Trends in Professional Education

Parent to Parent Programs: Parent Preferences for Supports

Parent to Parent programs facilitate connections between parents of young children with special needs by carefully matching an experienced parent in a one-to-one relationship with a parent newly referred to the program. Using national survey results, this article examines the preferences referred parents have for various kinds of emotional and informational supports provided through Parent to Parent programs and the impact of the child's age and severity of disability on parental preferences. Parents preferred a wide range of supports, with the child's age and severity of disability contributing to differences in their preferences. Supports to families must, therefore, be individualized and responsive to child and family characteristics. Key words: mentor parents, mutual support, parental preferences for support, parent support, Parent to Parent.

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As services for supporting families of young children with special needs are integrated into public policy and recognized as core components of quality early intervention programs, service providers are working to provide individualized responsive services for families with diverse needs. Parent to Parent programs offer a unique model for accomplishing this goal.1

Parent to Parent programs match a veteran parent—someone with experience as a parent of a child with a disability—with a referred parent—someone with a child with a similar disability who would like support from a parent who has "been there." Common experiences form the basis of a one-to-one relationship in which the referred and veteran parents arrange their own supportive interactions. The veteran parent typically shares information and provides emotional support to the referred parent in a one-to-one match that usually lasts from 1 to 6 months.1

The goals of the Parent to Parent match are:
- to provide parents with emotional support and understanding,
- to provide parents with informational support about disability and services.18

Well-established programs may offer additional services beyond the one-to-one match, such as a 24-hour telephone line, meetings for group support, training to become a veteran parent, social

1 Jef Young Children 1996, XI: 55-62

53
events, newsletters and brochures, advocacy training, and activities for other families members.

Parent to Parent programs, by their very nature, respond directly to individual needs, providing services that family members consider important, as one mother notes.

Parent to Parent has been my life line. When I first heard the diagnosis, I was devastated. Well-meaning doctors and nurses, as well as friends and families, simply did not understand. It was only when I talked with another parent through the Parent to Parent program that I could begin to hope for a future for us all. My veteran parent was gently there for me whenever I needed her.

Child and family characteristics, such as the age of the child who has a disability, the type and severity of the disability, family size and form, ethnicity, educational background, socioeconomic status, and geographic location, typically form the basis of the one-to-one match between veteran and referred parents. Child and family characteristics will also affect family needs and preferences for support. The age of the child with a disability may affect family preferences for support. Families who learn of their child's disability at or shortly after birth deal with a whole host of feelings. At the same time, these parents are having to deal with all of the new procedures and terminology related to their child's disability or special health need. Emotional and informational support for obtaining an accurate diagnosis, funding services, informing family and friends, addressing the stigma, and meeting the needs of siblings are frequently mentioned as needs by parents of young children with disabilities.

By the time a child with a disability is of school age, the issues that families face often relate to the individualized education plan (IEP) and inclusion in school and community activities. Parents at this stage may need informational support regarding their educational and legal rights, adaptations of the school curriculum, and strategies for their child to establish friendships. The severity of the family member's disability may also affect family preferences for support. Mothers of children with severe disabilities may prefer support related to meeting the needs of other family members, while mothers of children with mild or moderate disabilities prefer support directed to meeting the needs of the child with a disability. Some parents of children with mild disabilities may experience mixed feelings of confusion, frustration, relief, guilt upon receiving the diagnosis, and uncertainty about future dependency or independence when a child's progress is irregular, and they may prefer support to help them deal with their feelings.

There is a need for further information about whether and how preferred services vary with individual family characteristics and how services can be designed with parental preferences in mind.

The October, 1995 issue of Infants and Young Children included an article reporting the results of a national survey of Parent to Parent programs. Special focus was on veteran and referred parents who participate in Parent to Parent programs, the supports provided to referred parents, and the training provided to veteran parents. This article, an expansion of that Parent to Parent overview, provides answers to the following two questions:

1. What is the impact of the age of the child with a disability on parental preferences for support offered by Parent to Parent programs?
2. What is the impact of the severity of the child's disability on parental preferences for the supports offered by Parent to Parent programs?

