The impact of preterm birth <37 weeks on parents and families: a cross-sectional study in the 2 years after discharge from the neonatal intensive care unit

The Harvard community has made this article openly available. Please share how this access benefits you. Your story matters

Citation
Lakshmanan, Ashwini, Meghana Agni, Tracy Lieu, Eric Fleegler, Michele Kipke, Philippe S. Friedlich, Marie C. McCormick, and Mandy B. Belfort. 2017. “The impact of preterm birth <37 weeks on parents and families: a cross-sectional study in the 2 years after discharge from the neonatal intensive care unit.” Health and Quality of Life Outcomes 15 (1): 38. doi:10.1186/s12955-017-0602-3. http://dx.doi.org/10.1186/s12955-017-0602-3.

Published Version
doi:10.1186/s12955-017-0602-3

Citable link
http://nrs.harvard.edu/urn-3:HUL.InstRepos:31731888

Terms of Use
This article was downloaded from Harvard University’s DASH repository, and is made available under the terms and conditions applicable to Other Posted Material, as set forth at http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#LAA
The impact of preterm birth <37 weeks on parents and families: a cross-sectional study in the 2 years after discharge from the neonatal intensive care unit

Ashwini Lakshmanan1,2,3,4*, Meghana Agni5, Tracy Lieu6, Eric Fleegler7, Michele Kipke8, Philippe S. Friedlich1, Marie C. McCormick9,10,11 and Mandy B. Belfort12

Abstract

Background: Little is known about the quality of life of parents and families of preterm infants after discharge from the neonatal intensive care unit (NICU). Our aims were (1) to describe the impact of preterm birth on parents and families and (2) to identify potentially modifyable determinants of parent and family impact.

Methods: We surveyed 196 parents of preterm infants <24 months corrected age in 3 specialty clinics (82% response rate). Primary outcomes were: (1) the Impact on Family Scale total score; and (2) the Infant Toddler Quality of Life parent emotion and (3) time limitations scores. Potentially modifiable factors were use of community-based services, financial burdens, and health-related social problems. We estimated associations of potentially modifiable factors with outcomes, adjusting for socio-demographic and infant characteristics using linear regression.

Results: Median (inter-quartile range) infant gestational age was 28 (26–31) weeks. Higher Impact on Family scores (indicating worse effects on family functioning) were associated with taking ≥3 unpaid hours/week off from work, increased debt, financial worry, unsafe home environment and social isolation. Lower parent emotion scores (indicating greater impact on the parent) were also associated with social isolation and unpaid time off from work. Lower parent time limitations scores were associated with social isolation, unpaid time off from work, financial worry, and an unsafe home environment. In contrast, higher parent time limitations scores (indicating less impact) were associated with enrollment in early intervention and Medicaid.

Conclusions: Interventions to reduce social isolation, lessen financial burden, improve home safety, and increase enrollment in early intervention and Medicaid all have the potential to lessen the impact of preterm birth on parents and families.

Keywords: Impact on family, Impact on parents, Prematurity, High-risk infant, Post-discharge
Background
In the United States, nearly 500,000 infants, or 11.7% of all live births, are born preterm (<37 weeks’ gestation) each year [1, 2]. Preterm birth and the sometimes associated prolonged newborn hospitalization are great family stressors, and can lead to subsequent family dysfunction [3–5].

All preterm infants are at risk for re-hospitalization, as well as medical and neurodevelopmental complications, even moderate to late preterm infants (born at 32 to <37 weeks’ gestation) [6]. A particularly challenged sub-group is very low birth weight (VLBW) infants or those born < 1500 g. More than 90% of VLBW infants are discharged home from the neonatal intensive care unit (NICU). The burden of continued health and developmental problems faced by these infants is substantial [7–9]. For example, compared with normal birth weight children, VLBW children face a 2–3 fold greater risk for visual and hearing impairment, speech delays and attention disorders [10, 11]; may have poor feeding and growth, respiratory complications, and face neurocognitive difficulties [12–16].

Given these ongoing problems and risks, families of preterm children often must manage numerous medical and developmental needs above and beyond what is required for a healthy full term infant, for months or even years after the neonatal discharge. For example, during the first year of life, VLBW infants are prone to re-hospitalization and require increased outpatient care [17–19]. Parents must transport their child for medical appointments and therapies, communicate with the child’s pediatrician and other healthcare providers, and are often responsible for daily tasks, such as administering medications and monitoring chronic conditions.

