International Scholarly Research Network
ISRN Nursing
Volume 2011, Article ID 708318, 8 pages
doi:10.5402/2011/708318

Research Article

Mothers with Serious Mental Illness: Their Experience of “Hitting Bottom”

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Received 26 February 2011; Accepted 20 March 2011

Academic Editors: S. Keeney and B. Lundman

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This study sought to understand the experience of “hitting bottom” from the perspective of 32 mothers with serious mental illness. Secondary narrative analysis of 173 stories about experiences related to hitting bottom were identified. Enactment of their perceived mothering roles and responsibilities was compromised when confronted by the worst of illness. Subsequent to women’s descent to bottom was their need for a timely and safe exit from bottom. An intense experience in bottom further jeopardized their parenting and treatment self-determination and, for some, their potential for survival. The results suggest that prevention of bottom is feasible with early assessment of the diverse issues contributing to mothers’ vulnerabilities. Interventions to lessen their pain may circumvent bottom experiences. Healing necessitates purposeful approaches to minimize the private and public trauma of bottom experiences, nurture growth towards a future, and establish resources to actualize such a life.

1. Introduction

Women diagnosed with serious mental illness (SMI) are just as likely to parent as other adult women in the general population. Research shows that this group of women have normal fertility rates [1] and higher rates of induced abortion [2]. Studies, however, report conflicting findings about the mean number of offspring for women with SMI. While some comparison studies report that women with SMI bear fewer than the average number of children [3], others show the opposite result [1]. This difference may partially be explained by women’s age-at-first-diagnosis. That is, earlier age-at-first-diagnosis may contribute to a smaller mean number of offspring [4].

Globally, there is increasing awareness of the parenting status of mental health service users. It is estimated that approximately 60% of women with enduring mental health issues, internationally, have dependent children [5]. In both the United States and the United Kingdom, research shows that approximately one third of the female mental health services users are parents with dependent children [6–8].

In Canada, one in every ten children or 12.1% of all children under the age of 12 lives with a parent that had at least one psychiatric disorder in the previous 12 months [9]. Furthermore, only about one quarter of these parents reported receiving recent mental health services.

The expanding research on parents with SMI remains largely focused on maternal role challenges in relation to the presence of illness. Some researchers suggest that families led by these women are among the most vulnerable and unsupported in communities [10–13]. As a group, they often contend with a range of health and social disadvantages that are compounded by illness. Such challenging circumstances include comorbid conditions [14, 15], poverty [16], discrimination [17], lack of social support [18], lack of health and parenting information [19], inadequate housing [20], trauma [21], and constant fear of their children’s apprehension by Children’s Protection Services [22].

According to Hollingsworth [23], a diagnosis of SMI is a “fast track” to the termination of parents’ rights to the custody of their children. In a sample of 1954 mothers with SMI and other coexisting conditions, two-thirds were
separated from all their dependent children [24]. Of the women who lived with at least one of their offspring, they reported significantly less stressful life events. Due to the stigma of mental illness and its associated socioeconomic implications, the juxtaposition for mothers with SMI relates to the reality that requests for help heighten the risk of child removal. It is therefore understandable that the lack of services or the feared implications of exposure by illness may contribute to mothers parenting without professional intervention.

Without support, however, these mothers’ parenting abilities may be jeopardized by the complex interplay of mothering, mothering contexts, illness, services needs, and resources. Bybee and associates [25] conducted a correlation study to explore factors affecting the day-to-day functioning of a group of 379 mothers with SMI. For mothers with low service involvement levels, functioning was inversely related to social stress. These researchers suggest that intensive services mediate the effects of stress, thereby allowing mothers to better attend to their parenting responsibilities. This strength-based orientation to parenting identifies that access to adequate resources fosters parenting success, regardless of specific diagnostic label [16, 23]. According to Oyserman and associates [18].

Little research to date has sought out the voices of parents in disadvantaged circumstances… even though such research is critical if we are to understand the mechanisms underlying positive parenting (page 2505)

Although not all mothers with SMI experience negative parenting realities, for those that do, little is known about their subjective experiences. There are recent qualitative evidence reviews particular to parents with SMI [18, 26, 27]. The qualitative evidence presents emic perspectives from two groups: mothers and mental health service providers. From a maternal orientation, key findings include acknowledgement of role complexity, desire to minimize negative and optimize positive familial and individual outcomes, and identification of the need for significant trusted other [28–37]. A few studies present the perspective of those committed to servicing parents with SMI [38–41]. Overall, findings suggest the need for a range of individualized and purposeful material, emotional, educational, and relational supports to extend the benefits of existing treatment models.

