 1.

**Table 1** Individual child characteristics

<table>
<thead>
<tr>
<th>Parent ID number</th>
<th>Medical diagnoses</th>
<th>Age at diagnosis of dysphagia</th>
<th>Nutritional intake</th>
<th>Route to recruitment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CP epilepsy, GORD</td>
<td>Under 1 year</td>
<td>Oral diet</td>
<td>School</td>
</tr>
<tr>
<td>2</td>
<td>Lissencephaly</td>
<td>18 months</td>
<td>Oral diet</td>
<td>School</td>
</tr>
<tr>
<td>3</td>
<td>No diagnosis</td>
<td>8 months</td>
<td>Oral diet</td>
<td>School</td>
</tr>
<tr>
<td>4</td>
<td>CP, hydrocephalus, epilepsy, VI, HI</td>
<td>Under 1 year</td>
<td>PEG plus oral tastes</td>
<td>School</td>
</tr>
<tr>
<td>5</td>
<td>CP, chronic lung disease, GORD</td>
<td>1 year</td>
<td>Oral diet (previously PEG)</td>
<td>School</td>
</tr>
<tr>
<td>6</td>
<td>Alagille syndrome</td>
<td>Under 1 year</td>
<td>Half oral, half PEG</td>
<td>School</td>
</tr>
<tr>
<td>7</td>
<td>DS, CHD</td>
<td>Under 1 year</td>
<td>Oral diet</td>
<td>Charity</td>
</tr>
<tr>
<td>8</td>
<td>DS, GORD</td>
<td>5 years</td>
<td>Oral diet</td>
<td>Charity</td>
</tr>
<tr>
<td>9</td>
<td>Congenital cytomegalovirus, GDD</td>
<td>1 year</td>
<td>Oral diet</td>
<td>Family worker</td>
</tr>
<tr>
<td>10</td>
<td>DS, CHD, GORD, HI</td>
<td>Under 1 year</td>
<td>Oral diet</td>
<td>Charity</td>
</tr>
<tr>
<td>11</td>
<td>DS</td>
<td>Under 1 year</td>
<td>Oral diet</td>
<td>Charity</td>
</tr>
<tr>
<td>12</td>
<td>DS</td>
<td>Under 1 year</td>
<td>Oral diet (previously PEG)</td>
<td>School</td>
</tr>
<tr>
<td>13</td>
<td>DS, GORD, VI, HI</td>
<td>2.5 years</td>
<td>Oral diet</td>
<td>School</td>
</tr>
<tr>
<td>14</td>
<td>DS, Hirschprung’s disease, GORD</td>
<td>3 years</td>
<td>Oral diet</td>
<td>Charity</td>
</tr>
</tbody>
</table>

CHD, congenital heart disease; CP, cerebral palsy; DS, Down syndrome; GDD, global developmental delay; GORD, gastro-oesophageal reflux disease; HI, hearing impairment; PEG, percutaneous endoscopic gastrostomy; VI, visual impairment.

**Figure 1** Themes and subthemes generated from data.
We know our children better than [professionals] do as individuals; they have to take our views on board and respect our opinions.7

Facilitators and barriers
The facilitators and barriers to parents’ collaboration with the MDT were categorised into three separate subthemes: (1) accessing services, (2) professional knowledge and (3) professional skillset, as below:

1. Accessing services
Parents described three main issues around their ability to access services; time, staffing and flexibility of service delivery. When asked how satisfied they were with the support they had received for their child’s dysphagia, nine parents reported some degree of dissatisfaction with service accessibility and availability.

They won’t come and see him at home, which is a real shame.2

My daughter was ill for the whole winter. [The community Speech and Language Therapist] came as a favour to check her swallow, but there’s a fine line now she’s in school, with whose remit it is.12

They said they didn’t have enough staff [in hospital] to give support with her feeding.3

It was quite a few years before they had a permanent Speech and Language Therapist at school.4

Those who had felt largely satisfied with their support acknowledged the same qualities as important; reporting easier access to help and advice due to flexibility and availability of health and care staff.