The intensity of care and high level of vigilance required by families to meet the needs of their preterm child makes it likely that having a preterm child adversely affects the quality of life of the parents and the family overall. The 2006 Institute of Medicine’s (IOM) report on Preterm Birth: Causes, Consequences and Prevention stressed the importance of assessing aspects of family and parent quality of life and stress beyond maternal psychological well-being [20–22]. A better understanding of the impact of preterm birth on parent and family quality of life, as well as modifiable factors that predispose parents and families to greater or lesser impact would inform community-based and other structured assistance programs designed to lessen the impact.

Our main research question was, “Are modifiable characteristics (such as the use of community based and public assistance programs, financial burden, and health related social problems) associated with the impact of preterm birth on parents and families after NICU discharge?” The Anderson and Aday health utilization model [23, 24] provides a useful framework for addressing our research question because it uniquely captures the constructs of access, need, and quality of life. As presented in Fig. 1, we conceptualized potentially modifiable characteristics that influence parent and family impact as: (1) use of community based developmental services and public assistance programs; (2) financial burden; (3) health related social problems. We also specified predisposing characteristics (including socio-demographics and infant health characteristics) related both to modifiable characteristics and to outcomes that could act as confounders.

Our specific aims were (1) to describe the impact of preterm birth on parents and families and (2) to identify potentially modifiable determinants of parent and family impact. Specific variables of interest based on previous literature, were use of community-based resources, financial burden and health-related social problems [25].

Methods
Study design and participants
This was a cross-sectional study. We enrolled one parent (mother or father) of preterm (<37 weeks gestation) infants attending 3 outpatient clinics at a large tertiary children’s hospital. One clinic provides multidisciplinary medical and neurodevelopmental follow-up for infants with gestational age <32 completed weeks or birth weight <1500 g discharged from one of 3 large, academic NICU’s and affiliated community-based Level II nurseries, and for more mature or heavier preterm infants with severe medical conditions and/or social risk factors (101 participants enrolled). The second clinic provides pediatric pulmonary care for infants with lung disease that originates in the newborn period, predominantly bronchopulmonary dysplasia (57 participants enrolled). The third clinic provides follow-up care for infants who have suffered neurologic injury during the fetal or newborn period (38 participants enrolled). While some patients were seen at more than one clinic, they were only enrolled once in the study.

We included parents of preterm infants who were up to 24 months corrected age (age from term equivalent). Parents must have been able to answer questions in English or Spanish. If the infant was a multiple, only one response was collected from the family. Study staff provided eligible families with a letter describing the study. Consent was obtained when the parent agreed to complete the questionnaire, which was administered on a laptop (with privacy screens) in the clinic waiting room or examination room. Participants were provided a small incentive to complete the survey.

The Boston Children’s Hospital and Children’s Hospital Los Angeles human subjects committees approved the study protocol. Approximately 75% of preterm infants are
referred to high-risk infant follow up programs. [3] In this study, of the 239 eligible participants from October, 2011 to June, 2012, 196 completed the questionnaire (82% response rate). The questionnaire is available as Additional file 1.

Measurements
Measurements of primary outcomes, modifiable characteristics and potential confounders (predisposing characteristics) are summarized in Additional file 2: Table S1.

Impact on family
The Impact on Family (IOF) [26–28] measures the global impact of pediatric disability on the family and has been validated on samples of children with chronic health conditions, including preterm birth [25]. The IOF total score is derived from a 27-item questionnaire. For each item, parents indicate the extent to which they agree with a statement regarding the negative impact of the child on the family. Anchors for a 4-point Likert scale were: strongly agree; agree; disagree; and strongly disagree. Examples of IOF items are: “The illness is causing financial problems for the family” and “Our family gives up things because of illness.” IOF subscales include financial impact (8 points), disruption of planning (20 points), caretaker burden (12 points), and familial burden (16 points) for total possible score of 56 points. The total negative impact score served as our summary measure of family burden (higher scores indicate greater family burden).

In a previous study, internal consistency was high for the overall IOF Scale (Cronbach alphas for total impact, 0.83 to 0.89), but lower for financial (0.68 to 0.79) and coping (0.46 to .52) items [26]. High total scores on the IOF are associated with maternal psychiatric symptoms, poor child health, poor child adjustment, increased child hospitalizations, lower maternal education, and maternal receipt of public assistance [27–29], providing evidence for construct validity.