METHOD

Data were gathered through the national survey of Parent to Parent programs and participating parents conducted by The Beach Center on Families and Disability at The University of Kansas. The survey for referred parents targeted a random sample of parents from 190 of the Parent to Parent programs participating in the national survey. Seven hundred and four surveys were requested by referred parents, and 260 surveys, representing a conditional return rate of 34%, from 115 different Parent to Parent programs in 43 different states were returned.

The survey for referred parents provided demographic data and detailed information on the family
support and services received and preferred by referred parents. Referred parents responded to 15 questions, including three questions on the type of supports received and preferred. Survey respondents were provided with a listing of seven informational supports, 11 emotional supports, and 18 other supports. They were asked to rate which supports were provided to them and which of the supports they preferred for meeting their needs. Parents were instructed to write the item number corresponding to their preferred supports in boxes labeled “most important,” “second most important,” or “third most important.” For the purposes of this study, first, second, and third choices indicated a preferred support. A more detailed description of the questionnaire and sampling methodology for the current study can be found in the 1995 Infants and Young Children article.

RESULTS

Of the 340 referred parents returning surveys, 161 (69%) were parents of children aged birth to 5 years, while 179 (59%) referred parents had children 6 years and older (7 parents had missing data). Of the referred parents of children birth to 5, 111 (32%) reported their child to have a mild disability, 61 (18%) a moderate disability, and 39 (12%) a severe disability. Of the referred parents of children 6 and older, 98 (20%) reported their child to have a mild disability, 39 (8%) a moderate disability, and 23 (5%) a severe disability.

Survey results provided information about the types of emotional, informational, and other program supports that all referred parents preferred. For the current study, the preferences for support were divided into subgroups by the family characteristics of age of child and disability of the disability.

Preferred supports by age of child with a disability

Referred parent responses were grouped by the family characteristics of age of the child and disability of the child. Parental preferences of younger or older children were identified using a 9-point scale of importance. The scale was composed of 3 elements: 1) receive information about the disability, 2) living with and caring for the child, and 3) finding and getting the best possible help for the family member with a disability (advocate) and services for the family.

Group meetings, either for emotional or educational support, were preferred by a large percentage (at least 40%) of parents of both younger and older children with disabilities. Parental preferences for each of the other program activities beyond the math did not differ by more than 6 percentage points.

Preferred supports by severity of disability

Responses were grouped by the family characteristic of severity of disability into categories of mild, moderate, or severe, according to the severity of disability of the child as reported by the referred parents completing the survey. Table 2 presents parental preferences for emotional and informa-
Table 1. Parent preferences for supports by age of child