In an effort to contribute to the developing understanding of women’s experiences, this paper explores the specific experience of “hitting bottom.” To date, the experience and language of hitting bottom in the nursing literature has been associated primarily with substance-related disorders [42] and to a lesser extent life-threatening chronic illnesses [43]. Therefore, for this particular group of women with a primary diagnosis of SMI, further examination of their “hitting bottom” experience is warranted. Further, from a women’s agency and subjective orientation, knowledge about bottom experiences is imperative to the development of a strength-based, family-centered approach to well-being [16, 36].

2. Method

2.1. Design. The study design is a supplementary analysis, a type of secondary qualitative analysis [44, 45]. This approach is analogous to Thorne’s [46] retrospective interpretation and analytic expansion types of secondary analysis. Supplementary analysis involves an in-depth examination of a theme or subset of data for the purpose of extending the primary work. Hitting bottom experiences were initially identified in a grounded theory study [31]. Bottom involved mothers’ awareness of their distance from their children, compounded by an uncertainty of how to protect their children from witnessing intense illness. In two related studies about maternal health, mothers with SMI told unprompted, compelling stories that contained patterns revealing their human experience of “bottom”. Given that the data from these three studies, conducted between 2000 and 2008, were not analyzed to explore this phenomenon, secondary analysis offers a new layer of understanding of bottom as experienced by mothers. As Kohler Riessman [47] stated, the issue is not whether subsequent analysis of data is “truer,” rather, reinterpretation “illuminates a layered complexity…there is never a single authorised meaning” (page 321).

2.2. Setting and Sample. All of the ethically approved primary studies were conducted in northeastern Ontario. A variety of specialized mental health services were accessed since not one of them provided specific parenting programs for persons with SMI. Using theoretical and purposive sampling, a total of 37 inpatient and community women participated in the original studies. Study inclusion criteria were English-speaking, mentally competent, under a psychiatrist’s care for a SMI as defined by Ministry of Health [48], and a mother of at least one child between the ages of two and sixteen years, either living with or separate from them.

2.3. Data Collection and Analysis. One to six audio-taped semistructured interviews were the main data-gathering strategies in all studies. In this study, narrative analysis of stories rather than the original studies’ approach of thematic analysis was chosen. Narrative analysis preserves rather than fragments participants’ stories [49]. Further, stories are frequently identified in qualitative data as a way in which participants construct meaning. Through analysis of stories, understanding of the emic perspective is sought. To complete this study’s analysis, an eclectic narrative analysis approach was developed by the third author [50]. It combines functional analysis model of individual stories [51] with Agar and Hobbs [52] strategy of identifying story coherence across interviews.

Figure 1 outlines the six iterative steps of the analysis process. Data about hitting bottom was extracted from each of the verbatim, unmarked transcripts and placed in a separate file. For this aggregate data set, the researchers individually deconstructed each story of bottom into its six elements inclusive of abstract, orientation, complicating action, resolution, evaluation, and coda [51]. Each element was compared across all stories to determine common, content, structure, function, and meaning. In addition,
3. Results

3.1. Storytellers. All of the women told stories of mothering and illness. Thirty-two of them specifically described a hitting bottom experience. The age range of these mothers was 19 to 38 years. Seventeen women lived with the biological fathers of the children. Twenty-two mothers lived with their children, and of the 10 mothers separated from their children, all continued to keep contact with them. The women parented a total of 49 children ranging in age from two to 15 years. Each of the 32 mothers identified themselves as having a SMI diagnosis. During their length of illness, varying from three to twenty years, many reported receiving more than one mental health diagnoses. Several mothers acknowledged that they had been aware that “something wasn’t quite right” for years prior to seeking psychiatric services. Other mothers said they became “suddenly” ill after giving birth. The most common primary psychiatric diagnosis they received during their course of illness was a major mood disorder.