Impact on parent
The Infant Toddler Quality of Life Questionnaire™ (ITQOL) was developed in 1994 for use in children from 2 months to 5 years of age as a “profile measure” for health status and health-related quality of life. ITQOL adopts as its conceptual framework the World Health Organization’s definition of health as a state of complete physical, mental and social well-being and not just the absence of disease [30]. It has been used both in randomized clinical trials [31] and observational studies, and is accepted favorably for its ease of use and understandability [32].

In this study, we used the Family Burden scales of the ITQOL, which cover two parent-focused concept subscales, impact-emotion and impact-time, due to caring for their infant or toddler [30, 32, 33]. The parent impact-emotion domain consists of seven items in which the parent is asked to rate how much anxiety or worry each of the child characteristics described in the items has caused during the past 4 weeks (i.e., feeding/sleeping/eating habits; physical health, emotional well-being, learning abilities, ability to interact with others; behavior and temperament). The parent impact-time domain consists of seven items in which the parents is asked to rate how much of his/her time was limited for personal needs because of the problems with the child’s personal needs during the past 4 weeks. Internal consistency for the ITQOL parent-impact emotion and parent-impact time scales has been reported in three different populations, a Dutch general population sample (0.61, 0.64) [34], a functional abdominal pain sample (0.72, 0.73) [35] and a burn injury sample (0.79, 0.84) [30, 36].

Raw subscale scores are converted to standardized scores on a 0–100 continuum [37–40]. For each scale, higher scores indicate less emotional impact and fewer
time limitations on the parent (in other words, higher scores represent more favorable outcomes).

**Potentially modifiable characteristics**

**Use of community-based resources** Participants were asked yes/no questions about the use of community-based developmental resources (such as early intervention programs), use of social services such as food assistance programs, Supplemental Nutrition Assistance Program and the Women, Infant, Children’s program as well as energy assistance/disability programs such the Low Income Home Energy Assistance Program, Transitional Aid to Families with Dependent Children, and receipt of Supplemental Security Income (SSI).

**Financial Burden** In addition to questions about employment for the participating parent and his/her partner, we asked 6 yes/no questions from the 2007 Commonwealth Fund Biennial Health Insurance Survey [41–44] regarding unexpected costs, increased bills, increased out-of-pocket expenses and financial worry.

**Health-related social problems** HelpSteps.com is a survey designed to identify health-related social problems. Development of HelpSteps involved literature review and key informant interviews with health and social services experts, yielding an initial list of 25 social domains. Of those, the 5 most relevant domains were identified using a modified Delphi technique: (1) access to health care, (2) housing, (3) food security (4) income security and (5) intimate partner violence [45–48]. Most questions about these domains were adapted from previous surveys (e.g. National Health Interview Survey [49], the American Housing Survey [50], the Philadelphia Survey of Work and Family [51] and the Childhood Community Hunger Identification Project [52]), while a few newly written items were also incorporated into the final HelpSteps survey. In terms of content validity, the domains covered in HelpSteps are well-recognized as being closely tied to health outcomes and costs [53]. HelpSteps is highly effective in identifying problems that can be addressed by referral to appropriate social services [46, 47] and a qualitative study revealed that over 2/3 of participants found the HelpSteps questions to be highly relevant to their own problems [48].

**Predisposing characteristics (potential confounders)**

**Infant health and development** We obtained information from the medical record regarding delivery and complications during the neonatal hospitalization. We asked parents questions about their infant’s health status since discharge including the number of emergency department visits, monthly clinic appointments, and hospitalizations, immunizations, dependence on technology, and administration of prescription medications.

