Introduction

The intent of this article is two-fold: (1) to describe how a disability culture perspective has informed research and early intervention serving parents with physical disabilities and their babies, and (2) to articulate how this perspective is also guiding intervention for parents with cognitive disabilities and their babies. The term “disability culture” refers to the social model, civil rights or minority model of disability: disability as socially constructed, with an emphasis on its social meaning, and on social obstacles as the primary problem for people with disabilities and their families. This article identifies themes that have emerged from research and intervention in a disability culture-based organization, Through the Looking Glass (TLG), and which seem particularly salient for improving practice with parents with disabilities and their children.

Differentiating is an appropriate theme to consider initially. There has been a persistent problem in research and practice of blurring distinctions between parents with diverse disabilities. One can consider this over-generalizing a manifestation of the concept of “spread”\(^1\). Recently Olkin\(^2\) discussed the power of the negatively valenced disability characteristic to “spread” and evoke inferences about an individual’s other characteristics, leading to stereotyping: “A negative value attached to the fact of disability spreads to other unrelated...
aspects. Thus a person in a wheelchair is assumed to be cognitively impaired; a person with mild mental retardation is viewed as more profoundly retarded; people raise their voices to talk to a person who is blind. A deficit in one characteristic spreads such that similar deficits are ascribed to other characteristics.” (p.56). The process of “spread” also affects the perception of families with disabled members.

Negatively valenced “spread” appears to be one of the processes that has led to pathologizing parents with disabilities and their families in research and practice. That is, most of the over-generalizing about parents with disabilities has involved pathological assumptions about them, expressed in the emphases, language or hypotheses chosen, such as “The Mutative Impact of Serious Mental and Physical Illness in a Parent on Family Life” or the hypothesis that children of parents with multiple sclerosis would have damaged body images. A more recent article, “Child Abuse and Neglect by Parents with Disabilities” demonstrates both pathologizing and spread as it actually concerns only two families with mothers with cognitive disability rather than parents with disabilities in general. Buck and Hohmann, Cohen, Conley-Jung and Olkin have critiqued the methodology of the research literature that posits maladjustment in the children of parents with disabilities or pathologizes parents.

The pathological focus in research is a reflection of society’s particular stigma about parenthood by individuals with disability; that is, assumptions about disability commonly preclude parenting. Perhaps this attitude explains the persistent and potent tendency for parents with disabilities to be invisible and marginalized in society. Public systems, even in the disability community, do not tend to identify or gather information about parents with disabilities. As a result, these parents are not included in needs assessment, nor are funds earmarked for services for them. Invisibility results in a critical lack of resources for the growing numbers of families in
the community. People are often surprised to learn that there are approximately 8 million U.S. families with children under 18 who have one or both parents with a disability, or almost 11% of families\textsuperscript{9,10}.

This article describes an approach to research, resource development and early intervention that has evolved in response to the unmet needs and obstacles faced by parents with disabilities and their children. Material on parents with physical disabilities is presented first, identifying additional themes which are embedded in a disability culture perspective. Next, material on parents with cognitive disabilities is presented in relation to these themes. The integration of infant mental health and family therapy approaches with disability issues is discussed.

Parents with physical disabilities

The disability community has emphasized identifying and grappling with contextual, environmental, or social dimensions of disability. Parents with disabilities face numerous social obstacles, documented in a national survey of parents with disabilities. This national survey, conducted under the aegis of TLG\textsuperscript{10}, included more than 1200 parents, approximately 75% of whom had physical disabilities. Two out of five respondents reported facing attitudinal barriers as disabled parents, with one-third reporting discrimination as a parent with a disability. Practical obstacles to parenting included transportation (reported by four out of five), housing (40%), recreation access (66%), lack of access to baby care adaptations, and barriers to childcare. Attempts to take their children away were reported by 15%. Other significant issues included parents being told they could not use personal assistants to help with childcare and experiencing interference from assistants with their role as parents. More disturbing was the child abuse by personal assistants reported by 18 parents.
Costs of resources was another significant concern, even among the predominantly middle-class and well-educated participants in this survey. The average monthly household income for parents with disabilities is $1,000 less than that of non-disabled parents\textsuperscript{10}. Poverty is an especially crucial social obstacle among parents with disabilities because of the extra costs that parenthood brings and the lack of public funding of resources such as adaptive equipment and personal assistance for babycare. Unemployment and poverty are common in the disability community, with one out of three households having extremely low incomes\textsuperscript{11}.