To represent the subjective understanding of bottom as expressed by the storytellers, the findings are presented in two sections, stories of bottom and a case example. First, the content and structure components of the 173 stories of hitting bottom are identified as the worst of illness, the escape imperative, and the consequences. Then, the meaning and function of the stories are outlined separately. Each of these components is illustrated by story excerpts. The inclusion of the symbol “/” denotes pauses often between ideas or phrases. Each excerpt is coded by an assigned participant identifier, for example, P1. To present a unified account of bottom, a case example inclusive of all story components concludes the findings. This format of presentation explicates the researchers’ analytic logic, a reflection of fundamental credibility of this study [49, 54].

3.2. Stories of Bottom

3.2.1. Content and Structure. Across the data base, the women told stories of hitting bottom contextualized by overwhelming illness that impeded enactment of their perceived roles and responsibilities as mothers. Three common content and structural components were identified in these stories. Mothers consistently talked of bottom as what they conceived to be the worst of illness. Second, mothers described their private internal dialogue as they searched for an exit from bottom with a limited range of possibilities. This structural and content component of the bottom stories was labelled as the escape imperative. Finally, these stories also included retrospective descriptions of the implications of being at bottom for themselves and their children. This final structural and content constituent was labelled as the consequences.

3.2.2. The Worst of Illness. Women told stories characterizing the bottom experience as the worst possible illness reality. This story structure included three content areas: imposed kinetic force, unrecognizable self, and disaffiliation.

3.2.3. Imposed Kinetic Force. Mothers attributed an actual or metaphorical motion to the worst of illness. As such, they explained an inescapable, often unexpected, and rapid descent to bottom as they found themselves “sliding” or “spiralling” out of control, “rolling down a hill”, and “falling, falling, falling”. This motion precipitated “losing touch”, “hitting hard” and finding themselves “at the end of their rope” in a “tunnel”. Excerpts of complete bottom stories which represent the kinetic experience of the worst of illness are as follows.

P2 I started sliding and sliding and I knew I won’t be able to last much longer.
3.2.5. Disaffiliation. Women’s accounts included the interplay of multiple familial, economic, health, and social service stressors. These external pressures combined with their SMI trajectory exacerbates real or perceived dissociation from others during the worst of illness. This disconnectedness from significant others and loss of typical role functioning served to “isolate” and “embarrass” them.

P19 Like they (Children’s Services) just took the kids away/My mom did not want me to live with her because she had my (children) and we were fighting all the time.

P29 It is more about lost. I was just lost. Just feeling lost and not knowing where I belonged or who loved me.

3.2.6. The Escape Imperative. Within the bottom stories, all women spoke about an intensive internal dialogue about an escape from bottom. This self-talk focused on their critical need to alleviate their suffering at bottom. Determined to search for an exit, they identified that there were “no easy answers” especially in the context of their diminishing physical, emotional, cognitive, and social resources. In such “devastating,” “discouraging,” “destructive,” and “cruel” circumstances many women considered the merit, pragmatics, and implications of suicide for themselves and significant others, most importantly their children as a way out of bottom.

P4 If want the best for my loved ones than maybe I need to disappear in the sense that yeah/ok/finish it off then. I may be doing more harm to my children and to my loved ones by being here than not being here at all. So basically that was the black and white of the situation at that point. I was incessantly tired, and started thinking very negatively and not being able to see beyond it you know. I want the best for my family.

P23 I’d just be sitting here and just doing nothing and all of a sudden-boom. And it snowballed down from there.

P21 It’s emotional torture/I know that/I equate it to being a homeless person living out of a cardboard box in the middle of a winter. It’s minus 30. I have no shoes. No coat. This is how I felt at bottom-naked.

P24 Oh I was terrified. I did not know where I was or what was there. There was no help. Oh my God, my mind went insane. I got so confused.

P16 You do not think. You do not realize. You do not think you’re broken in it.

3.2.7. The Consequences. The final structural component of the bottom stories was “the consequences” of living through this experience for their children and themselves as both a person and a mother. As mothers, they shared a primal sense of parental responsibility to protect their children regardless of the circumstances. Many women shared recollections of their struggle to fulfill this obligation in the midst of bottom. Although they expended effort to “pretend that things were OK,” they spoke of “tremendous guilt” associated with the “kids seeing suffering,” and “illness invading their (children’s) lives.” Many women spoke of being ashamed that they experienced bottom and were displaced from their responsibilities as mother. Subsequent to the bottom experience, many women expressed a perceived sense of heightened familial and community surveillance and scrutiny of their mothering competency. For some women, loss of temporary or permanent custody was an imposed outcome of the bottom.

