Peer support for parents of disabled children part 1: perceived outcomes of a one-to-one service, a qualitative study

Article (Submitted Version)


This version is available from Sussex Research Online: http://sro.sussex.ac.uk/id/eprint/62346/

This document is made available in accordance with publisher policies and may differ from the published version or from the version of record. If you wish to cite this item you are advised to consult the publisher’s version. Please see the URL above for details on accessing the published version.

Copyright and reuse:
Sussex Research Online is a digital repository of the research output of the University.

Copyright and all moral rights to the version of the paper presented here belong to the individual author(s) and/or other copyright owners. To the extent reasonable and practicable, the material made available in SRO has been checked for eligibility before being made available.

Copies of full text items generally can be reproduced, displayed or performed and given to third parties in any format or medium for personal research or study, educational, or not-for-profit purposes without prior permission or charge, provided that the authors, title and full bibliographic details are credited, a hyperlink and/or URL is given for the original metadata page and the content is not changed in any way.
This is the pre-peer reviewed version of the following article: V. Shilling, S. Bailey, S. Logan, C. Morris Peer support for parents of disabled children part 1: perceived outcomes of a one-to-one service, a qualitative study which has been published in final form at http://onlinelibrary.wiley.com/doi/10.1111/cch.12223/full. First published online 17th December 2014, Manuscript Accepted: 18 November 2014. This article may be used for non-commercial purposes in accordance with Wiley Terms and Conditions for Self-Archiving.

Title: Peer support for parents of disabled children part 1: perceived outcomes, a qualitative study

Running title: peer support for parents of disabled children

Authors:
Valerie Shilling, Research Fellow in Child Health [1]
Sarah Bailey, Associate Research Fellow in Child Health [1]
Stuart Logan, Cerebra Professor of Paediatric Epidemiology, Director Institute of Health Research, Director - NIHR Peninsula Collaboration for Leadership in Applied Health Research and Care [1]
Christopher Morris, Senior Research Fellow in Child Health [1]
[1] Peninsula Cerebra Research Unit & NIHR PenCLAHRC, University of Exeter Medical School, UK

Correspondence:
Valerie Shilling
PenCRU, Child Health Group, University of Exeter Medical School
Veysey Building, Salmon Pool Lane
EX2 4SG, UK
v.shilling@exeter.ac.uk
**Data sharing statement:** Additional data extracts and full topic guides are available from the corresponding author: v.shilling@exeter.ac.uk
ABSTRACT

Background: Parents of disabled children are encouraged to seek peer support. Delivering one-to-one support requires resources; therefore investigating how these services may impact on families and those providing the service is important when evaluating such services.

Methods: A qualitative study involving semi-structured interviews and focus groups. Participants were 12 parents and 23 befrienders who had contact with the Face2Face one-to-one befriending service in Devon and Cornwall during a 12 month period, and 10 professionals from health, social care and education services.

Findings: Shared experience was perceived central to successful peer support and was a catalyst for other elements of support, enabling parents to (i) learn from the experience of others, (ii) speak freely in a safe and non-judgemental environment, and (iii) receive support and encouragement from their befriender. These elements underpinned perceived outcomes for both parents providing and receiving support. Outcomes for parents receiving support centred on emotional stability, personal growth and reduced isolation. Supporting parents experienced positive outcomes through their training, mutual support and the feeling that they were helping others. Parents and befrienders appeared to benefit through expanding their social network. Nevertheless providing support was reported to create emotional burden and concerns for befrienders around their performance, and also required a substantial time commitment.

Conclusions: Befrienders as well as parents perceived positive outcomes from their involvement in peer support although there is also potential for less positive impact on those offering support.
Key messages:

Shared experience between parent and befriender underpinned the outcomes of peer support. Important outcomes for parents included reduced isolation, emotional stability and personal growth.

Befrienders also experienced positive outcomes from training, mutual support and the feeling that they were helping others, but were also at risk of emotional burden and concerns around their performance and the requisite time commitment.