Despite these social obstacles, parents with physical disabilities have been applying their expertise in problem-solving to the realm of parenting for generations. The scant nonpathologically focused research documents positive outcomes for these parents and their children\textsuperscript{6,7,8,12,13}. Usually these outcomes have occurred in the absence of specialized resources or early intervention. Given the social obstacles these positive outcomes are a testimony to the resilience of parents with physical disabilities and their children. Parents with disabilities have expressed concerns about generalized stigmatization of their families as being particularly needy, and it is important to clarify that many parents with physical disabilities can manage with no intervention or short-term or periodic services. Services or resource requirements for families of parents with physical disabilities would be dramatically reduced if there were fewer social obstacles.

Elsewhere\textsuperscript{14} I have described the cumulative effect of repetitive negative social suggestions or messages on our families with disabilities. Social obstacles and lack of adaptations not only exclude our families and complicate our daily lives; they are dismissive and devaluing of our families. They are subtly wounding, re-occurring through the course of every day life. They reify stigma. So it is understandable that a theme in the disability community has been an emphasis on
universal design and access: the elimination of barriers that are due to attitudinal bias and lack of expertise, as well as barriers which are physical or communicative in nature.

While the disability community advocates for universal access, making life work in the face of obstacles--pragmatic ingenuity regarding adaptation is another theme. Experiences with disrespectful services and information which connoted inferiority, deficit or pathology in people with disabilities, has led to an emphasis within the disability community on respect for expertise and adaptations derived from personal disability experience.

Therefore, as a disability community-based agency approaching the development of services and research with parents with disabilities and their children, TLG emphasized learning from our community’s families, observing and documenting how parenting works. When parents complained that professionals questioned their ability to competently care for their babies, TLG conducted a research project from 1985-88 that videotaped how mothers with physical disabilities cared for their babies and toddlers. Without intervention or baby care adaptations, most mothers developed ingenious solutions to disability obstacles. Babies adapted to their mothers’ disabilities as early as one month of age, e.g., holding still and compact (like a kitten) when lifted. There was a natural reciprocal adaptation process that developed over time.

In subsequent work with particularly stressed families, TLG staff were startled by social services and mental health practice that was pathological and uninformed about adaptations or disability culture norms. A parent with significant cerebral palsy was videotaped by child protection workers while diapering her baby--without any adaptations being provided and after a long period of out-of-home placement which interfered with the natural reciprocal adaptation process between parent and child. This tape was cited in court as evidence for her parental incapability, despite the similarity to long diaperings by high functioning mothers with cerebral
palsy in the community\textsuperscript{15,16}. In another instance it was assumed that gaze between a baby and his mother with significant cerebral palsy had not been established because of the (hypothesized) intrapsychic pathology of the mother. Actually the problem was that no one had provided a way to make mutual gaze feasible and comfortable. With adaptive positioning, gaze was promptly established between mother and baby. At TLG we concluded that one cannot assess the potential of a relationship between a parent with a significant physical disability and a baby without first providing whatever adaptive techniques and equipment make it possible for interaction to occur and the infant/parent relationship to develop\textsuperscript{16,17}.

Building on the solutions of the pioneering disabled parents in our community, TLG conducted a series of research projects to design and provide individualized baby care adaptations in order to ease the number of obstacles at the outset of parenting\textsuperscript{18-22}. The disability community’s emphasis on empowerment was inherent to this process, as it involved mutual problem-solving and brainstorming between parent and occupational therapists (one of whom was a mother with cerebral palsy herself). Another example of the disability community orientation was the non-pathological emphasis on “environmental mismatch”, e.g., “the barriers or physical elements in the environment which fail to match the functional abilities of the parent are seen as the problem rather than the parent’s physical limitations\textsuperscript{20(p72)}.” This approach is more respectful and also more conducive to change, e.g. one can focus on the problem being how to set up a diapering surface which accommodates a wheelchair rather than the problem being a mother who can’t stand. Our research on the impact of adapted baby care equipment found it to be inherently empowering, to decrease environmental barriers and increase parents’ functional baby care abilities and involvement, and to decrease fatigue and pain and seemingly prevent secondary injury. We observed that as baby care tasks became easier some parents
became less focused on the physical demands of the task and engaged in more positive interactions with their babies\textsuperscript{20-22}.