The [respiratory] physios are really good. They come in [if she is unwell] and monitor her sats. It saves going to A&E and they take swabs that can be sent off.5

When we went for the Videofluoroscopy the Speech and Language Therapist came with us...Every six weeks she comes here to see him.7

We’re lucky that we have had support. They’re always at the other end of the phone if we need it.11

We have much more support now.10

2. Professional knowledge
Parents attributed their health and care professionals with a level of expertise and specialist knowledge; however, some parents felt that this knowledge was poorly demonstrated by professionals at crucial time points when it was needed; particularly in the early stages of seeking support and specialist advice for their child. Five parents received misdiagnoses of their child’s dysphagia, which resulted in a wait of up to 5 years for an accurate diagnosis and to receive specialist intervention.

I kept showing her feeding to so many different consultants and no one spotted it ...they kept saying ‘it’s a virus, she’s fine’. Thirteen times in her first year she was on antibiotics for chest infections.4

I’m surprised the paediatrician didn’t signpost [specialist assessment] more clearly; she knew what [Lissencephaly] was and its implications.2

I just couldn’t believe it took them so many years to take it seriously; it amazed me.5

Those with positive experiences also highlighted the importance of well-demonstrated expert knowledge. These families reported an increase in their knowledge of their child’s condition as a result.

[The specialist’s] advice, which I found useful was that there were three things to look at; if he was getting pneumonias...if he was failing to thrive...or if it as taking a very long time to feed him.2

[The specialist] had lots of questions; over the phone I was talking to her and she said my daughter had reflux and dysphagia, just from the phone call.8

3. Professional skillset
Parents provided a list of qualities they value in healthcare professionals (table 2), placing the greatest importance on communication skills. Parents valued professionals who demonstrated a willingness to listen and who actively sought their opinions.

Parents have the answers even if we don’t have the terminology.12

More the medical side than the community side have no respect or value for what the parents have to say...it’s quite nice when you find people who actually listen.10

The interpersonal (skills) are as important as the clinical (skills).13

Parents also described the importance of a positive working relationship between healthcare professionals, themselves and their child:

In the medical profession it’d hurt me when they wouldn’t even address my son at appointments; they would talk to me about him, in front of him, without even saying hello.2

[I value] people that are good at interacting with my daughter...people who work with her rather than just talking to mum.9

Every six weeks [our Speech and Language Therapist] comes here to see [my son] ... he really likes her.7

Parents with negative experiences described times when they felt unheard and ‘on a different page’ to professionals. Parents raised concerns that too strong a
focus was placed on health and risk, at times to the detri-
ment of their child’s emotional well-being and QoL.

I felt that [the professionals] were very risk-averse…their
first concern was to ensure the risk was minimized com-
pletely and utterly, rather than recognising that there was
a risk but that it wasn’t too great yet.2

We told school that if the children are having any treats,
our daughter can have them too…There was an event
where all the children were having hot chocolate with
marshmallows. They wouldn’t let our daughter have one
and she got upset.8

Parents discussed the importance of information
sharing, identifying a need for up-to-date, jargon-free
information to maximise their understanding and
involvement. Parents reported a similar need for more
effective information sharing between professionals
within the MDT. Parents identified times when poor col-
laboration had negatively impacted on the well-being of
their child or on themselves as caregivers.

[The professionals] don’t liaise with each other at all…
although they are all in the same building.1

I repeat myself all the time…it’s hard to keep bringing
up the past.5

In school there is a lack of communication…at one time
they had stopped feeding her. I felt so upset by it.6

Responses to barriers
Parents’ responses to the challenges they have faced
were summarised in the following themes: (1) reacting
emotionally, (2) seeking solutions and (3) making
decisions.

1. Reacting emotionally
Parents who encountered barriers to support had
reported high levels of anxiety, frustration and distress
and feelings of being unheard. Parents who experienced
difficulties at the early stages of their child’s care
described associated feelings of guilt and helplessness at
their child’s failure to thrive, and frustration in their
search for answers. All parents with these experiences
reported a significant impact on their emotional
well-being.