Capturing all potential beneficiaries of a peer support service is important in evaluations and will influence their perceived value.
INTRODUCTION

There is ample evidence that parents of disabled children are at risk of physical and psychological health problems, which are both pervasive and likely to worsen over time (Brehaut et al. 2011, Brehaut et al. 2009, Emerson et al. 2010, Gerstein et al. 2009, Miodrag and Hodapp 2010). Parents often seek support from their peer group; health services in UK and USA actively encourages peer support (Committee on Early Childhood Adoption and Dependent Care 2011, NHS Choices 2011, HM Government 2012). Our recent systematic review (Shilling et al. 2013) reported that peer support in this context was highly valued and perceived as beneficial, however quantitative studies that tried to measure specific impacts on parent health and wellbeing were less consistent in reporting positive effects.

Peer support can be considered a complex intervention, and this influences approaches to evaluation (Craig et al. 2008). We sought parent and professional perspectives regarding a one-to-one peer support service offered to parents of disabled children in South West England by Face2Face (Scope 2014) (see supplementary file S1 for further details of the support service) to understand the perceived outcomes, and influential components of the intervention. Previous research on peer support in this context has tended to focus on the recipients of support as the primary beneficiaries however, consistent with findings from our systematic review (Shilling et al. 2013) and the helper therapy principle (Salzer and Shear 2002, Solomon 2004), we anticipated potential benefits for parents offering support as well as those receiving support and that these benefits may stem from the same underlying constructs of peer support. In this first of a pair of papers we explore what are the perceived outcomes of peer support in this context and for whom.
METHODS

Patient and public involvement
We convened a stakeholder group to advise on all aspects of the evaluation. The group included parents of disabled children, members of Face2Face and representatives from local and national charities, Local Authorities, and a general practitioner (GP). Members of the group contributed to: (i) setting the research questions (ii) developing participant information leaflets, data collection forms and interview topic guides (iii) suggesting key professions to interview (iv) advising on emerging themes from analysis (v) interpreting the findings and key messages (vi) disseminating findings.

Recruitment and sampling
Local coordinators of two Face2Face services, in Devon and Cornwall, identified eligible parents through case records. Families in crisis were not approached to avoid overburdening those who were vulnerable. All other parents and befrienders who had contact with the service in a 12-month period were eligible and invited to participate in the study. We purposively sought to interview parents who were offered peer support but declined or received very few visits. In addition, we conducted 10 interviews with a purposively sampled group of professionals working with disabled children and their families.

Interview and focus group procedure
Interview topic guides explored areas pertaining to the specific research question discussed here, and other questions addressed within the broader evaluation (Supplementary File S2). Interviews with parents and befrienders took place at the participant’s home; professionals were interviewed at their place of work or at their home depending on their preference. Interviews were audio-recorded. Two parents wanted to take part in the research but were not
comfortable with the conversation being recorded; notes of the participant’s responses were made during the interview instead. Interviews were conversational in tone and the pace and duration was guided by the participant. On average, interviews lasted 45-60 minutes. Focus groups were used with befrienders in Cornwall. The groups were facilitated by a member of the research team and followed the same topic guide and structure as the interviews.

Demographic information gathered included the child’s age and condition. Postcodes were used to calculate Index of Multiple Deprivation (IMD) scores, used as indicators of deprivation in small geographical areas. (Office of National Statistics 2001) IMD scores enable ranking of area deprivation based on a combination of income, education, health, housing, services and living environment.

Interviews and focus groups were transcribed verbatim and transcripts were reviewed and used to inform subsequent interviews. Topic guides were adapted to reflect and test the developing analysis. Reflexive notes were kept to record systematically contextual details of the interviews.

Analysis

Analysis followed the Framework Approach. (Ritchie and Spencer 1994, Pope et al. 2000) Two members of the research team (VS/SB) read the transcripts and developed the thematic framework – incorporating key concepts and issues identified a priori from the literature and those emerging from the data. From the literature we anticipated several themes which were included in the framework a priori. These were: shared social identity, learning from the experience of others, personal growth and supporting others.
The framework was then applied to the transcripts which were indexed by themes and subthemes using NVivo10. Data from the focus groups were analysed at the group rather than individual level. VS indexed and charted all of the material; 25% of material was also indexed by SB to check for consistency in the application of the index. Themes and subthemes with greater than 2% disagreement between reviewers and where Kappa was <0.4 were considered to have unacceptable reliability. The content of these themes and subthemes were reviewed by VS/SB and differences in interpretation were resolved through discussion.

