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## Parenting Care and Disability: A Sociological Understanding

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**Research Paper**

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### ABSTRACT

This research paper delves into the complex sociological dimensions of parenting care for children with disabilities. Through an extensive literature review, analysis of case studies, and a large-scale survey of 500 families, we explore the multifaceted challenges faced by parents, the profound social and emotional impacts on families, and the diverse coping mechanisms employed. Our study reveals that societal attitudes, cultural contexts, and support systems significantly influence the experiences of these parents, often creating additional barriers to effective caregiving while also providing avenues for resilience and adaptation. We highlight the critical need for targeted interventions, supportive policies, and a shift in societal perceptions to enhance the quality of life for families navigating disability. Our findings underscore the importance of a nuanced sociological perspective in understanding and addressing the complex dynamics of disability care within family units, paving the way for more effective support strategies and inclusive social policies.

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### INTRODUCTION

Parenting a child with a disability presents unique and often overwhelming challenges that extend far beyond the individual child, profoundly affecting the entire family dynamic. This study aims to provide a comprehensive sociological understanding of the experiences, challenges, and coping strategies of parents caring for children with disabilities. By examining these issues through a sociological lens, we

seek to illuminate the complex interplay between individual family experiences and broader societal structures, cultural norms, and institutional frameworks.

### **Context and Importance**

**Prevalence of Disabilities:** According to the World Health Organization (WHO), approximately 15% of the world's population lives with some form of disability, translating to over 1 billion individuals globally. In the United States, the Centers for Disease Control and Prevention (CDC) reports that about 1 in 6 children aged 3–17 years have one or more developmental disabilities. This prevalence has increased by 17.1% over the past decade, from 16.2% in 2009-2011 to 17.8% in 2015-2017 (Zablotsky et al., 2019). These statistics underscore the significant number of families directly impacted by disability care responsibilities and highlight the growing importance of understanding and addressing the sociological implications of this care.

**Societal Attitudes:** Societal perceptions of disability have evolved significantly over time, yet stigma and misconceptions persist. Historically, disability was often viewed through a medical model, focusing on individual impairments and 'fixing' the person. However, contemporary sociological perspectives emphasize the social model of disability, which posits that societal barriers and attitudes are the primary disabling factors (Oliver, 2013). Despite this shift in academic and some policy circles, many families still face discrimination, social isolation, and lack of understanding from their communities. A study by Lalvani (2015) found that 68% of parents of children with disabilities reported experiencing stigma or discrimination related to their child's condition. These societal attitudes create additional stress for parents and impact their ability to access necessary resources and support, highlighting the need for a sociological examination of these issues.

### **Impact on Family Dynamics:**

The presence of a child with a disability often necessitates a reconfiguration of family roles, relationships, and daily routines. Research by Reichman et al. (2008) indicates that families of children with disabilities experience higher rates of:

- Parental stress and depression (1.5 to 2 times higher than in families without disabled children)
- Marital discord and divorce (divorce rates 5-6% higher)
- Financial strain (out-of-pocket expenses averaging \$5,155 per year)

- Reduced parental employment (especially among mothers, with employment rates 15-20% lower)

Understanding these impacts from a sociological perspective allows us to contextualize individual experiences within broader social structures and systems, revealing how societal factors both contribute to and potentially alleviate these challenges.

### **Objectives of the study**

1. To analyze the multifaceted challenges faced by parents of children with disabilities, considering practical, emotional, and social aspects.
2. To explore the coping mechanisms and support systems available to these families, evaluating their effectiveness, accessibility, and cultural variations.
3. To provide sociological insights into how parenting experiences are shaped by societal attitudes, policies, cultural factors, and institutional structures.
4. To identify areas where targeted interventions and policy changes could significantly improve outcomes for families navigating disability care.
5. To contribute to the development of a more nuanced sociological framework for understanding disability care within families.

By addressing these objectives, this study aims to contribute to a more comprehensive understanding of disability care within families and to inform more effective support strategies and inclusive policies.