P1 I lost my joint custody.

P2 Ah (my child) feels abandoned/feels rejected/unloved and all it does to me is just/it adds more/I feel guilty as it is/it just adds to the guilt.

P5 You are second judging yourself and then you have others second judging you.

P6 I do not want the kids to see me suffer and worry about what effect it will have. I do not want them to see that or to know that. Why should, why should it invade their lives? (silence) I do not think it should. The kids they need just to be kids and then there’s a need to grow up.

P27 (My child) is my main priority and God knows how long it is going to take me to get well in the hospital. There is no way I want (my child) in the hands of (child services). Because once it goes that far/there is more chance/more likely that you’ll never get them back.

3.2.8. Meaning. All of the women spoke of the importance and value of their role as mothers. In the context of bottom, the mother-child relationship became increasingly tenuous. The heightened intensity of the illness circumstances jeopardized their self-care practices and frequently compromised their ability to mother. Insight regarding the complexity of their descent to bottom, and their experiences while at bottom began to emerge only after escape from bottom. Mothers acknowledged that despite their best efforts to protectively conceal their illness circumstances, this was no longer possible. This illness-imposed impediment to their
mothering role was associated with an intense private sense of guilt and contributed to worsening symptoms.

P11 You have so much pain you do not know where it goes so you turn it inward on yourself.

At bottom, these women knew that others questioned their fitness to mother. They feared losing their parental right to custody as a result of their illness-imposed bottom-experience. Their bottom stories illustrated their sense of powerlessness as both a person and a mother.

P2 (My child) could do a lot better. (My child) deserves a lot better.

P28 It’s the people you know who try to get help that get their kids taken away from them.

A prolonged period or intense experience at bottom threatened women’s self-determination in treatment, parenting involvement, and potentially survival. A timely and dignified escape from bottom was desired. For women survival through and from bottom was often influenced by perceived implications on their children.

P6 It’s too hard/too hard. I feel sometimes there is no hope/You know I still feel that I will be around. That’s a scary thought/very very scary thought.

P17 If my (child) is not ok then I’m not.

P18 I love my children too much/they were the only reason I wanted to live.

Being caught in the “horror” of bottom, often void of resources, however, they were unable to actualize a “life without suffering” for themselves with their children. A reality sought but not achievable in the context of bottom.

3.2.9. Function. The bottom stories demonstrate the impossibility of balancing responsibilities for self and others in the context of one’s worst illness reality. Although bottom compromised the women’s preferred mothering for a circumscribed period of time, it also sustained self and public doubt regarding their credible identity and capacity as mothers beyond bottom. These stories suggest that there is a need to create a safe context in which women may seek and receive early supportive care with a heightened likelihood of retaining their status as mother. Identifying and addressing the range of health and social circumstances that contributed to their vulnerabilities may lessen women’s fragility and despair. Such issues, unattended at both a personal and social level, can accelerate movement to bottom, suspend women in bottom, and impede their safe exist from bottom.

3.3. Case Example. The case example outlined in Table 1 demonstrates inclusiveness of the above content, structure, meaning, and functional components of the bottom stories. P22 is a divorced mother living with her 12-year-old child. P22 had been diagnosed with schizophrenia 11 years prior to the initial data collection interview. The following narrative was told by P22 in response to the interviewer’s request to elaborate on her expressed “not wanting or wishing to wake up in the morning” in relation to her discussion about bottom. Her expanded response acknowledged her bottom experience as distinct from suicidal ideation through her intent to not only escape her immediate circumstances, but re-establish agency.
4. Discussion

The bottom stories shared by the women in this study emphasized mothering in the context of overwhelming circumstances. When the “uncontrollable” force of illness disrupted their relationship with their children, their perceived experience of psychological pain was maximized. Within this context, hitting bottom was inevitable especially in the absence of comprehensive and appropriate mitigating resources. As such, there is an obligation for others, including health care professionals, to develop an awareness of each woman’s unique indicators that may precipitate a descent to bottom. An informed understanding of such cues may provide an opportunity to develop an individualized woman-centered purposeful plan to prevent the intensity of the “fall” to bottom or ultimately, a complete avoidance of a bottom experience. Recent evidence [16, 29, 30, 41] reinforces that strengths-based, interagency support has the potential to make a positive difference for maternal mental health. By extension, this emphasizes the need for psychiatric mental health nurses to enhance their competency in delivering health-promoting family-focused interventions [11] as early as the prenatal period [55].