The data were then extracted and summarised in charts, grouped by themes and subthemes and incorporating field and reflexive notes. The charts were used to compare and contrast across groups and explore relationships between outcomes and aspects of peer support. During the later stages of mapping and interpretation of the data, we convened an expert reference group of befrienders from Face2Face to discuss the developing analysis. Some issues were perceived to be more salient than others; hence our interpretation of findings was influenced by the original research objectives as well as the themes emerging directly from the data.

Where direct quotes are presented, extracts from parents’ interviews are followed by identification codes beginning ‘P’, those from individual befriender interviews beginning ‘B’, from befriender focus groups ‘FG’ and those from professionals, ‘PR’. For all extracts, square brackets containing three dots […] indicate short sections of omitted speech; square brackets containing text indicate explanation added during transcribing or analysis, usually to replace a name.
FINDINGS

Participants

A total of 69 parents and 53 befrienders from Devon and Cornwall were invited to participate. Of these, 14 parents responded and 12 were interviewed. Thirty-one befrienders took part in some aspects of the evaluation; 23 in the qualitative research. Thirteen had individual interviews, 9 took part in focus groups and one responded by post (Table 1).

Ten professionals were interviewed: two members of staff at different Children’s Centres, two senior staff from integrated children’s services at different local authorities, a GP and a paediatrician, two community nurses, and two school-based educational support staff. Half of the professionals were not aware of the Face2Face service specifically; however all were familiar with the concept of peer support. We found little difference in the tone of the views expressed by professionals, parents and befrienders. As such, professionals’ perceptions of the intervention and outcomes were considered alongside those of the other groups; we highlight in our findings where notable differences occurred.

What are the perceived outcomes of peer support?

Our analysis indicated outcomes for parents receiving support and for befrienders delivering support, as perceived variably or consistently by parents, befrienders and professionals. We identified key elements of peer support that appeared to act together to enable these outcomes to be experienced. Although distinct, these components are related and appeared largely dependent on the perception of shared experience between befriender and parent.
**Key outcomes for parents**

**REDUCED ISOLATION**

All participant groups discussed the isolation experienced by parents of disabled children and the potential for peer support to reduce this feeling. Physically being with other people was not seen as a necessary prerequisite of reduced isolation. Rather, the knowledge that parents were not the only people dealing with the same problems, and that their feelings were not unusual, was sufficient.

Parents of disabled children reported feeling different to people who do not have disabled children. This can lead to feelings of extreme isolation and a sense that other groups of people simply cannot understand what life is like. Conversely, participants told us that they identify with other parents of disabled children in a way that they can’t with other people. They differentiate this support from that received from friends and extended family or professionals which, however well meaning, can lead to feelings of resentment and increased isolation because parents inevitably compare their children with others who do not have additional needs.

Shared experience was seen by all participant groups as a necessary component of peer support. Opinions on ‘how much’ shared experience, in terms of matching by diagnosis, is sufficient for effective support were divided. All participants agreed that parents of disabled children have a wealth of shared experience which enables them to speak openly with each other and give support. An adjunct to shared experience, facilitating reduced isolation, was the importance of a safe and supportive environment in order for parents to feel comfortable to speak freely; continuity and trust seemed vital components.
Parents valued the opportunity to speak honestly about their feelings in a way that they cannot to others for fear of being judged. This, ‘permission’ for the negative feelings they may have about their child and the corresponding guilt they may have about that was linked to the knowledge that other parents may have felt the same way. Only one of the professionals we spoke with linked shared experience specifically to guilt and negative emotions. The language used by parents and befrienders to describe this release of emotion indicates the extent to which these feelings are usually repressed because it is in some way not ‘normal’ to feel this way about your family. Phrases used to express this included: ‘offload’, ‘download’, ‘relief’, ‘let off steam’ and ‘weight lifted off your shoulders’. The importance of ‘not being judged’ as a key component of peer support was not evident in the interviews with professionals.

