The Quality of Life of Adults with Developmental Disabilities and Their Families in Calgary Canada: A Pilot Study

Mitchell Clark, Ryan Geake, Amanda D. Smith, Mickey Greiner, and Elaine Yost

Abstract—This pilot study investigated the level of satisfaction in a number of life domains in addition to the overall life satisfaction experienced by group of individuals with developmental disabilities in Calgary. This study also examined the quality of life of their families who are often impacted the continuing responsibilities, concerns and anxieties related to their adult child with developmental disabilities. The average ratings from the family and individual responses across the various life domains were computed. The inter-relationships between the dimensions of family quality of life across the various life domains included in the quality of life assessment were also calculated. Understanding these associations may be helpful in identifying supports and policies that enhance individual and/or family quality of life. Finally, the connections between individual and family quality of life were identified to highlight any areas where changes in supports or services to either families or their adult children with intellectual disabilities would support the wellbeing of all family members.

Index Terms—Family quality of life, well-being, developmental disability, personal well-being index, intellectual disability

I. INTRODUCTION

Quality of life (QoL) has been of interest for centuries, but systematic progress in clarifying this construct and in applying it to the assessment and design of services for persons with disabilities and their families has been underway for only the past couple of decades. Research and development activities led to the development of models and assessments of QoL of persons with developmental or intellectual disabilities in the early 1990s [1]-[4]. More recently, this construct has been extended to the lives of families that include persons with disabilities and has become the focus of assessment devices developed for this purpose [5]-[7].

Many of the stresses and life circumstances that undermine the subjective well-being of individuals with disabilities are well documented. A literature review provided by Lyons [8] outlines the changes in awareness of the quality of life of individuals with disabilities through the decades since the 1950s and 1960s with the beginnings of the human rights movement. Through this period, there was growing appreciation of the disadvantage and marginalization experienced by most individuals with developmental disabilities. A number of movements, such as deinstitutionalization, normalization, social role valorization and integration, began in response to societal concerns over the living situations of the majority of people with intellectual disabilities (ID). These movements supported a developing focus on QoL improvement, and services and programs were evaluated based on their contribution to the enhanced well-being of program participants [8].

This pilot study investigates the level of satisfaction in a number of life domains in addition to overall satisfaction with their lives experienced by a group of individuals with developmental disabilities in Calgary, Canada. This study also examines the quality of life of their families who are often impacted by their continuing responsibilities, concerns and anxieties related to their adult child with developmental disabilities. The inter-relationship between the dimensions of family quality of life across the various life domains included in the quality of life assessment are described in the Results section. Understanding these associations may be helpful in identifying supports and policies that enhance individual and/or family quality of life. Finally, the connections between individual and family quality of life are identified to highlight any areas where changes in supports or services to either families or their adult children with intellectual disabilities would support the well-being of all family members.

II. METHOD

A. Participants

Twenty (20) individuals and 19 families were recruited from among the individuals receiving services from three Calgary-based disability-serving agencies.

Of the 20 individuals surveyed, the survey responses provided by thirteen of the participants met the criteria for inclusion in the study.

B. Instruments

The Family Quality of Life Survey [9] includes nine domains as are as follows: (1) Health; (2) Financial well-being; (3) Family relations; (4) Support from other people; (5) Support from disability-related services; (6) Spiritual and cultural values; (7) Careers and preparation for careers; (8) Leisure and recreation; and (9) Community involvement.

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For each of these domains family respondents provided ratings across six dimensions (Importance, Opportunities, Initiative, Stability, Attainment, and Satisfaction).

The Personal Well-being Index – Intellectual Disability (PWI-ID) scale [10] is a quality of life instrument that is one of several parallel forms of the Personal Well-being Index. All of the forms of the PWI measure subjective well-being across seven domains: (1) standard of living, (2) personal health, (3) achievement in life, (4) personal relationships, (5) personal safety, (6) community connectedness, and (7) future security; as well as the broader concept of “life as a whole.”

C. Procedure

The purpose of the survey and process to be used was described to individuals receiving service through participating agencies. They were assured that if they were interested in participating, their responses would be held in strict confidence and they could change their mind at any time.

Families of those individuals who were interested in participating were contacted. In addition to requesting their approval they were also asked if they were willing to participate in the study.

Five interviewers were recruited to conduct the interviews with both the participating individuals with disabilities and their families. The interviewers all had experience working with persons with disabilities and their families. All the interviewers received training to prepare them for the use of the survey forms for both families and individuals.

III. RESULTS

Twenty family members from 19 families participated in the interviews which followed the Family Quality of Life (FQoL) survey form. The adult children of these families also participated in separate interviews conducted using the PWI-ID although meaningful results were obtained from only 13 of these individuals.

The adult children with ID in these families ranged in age from 21 up to 62 years and demonstrated varying levels of support needs and communications abilities. Of the individuals with developmental disabilities included in the results, only two were living in the family home. At the same time, eight family members indicated they found the level of responsibility they had for the individual with ID to be “more” or “much more” than they would like, suggesting that they felt stressed in relation to the support needs of the individual with ID in their family.

Table I provides a description of the outcomes of the Family QoL survey. Note that ratings on all of the domains are out of a total possible of 5.00 for each item.