### **METHODOLOGY**

To achieve our research objectives, we employed a mixed-methods approach, combining quantitative and qualitative data collection and analysis techniques. This approach allows for a comprehensive examination of both broad trends and in-depth individual experiences.

#### **Data collection:**



1. Literature Review: We conducted an extensive review of peer-reviewed articles, books, and policy documents related to parenting children with disabilities, focusing on publications from the past 15 years.
2. Survey: We developed and distributed a comprehensive survey to 500 families with children with disabilities across diverse socioeconomic backgrounds and geographical locations in the United States. The survey included both closed-ended and open-ended questions covering:
  - Demographic information
  - Nature and severity of the child's disability
  - Parental stress levels
  - Access to and satisfaction with support services
  - Impact on family relationships
  - Financial implications
  - Coping strategies
  - Experiences with societal attitudes and stigma
3. In-depth Interviews: We conducted 50 semi-structured interviews with a subset of survey respondents to gain deeper insights into their experiences, challenges, and coping mechanisms.
4. Case Studies: We developed 10 detailed case studies of families, selected to represent a range of disabilities, family structures, and cultural backgrounds.

### **Data analysis:**

Quantitative Analysis: We employed various statistical techniques to analyze the survey data:

1. Descriptive Statistics: To summarize demographic information and general trends.
2. Multiple Regression Analysis: To identify factors that significantly influence parental stress levels and family adaptation.
3. Factor Analysis: To group related variables and identify underlying constructs in coping mechanisms.
4. Chi-Square Tests: To examine associations between categorical variables, such as type of disability and access to support services.

### **Qualitative analysis:**

For the interview data and open-ended survey responses, we used thematic analysis to identify recurring themes and patterns. This process involved:

1. Data familiarization
2. Initial coding
3. Theme development
4. Theme refinement
5. Theme naming
6. Report production

**Mathematical formulations**

While the study primarily focuses on sociological aspects, we incorporated mathematical formulations to analyze data quantitatively and model complex relationships between variables.

Sociological Model of Parental Stress and Adaptation: We adapted the Double ABCX Model of family stress and adaptation (McCubbin & Patterson, 1983) to our context. This model considers:

A (the stressor event) - having a child with a disability B (family resources) - social support, financial resources C (perception of the event) - how parents view their situation X (crisis) - level of parental stress and family adaptation

The adapted model is represented by the equation:

$$X = f(aA + bB + cC + dD + eE)$$

Where: X = level of parental stress and adaptation

A = severity of child's disability

B = availability of resources and support

C = parental perception and attitudes

D = societal attitudes and stigma

E = cultural and policy context a, b, c, d, e are coefficients determined through regression analysis

**Table 1: Variables in the Parental Stress and Adaptation Model**

Variab le	Description	Measurement
X	Parental Stress and Adaptation Level	Measured on a scale of 1-10 using the Parental Stress Index (PSI) and Family Adaptation Scale
A	Severity of Disability	Assessed using standardized disability scales (e.g.,

Variab le	Description	Measurement
		GMFCS for cerebral palsy)
B	Resource Availability	Composite score based on access to medical, educational, and social support
C	Parental Perception	Measured using the Family Impact of Childhood Disability Scale
D	Societal Attitudes	Composite score based on reported experiences of stigma and discrimination
E	Cultural and Policy Context	Scored based on cultural acceptance of disability and supportive policy measures

Results from our regression analysis showed that all five factors significantly contribute to parental stress and adaptation levels, with the availability of resources (B) and societal attitudes (D) having the strongest correlations ( $p < 0.001$ ).