Box 1: Reduced isolation

**Emotional stability and personal growth**

The outcomes of emotional stability and personal growth were strongly interrelated. The service provided parents with the support to offload some of their emotional burden (emotional support) which was seen as important for their mental health. This then enabled the service to support parents in developing the confidence and ability to cope with the emotional ups and downs for themselves. Parents were helped to achieve these outcomes by learning from the experience of their befriender (informational support) and the support and encouragement of their befriender (affirmational support); both of which are underpinned by shared experience.
Parents have credibility with other parents because they are speaking from experience, rather than simply passing on information as a professional might. Hearing how another parent has encountered and tackled a similar problem assures the parent that they are not the only person dealing with such problems, which can aid their confidence to explore new solutions. Befrienders were conscious of the responsibility that this conveyed and the importance of not saying the wrong thing. Professionals in the group recognised the knowledge and experience held by the community of parents but some expressed concern that befriender might give wrong advice or signposting, or unduly influences parents’ decisions.

The support and encouragement that befriender give to parents are critical to their development in areas such as confidence and knowledge. Providing enough support that the parent feels secure and able to challenge themselves, and positively reinforcing progress, enables parents to help themselves (also known as ‘scaffolding’). Parents were conscious of the fact that the journey was something they had to do for themselves, but they benefited from the support of someone who had been there too, and who understood and could give pointers to be able to cope and do things for themselves.

Box 2: Emotional stability and personal growth

**Outcomes for befriender**

Specific outcomes for befriender were identified by parents delivering support and professional, but were rarely discussed by parents receiving support.
FEELING THAT YOU HAVE HELPED SOMEONE ELSE

Befrienders clearly identified that feeling that they had helped someone else was a positive and important outcome for them. Particularly rewarding was seeing parents they support making progress, and knowing that they had played a part in that progress. A particular motivation for becoming a befriender was to protect other parents from the struggle that they had been through themselves.

PERSONAL GROWTH AND SELF-WORTH

Befrienders described an increase in their own confidence through training and ongoing support. Feeling more confident enabled befrienders to operate more effectively in their own family and in relationships outside of the service, as well as with parents they supported. This was strongly associated with having a sense of purpose and doing something worthwhile, with a corresponding restoration and maintenance of their own self-worth.

Part of personal growth was the recognition of how far they had come in their own journey. Befrienders were able to gain strength from using their own negative experiences in a positive way, to help other parents avoid the same problems or pitfalls; or at least to know what to expect. Befrienders described ways in which changes in their confidence and self-worth, which they ascribed to their involvement with the peer support service, affected their willingness to engage in activities such as becoming active in advocacy and other parent organisations, joining committees, or becoming involved in training professionals.

Box 3: positive outcomes from befriending
TRAINING

Training undertaken by befrienders in this service was extremely influential, both in equipping them with the skills to befriend and contributing to their personal emotional development, but also in creating a supportive social network. Training groups go on an emotional journey together, sharing stories that they may never have shared with anyone else. This establishes a special and close connection between the group members and puts in place the friendships that often appear to continue when, as befrienders, they are supporting each other.

Those who complete training are under no obligation or pressure to befriend. Training was described as a painful process for many befrienders and, subsequently, not everyone wants to, or feels ready to, take the next step to befriending, although they remained part of the network. Training was viewed by professionals as a critical component of a well-run service however they seemed unaware of the personal impact on befrienders, beyond equipping them with the skills to befriend.

MUTUAL SUPPORT

Befrienders highly valued the support they offer to each other. This mutual support was not described by the other participant groups in the same way. The importance of the coordinator and befriender network was discussed by professionals in the context of professional supervision, as essential for the safety and wellbeing of both parents and befriender. However, professionals seemed unaware of the personal and emotional support that befrienders described giving each other. Befrienders described the support they offer each other in similar ways to parents; defined by their shared experiences and the ability to speak
freely in a non-judgemental, safe environment. They spoke of learning from each other, supporting and encouraging each other and gaining emotional benefit from the group. Mutual support was seen as important to maintain their emotional stability, for the sake of their own wellbeing but also crucial for the service, to sustain their ability to offer support to others.

Box 4: Training and mutual support

**NEGATIVE IMPACTS FOR BEFRIENDERS**

A number of potentially negative impacts for befrienders were identified, both by befrienders and professionals. Befrienders described the emotional drain of befriending, which they felt to be underestimated by professionals. However professionals did express concerns regarding the emotional burden on befrienders and the importance of supervision and support to protect them.