**Power differential issues** and **empowerment**, are key constructs in the disability community. Services, such as personal assistance or assistive technology, are viewed as enhancing independence as long as the individual with disabilities has the decision-making authority to orchestrate them. There is a strong preference for service provision by individuals with personal disability experience. This value is reflected in disability community agencies such as TLG being staffed predominantly by individuals with personal disability experience.

**Disability culture as support.** In *What Psychotherapists Should Know About Disability*, Olkin has discussed the power of disability culture inclusion for individuals with disabilities. It is especially informative to consider the role of disability culture for parents with physical disabilities. Since parenthood by individuals with disabilities is particularly stigmatized, the disability community can provide a buffer and an antidote to social stigma, reframing the meaning of disability. The community carries practical problem solving strategies (such as baby care adaptations) and is a source of role models for people who were not socialized to expect parenthood.

**Interdependence** is a related theme. A number of TLG research projects have documented the role of parental teamwork in disability community families\textsuperscript{15,20,23}. TLG conducted a second national survey, of couples with young children in which one partner was a parent with a physical disability\textsuperscript{23}. This study detailed household division of work and decision-making, comparing these couples to couples in which both partners were non-disabled. The first analysis, of couples in which only the mother has a disability, found that the able-bodied fathers did a little more housework and the mothers with disabilities did a little more childcare, suggesting
couples were making satisfactory disability adaptations, as they were more satisfied with their childcare role division than were non-disabled couples.

Many parents are sensitive to the stereotype about parents with disabilities over-burdening or parentifying their children. A number of TLG studies suggest that there is a tendency for mothers with physical disabilities to actually avoid placing their children in helping roles, even the usual household chores\textsuperscript{7,20-23}. In the absence of babycare adaptations, mothers tend to over-use their own bodies, minimizing their babies’ need to adapt\textsuperscript{21-23}.

The disability community’s familiarity with the patterns and norms of our families enhances the ability to differentiate between situations. In our research\textsuperscript{21} the occupational therapist who was a mother with hemiplegia cerebral palsy interpreted a parent’s diapering with one hand as the least demanding. The occupational therapist who was the least experienced in observing baby care by parents with disabilities interpreted tasks as more demanding. Adequate familiarity with parents with disabilities helps one neither exaggerate nor neglect a need for intervention.

Drawing from experience with solutions of parents with disabilities, one can identify applicable solutions and tailor them for diverse families. One can differentiate between what is common and readily adjusted to by babies,(e.g. slow diapering), and what is unusual and worrisome,(e.g. a toddler purposefully knocking down a parent with balance difficulties). One can differentiate between characteristics of babies that may present particular challenges for a particular disability situation of a parent. With experience one is less likely to over-generalize about disability or physical disability in a parent: recognizing subtle differences in functioning; determining crucial distinctions between progressive or relatively stable disabilities; assessing whether disability is long-term or recently acquired or worsened; evaluating whether physical disability is complicated by a cognitive or psychiatric component or a trauma history. One can also
differentiate between parents who do not or would not identify as having a disability or being part of disability culture, versus parents who would be much more open to a disability culture insider as an intervenor.

**Integrating infant mental health and family therapy approaches**

The disability community’s emphasis on contextual and environmental factors is consistent with considering the family system and interaction in the relationships between parents and children. Understanding the experience of families of people with disabilities means considering the perspectives and experiences of all family members as they are affected by the social context.