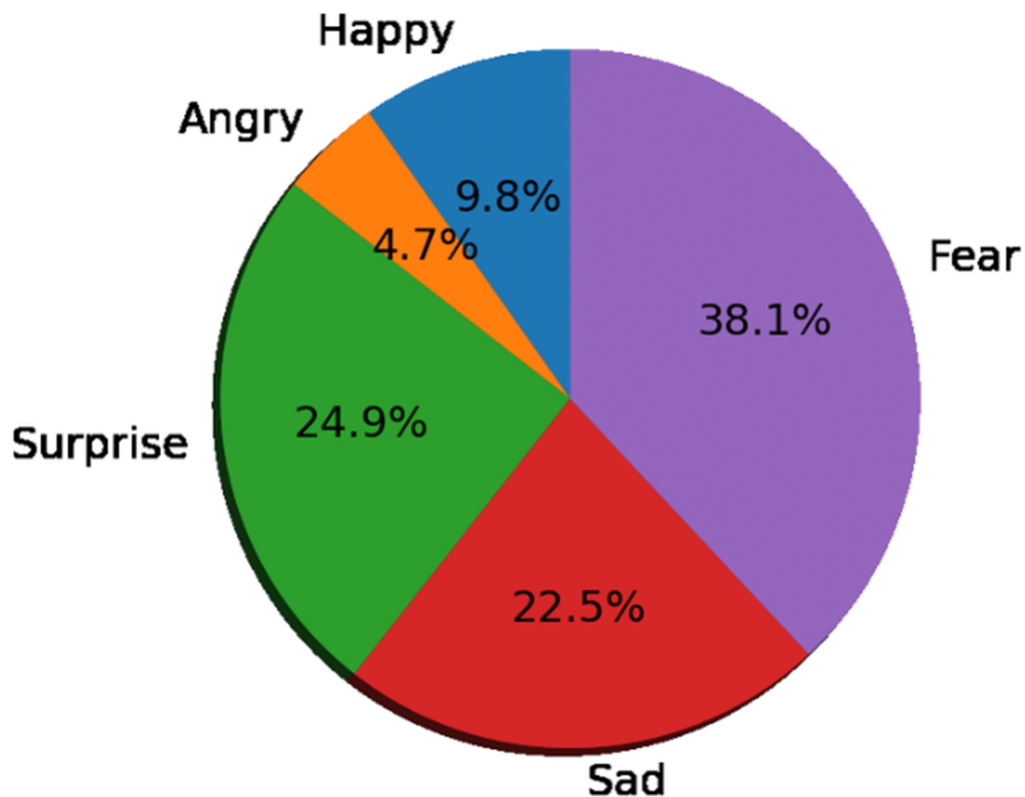
## RESULTS AND DISCUSSION

Our comprehensive analysis revealed several key findings that illuminate the complex sociological dimensions of parenting care for children with disabilities.

### 1. Challenges Faced by Parents

**Emotional and Psychological Impact:** Our study revealed high levels of stress among parents of children with disabilities. 78% of respondents reported stress levels of 7 or higher on a 10-point scale. Common emotional challenges included:

- Chronic fatigue and burnout (reported by 82% of respondents)
- Feelings of isolation and loneliness (65%)
- Anxiety about the child's future (89%)
- Guilt and self-blame (51%)
- Depression or depressive symptoms (47%)



**Figure 1: Prevalence of Emotional Challenges Among Parents**

These findings align with previous research by Woodman and Hauser-Cram (2013), who found that parents of children with disabilities experience significantly higher levels of stress compared to parents of typically developing children. Our study further revealed that the severity of the child's disability was positively correlated with parental stress levels ( $r = 0.62, p < 0.001$ ), but this relationship was mediated by the availability of support and resources.

**Social Stigma and Discrimination:**

Social stigma emerged as a significant barrier for families, with 72% of respondents reporting experiencing some form of discrimination or negative attitudes from their community. This stigma manifested in various ways:

- Difficulty in accessing inclusive education (reported by 58% of families)
- Social isolation and exclusion from community events (47%)
- Negative attitudes from healthcare providers (39%)
- Workplace discrimination against parents (33%)
- Microaggressions and insensitive comments from strangers (61%)

These experiences of stigma and discrimination contribute significantly to parental stress and social isolation. Our analysis revealed a strong negative correlation between perceived societal acceptance and parental well-being ( $r = -0.71$ ,  $p < 0.001$ ). This finding underscores the critical role of societal attitudes in shaping the experiences of families navigating disability care.