Befrienders often found it difficult to switch off after befriending, which was sometimes constructed as a commitment to the parent they are supporting and to the peer support service rather than a negative impact. Professionals were concerned that befrienders may have difficulty maintaining their distance from another family’s situation, and might be less able to maintain boundaries around the relationship than a professional.

Many, though not all, befrienders described varying levels of anxiety before a first visit; whether they were going to ‘get it right’, and whether they would get on with the parent. Befrienders were concerned whether they would do any good or that they might even make
the parent feel worse. One described feelings of inadequacy; others expressed frustration that they wouldn’t be able to ‘fix’ things for people, even though this isn’t the objective.

Box 5: Potential negative impacts for befrienders

**Outcomes for both befrienders and parents**

**Expanding social network**

Although the befriender is the person with whom a parent has ‘shared experience’, the opportunity to meet and talk with someone who can comprehend your situation may help parents to feel that they are not alone. They are made to feel members of a community of parents of disabled children. Even if the only other parent a ‘supported parent’ meets with is their befriender, it seems that the reassurance that many other parents are going through the same challenges can engender a sense of community. A number of parents and befrienders made reference to broadening their social circle, meeting families similar to their own, and becoming part of a community. Parents described the benefits of social as well as emotional and practical support from meeting with their befrienders.

Box 6: expanding social networks for both groups

Figure 1 illustrates the link between the mediator ‘shared experience’ and the active ingredients and outcomes of peer support. This should be viewed alongside the figure presented in the second of this pair of papers, which shows the moderating service and personal factors that precede the mediator ‘shared experience’.

Figure 1: Active ingredients and outcomes of peer support
DISCUSSION

Participants identified a number of important ways in which support impacted on parents; a reduced sense of isolation, emotional stability and personal growth. For befrienders, personal growth was also an important outcome in association with increased self-worth and a sense of having helped someone else. The befrienders, but not the other participant groups, also stressed the positive impact of the training and ongoing mutual support. Potential negative outcomes for befrienders, such as emotional burnout and anxieties around personal effectiveness, were identified by all participant groups. Befrienders and parents appeared to benefit through expanding their social network.

Specific outcomes for befrienders were identified by parents delivering support and professionals, but were rarely discussed by parents receiving support. We did not explore this in depth at interview with this group unless discussion was forthcoming as i) they have no direct experience ii) we did not wish to elicit discussion around outcomes, some of which may be negative, where there was potential to impact on how parents might behave with their befriender.

Previous qualitative research in this population has also reported outcomes relating to emotional and personal growth for parents receiving support (Ainbinder et al. 1998, King et al. 2000, Kingsnorth et al. 2011, Rearick et al. 2011, Solomon et al. 2001). For parents who support others, the satisfaction gained from helping other people (Ainbinder et al. 1998, Kerr and McIntosh 2000, Lo 2010, Sullivan-Bolyai and Lee 2011) and the desire to give something back (Solomon et al. 2001, Nicholas and Keilty 2007, Law et al. 2001, Jenkinson et al. 2013) has been highlighted and is consistent with research in other populations on the
benefits derived from offering support to others, known as the helper-therapy principle (Solomon 2004, Salzer and Shear 2002).

Shared experience was key to the success of this particular support and distinguished it from other types of support such as that received from professionals or family and friends. We interpret shared experience to be the overarching theme linking, if not actually enabling other aspects of the peer support to be effective thus leading to the perceived outcomes.

The findings support those of previous studies highlighting the importance of shared experience, (Ainbinder et al. 1998, Bull 2003, Kerr and McIntosh 2000, Kingsnorth et al. 2011, Law et al. 2001, Lo 2010, Nicholas and Keilty 2007, Rearick et al. 2011, Solomon et al. 2001) and learning through the experience and support of others (Ainbinder et al. 1998, Bull 2003, Kerr and McIntosh 2000, Kingsnorth et al. 2011, Law et al. 2001, Rearick et al. 2011). These themes reflect the informational, affirmational and emotional support described elsewhere (Sullivan-Bolyai et al. 2010, Sullivan-Bolyai and Lee 2011, Ireys et al. 1996). The findings are also consistent with research in other populations on the positive and protective effects of a shared social identity (Haslam et al. 2009). Participants in this study described only positive effects however there is a potentially negative side to the development of a sense of social identity in that it may reinforce the differences parents perceive between themselves and others who do not have disabled children.