The experience for these women became increasingly complicated as their illness circumstances reciprocally exacerbated the challenges inherent in the mothering role and at bottom. Health professionals who focus exclusively on acute symptoms without consideration of the mothering role may not appreciate the depth of the pain nor the potential strengths and motivations that will assist the woman to exit bottom safely. Hitting bottom may be a key point for intervention since these women described exhaustion and lack of trust regarding their internal resources or interpersonal connections to improve their situation. Unable to advocate for themselves at bottom may be liken to Broucek’s [56] self-objectification in shame.

...is as if the ground under one’s feet were giving way; depth and spatial relationships may have altered and one’s place in space uncertain, resulting in a kind of vertigo. Such experiences dramatically illustrate to what extent the reality of our everyday phenomenal world is dependent on an intact sense of self and an intact set of interpersonal coordinates and not simply on an intact brain (page 40).

In the event that a bottom experience is not mitigated, there is an urgent need to minimize the trauma of a woman’s experience. It is essential to provide guidance toward a safe exit that is not only desirable but achievable. A woman’s contemporaneous planning for exit from bottom is difficult given that agency may be affected by the presence of acute symptoms. Early acknowledgement and ongoing validation of a woman’s needs as mother is important to the creation of a genuine supported passage out of bottom. Innovative interventions to preserve agency and sense of meaningful connections to family involves affirming and strengthening women’s competencies while cognizant of the oppressive presence of socially constructed unrealistic ideals of motherhood. [10, 19, 35] Such approaches may lessen the time and acuity of distress associated with bottom.

Once exit has been achieved, it is essential that women have an opportunity to talk through their experience in the context of a safe receptive relationship. The processing of the bottom experiences may allow for an increased understanding concerning the complexity of their descent to bottom and decrease their sense of guilt. Further, a purposeful debriefing may be conducive to healing thereby enhancing capacity for coping with the precipitating events and emerging consequences of bottom. The integration of a realistic insight about bottom is fundamentally linked to establishing a future characterized by well-being. Noh [57] asserted that responsibility in healing involves the acknowledgment of a continuum of women’s competencies shaped by social, cultural, and familial supports. In the void of bottom, the women in this study emphasize the need to recognize their competency with appropriate resources.

A commonly identified limitation associated with secondary analyses relates to the assertion that qualitative data is co-constructed during the original study and thereby, limiting reuse for other purposes [45, 46]. The concerns about data reuse may be countered in this study given women’s spontaneous telling of bottom stories in the original investigations. This emergent complex phenomenon warranted focused exploration by the primary researchers to more fully identify the salient features of bottom for mothers with SMI [45]. This secondary analysis generated a refined understanding of bottom that remains coherent with the preliminary conception in the original studies [31].

5. Conclusions

Thirty-two women with SMI, involved in three qualitative studies, told a total of 173 stories about experiences related to hitting bottom. These stories suggest that there is a need to create a safe context for women to receive early intervention to maximize their capacities and stability. Further, the results suggest that mental health providers must identify the diverse issues contributing to mothers’ vulnerabilities in an effort to lessen their fragility and pain. Healing necessitates multiple approaches to address the private and public trauma of bottom experiences while nurturing their development. Preventative supportive family approaches are recommended given the burdensome legacy of bottom.