Addressing disability obstacles can clarify and uncover issues in the infant/parent relationship that can benefit from intervention. With one mother, providing a way for her to carry her baby revealed issues with physical closeness, eventually found to be associated with her own history of childhood abuse. Baby care adaptations can produce rapid change; some individuals, couples or intergenerational families may have difficulty tolerating the sudden increase in functioning, especially at the change-laden time of early parenthood, when roles are renegotiated in family systems\(^{17,24}\). One mother rejected babycare adaptations when use of them meant that the grandmother felt hurt that her helping role was lessened\(^{20}\). Rapid change resulting from adaptive equipment may be especially problematic if there is also a disability change at this point. Adaptive equipment may have an intolerable negative connotation for a parent experiencing a new or worsened disability; for instance, the concrete need for an adapted rather than “normal” crib can represent a painful acknowledgment of loss.

Research on women with physical disabilities has pointed out how some women with disabilities remain with abusive partners because they are physically dependent on these partners and concerned about losing their children because of their disabilities\(^{25}\). This pattern is more
common among women who are isolated from the disability community and its resources, and who do not have the benefit of adaptations that can decrease dependency on assistance.

Integrating disability culture expertise and awareness of adaptations with infant mental health and family therapy/family systems knowledge has been effective in intervention with particularly stressed families. A mother with a post-natal exacerbation of multiple sclerosis became extremely depressed. She relied on her able-bodied husband to provide the infant care to such an extent that the baby was not forming a relationship to the mother and the father was becoming overwhelmed. Baby care adaptations helped alleviate her depression, increasing a balance of functioning in the couple and allowing the relationship between mother and infant to flourish.

Parents with cognitive disabilities

There are many clear differences between parents with physical disabilities and parents with cognitive disabilities. For instance, most parents with cognitive disabilities have a need for long-term intervention in which the change process is slow, and these parents are limited in their ability to initiate adaptations. In contrast to parents with physical disabilities, parentification is often an issue for the children of parents with cognitive disabilities. Yet it is informative to apply the previously discussed disability community themes to a consideration of parents with cognitive disabilities. The themes of spread and differentiation are particularly relevant. There are significant differences, often blurred in practice and research, between cognitive difficulties associated with disabilities such as head injury, stroke, multiple sclerosis, or developmental disability. This discussion is focused primarily on early intervention issues that are applicable to parents with developmental disabilities. However, “developmental disability” or “mental retardation” are labels which are applied to parents with diverse functional abilities as well. Advocates have alleged discriminatory practice when legal and social services systems presume
parental incompetence and the inability to benefit from reunification services, based on the
categorical diagnosis of “mental retardation” or “developmental disability”, rather than on
individual functioning and behavior of a parent with his or her child26.

The need for familiarity is another disability community theme. One needs extensive
experience with parents with cognitive disabilities in order to evaluate parental capability. As
with parents with physical disabilities one cannot discern the full potential in parents with
cognitive disabilities without providing adaptations that are individualized to the parent’s
functioning. Many problems with current practice are related to this issue. For instance, many
children’s protection services departments send parents with cognitive disabilities to generic
parenting classes, which are more likely to undermine their self-esteem than to be helpful. If
parents do not benefit from generic intervention, they are typically portrayed as incapable --
rather than questioning the appropriateness of the intervention. Even curriculum-based
approaches to intervention that are developed specifically for parents with cognitive disabilities
are inherently limited in their responsiveness to the wide variations in functioning of this
population of parents.

As Jeree Pawl has said, “Pulling together the threads of hope and the evidence of possibility is
our task. Often it is not easy. But without real trust, we convey despair --or worse. This
undermining message--which parents will apprehend--interferes with whatever positive
possibilities we might create27(p5). In the case of parents with disabilities, positive possibilities
are enhanced by adaptations.