<table>
<thead>
<tr>
<th>Supports</th>
<th>Birth-5 years</th>
<th>5-6 years</th>
<th>6 years and older</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emotional support</td>
<td><strong>N = 155</strong></td>
<td><strong>N = 70</strong></td>
<td></td>
</tr>
<tr>
<td>Suggest to listen and understand</td>
<td>60%</td>
<td>60%</td>
<td></td>
</tr>
<tr>
<td>Knowledge of others who are doing OK</td>
<td>59%</td>
<td>50%</td>
<td></td>
</tr>
<tr>
<td>Help for the future</td>
<td>53%</td>
<td>50%</td>
<td></td>
</tr>
<tr>
<td>Help in being less alone</td>
<td>36%</td>
<td>42%</td>
<td></td>
</tr>
<tr>
<td>Ways to deal with stress</td>
<td>25%</td>
<td>28%</td>
<td></td>
</tr>
<tr>
<td>Dealing with the diagnosis</td>
<td>20%</td>
<td>24%</td>
<td></td>
</tr>
<tr>
<td>Problem-solving support</td>
<td>16%</td>
<td>27%</td>
<td></td>
</tr>
<tr>
<td>Informational support</td>
<td><strong>N = 157</strong></td>
<td><strong>N = 69</strong></td>
<td></td>
</tr>
<tr>
<td>Disability information</td>
<td>65%</td>
<td>59%</td>
<td></td>
</tr>
<tr>
<td>Going for child</td>
<td>58%</td>
<td>50%</td>
<td></td>
</tr>
<tr>
<td>Ways to find and get services</td>
<td>54%</td>
<td>55%</td>
<td></td>
</tr>
<tr>
<td>Community resources</td>
<td>48%</td>
<td>56%</td>
<td></td>
</tr>
<tr>
<td>Financial information</td>
<td>19%</td>
<td>24%</td>
<td></td>
</tr>
<tr>
<td>Support care information</td>
<td>19%</td>
<td>20%</td>
<td></td>
</tr>
<tr>
<td>Other program support</td>
<td><strong>N = 126</strong></td>
<td><strong>N = 58</strong></td>
<td></td>
</tr>
<tr>
<td>Group meetings for emotional support</td>
<td>48%</td>
<td>50%</td>
<td></td>
</tr>
<tr>
<td>Group meetings for education</td>
<td>44%</td>
<td>50%</td>
<td></td>
</tr>
<tr>
<td>24-hour warm line</td>
<td>25%</td>
<td>29%</td>
<td></td>
</tr>
<tr>
<td>Social events</td>
<td>26%</td>
<td>22%</td>
<td></td>
</tr>
<tr>
<td>Activities for other family members</td>
<td>12%</td>
<td>7%</td>
<td></td>
</tr>
</tbody>
</table>

- Emotional supports and the program supports beyond the match, according to the level of severity of the disability of the child.
- At least three-fourths of all parents ranked having someone to listen and understand as important, regardless of the severity of their child’s disability; and at least one-third of all parents ranked help in feeling less alone as important regardless of the severity of their child’s disability. As can be seen in Table 2, receiving information about the disability was important to the greatest percentage (74%) of parents whose children have mild disabilities and the smallest percentage (62%) of parents of children with moderate disabilities. The greatest percentage of parents of children with severe disabilities (69%) ranked information about living with and caring for the child as the most preferred type of support. Information about respite care and financial assistance were preferred by 32% of parents of children with severe disabilities, compared to less than 15% of parents of children with mild disabilities.

- The greatest percentage of all parents, regardless of the severity of their child’s disability, reported that group meetings for emotional or educational support were important. More than twice as many parents of a child with a severe disability ranked a telephone warm line as important, as compared to parents of children with mild disabilities (38% and 18%, respectively).

- Preferred supports by age of child and severity of disability

Table 3 presents information on the parental preferences for emotional and informational supports and other program supports beyond the match, as determined both by the age of the child and the severity of the disability. As can be seen in Table 3, over half of all parents ranked the emotional support of having someone to listen and understand as important, regardless of the age of their child and the severity of their child’s disability. The greatest differences in how parents ranked the emotional supports provided by Parent-Parent programs often existed between parents of young children with mild disabilities and parents of young children with severe disabilities. Two emotional supports—dealing with stress and problem-solving support—were ranked as important by almost twice as many parents of young children with severe disabilities as by parents of young children with mild disabilities.