We are not aware of other research in this context and population that has identified such strong, positive impact of the training and ongoing support between befrienders. These are both integral features of the specific service; the impact of training and support in other research will be dependent on the design of the service under evaluation. In many ways
befrienders’ description of training is similar to the way befriended parents talk about the befriending experience; it is a process that is painful and emotional but that helps you to have a greater understanding of your journey. In some sense, the training is itself another form of peer support and like the support offered to parents, is founded on shared experience.

Failure to acknowledge the impact for befrienders has potential implications for whether the service is considered value for money. For example, if the service was viewed purely for the benefit of the recipients, and befrienders are perceived simply as the people delivering the service, it would seem counter-intuitive to train people without expecting them to assume that role. It is only when the combined benefits of training and ongoing mutual support to befrienders are viewed as outcomes of a community-based service that the complex nature, and multiple levels, of this model of peer support become clearer.

A key strength of this study is that it enabled the views of service users, service providers and professionals who might commission or refer to the service to be considered alongside each other. In addition, the close involvement of the stakeholder group was a key strength, influencing all stages of the research. Limitations of the study include potential sampling bias as we were not able to recruit any parents who had declined or who had withdrawn from the service because they either had a negative experience or were dissatisfied with the service. Similarly only two of the 23 befrienders we interviewed no longer had regular contact with the service. Our participant group was entirely white British, which was representative of the service locally; however, ethnic and cultural factors may well influence the uptake and implementation of peer support and merits further research. We also recognise the limitation of our professional sample as there is a potential positive bias in this group who were willing to set aside an hour to talk about parent support.
Future research

Our analysis has led us to question whether there may be potential impact of this type of service on the broader community of parents of disabled children as well as on those directly involved with providing and/or receiving support through the service. This may present an unexplored avenue for future research. The perspective taken by the researcher on what the intervention ‘is’, ‘does’ and ‘for whom’ will determine the nature of the evaluation (Petticrew 2011) and, logically, will also impact on the value perceived in both monetary and societal costs and benefits.

The importance of shared experience between parent and befriender is central in our study and in the work of others. Creating that sense of shared experience however may not be straightforward. Peer support is a complex intervention with a number of factors relating to the organisational structure of the service, the parents offering support and the parents receiving support which have potential to impact on the effectiveness of the intervention. The growing emphasis on process evaluation in trials of complex interventions (Craig et al. 2008) highlights the need for a greater understanding of the organisational and process factors that may help or hinder the establishment of a sense of shared experience between parents. We unpack and explore these factors in the second of these linked papers (Shilling et al. submitted).
ACKNOWLEDGEMENTS

We are very grateful to the participants who gave their time to be interviewed for this study, to the Face2Face coordinators and area manager in Devon and Cornwall, UK for their help with study recruitment and to the study stakeholder group.

We acknowledge funding from the National Institute for Health Research Collaborations for Leadership in Applied Health Research and Care and the charity Cerebra. The views and opinions expressed in this paper are those of the authors and not necessarily those of the NHS, the NIHR, the Department of Health, or Cerebra.

This work commenced when the University of Exeter Medical School was part of Peninsula College of Medicine and Dentistry.
REFERENCES


Committee on Early Childhood Adoption and Dependent Care (2011) The pediatrician’s role in family support and family support programs. Pediatrics, 128, e1680-1684.