References

[1] C. T. Mowbray, D. Oyserman, D. Bybee, P. Macfarlane, and A. Rueda-Riedle, “Life circumstances of mothers with serious mental illnesses,” Psychiatric Rehabilitation Journal, vol. 25, no. 2, pp. 114–123, 2001.
[2] T. M. Laursen and T. Munk-Olsen, “Reproductive patterns in psychotic patients,” Schizophrenia Research, vol. 121, no. 1–3, pp. 234–240, 2010.
[3] J. J. McGrath, J. Hearle, L. Jenner, K. Plant, A. Drummond, and J. M. Barkla, “The fertility and fecundity of patients with psychoses,” Acta Psychiatrica Scandinavica, vol. 99, no. 6, pp. 441–446, 1999.
[4] J. Earle and J. McGrath, “Motherhood and schizophrenia,” in Women and Schizophrenia, D. J. Castle, J. McGrath, and J. J. J. Kulkami, Eds., pp. 79–94, Cambridge University Press, London, UK, 2000.

[5] P. Reeder, M. McClure, and A. Jolley, Family Matters: Interfaces between Child and Adult Mental Health, Routledge, London, UK, 2000.

[6] T. Craig and E. J. Bromet, “Parents with psychosis,” Annals of Clinical Psychiatry, vol. 16, no. 1, pp. 35–39, 2004.

[7] C. Handley, G. A. Farrell, A. Josephs, A. Hanke, and M. Hazeldon, “The Tasmanian children's project: the needs of children with a parent/carer with a mental illness,” The Australian and New Zealand Journal of Mental Health Nursing, vol. 10, no. 4, pp. 221–228, 2001.

[8] M. Östman and L. Hansson, “Children in families with a severely mentally ill member. Prevalence and needs for support,” Social Psychiatry and Psychiatric Epidemiology, vol. 37, no. 5, pp. 243–248, 2002.

[9] D. G. Bassani, C. V. Padoin, D. Philipp, and S. Veldhuizen, “Estimating the number of children exposed to parental psychiatric disorders through a national health survey,” Child and Adolescent Psychiatry and Mental Health, vol. 3, article 6, 2009.

[10] A. Blanch, K. J. Nicholson, and J. Purcell, “Parents with severe mental illness and their children: the need for human services integration,” in Women's Mental Health Services: A Public Health Perspective, B. Lubotsky Levin, A. K. Blanch, and A. Jennings, Eds., pp. 201–214, Sage, Thousand Oaks, Calif, USA, 1998.

[11] D. Maybery and A. Reupert, “Parental mental illness: a review of barriers and issues for working with families and children,” Journal of Psychiatric and Mental Health Nursing, vol. 16, no. 9, pp. 784–791, 2009.

[12] A. Monds-Watson, R. Manktelow, and M. McColgan, “Social work with children when parents have mental health difficulties: acknowledging vulnerability and maintaining the "Rights of the child,"” Child Care in Practice, vol. 16, no. 1, pp. 35–55, 2010.

[13] J. Nicholson and K. Biebel, “Commentary on “Community mental health care for women with severe mental illness who are parents”—the tragedy of missed opportunities: what providers can do,” Community Mental Health Journal, vol. 38, no. 2, pp. 167–172, 2002.

[14] L. M. Howard and K. Hunt, “The needs of mothers with severe mental illness: a comparison of assessments of needs by staff and patients,” Archives of Women's Mental Health, vol. 11, no. 2, pp. 131–136, 2008.

[15] M. L. Zilberman, H. Tavares, S. B. Blume, and N. El-Guebaly, “Towards best practices in the treatment of women with addictive disorders,” Addictive Disorders and Their Treatment, vol. 1, no. 2, pp. 39–46, 2002.

[16] A. Reupert and D. Maybery, “Families affected by parental mental illness: a multiperspective account of issues and interventions,” American Journal of Orthopsychiatry, vol. 77, no. 3, pp. 362–369, 2007.

[17] K. Kaplan, P. Kottsieper, J. Scott, M. Salzer, and P. Solomon, “Adoption and safe families act state statutes regarding parents with mental illnesses: a review and targeted intervention,” Psychiatric Rehabilitation Journal, vol. 33, no. 2, pp. 91–94, 2009.

[18] D. Oyserman, C. T. Mowbray, P. A. Meares, and K. B. Firminger, “Parenting among mothers with a serious mental illness,” American Journal of Orthopsychiatry, vol. 70, no. 3, pp. 296–315, 2000.

[19] C. Alakus, R. Convell, M. Gilbert, A. Buist, and D. Castle, “The needs of parents with a mental illness who have young children: an Australian perspective on service delivery options,” International Journal of Social Psychiatry, vol. 53, no. 4, pp. 333–339, 2007.