- Whether the child’s disability is mild or severe appears to have less of an effect on the preferences of parents of older children for emotional supports. With the exception of the emotional support of giving an awareness that there are other families who are doing OK (ie, 15% of parents of children...
<table>
<thead>
<tr>
<th>Supports</th>
<th>Mild N = 69</th>
<th>Moderate N = 50</th>
<th>Severe N = 66</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emotional support</td>
<td>72%</td>
<td>70%</td>
<td>65%</td>
</tr>
<tr>
<td>Someone to listen and understand</td>
<td>55%</td>
<td>50%</td>
<td>56%</td>
</tr>
<tr>
<td>Knowledge of others who are doing OK</td>
<td>30%</td>
<td>30%</td>
<td>33%</td>
</tr>
<tr>
<td>Hope for the future</td>
<td>33%</td>
<td>33%</td>
<td>33%</td>
</tr>
<tr>
<td>Help in feeling less stressed</td>
<td>22%</td>
<td>22%</td>
<td>21%</td>
</tr>
<tr>
<td>Help to deal with stress</td>
<td>34%</td>
<td>32%</td>
<td>34%</td>
</tr>
<tr>
<td>Dealing with the diagnosis</td>
<td>25%</td>
<td>25%</td>
<td>25%</td>
</tr>
<tr>
<td>Problem-solving support</td>
<td>17%</td>
<td>17%</td>
<td>17%</td>
</tr>
<tr>
<td>Informational support</td>
<td>77%</td>
<td>77%</td>
<td>80%</td>
</tr>
<tr>
<td>Disability information</td>
<td>51%</td>
<td>50%</td>
<td>52%</td>
</tr>
<tr>
<td>Caring for child</td>
<td>39%</td>
<td>37%</td>
<td>35%</td>
</tr>
<tr>
<td>Ways to deal and get services</td>
<td>57%</td>
<td>57%</td>
<td>57%</td>
</tr>
<tr>
<td>Community resources</td>
<td>52%</td>
<td>52%</td>
<td>52%</td>
</tr>
<tr>
<td>Financial information</td>
<td>19%</td>
<td>20%</td>
<td>20%</td>
</tr>
<tr>
<td>Respite care information</td>
<td>11%</td>
<td>11%</td>
<td>12%</td>
</tr>
<tr>
<td>Other program supports</td>
<td>60%</td>
<td>59%</td>
<td>61%</td>
</tr>
<tr>
<td>Group meetings for emotional support</td>
<td>70%</td>
<td>69%</td>
<td>68%</td>
</tr>
<tr>
<td>Group meetings for education</td>
<td>70%</td>
<td>70%</td>
<td>70%</td>
</tr>
<tr>
<td>Social event</td>
<td>35%</td>
<td>35%</td>
<td>36%</td>
</tr>
<tr>
<td>Activities for other family members</td>
<td>22%</td>
<td>24%</td>
<td>25%</td>
</tr>
</tbody>
</table>

Table 2. Parent preferences for supports by severity of disability of child

with mild disabilities ranked this support as important, whereas 22% of parents of children with severe disabilities preferred this support and the emotional support of finding ways to deal with stress (53% of parents with mild disabilities ranked this support as important, whereas 44% of parents with severe disabilities preferred this support), there were no other supports that had a greater than 10 percentage point discrepancy in rankings between parents of children with mild and severe disabilities.

The data reveal some interesting similarities and differences in parental preferences overall. As seen in Table 3, some supports are preferred regardless of the age of the child and the severity of the child's disability (i.e., having someone to listen and understand, information about the disability and how to care for the child, as well as information about services and community resources). Group meetings for emotional and/or educational support are valued by a significant percentage (in at least 50% of all parents regardless of the age and severity of disability of their child. It is also interesting to note that at least one-third of the parents who have a child with a severe disability ranked information about respite care and financial assistance as preferred regardless of the age of their child.

While some preferences appear to remain relatively constant over time, it is interesting to note in Table 3 that parents who have children with mild disabilities generally reported greater differences in their preferences as the child gets older than did parents of children with severe disabilities, whose preferences remained more constant over time.

**Discussion**

The wide range of supports considered important by referred parents suggests that when Parents:10 Parent programs (and other family support programs) offer a continuum of supports, parents are able to individualize their Parent to Parent experience. Understanding the differences in their preferences for support based upon age and severity of disability
Table 3. Parent preferences for supports by age and severity of disability of child