To assess infant development, we used the Motor and Social Development (MSD) scale [54], which was developed by the National Center for Health Statistics to measure motor, social and cognitive development of young children. Of 48 items derived from standard measures of child development, including the Bayley Scales, Gesell Scale, and Denver Developmental Screening Test, parents complete 15 age-specific items, which ask about specific developmental milestones such as laughing out loud, pulling to stand, and saying recognizable words [55]. We selected the MSD because it is brief and allows for scoring based on a large, national sample [56] with a normative mean of 100 and standard deviation 15, similar to other developmental tests. Higher scores indicate better development. In a previous study of former preterm infants, we showed that the MSD has good internal consistency (Cronbach alpha 0.65-0.88) and is modestly correlated with Bayley alpha 0.45-0.88 and is moderately correlated with Bayley Scales of Infant and Toddler Development, 3rd edition, a gold standard professionally administered neurodevelopmental test [56]. Another study reported that infants with lower gestational age at birth have lower scores on the MSD [54], although the MSD includes a cognitive, motor, and social subscales, the degree to which it is sensitive to language/communication delays is unknown, which is a potential limitation.

**Statistical analysis** Our main outcomes were: (1) impact on family total score; (2) impact on parent score determined by the concept of emotion; and (3) impact on parent score determined by the concept of limitation of time. Potentially modifiable determinants included the use of community-based resources, financial burden, and health-related social problems. Potential confounders (predisposing characteristics) were socio-demographics and infant pre-disposing and post-discharge characteristics.

In bivariate analyses, we compared outcome scores across categories of predisposing characteristics (potential confounders) and potentially modifiable determinants. We calculated p-values using non-parametric tests (Wilcoxon Rank Sum or Kruskal Wallis). To identify potentially modifiable determinants independent of confounders on our primary outcomes, we created parsimonious multivariable models, adjusting for variables of a priori interest and for other characteristics found to be significant at p < 0.1. We also examined each model using variance inflation factors (VIF) and did not detect significant collinearity (VIF ≤ 2 for all models).

We used SAS version 9.4 (SAS Institute Inc., Cary, NC) for analyses.
### Table 1: Pre-disposing characteristics and unadjusted associations with impact on family and impact on parent scores (n = 196)\(^a\)

| Characteristic | Time from discharge (months) | Total cohort | Impact on family | Impact on parent |
|----------------|-----------------------------|--------------|------------------|------------------|
|                |                             | % of sample  | Median (IQR)     | Median (IQR)     | Median (IQR) | p-value | Median (IQR) | p-value | Median (IQR) | p-value |
| Time from discharge (months) | < 6 | 31 | 27 (21–35.5) | <0.04 | 78 (57–93) | <0.04 | 76 (67–95) | <0.04 |
|                            | ≥ 6 to < 12 | 30 | 20.5 (17–30) | 86 (64–96) | 90 (81–100) | 0.12 |
|                            | ≥ 12 to < 18 | 24 | 22 (18–31) | 86 (61–93) | 90 (71–100) | 0.36 |
|                            | ≥ 18 | 15 | 23 (16–31) | 82 (64–96) | 90 (67–100) | 0.89 |
| Socio-demographics | Person completing survey | | | | | | | |
| Person completing survey | Mother | 77 | 23 (17–30) | 0.07 | 85 (68–96) | <0.01 | 90 (74–100) | <0.01 |
|                            | Father | 23 | 27.5 (20–36) | 69.5 (53–86) | 86 (52–90) | 0.97 |
| Race/ethnicity | White non-Hispanic | 67 | 24.5 (18–33) | 0.2 | 82 (61–93) | 0.6 | 90 (71–100) | 0.1 |
| Hispanic | 7 | 20 (17–23.5) | 89 (85.5–93) | 100 (88–100) | 0.04 |
| Black non-Hispanic | 11 | 26 (18–41) | 81 (61–100) | 78 (57–100) | 0.01 |
| Other | 15 | 22 (17–36) | 88 (71–98) | 87 (70–94) | 0.02 |
| Income ($/Year) | Income ($/Year) | 24 | 24 (19–38) | 0.6 | 86 (69–93) | 0.4 | 90 (71–100) | 0.4 |
| Income ($/Year) | 15 | 23.5 (17.5–35) | 78 (66–96) | 90 (52–100) | 0.08 |
| Income ($/Year) | ≥ 80,000 | 52 | 23 (17–30) | 82 (61–93) | 90 (76–95) | 0.06 |
| Missing | 9 | | | | | | |
| Highest level of education (either parent) | High school | 32 | 26 (19–39) | 0.2 | 83.5 (61–93) | 0.9 | 90 (62–100) | 0.4 |
| Highest level of education (either parent) | At least some college | 68 | 23 (18–30) | 0.05 | 82 (64–93) | 0.01 | 90 (76–100) | 0.01 |
| Language | Non-English | 6 | 26.5 (20–34) | 0.5 | 85.5 (57–89) | 0.9 | 85 (57–100) | 0.8 |
| Language | English | 94 | 24 (18–32) | 0.05 | 82 (64–93) | 0.05 | 90 (71–100) | 0.05 |
| Infant characteristics | Birthweight (grams) | 53 | 25 (18–33) | 0.06 | 75 (61–82) | 0.2 | 100 (95–100) | 0.2 |
| Birthweight (grams) | ≥ 1000 to < 1500 | 24 | 24 (17–30) | 86 (69–96) | 84 (62–100) | 0.02 |
| Birthweight (grams) | ≥ 1500 to < 2500 | 17 | 18.5 (15–23.5) | 86 (76–93) | 90 (86–100) | 0.03 |
| Birthweight (grams) | ≥ 2500 | 6 | 27 (20–36) | 61 (18–90) | 77 (33–100) | 0.02 |
| Gestational age (weeks) | < 24 to < 28 | 41 | 26 (18–33) | 0.07 | 82 (66–93) | 0.2 | 88 (67–100) | 0.8 |
| Gestational age (weeks) | ≥ 28 to < 32 | 37 | 21 (17–28) | 85 (66–96) | 90 (71–100) | 0.02 |
| Gestational age (weeks) | ≥ 32 to < 34 | 9 | 26 (18–30) | 81 (49–91) | 81 (69–95) | 0.02 |
| Gestational age (weeks) | ≥ 34 to < 37 | 13 | 29 (20–41) | 71 (35–93) | 81 (57–100) | 0.02 |
| Multiple Birth | Yes | 15 | 25 (17–37) | 0.44 | 77 (34–91) | 0.7 | 81 (70–96) | 0.7 |
| Multiple Birth | No | 85 | 23 (18–33) | 0.89 | 81 (60–92) | 0.89 | 89 (70–99) | 0.89 |
Results