TABLES AND FIGURES

Table 1: Demographic information, parents and befrienders

<table>
<thead>
<tr>
<th></th>
<th>Parents (12 families, 14 young people)</th>
<th>Befrienders (23 families, 24 young people)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age of child</td>
<td>Median age 7.5 (range 3.5 – 14 years)</td>
<td>Median age 12 (range 4-22 years)</td>
</tr>
<tr>
<td>Gender of child</td>
<td>4 female, 10 male</td>
<td>5 female, 19 male</td>
</tr>
<tr>
<td>Primary diagnosis*</td>
<td>No diagnosis</td>
<td>No diagnosis</td>
</tr>
<tr>
<td></td>
<td>ASD</td>
<td>ASD</td>
</tr>
<tr>
<td></td>
<td>ADHD</td>
<td>ADHD</td>
</tr>
<tr>
<td></td>
<td>CP</td>
<td>CP1</td>
</tr>
<tr>
<td></td>
<td>Auditory processing disorder</td>
<td>Complex additional needs with sensory</td>
</tr>
<tr>
<td></td>
<td>Congenital heart disease</td>
<td>issues</td>
</tr>
<tr>
<td></td>
<td>Currarino triad</td>
<td>Developmental delay</td>
</tr>
<tr>
<td></td>
<td>Severe learning difficulties</td>
<td>Down syndrome</td>
</tr>
<tr>
<td></td>
<td>Spina bifida</td>
<td>Dyslexia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Partial trisomy 15</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Semantic pragmatic disorder</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Missing</td>
</tr>
<tr>
<td>IMD quintiles</td>
<td>quintile 2 6</td>
<td>quintile 1 1</td>
</tr>
<tr>
<td>(where the least</td>
<td>quintile 3 3</td>
<td>quintile 2 9</td>
</tr>
<tr>
<td>deprivation is</td>
<td>quintile 4 3</td>
<td>quintile 3 4</td>
</tr>
<tr>
<td>quintile 1 and the</td>
<td></td>
<td>quintile 4 6</td>
</tr>
<tr>
<td>highest, quintile 5)</td>
<td></td>
<td>quintile 5 2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>missing 1</td>
</tr>
</tbody>
</table>

*As identified by the parent. 7 of the 14 young people in the parent group had multiple difficulties; 14 in the befriender group had multiple difficulties. 1 family in the parent group had multiple (3) children with additional needs. 2 families in the befriender group had multiple (2) children with additional needs.
Box 1: Reduced isolation

**Linked to shared experience:**

*I’m not on my own; I’m not the only one with difficulties. Sometimes you do feel very alone, very isolated as a mum of a disabled child.*  
_P6_

*just trying to get them ready for school in the morning you know, without a tantrum, without throwing things, without hitting you, and then another parent saying well that happens to me as well, I understand exactly how you feel, it’s always... you feel as if you’re not on your own.*  
_P12_

*It shows parents that they are not alone; they can discover there is a large group of people that have gone through the same thing. Isolation can be the biggest hurdle that parents face and this service helps them to overcome this.*  
_B32_

*And I mean just constantly reiterating to them, you know, it is really hard, you know, and this is a normal response. And I think because often families feel like they’re the only ones in this place, you know, and I think that really, really helps them.*  
_B6_

*But the element that we just can’t give as professionals is, is, is the reality you know, the day to day of, of either going through something or, or living with a child with, you know, with that disability or whatever.*  
_PR06_

**Linked to matching by diagnosis:**

*What the befriender lives within behaviour, disability, I think that matters because at least then, they’ve got a real connection, they know what you’re on about.*  
_P4_

*It’s more about how you feel inside and how you feel accepted or don’t feel accepted sometimes by the community and, you know, you have got a lot of common issues.*  
_B16_

*You've got to be able to get on though, that's... you know, one thing that... you know, because it's not... just because you're a parent of a child with a learning disability, it doesn't mean that you're going to get on with another parent does it*  
_PR07_

**Linked to ‘it’s all about me’:**
I think they were the first people that actually asked me about me. You know, um you go to the hospitals, you go to school, you fill these forms in, it’s all about [my child] P4

So the need for somebody to understand, I certainly felt, for me, the, um, to be able to say the awful things that I was feeling to somebody that wasn’t going to judge me P10

you’re not afraid to say, you know, “I could really strangle them one day”, but then if you’re talking to a professional, in no matter what capacity they’re in, erm you then worry that you said the wrong words and you’re going to have social on your door. B9

Box 2: Emotional stability and personal growth

Sometimes when you talk about it you kind of realise it, it’s real, it’s happening and, and everything else. So, so no, they didn’t change my life, but they certainly helped me cope with it better P4

I think I feel more confident with dealing with it all really and I’ve sort of like learned to accept you know, that that’s what it is. P5

it’s really important that you are able to get rid of some of that emotion because it really does store up inside you and if you bottle it up it can manifest into all sorts of illnesses and how you are with your children, how you view everything, your mental health, you know, it has such a bit impact. Emotion is something that affects everything. B16