<table>
<thead>
<tr>
<th>Supports</th>
<th>Mild Birth-5 years N = 52</th>
<th>Moderate Birth-5 years N = 61</th>
<th>Severe Birth-3 years N = 39</th>
<th>Mild 6 years and older N = 29</th>
<th>Moderate 6 years and older N = 29</th>
<th>Severe 6 years and older N = 23</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emotional supports</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Someone to listen and understand</td>
<td>74%</td>
<td>60%</td>
<td>58%</td>
<td>70%</td>
<td>69%</td>
<td>68%</td>
</tr>
<tr>
<td>Knowledge of others who are doing OK</td>
<td>57%</td>
<td>51%</td>
<td>34%</td>
<td>15%</td>
<td>33%</td>
<td>23%</td>
</tr>
<tr>
<td>Hope for the future</td>
<td>59%</td>
<td>49%</td>
<td>29%</td>
<td>29%</td>
<td>35%</td>
<td>33%</td>
</tr>
<tr>
<td>Help in being less alone</td>
<td>55%</td>
<td>30%</td>
<td>29%</td>
<td>40%</td>
<td>45%</td>
<td>35%</td>
</tr>
<tr>
<td>Ways to deal with stress</td>
<td>50%</td>
<td>32%</td>
<td>27%</td>
<td>21%</td>
<td>27%</td>
<td>26%</td>
</tr>
<tr>
<td>Dealing with the diagnosis</td>
<td>46%</td>
<td>27%</td>
<td>23%</td>
<td>24%</td>
<td>20%</td>
<td>21%</td>
</tr>
<tr>
<td>Respite care support</td>
<td>25%</td>
<td>29%</td>
<td>17%</td>
<td>15%</td>
<td>13%</td>
<td>17%</td>
</tr>
<tr>
<td>Informational supports</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Disability information</td>
<td>78%</td>
<td>68%</td>
<td>60%</td>
<td>65%</td>
<td>65%</td>
<td>65%</td>
</tr>
<tr>
<td>Caring for child</td>
<td>60%</td>
<td>63%</td>
<td>61%</td>
<td>60%</td>
<td>60%</td>
<td>60%</td>
</tr>
<tr>
<td>Ways to find and get services</td>
<td>63%</td>
<td>55%</td>
<td>45%</td>
<td>55%</td>
<td>61%</td>
<td>60%</td>
</tr>
<tr>
<td>Community resources</td>
<td>45%</td>
<td>55%</td>
<td>39%</td>
<td>35%</td>
<td>46%</td>
<td>39%</td>
</tr>
<tr>
<td>Financial information</td>
<td>25%</td>
<td>24%</td>
<td>23%</td>
<td>25%</td>
<td>15%</td>
<td>10%</td>
</tr>
<tr>
<td>Respite care information</td>
<td>20%</td>
<td>17%</td>
<td>17%</td>
<td>16%</td>
<td>12%</td>
<td>39%</td>
</tr>
<tr>
<td>Other program supports</td>
<td>N = 39</td>
<td>N = 46</td>
<td>N = 33</td>
<td>N = 33</td>
<td>N = 23</td>
<td>N = 18</td>
</tr>
<tr>
<td>Group meetings for emotional support</td>
<td>61%</td>
<td>44%</td>
<td>53%</td>
<td>70%</td>
<td>53%</td>
<td>50%</td>
</tr>
<tr>
<td>Group meetings for education</td>
<td>41%</td>
<td>24%</td>
<td>39%</td>
<td>44%</td>
<td>57%</td>
<td>50%</td>
</tr>
<tr>
<td>24-hour warm line</td>
<td>18%</td>
<td>19%</td>
<td>43%</td>
<td>18%</td>
<td>33%</td>
<td>33%</td>
</tr>
<tr>
<td>Social events</td>
<td>28%</td>
<td>18%</td>
<td>41%</td>
<td>28%</td>
<td>30%</td>
<td>11%</td>
</tr>
<tr>
<td>Activities for other family members</td>
<td>15%</td>
<td>7%</td>
<td>18%</td>
<td>6%</td>
<td>13%</td>
<td>6%</td>
</tr>
</tbody>
</table>
of the child will help individuals who are developing
a Parent to Parent program and other family
organizations in planning and modifying the services
provided. A discussion of these differences and some
implications for parents and service providers of
these parent preferences are presented.

Preferred support by age of child with disability

A one-to-one Parent to Parent match provides a
served parent with an opportunity to talk with a
veteran parent—someone who can listen and
understand based upon their own similar family
and disability experiences—and this type of support
may be particularly helpful for the preferred type of emotional support
available in Parent to Parent programs regardless of
the age of the child. This reinforces the findings of previous studies of Parent to Parent pro-
grams.13,14 and suggests that Parent to Parent pro-
grams are indeed meeting a preferred need.