[20] P. Montgomery, S. Brown, and C. Forchuk, “A comparison of individual and social vulnerabilities, health, and quality of life among Canadian women with mental diagnoses and young children,” Women's Health Issues, vol. 21, no. 1, pp. 48–56, 2011.

[21] M. D. McPherson, J. Delva, and J. A. Cranford, “A longitudinal investigation of intimate partner violence among mothers with mental illness,” Psychiatric Services, vol. 58, no. 5, pp. 675–680, 2007.

[22] J. M. Park, P. Solomon, and D. S. Mandell, “Involvement in the child welfare system among mothers with serious mental illness,” Psychiatric Services, vol. 57, no. 4, pp. 493–497, 2006.

[23] L. D. Hollingsworth, “Child custody loss among women with persistent severe mental illness,” Social Work Research, vol. 28, no. 4, pp. 199–209, 2004.

[24] J. Nicholoson, N. Finkelstein, V. Williams, J. Thom, C. Noether, and M. DeVilbiss, “A comparison of mothers with co-occurring disorders and histories of violence living with or separated from minor children,” Journal of Behavioral Health Services and Research, vol. 33, no. 2, pp. 223–243, 2006.

[25] D. Bybee, C. T. Mowbray, D. Oyserman, and L. Lewandowski, “Variability in community functioning of mothers with serious mental illness,” Journal of Behavioral Health Services and Research, vol. 30, no. 3, pp. 269–289, 2003.

[26] E. A. Craig, “Parenting programs for women with mental illness who have young children: a review,” Australian and New Zealand Journal of Psychiatry, vol. 38, no. 11–12, pp. 923–928, 2004.

[27] P. Montgomery, “Mothers with a serious mental illness: a critical review of the literature,” Archives of Psychiatric Nursing, vol. 19, no. 5, pp. 226–235, 2005.

[28] L. Wilson and M. Crowe, “Parenting with a diagnosis bipolar disorder,” Journal of Advanced Nursing, vol. 65, no. 4, pp. 877–884, 2009.

[29] M. J. Venkataraman and B. Ackerson, “Parenting among mothers with bipolar disorder: strengths, challenges, and service needs,” Journal of Family Social Work, vol. 11, no. 4, pp. 389–408, 2008.

[30] B. Davies and D. Allen, “Integrating “mental illness” and “motherhood”: the positive use of surveillance by health professionals. A qualitative study,” International Journal of Nursing Studies, vol. 44, no. 3, pp. 365–376, 2007.

[31] P. Montgomery, C. Tompkins, C. Forchuk, and S. French, “Keeping close: mothering with serious mental illness,” Journal of Advanced Nursing, vol. 54, no. 1, pp. 20–28, 2006.

[32] A. Diaz-Caneja and S. Johnson, “The views and experiences of severely mentally ill mothers—a qualitative study,” Social Psychiatry and Psychiatric Epidemiology, vol. 39, no. 6, pp. 472–482, 2004.

[33] R. G. Sands, N. Koppelman, and P. Solomon, “Maternal custody status and living arrangements of children of women with severe mental illness,” Health and Social Work, vol. 29, no. 4, pp. 317–325, 2004.

[34] B. J. Ackerson, “Coping with the dual demands of severe mental illness and parenting: the parents’ perspective,” Families in Society, vol. 84, no. 1, pp. 109–118, 2003.

[35] I. Savvidou, V. P. Bozikas, S. Hatziigeleki, and A. Karavatos, “Narratives about their children by mothers hospitalized on...
a psychiatric unit," *Family Process*, vol. 42, no. 3, pp. 391–402, 2003.

[36] B. Hinden, T. Wolf, K. Biebel, and J. Nicholson, “Supporting clubhouse members in their role as parents: necessary conditions for policy and practice initiatives,” *Psychiatric Rehabilitation Journal*, vol. 33, no. 2, pp. 98–105, 2009.

[37] R. Ueno and K. Kamibeppu, “Narratives by Japanese Mothers with chronic mental illness in the Tokyo metropolitan area: their feelings toward their children and perceptions of their children’s feelings,” *Journal of Nervous and Mental Disease*, vol. 196, no. 7, pp. 522–530, 2008.