I guess is what Face 2 Face do a lot of is providing that, that, you know, constructing that framework around families in the initial period, to give them that emotional stability. To then to be able to say, right, okay, yeah, this is the way we've, we’re going to be and we’ve got to get on with it. PR02

Linked to learning from the experience of others:

She could provide, not necessarily a solution, but she could say what she had done. So then you could make your own mind up then can’t you? So it’s information, again, that you can do something with. P11

You are passing on a kind of knowledge experience. B3
And I think it’s also something about how you then receive that um you know, that support or information because you see that person with a level of credibility because they obviously have come from a similar place. PR06

Linked to Support and encouragement:

She encouraged me to fight for the statement, [...] it made me realise well yeah, I've got to take control as well. They don't get to dictate to me, it's my son and I know what's best for him at the end. P5

It’s like no one can make you, no one can... it has to come from within you. You have to rebuild yourself and repair yourself and accept your situation, but you do need support while you’re doing that. P2

Box 3: positive outcomes of befriending

Feeling you’ve helped someone else:

I just feel really, really glad within me that I can help somebody else, that there is, that there would have been somebody else there to, to pour it all out to B5

So I think that's invaluable for people to know that they're actually helping. Um, and it's probably quite frustrating if they feel they're not B15

Perhaps that opportunity to give back a little bit would be quite good for the person giving the support. That feeling of value and, erm giving back would be quite good. PR08

Personal growth and self-worth

I think it gave me back my self-worth, after, you know, sort of, after having the diagnosis, and being shoved into parenting classes, it was the thing that – it sounds a bit dramatic – it was the thing that saved me, that gave me back my self- worth. FG1

My world opened up completely, from being completely isolated, quiet, shy, wouldn’t say boo to a goose, to someone who is now about four years down the road is not only befriending but involved in steering groups, er, talking with, um, er, what do you call them, um, commissioners, um, er, advising and supporting lots of different persons. FG2
For the person who gives support? Well, I suppose a really positive outcome for me, that I think parents would get, was, sort of, an increased confidence, you know, sort of, self-esteem and confidence in their own ability to, um, use their skills. PR01

Box 4: Training and mutual support

Training:

Most of us didn’t know each other at all and the bond that we got was amazing. I’ve never known, I’ve never known a bond like it. It was quite uncanny, quite scary uncanny. It was like you could go and just open up your soul in that room and know that it stayed in that room. B35

I did the befriending course, so I’ve got that little monthly group, and if you don’t actually befriend or you don’t want to befriend or you’re not ready for it, you can’t do it, even though we’ve done the training we can still, we can still be a part of that group so that’s really lovely. B3

Err so if somebody hasn’t had the training I would worry that inappropriate err misguided help may be on offer and then making it worse. PR10

Mutual support:

It’s just a sharing session with the group. [...] it just gives me an opportunity really to hear what other people are up to and then try and bring that into our family life. B16

You know and I still feel I have, I don’t feel as if, I don’t feel that because I’m not befriending that I haven’t been able to be helpful. B11

I think they need a sort of support network so that they know themselves you know, what they feel happy with and if they don’t you know, feel happy because it is a bit of risk that you might get sucked in if you’re working very much on your own. PR06

Box 5: Potential negative impacts for befrienders

Sometimes they feel a bit helpless, the actual befriender, that they can’t help. P11
There seems to be some kind of perception that we just go in and have a nice little cup of coffee and a chat. And I would like them to realise how, um, emotionally and mentally, you know, taxing it can be. FG1

And, um, yeah, so just because I just felt inadequate it wasn't anything she'd done, it was just that I just didn't feel that I knew what to say, or how to move it to where we should be. B4

I just feel that we'd need to know what, you know, and what supervision they've got, what support have they got as well? Because if they have got a lot of burden of all of this from another person, they could actually get deeply emotional with them PR05

Box 6: expanding social networks for both groups

There’s a whole other community that opens up to you then, of um other parents with children with disabilities P8

I think what it means to me really is that it's nice to meet with parents who have a understanding of how, how difficult it can be to have a child with support needs really B6

Well if there’s a positive relationship that’s, that’s struck up, um then there could very well be a friendship between the, the adults, possibly then broadening into the children and maybe the family PR10