One needs considerable experience in order to provide effective adaptations for parents with
diverse cognitive limitations. Such adaptations are often neglected and, instead, the parent is
characterized as non-compliant or unable to benefit from services. In contrast to parents with
physical disabilities, parents with cognitive disabilities are much less likely to self-initiate adaptations. Thus, there is a far greater need for adaptations to be introduced by intervenors. Yet professionals lack training and information regarding adaptive strategies, and this is reflected in current problems in evaluation and intervention. The misuse of generic unadapted parenting classes is one example of such poor practice. Evaluation of parenting capability often relies on measures which have not been normed with parents with cognitive disability or which preclude success by those without high verbal and cognitive functioning. Observation of actual parent/child interaction during evaluation may be limited, absent or in an inappropriate setting such as an office. Contextual approaches, such as observation in the home and community, are often neglected. Too often evaluations are conducted by professionals with inadequate familiarity with parents with cognitive disability.

Social stigma and obstacles are issues emphasized in a disability culture perspective. Poverty is a common stressor in the lives of parents with cognitive disabilities and over the years a number of researchers have examined its impact28-32.

Expertise and adaptation during evaluation and intervention are especially crucial to counteract the social stigmatization regarding cognitive disabilities affecting parents as well as professionals. In comparison to physical or sensory disabilities, cognitive impairments tend to be particularly stigmatized or ranked as less acceptable2. The qualitative “life narrative” research of Tim and Wendy Booth28,29 documents the on-going, intense and pervasive effect of stigma and discrimination in the lives of parents with cognitive limitations and their children.

The Booths’ study of adult children of parents with cognitive disability identified the effects of social exclusion as a major problem in the families’ lives. They suggested that this argued for the applicability of the social model of disability to this population of people with disabilities:
“When problems are seen as rooted in people’s personal deficits and limitations they may seem intractable and out of reach. Shifting the focus onto features of people’s lives that can and should be changed challenges the negative stereotypes that inform such thinking and opens up possibilities of social action in support of families”29(p38).

Issues of power differential and empowerment are key concerns in disability culture; they are issues that arise in the face of social stigmatization and exclusion. Parents with cognitive disabilities often have a lifetime of being “one down”, of feeling powerless. Intervenors are inherently “one up” in a power position relative to them. So, it can be effective to intervene in such a way that the power differential is softened or counteracted rather than accentuated. A very respectful approach, eliciting the parent’s goals and ideas and acknowledging one’s own limitations, mistakes, or problem-solving process, can be helpful.

Teaching needs to be handled very sensitively with many parents, particularly those with mild cognitive limitations and long-term issues of “passing” as nondisabled. Many parents have been scarred by disrespectful treatment, including teaching or behavioral intervention that has felt demeaning. Parenthood may be one of the first experiences that has implied normalcy--and teaching that challenges their competence as parents is likely to mobilize resistance or opposition33. Defenses are manifest in a variety of ways, including a tendency to withhold problems or questions, a hypersensitivity to any intervention implying deficits, a rejection of even critically needed supports, polarizing or withdrawing in reaction to didactic approaches.

Adaptation has been described as a central issue in disability culture, related to expectations of a full life despite social obstacles. A respectful orientation is particularly crucial for intervenors working with parents with cognitive disabilities; it is an underlying adaptation needed in the face of pervasive stigmatization. A gradual process of attunement allows one to
discern how direct or indirect one needs to be at a particular point in the relationship with a parent. That is, some parents can tolerate teaching from the outset, others only after a respectful working relationship has been established. An effective approach has been a slow process of observing and building upon concrete moments that emerge from the parent and parent/child interactions in the home setting. In an ongoing way, one can evaluate the interaction between parent and child as well as the impact of one’s interventions on the parent and on parent/child interaction. Over time, the intervenor must adjust to the parent’s particular abilities and limitations in order to be effective yet avoid being patronizing. With experience one learns to intuitively discern and attune to the individual’s sense of time, short and long-term memory (and for what), whether reading is a workable modality for conveying information, processing abilities regarding lengths of sentences, pauses between sentences, series of questions, sequencing, etc. This attunement process involves trial and error, missteps and repair. Particularly around issues of protection, one must sometimes intervene in direct ways that predictably may offend and necessitate repair of the working relationship. One needs to voice concerns along the way and not collude with hazardous or hurtful behavior toward a child. One might decide to facilitate a safety scenario, such as “What would you do if your baby started choking on some food?”, and find that the parent feels disrespected by this. Eliciting parental problem-solving around safety, when possible, is less likely to mobilize resistance and more likely to result in the parent owning the solutions. Over time, as limitations are clarified, one needs to explore openness to setting up the environment to enhance functioning, such as offering or suggesting concrete adaptations to bypass problems like digital clocks, digital thermometers, a premeasured dose of Tylenol, calendars with appointments, watches with alarms, feeding or medication charts, or premixed formula. The need for repetition and problems with
generalization can be dealt with, without nagging or demeaning, by presenting concepts via multiple modalities in varied situations, e.g. video, books, charts, and varied ways of talking about the issue from the perspectives of different intervenors. When parents have intense polarizing or oppositional patterns it can be effective to intersperse important suggestions into other comments, sandwich critical or sensitive material with positive comments, or use a “lightning rod” preface to the suggestion such as “this may seem like a dumb idea but...”. It can be helpful to even the power differential by normalizing not knowing, e.g., “I used to do this with my baby until my neighbor gave me this idea”.