I struggled with it; it was awful. It made me extremely
depressed because I just didn’t know what was going on,
and no one seemed to be paying any attention.3

I felt that this was very poorly handled in terms of the
emotional impact it can have on a parent, to say you
might not be able to feed your child yourself.2

[The Speech and Language Therapist] was only focussed
on communication. She asked us every time how feeding
was going; I dreaded her asking me that…I hated telling
her he just couldn’t eat, but she just said to keep trying.
It was so demoralising…13

2. Seeking solutions
All parents described times when they had adopted a
proactive approach towards seeking and receiving help:
researching into their child’s medical conditions, recom-

dended treatment, medication and available support.
Parents report using the knowledge they had gained to
actively request or refuse specific services or
interventions.

It’s been more parent-driven; no one takes ownership of
the problems.13

<table>
<thead>
<tr>
<th>ID number</th>
<th>Qualities listed</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Accessible; two-way communication; being honest without being too negative</td>
</tr>
<tr>
<td>2</td>
<td>Flexibility in appointments; listening and walk alongside you; treating you as a thinking person; addressing my son at appointments</td>
</tr>
<tr>
<td>3</td>
<td>Interacting with my daughter; working well with other professionals; following through with what they say; talking to who they say they will; when you feel they want to help you and are on your side; making notes and giving you copies; being contactable; being approachable; being honest; not being overworked</td>
</tr>
<tr>
<td>4</td>
<td>Being there; being contactable; liaising with each other; being trustworthy</td>
</tr>
<tr>
<td>5</td>
<td>Being caring, friendly; knowing my child so I don’t need to repeat myself; record keeping</td>
</tr>
<tr>
<td>6</td>
<td>Good people skills, sensitive to my child’s needs and my concerns; good communication; sympathy; seeing my child as an individual</td>
</tr>
<tr>
<td>7</td>
<td>Listen and take the parents’ views seriously; communication with parents; explaining jargon; sharing information</td>
</tr>
<tr>
<td>8</td>
<td>Showing expertise in their field; treating children as individuals; not labelling children</td>
</tr>
<tr>
<td>9</td>
<td>Professionals working together; putting my child first; sharing information</td>
</tr>
<tr>
<td>10</td>
<td>Listen to parents; being reliable</td>
</tr>
<tr>
<td>11</td>
<td>Approachability; being able to explain things</td>
</tr>
<tr>
<td>12</td>
<td>Honesty; encouragement; new ideas; continuity</td>
</tr>
<tr>
<td>13</td>
<td>Approachability; friendliness; warms; engaging our child; active, unhurried listening; empathy</td>
</tr>
<tr>
<td>14</td>
<td>Asking questions of me; active listening; acting on what we say; not labelling our child</td>
</tr>
</tbody>
</table>

Table 2 Valued professional qualities

Parents reported the value of peer support and identified times when they have felt empowered by the knowledge they have gained from other parents and carers in a similar situation.

At first I didn’t want a PEG but...I spoke to other parents who said they’re really good.5

It’s been useful meeting other parents, to talk about things with them and share experiences.11

I wrote to the GP and asked [for thickener], because somebody else used it and gave me some sachets.1

Talking to other parents [helps] too, not just professionals.4

Some parents used the knowledge they had gained to directly challenge professional advice or seek second opinions; six parents contacted private professionals for second opinions and two families contacted overseas specialists for advice.

I see a private person because I’m unhappy with the support I get.3

I discovered a Speech and Language Therapist in Brazil... she said my daughter has [dysphagia]...I was shocked; she’d had problems all that time and we had been told it was nothing.8

3. Making decisions
Parents described times when they had taken matters ‘into their own hands’ in the day-to-day management of their child’s difficulties. Parents reported developing their own strategies based on a ‘trial and error’ approach with their child; employing a range of techniques to maximise their child’s swallowing safety. Techniques included postural adaptations, changes to mealtime utensils and environmental strategies.

We had been spoon-feeding him water...he was alright with spoonfuls; any more and he coughs.5

She would have rice pudding and I would give her a dummy after to help her swallow...when we put the dummy in then she would initiate the swallowing action.6

I take her drink away for a second [between sips] to give her a chance to breathe before carrying on.10

A similar approach was used to manage difficulties encountered with fluid thickeners. Parents reported a range of difficulties using thickeners on an everyday basis including lumpiness; problems maintaining fluid consistency over time; managing liquid medications and modifying their child’s diet or fluid consistency when unwell or tired. Parents described a number of techniques that they developed using their knowledge and experience of their child, in order to avoid such difficulties.