[38] H. Khalifeh, C. Murgatroyd, M. Freeman, S. Johnson, and H. Killaspy, “Home treatment as an alternative to hospital admission for mothers in a mental health crisis: a qualitative study,” *Psychiatric Services*, vol. 60, no. 5, pp. 634–639, 2009.

[39] M. W. Wan, S. Moulton, and K. M. Abel, “The service needs of mothers with schizophrenia: a qualitative study of perinatal psychiatric and antenatal workers,” *General Hospital Psychiatry*, vol. 30, no. 2, pp. 177–184, 2008.

[40] J. Nicholson, B. R. Hinden, K. Biebel, A. D. Henry, and J. Katz-Leavy, “A qualitative study of programs for parents with serious mental illness and their children: building practice-based evidence,” *Journal of Behavioral Health Services and Research*, vol. 34, no. 4, pp. 395–413, 2007.

[41] B. R. Hinden, K. Biebel, J. Nicholson, and L. Mehnert, “The invisible children’s project: key ingredients of an intervention for parents with mental illness,” *Journal of Behavioral Health Services and Research*, vol. 32, no. 4, pp. 393–408, 2005.

[42] M. K. Brewer, *Leaving the abyss: a phenomenological study of women’s recovery from alcoholism*, Doctoral thesis, 2000.

[43] B. Curtin, R. Mapes, D. Petillo, and E. Oberley, “Long-term dialysis survivors: a transformational experience,” *Qualitative Health Research*, vol. 12, no. 5, pp. 609–624, 2002.

[44] J. Heaton, *Reworking Qualitative Data*, SAGE, London, UK, 2004.

[45] J. Heaton, “Secondary analysis of qualitative data,” in *The SAGE Handbook of Social Research Methods*, P. Alasuutari, L. Bickman, and J. Brannen, Eds., pp. 506–519, SAGE, Los Angeles, Calif, USA, 2008.

[46] S. E. Thorne, “Secondary analysis in qualitative research: issues and implications,” in *Critical Issues in Qualitative Research Methods*, J. M. Morse, Ed., pp. 263–279, Sage, London, UK, 1994.

[47] C. Kohler Riesmann, “A thrice-told tale: new readings of an old story,” in *Narrative Research in Health and Illness*, B. Hurwitz, T. Greenhalgh, and V. Skaltans, Eds., pp. 309–324, Blackwell Publishing, Oxford, UK, 2004.

[48] Ontario Ministry of Health and Long-Term Care, *Making it Happen. Operational Framework for the Delivery of Mental Health Services and Supports*, Queen’s Print for Ontario, Ottawa, Canada, 1999.

[49] C. Kohler Riesmann, *Narrative Methods for the Human Sciences*, Sage, Thousand Oaks, Calif, USA, 2008.

[50] P. H. Bailey and S. Tilley, “Story telling and the interpretation of meaning in qualitative research,” *Journal of Advanced Nursing*, vol. 38, no. 6, pp. 574–583, 2002.

[51] W. Labov and J. Waletzky, “Narrative analysis: oral versions of personal experience,” in *Essays on the Verbal and Visual Arts*, J. Helms, Ed., pp. 12–44, University of Washington Press, Seattle, Wash, USA, 1972.

[52] M. Agar and J. R. Hobbs, “Interpreting discourse: coherence and the analysis of ethnographic interviews,” *Discourse Processes*, vol. 5, pp. 1–32, 1982.

[53] C. Antaki, *Exploring and Arguing: The Social Organization of Accounts*, Sage, London, UK, 1994.

[54] S. E. Thorne, *Interpretive Description*, Left Coast Press, Walnut Creek, Calif, USA, 2008.

[55] Y. Hauck, D. Rock, T. Jackiewicz, and A. Jablensky, “Healthy babies for mothers with serious mental illness: a case management framework for mental health clinicians,” *International Journal of Mental Health Nursing*, vol. 17, no. 6, pp. 383–391, 2008.

[56] F. V. Broucek, * Shame and the Self*, Guilford Press, New York, NY, USA, 1991.

[57] E. Noh, “Asian American women and suicide: problems of responsibility and healing,” *Women and Therapy*, vol. 30, no. 3–4, pp. 87–107, 2007.
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