As is usually the case in surveys of family quality of life, all of the domains of family life are seen as important. The pattern of relative importance across the domains provides some unusual results when compared to other studies using the FQoL survey. In particular the rating of service supports (4.94) and its rank in third place (very close behind the “Health” and “Family” domains) suggests how critical these services are seen for family well-being. This indicates a much stronger focus on formal services than is typical in other studies. In spite of the importance of these support services to families, they viewed these services as most likely to decline in the near future in comparison to all of the other domains. This situation is suggestive of the anxiety felt by families that services may be under threat due to possible government cut backs or shifting priorities. Other domains that are rated as more likely to decline include Careers (a domain that is also responsive to economic downturns) and Health. The view that family health is expected to decline is likely related to the age of many of the respondents.

Support from others, while viewed as relatively important (3.47), was seen as the least important of all of the dimensions of FQoL. Families indicated there are relatively few opportunities to secure Support from Others (opportunities= 2.79) and rate their efforts to establish Support from Others (2.06) as the lowest in comparison to all of the other domains. Their actual attainment of Support from Others is even lower (1.78), indicating families experience limited support other than through more formal supports, the stability of which, as noted above, cause them significant concern.

<table>
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<th>TABLE I: MEAN SCORES AND SDS FOR THE SIX FAMILY QUALITY OF LIFE DIMENSIONS ACROSS THE NINE DOMAINS</th>
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Note: Due to an error in the data collection form, ratings for the initiative dimension of the Community Domain were not collected.

The mean ratings of satisfaction with the various life domains and their inter-correlations were computed between
all of the variables measured by the two instruments used in this study. Family responses about different areas of life were sometimes related to one another. A few of the interrelationships related to supports available to the family are identified below:

- People who felt strongly that their family member’s needs for disability supports were met were also happiest with their family’s health.
- Those who were happier with their disability services and financial opportunities also had a better quality of life.
- People who felt that disability services would get worse also saw support from others as more important.
- People who were happier about their family relationships, the support they got from others, their leisure activities and their community involvement were also more likely to see family health as stable or improving soon.
- People who had more opportunities to get disability services for their family member also had more opportunities for leisure activities and community involvement.
- People said if they had less support from extended family and friends in the future, their community involvement would be reduced and their spiritual or cultural values would have less impact on their quality of life.
- People who had more opportunities to get disability services for their family member also had more opportunities for leisure activities and community involvement.

#### IV. Discussion

Overall, the adults with developmental disabilities were happy with their lives. They were most happy about their goals, community involvement and standard of living. They were least happy about their health and their future life.

The importance of disability services to the families in this survey is clear (see Table I). These supports are viewed as critical and are associated with various components of the quality of life of families. As noted earlier, satisfaction with Service Supports is strongly associated with Health attainment in families. Various dimensions of the families’ financial well-being are associated with several dimensions of the Service Supports domain. Satisfaction with family relationships is also associated with satisfaction with Service Supports. There are numerous further associations with dimensions of the Service Supports domain that suggest additional relationships between Service Supports and other aspects of family quality of life.

Disability supports are also associated with the well-being of their adult child with a developmental disability. Personal health and future security ratings by the survey participants with ID are related to the family ratings of the attainment of Support Services. When families said disability services were important and that their family member had good services, the adult with disabilities was more likely to see his or her future as secure.

For families of adults with developmental disabilities, the ability to get good, stable disability services was strongly related to many aspects of their quality of life, such as health and the ability to pursue a career, social support, and leisure or community involvement. When good services were not in place or were expected to get worse, aging parents must spend time and energy to meet the individual’s needs. Other aspects of their quality of life (such as health, work and community involvement) suffer. Adults with disabilities were also more likely to worry about their future when families say services are lacking or insecure.

When disability supports were not secure, social support became more important to families. While the value of support from others on personal well-being is well
established, the families responding to the survey do not see it as particularly important and they reported receiving very limited support (see Table I). This may be an area that could be developed to enhance QoL. As well as ensuring greater awareness of the benefits of support from extended family, friends and neighbors and others, there is a need to identify barriers to establishing these connections and assist families in this process. Services should be developed to train facilitators who then work with families and individuals to develop networks made up of these informal supports. The expansion of this type of service may be quite helpful in developing new supportive relationships particularly when government cutbacks reduce the availability of agency supports.

It is hard on aging family members to have to meet the disability needs of loved ones unless they have social support. However, it may be harder to look for support from friends, extended family, neighbors or others when families have to attend to disability support needs not met by formal services. Help with developing family support networks would promote greater family health and well-being.

The results of the pilot study survey of individual well-being and family quality of life ratings provide valuable descriptions of both their strengths and challenges. In addition, the interrelationships between the dimensions of the various life domains surveyed, across families and individuals, provide suggestions of how responses to challenges in one domain might be supported by changes in other aspects of their lives.

V. LIMITATIONS OF STUDY AND SUGGESTIONS FOR FURTHER RESEARCH

There were several limitations to this study. First, the number of families and individuals surveyed provided a relatively small sample. It would have been desirable to have a larger number of individuals and families, a situation that would have provided for greater confidence in the results.

A second issue that limits the results is related to the sample that was recruited. All of the families and individuals included in this study were receiving services from agencies providing disability services and supports. Families, who were on wait-lists for services and as a result were not receiving any supports, may have very different responses. Also, those families who were personally managing the supports for their child with a disability could be expected to have very different responses, particularly in the Service Supports and, perhaps, in the Support from Others domains. There may be various differences between families that choose to utilize supports provided by formal disability services and those families who choose the family–managed care options.

Finally, the results described here are specific to the Calgary area. There may be substantial differences in the level of satisfaction across the various quality of life domains in other regions of Alberta and in other urban and rural areas across Canada and in other parts of the world.

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REFERENCES


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