We use a lower dose [of thickener] in milk, it’s just enough to slow it down. In his juice we use slightly more because the cups have bigger holes in.7

We recognise when it needs to be thicker, like when she’s poorly.13

We do thicken some of his medication...we give medication in a syringe too so we can control how he has it.11

DISCUSSION
This study proceeded with explorative aims to seek parents’ experiences of their involvement with their MDT. As a qualitative study this does not seek to generalise findings to the wider population, particularly considering the age ranges of children involved and the geographical spread. Qualitative methodologies employed were deemed appropriate for a study of its size and nature.23 27 34 28 Reliability and validity were established during analysis through inter-rater agreement and participant validation.

Participants acknowledged the need for multidisciplinary working to ensure holistic management of their child’s difficulties, as reported elsewhere within the literature.23–27 35 Parents encountered a number of barriers to accessing appropriate services in the early stages of their child’s life and described a range of negative physical and emotional effects on themselves and their children. While resources may be limited, improved staffing and time for direct family support may have reduced the impact of these barriers. Shared resources such as shared workspaces for health professionals may serve to improve collaboration and reduce carer burden. More flexible models of service delivery, such as improved crossover between school and community services, may have enabled some families to access support quicker and encounter fewer moments of delay and procedural ‘red tape’.

Families who received repeated misdiagnoses of their child’s condition described higher levels of dissatisfaction in the MDT and negative impacts on child and family health and well-being. Participants reported their frustration and upset in knowing there was ‘something wrong’ with their child’s health yet failing to receive appropriate support and diagnostics. It could be argued that the parents who sought second opinions from private and overseas professionals, did not receive advice that could not have been provided by their child’s MDT.

Participants expressed their need for healthcare professionals to work proactively, particularly in the early stages of seeking and receiving a diagnosis of dysphagia. Previous work has shown that professionals who work holistically and proactively are better able to anticipate a
child's needs, referring for specialist intervention before a critical need arises. Parents acknowledged the importance of communicative and interpersonal skills in facilitating a trusting relationship between family members and professionals, particularly when balancing considerations around risk and QoL. This issue was particularly pertinent for parents caring for children considered for percutaneous endoscopic gastrostomy feeding; an issue requiring great sensitivity and discussed elsewhere in the literature. The need for good-quality information was highlighted to enable informed decision-making and effective communication was considered to reduce the burden on parents to repeat their child's medical information to different members of the MDT. Keyworking schemes such as Early Support and shared documentation may be one step towards this end in order to streamline communication and to provide a single point of contact for families caring for children with complex needs; however with current financial and structural changes to the healthcare and social-care system such resources may be limited.

The emotional, financial and physical burdens of caring for a child with a disability are well reported. Parents reported a significant impact on the emotional and physical well-being of themselves and their children, acknowledging a need for proactivity on the part of the MDT, particularly in the early stages of receiving diagnoses and requesting specialist support. Parents utilised a range of creative approaches in managing the everyday difficulties of using thickening agents and drew on the experiences of themselves and others to develop solutions that worked for them.

**Limitations:** As a self-selecting cohort the sample in this study is open to a degree of bias. Because other agencies such as schools, charities and support groups were involved in disseminating the information a response rate could not be calculated as it was unclear how many parents had come into contact with the initial information regarding the study. Social demographics were not taken for this study; further research exploring the experiences of socially isolated families would be beneficial. Parents were invited to check the accuracy of transcripts; this feature of the study was not accounted for in the initial stages and as such may have been considered an additional burden which had not been anticipated by parents at the recruitment stage.

Further developments of this research exploring in more depth the different types of service provided to families with children with dysphagia would be beneficial. Similarly although social demographics were not taken for this study's further research, this study could be expanded by studying socially isolated or ‘hard to reach’ families.

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**Contributors** EC and CHS accessed the data for the study in its entirety prior to and during write-up; BH contributed to the conception and design of the study, critical revision of draft versions of manuscript and approval of final version. EC was responsible for participant recruitment, data collection and analysis, write-up of manuscript and had access to all data. CHS contributed towards research design, provided advice and supervision on research processes and data write-up, and critically revised and approved the manuscript (draft and final versions).

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