While the literature suggests that feelings of
isolation and loneliness often accompany the birth
of a child with a disability,17,18 it is interesting to
note that, while parents of younger children fre-
quently mention their preference for help in feeling
lost alone, parents of older children actually have
greater preferences for this type of support. Dealt-
ing with feelings of isolation apparently converge
to be an issue for these parents of older children,
and professionals providing services to families
who have older children with special needs should
be aware of that long-term need.19 A single parent
of a 9-year-old boy with autism explains:

Even though I have received lots of information about autism from many different professionals, until I con-
ected with Parent to Parent I had not had an opportunity to talk to another parent who shared my experiences, my
frustrations, and, yes, my special joys. I felt very alone with my feelings.

As children with disabilities move into the school system, parents often face a whole host of new
decisions in a system that is not as family-centered as an early intervention program.20 Given the
expansion of the family’s environment and less
support from the system, it is perhaps not surpris-

ing that more parents of older children preferred
problem-solving support than did parents of
younger children.

Parents, regardless of the age of their child with a
disability, are searching for information about the
disability, community resources and services, how
to find and get help, and living with and caring for
their child. The fact that these four different kinds of
informational supports were ranked as preferred
by almost half of all parents suggests that the need for
these types of information remains across the lifespan.

The importance of group meetings, both for
emotional and educational support, for parents of
both younger and older children with disabilities
suggests that some parents appreciate opportuni-
ties to connect with other parents in a group setting,
regardless of the age of their child. While Parent to
Parent programs are founded on the basis of the
one-to-one match between two parents, 84% of the
programs also provide group activities for support
and to respond to these parental preferences.14

Preferred support by severity of disability

Just as the age of the child with a disability has
very little impact on how strongly parents seem to
prefer the emotional support of having someone to
listen and understand, so, too, the severity of the
child’s disability does not seem to influence parent-
al preferences for this support. While some sup-
port exists in the literature that parents of children
with severe disabilities experience higher levels of
stress,14 and it might, therefore, be assumed that
parents of children with severe disabilities would
have a greater need for this type of support than
would parents of children with mild disabilities, the
current findings do not support this assumption.

Parents of children with disabilities of all severity
levels value the opportunity to talk with someone
who shares their experiences.

With respect to preferences for informational
supports, it is interesting to note that, while a large
percentage of all parents regardless of the severity
of their child’s disability prefer receiving informa-
tion about
• the disability,
• finding and getting help, and
• community resources.
Fewer parents of children with severe disabilities preferred these kinds of information. These differences may again be related to the fact that a more severe disability may come with more information for the parents about the disability itself and about available services.
Living with and caring for a child with a severe disability often requires expensive equipment and monitoring and caring 24 hours every day. It is, therefore, not surprising that there are significant differences in the preferences of parents for information about special care and financial assistance based on the severity of the child’s disability.
Information about how to get some away from childcare responsibilities and money to pay for needed services and equipment is an important support, especially for parents of children with severe disabilities. A father of a 5-year-old daughter whose disability requires total care-related support:

It is very difficult for us even to find a chance to spend some time with each other—she is always on duty. And although none of our children are chronically ill, the additional caring responsibilities for our 5-year-old are too overwhelming. Report cards are an option, but it is in very short supply, and you have to plan for this in advance.
The support we need is hard to find.

Just as parents of children of all ages mentioned their preferences for group activities in large numbers, so did group activities were also preferred by large percentages of parents regardless of the level of severity of the child’s disability.
Preferred supports by age of child and severity of disability

As might be expected, when age and severity of disability are combined, many of the same parental preferences are apparent when age and severity of the disability are considered separately. By noting which emotional, informational, and other program supports are preferred by all parents regardless of the age and severity of the disability of the child, coordinators of newly developing Parent to Parent programs, as well as other service providers building family support programs, will be able to prioritize program components to be developed.