Identifying parent qualities that you admire or can learn from is especially conducive to respectful and effective intervention. Activities that the parent particularly enjoys can be focused on to enhance the role of the parent in interaction with the child. Videotaping can be used to reinforce the strengths of a parent and to enhance the ability of the parent to observe the baby and wonder about his or her inner experience. It can also be used to enhance the parent’s ability to be assertive, in this case about reactions to the intervenor’s actions--as a way of discussing their working relationship. The parent can observe the videotaped mistakes or insensitivities of the intervenor and these can be discussed and repaired. The continuity of this intervenor/parent relationship in the face of negative moments provides a model for the relationship between parent and child. Jeree Pawl has said: “We learn over time that everything we think we know is a hypothesis; that we have ideas, but that we don’t have truth. We learn that those with whom we work have all the information we need, and that this is what we will work with. When we know this, our attitude conveys it; and the child and the parent sense themselves as sources, not objects....In this context, they become aware of a mutual effort--one in which a sense of
partnership can be maintained much of the time. They do not feel weighed, measured, or judged. They do feel listened to, seen and appreciated.\textsuperscript{26}(p.5)

**Integrating infant mental health approaches**

The literature on parents with cognitive disabilities is seldom informed by current mental health perspectives. This is ironic given the degree of trauma in the past and current lives of so many women labeled with developmental disability\textsuperscript{35}. Programs serving mothers with cognitive disabilities report a startling prevalence of trauma histories\textsuperscript{33}. In 1999, 77% of the parents in TLG’s program for parents with cognitive disabilities had personal histories of trauma or abuse.

Infant mental health knowledge has been particularly salient, with its expertise about helping parents develop new models of attachment in which others are experienced as caring and reliable and themselves experienced as worthy of care and capable of nurturing\textsuperscript{36}. Identifying and eliciting the “ghosts in the nursery”\textsuperscript{37} has been effective with many parents with cognitive limitations. Eliciting the meaning of behavior is often effective in producing change that is not achieved by more educative or behavioral approaches. One mother adamantly refused to allow her toddler to attend a childcare center until early memories of school maltreatment and taunting were surfaced. Another mother curtailed the mobility of her baby for long hours while she fastidiously cleaned, continuing this until she was helped to re-process a removal of previous children that she had incorrectly attributed to her messy household. Understanding the meaning and history of behavior can lessen an intervenor’s tendency to judge behavior and therefore enhance the ability to form a positive relationship with the parent in which more positive parental behaviors can be elicited.

Cognitive limitation can mean that verbal working through of past trauma is less feasible than it is with other parents. The nature of the containing relationship with the parent becomes
even more important because it enacts and puts into practice concretely and understandably what we want to be reflected in the infant/parent relationship--e.g., kindness, consistency, responsiveness, respect, handling of change and transitions, limitsetting, exerting influence, negotiating, problem-solving, tolerance of different perspectives, awareness of others’ experience. This kind of relationship between parent and intervenor provides a model for secure attachment for parents who often didn’t have this experience in their own early childhoods. Unfortunately parents with cognitive limits are especially likely to have multiple superficial and short-term relationships with service providers--the exact opposite of what they need.