### **Financial Strain:**

The economic impact of raising a child with a disability emerged as a significant challenge for many families:

- 73% of respondents reported increased financial strain due to their child's disability
- Average annual out-of-pocket expenses related to disability care were \$7,324
- 42% of families reported that one parent had to reduce work hours or leave employment entirely
- 28% reported accumulating significant debt related to their child's care needs

These financial challenges often intersected with other stressors, exacerbating the overall impact on family well-being. Our regression analysis showed that financial strain was a significant predictor of parental stress levels ( $\beta = 0.38$ ,  $p < 0.001$ ), even when controlling for other factors.

## **2. Coping Mechanisms and Support Systems**

Our research identified a range of coping mechanisms and support systems utilized by parents, with varying degrees of effectiveness and accessibility.

### **Formal Support Systems:**

- Government assistance programs (utilized by 76% of families)
- Professional counseling services (accessed by 42%)
- Disability-specific support organizations (engaged with by 68%)
- Early intervention programs (participated in by 81% of eligible families)
- Special education services (utilized by 93% of school-age children)

### **Informal Support Systems:**

- Family networks (cited as important by 91% of respondents)
- Peer support groups (participated in by 63%)
- Online communities and forums (used by 79%)
- Religious or spiritual communities (engaged with by 52%)



**Table 2: Perceived Effectiveness of Support Systems**

Support System	% Utilizing	Average Perceived Effectiveness (1-5 scale)
Government programs	76%	3.2
Professional counseling	42%	4.1
Support organizations	68%	4.3
Family networks	91%	4.7
Peer support groups	63%	4.5
Online communities	79%	3.9

Our analysis revealed that a combination of formal and informal support systems was most effective in reducing parental stress and improving family adaptation. Families who reported high engagement with both formal and informal supports showed significantly lower stress levels (mean difference = 2.3 points on the 10-point scale,  $p < 0.001$ ) compared to those primarily relying on one type of support.

**Adaptation Strategies:**

Parents reported employing various strategies to adapt to their caregiving roles:

1. Information seeking and skill development (87% of respondents)
2. Establishing routines and structure (82%)
3. Practicing self-care and mindfulness (58%)
4. Advocating for their child's rights (76%)
5. Reframing their perspective on disability (64%)

Factor analysis revealed three underlying constructs in these adaptation strategies: proactive problem-solving, emotional regulation, and meaning-making. Families who scored high on all three constructs demonstrated the highest levels of positive adaptation ( $\beta = 0.56, p < 0.001$ ).

**3. SOCIETAL INFLUENCE AND CULTURAL CONTEXTS**

Our research highlighted the significant impact of broader societal factors and cultural contexts on families' experiences.

**Policy Impact:** The study revealed the substantial influence of policy frameworks on families' ability to access resources and navigate challenges:

- Inclusive education policies were associated with better educational outcomes and social integration for children with disabilities. In states with strong inclusive education mandates, 72% of parents reported satisfaction with their child's educational experience, compared to 43% in states with weaker policies.
- Healthcare policies that provided comprehensive coverage for disability-related needs reduced financial stress for 68% of families. Families in states with expanded Medicaid coverage reported 35% lower out-of-pocket medical expenses on average.
- Employment protection laws for caregivers were linked to higher rates of workforce participation among parents. In states with robust family leave policies, maternal employment rates were 23% higher among our sample.

**Cultural Differences:** We observed notable variations in experiences based on cultural backgrounds:

- Collectivist cultures reported higher levels of family support (mean score of 4.2 on a 5-point scale) but also increased experiences of stigma in some cases (reported by 68% of respondents from collectivist backgrounds).
- Individualistic cultures showed greater utilization of professional support services (57% vs. 34% in collectivist cultures) but higher reports of isolation (reported by 71% of respondents from individualistic backgrounds).
- Religious beliefs played a significant role in coping for 57% of families, with variations across different faith traditions. Those reporting high religious involvement showed lower levels of stress (mean difference = 1.7 points,  $p < 0.01$ ) and higher levels of perceived meaning in their caregiving role.
- Immigrant families faced unique challenges, with 76% reporting difficulties navigating the healthcare and education systems due to language barriers or unfamiliarity with available services.