These program supports that were mentioned as preferred by at least 50% of all parents regardless of the age and severity of the disability of the child include:
• emotional support—having someone to listen and understand.
• information support—having information about the disability, finding and getting services, living with and caring for the child, accessing community resources, and
• other program supports—having group meetings for either emotional or educational support.

Parents and/or professionals interested in establishing a Parent to Parent program and service providers working with families who have children with special needs may want to be sure that these programs support the existence of these particular supports for parents.

It is also interesting to note that many of the informational supports received similar rankings over time by parents, with the severity of the disability having only a minimal impact—suggesting that parents need for information remains relatively stable. Those developing Parent to Parent programs and other family support programs should be prepared to provide basic information about a wide range of disabilities and to access sources that provide more detailed disability information for those families facing this depth of information. Providers of services to children with special needs and their families should also acquire themselves with other community services and agencies providing services for persons with disabilities, as well as general community programs for all community residents. Contact with a parent who has first-hand experience including a child with a severe language disability into a museum
workshop, knowledge about which fast food res- taurants have the least accessibility for a wheelchair, or a tip about a gymnastics instructor who knows sign language may encourage parents to involve their children in community programs.

More than twice as many parents of children with severe disabilities ranked information about financial assistance and information about respite care as preferred supports than did parents of children with mild disabilities, regardless of the age of the child. Families with children who have severe disabilities often deal with expensive and extensive care-taking responsibilities, thus making information about financial assistance and respite care opportunities particularly valuable to them. While these sorts of information were ranked as preferred by much smaller percentages of parents of children with mild disabilities, Parent to Parent and other family support programs that seek to be cross-disability will need to have access to this information in order to meet the needs of all parents.

Parents' preferences for emotional support showed some variability, both over time and between levels of severity of the disability. This finding is, perhaps, not surprising given that parents bring their own set of emotional needs and coping styles to their family situation. Help in dealing with the diagnosis is the one emotional response that does appear to remain relatively stable across time and across severity of the disability. It will be important for providers involved in parent support to recognize that dealing with the diagnosis is not a single effort in time for parents, but rather that parents continue to deal with the diagnosis and its implications throughout the life of the child.

As Parent to Parent programs grow in response to the needs and preferences of participating parents in their communities, many programs have added a variety of other supports and services. Although Parent to Parent programs revolve around the one-to-one match between veteran and referred parent, group meetings for emotional or educational support were frequently ranked as preferred supports regardless of the age and severity of the disability of their child. Apparently, the sharing of common experiences is also appreciated in a group setting. Contact with several veteran parents and other referred parents may offer some advantages such as exposure to a variety of experiences as a parent of a child with a disability, guidance from parents who have "been there," or empowerment and empathy from other parents whose children are age-mates.

In addition to emotional and educational group support, access to a 24-hour telephone warm line was ranked as a preferred support by at least one-third of the parents of children with severe disabilities regardless of the age of their child. It may be that parents of children with severe disabilities prefer a "way at home" style of support rather than such activities as social events that require all of the logistical arrangements of preparing and transporting family members. As Parent to Parent programs mature, the parental preferences for these various other program supports will provide helpful information in prioritizing the development of new program components.

While it is helpful to observe which supports are preferred by the largest percentages of referred parents, an individualized family support program will also consider the supports ranked by a lesser number of parents. Virtually all of the emotional, informational, and other program supports were ranked by some parents as one of the three most important supports offered by Parent to Parent programs. To these parents, these less popular supports were significant.

Information about the types of support preferred by referred parents in Parent to Parent programs will aid in the planning stages of such programs and in the development of family support activities for parents. While certain types of support are considered by a large number of families to be important, other supports are just as important to a smaller number of families and, thus, should be taken into...
consideration. The wide range of supports that
referral parents considered important to their
individual situations required to the scope of sup-
port families need to receive from involvement in
Parent to Parent programs and other family support
efforts. The information gained from this study
reinforces the belief that the process of supporting
families must be individualized by offering a variety
of support options responsive to child and family
characteristics.

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