Early intervention practitioners are often taught to use role modeling to teach parenting skills. Infant mental health experience suggests that this approach should be used cautiously and selectively. Modeling skills may “out-parent the parent” and undermine vulnerable parental self-esteem. They may contribute to the more pervasive problem of the baby being drawn to the practitioner during home-based intervention. A more appropriate stance is to be the intervenor for the relationship between the parent and baby--facilitating and reinforcing positive aspects of the relationship. Recent infant mental health discussions\(^\text{38}\) describe this as “inclusive interaction”. It appears to be even more important to establish inclusive work from the outset with this population of parents, primarily due to the parental performance anxiety issues that arise when the intervenor is too centrally involved with the child. The sort of therapeutic relationship inspired by infant mental health ideas provides modeling at a deeper level than that of skills; one’s relationship with a parent is a model for the infant/parent relationship and is a laboratory for developing abilities that contribute to that relationship.

Infant mental health experience conveys a need for modest goals and a belief that even modest improvements in a baby/parent relationship can have a profound and lasting impact on a
child. Intervention with these families can require considerable patience and necessitates supervision and support for workers so they can support the parent and the parent in turn can nurture the baby. One needs to provide models for attachment on all levels.

**Considering the family context**

An inclusive approach should not only focus on the infant/parent dyad; the family system needs to be included as well. One needs to consider the family’s ambivalence about an increased role for the parent with a cognitive limitation. Birth is a developmental point in the family life cycle when roles are renegotiated and an outsider facilitating even more change may not be well received. A respectful orientation to the family, not just the parent, can be crucial. There is evidence of improved outcome when families provide consistent support that complements the abilities of the parent. Yet the ongoing need for family support can be wearing for families. Particular tensions tend to arise in intergenerational households. Family therapy expertise can be an essential part of intervention and can help the family system sustain positive and respectful support. This is especially crucial because of the social exclusion experienced by adults with cognitive disabilities and the centrality of the family in their social life.

**Service systems instead of disability culture.** Disability culture has seldom functioned as a support or buffer for this population of families. In the absence of positive ongoing disability culture or family support it is especially crucial that services simulate nurturing and practical assistance provided by long-term family involvement. Unfortunately services systems may carry stigma themselves or even be abusive, disempowering or otherwise contribute to the problems of parents. There tends to be inadequate training and supervision, poor reimbursement and high turn-over of providers working with this population. Lack of continuity, patterns of excessive rescuing and subsequent burn-out, judgmental and negative approaches, and interference with
the infant/parent relationship are all too common. Family or individual emotional patterns can be reflected in the service system, e.g., “splitting”, and workers can get in conflicts that reflect and perpetuate clients’ difficulties. There is a strong need for coordinated efforts and interagency teamwork. Though home-based intervention is crucial, services offering peer contact, such as parent support groups, need to be more available.

**Interdependence.** Research on adult children of parents with cognitive limitations found that the strength of the parents’ support system was important to their children’s experience.\(^{29}\)

Current research at TLG is investigating the perceptions of mothers with cognitive disabilities regarding the nurturing versus interfering aspects of their family, community and therapeutic support systems.\(^{41}\) The Booths are particularly eloquent about the problematic aspects in the support networks of parents with cognitive disabilities and present a normalized view of their interdependence during parenthood: “Competence may more properly be seen as a feature of parents’ social network rather than as an individual attribute. The notion of what might be termed ‘distributed competence’ underlines the fact that parenting is mostly a shared activity and acknowledges the interdependencies that comprise the parenting task.\(^{30}\) This is consistent with the disability culture’s contextual view of parenting that refocuses on the elements in the social network and environment that are compensatory and nurturing versus the elements that are undermining and stressful.

We as intervenors need to be self-reflective and vigilant about our own roles in the lives of these parents, ensuring that we are truly contributing to positive outcomes. As respectful intervenors we need to recognize the commonalities as well as the differences between our families. It is hoped that this discussion has increased familiarity with disability culture.
perspectives and has helped practitioners who are crossing the “lines of difference” presented by parental disability.


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