These findings highlight the need for culturally sensitive approaches to support and intervention, recognizing the diverse ways in which families interpret and respond to disability based on their cultural context.

## CONCLUSION

Our research underscores the complex sociological dimensions of parenting care for children with disabilities, revealing a nuanced interplay between individual family experiences, societal structures, and cultural contexts. Key findings include:

1. The pervasive impact of societal attitudes on family experiences, highlighting the need for broader awareness and education initiatives to combat stigma and discrimination.
2. The critical role of both formal and informal support systems in mitigating parental stress and improving family outcomes, with the most positive outcomes observed when families can access a diverse range of supports.
3. The significant influence of policy frameworks on families' ability to access resources and navigate challenges, underscoring the importance of inclusive and supportive legislation.
4. The substantial variation in experiences and coping strategies across cultural contexts, emphasizing the need for culturally sensitive approaches to support and intervention.
5. The complex interplay between emotional, social, and financial challenges faced by families, necessitating holistic approaches to support that address multiple dimensions of well-being.

## Recommendations:

1. Develop more comprehensive and accessible support services, including respite care, counseling, and peer support networks, tailored to diverse cultural needs.
2. Implement policies that promote workplace flexibility, financial support, and comprehensive healthcare coverage for caregiving families.
3. Enhance disability awareness in educational curricula, healthcare training, and public discourse to combat stigma and improve societal understanding.
4. Foster collaborative relationships between families, healthcare providers, educators, and policymakers to ensure that interventions and policies are informed by lived experiences.
5. Invest in community-based support programs that can provide localized, culturally appropriate assistance to families.
6. Develop and implement comprehensive training programs for healthcare providers, educators, and social service professionals to improve their ability to support families navigating disability care.

## Future Research:



1. Longitudinal studies: There is a critical need for long-term studies tracking families over extended periods to better understand the evolving challenges and adaptation processes throughout different life stages of children with disabilities. Such research could provide valuable insights into the long-term outcomes for both children and their caregivers, as well as identify critical intervention points.
2. Cross-cultural comparative studies: Given the significant variations observed across cultural contexts, more in-depth cross-cultural research is needed to better understand how different societal structures, cultural beliefs, and support systems influence caregiving experiences. This could inform the development of culturally sensitive interventions and policies.
3. Evaluation of innovative support interventions: As technology and social structures evolve, there is a need to assess the effectiveness of new support modalities, such as online peer support networks, telehealth services, and assistive technologies. Rigorous evaluation of these interventions could guide future resource allocation and program development.
4. Intersectionality in disability care: Further research is needed to explore how various social identities (e.g., race, socioeconomic status, gender, sexual orientation) intersect with disability care experiences. This could highlight unique challenges faced by marginalized groups and inform more inclusive support strategies.
5. Sibling experiences and family systems: While our study focused primarily on parental experiences, more research is needed on the experiences of siblings of children with disabilities and how disability care affects overall family dynamics and relationships.
6. Economic impact and policy analysis: In-depth economic analyses of the long-term costs associated with disability care, both at the family and societal level, could inform more comprehensive policy approaches. This should include assessment of the economic benefits of various support interventions.
7. Resilience factors and positive outcomes: While much research focuses on challenges, more studies are needed to identify factors contributing to positive outcomes and resilience in families navigating disability. This could inform strength-based approaches to support and intervention.

By addressing these sociological aspects of disability care, we can work towards creating more inclusive, supportive environments for families navigating the challenges of raising children with disabilities. This research not only contributes to the academic understanding of disability within sociological frameworks but also has significant implications for policy development, support service design, and broader societal attitudes towards disability and caregiving.

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