Participant characteristics and outcomes

Predisposing characteristics of study participants and outcome measures are shown in Table 1. A majority of participants were white, non-Hispanic (67%). 52% reported an annual household income of ≥ $80,000 and 68% of mothers had attended at least some college. The median (IQR) gestational age of infants at birth was 28 weeks (26–31). The median (interquartile range, IQR) chronologic age of infants at the time of study participation was 10.4 months (7.5-17.2).

Unadjusted associations of predisposing characteristics (potential confounders) with outcomes

As shown in Table 1, among pre-disposing characteristics, the use of medical technology, receipt of at least one prescription medication daily, one or more readmission or emergency department visit after neonatal discharge, and 2 or more clinic appointments per month were all associated with greater impact on family, parents, or both. Additionally, having an infant with a low developmental score (MSD < 85) was associated with greater impact on the family and parent emotion. Fathers who completed the survey had higher impact scores than mothers on the parent-focused domains of emotion and time limitations. Of note when we performed an additional sensitivity analysis by running our multivariate models for mothers only, we found our multivariate model estimates were similar in magnitude and direction as the full models that included fathers.

Potentially modifiable characteristics and unadjusted associations with outcomes

Table 2 shows that use of public housing and public assistance program were associated with greater impact on family. Compensation for time taken off from work was associated with a lower parent emotional score (less parental impact) while use of social services, public housing, enrollment in Medicaid and an unsafe home environment were associated with a higher IOF score (greater family impact). Markers of financial burden (including unpaid time off work, increased out-of-pocket
| Characteristics                                      | % of sample | Impact on family | Impact on parent | Time limitations score |
|------------------------------------------------------|-------------|------------------|------------------|-----------------------|
|                                                      |             | Total score      | Emotion score    | Time limitations score |
|                                                      |             | Median (IQR)     | p-value          | Median (IQR)          | p-value |
| Use of community-based services                      |             |                  |                  |                       |         |
| Use of a community-based developmental program (early intervention) |             |                  |                  |                       |         |
| Yes                                                  | 92          | 23 (18–31)       | 0.9              | 82 (61–93)            | 0.9     | 90 (71–100) | 0.2 |
| No                                                   | 8           | 28 (17–36)       |                  | 82 (64–93)            |         | 83 (33–95) |     |
| Use of social services                               |             |                  |                  |                       |         |           |     |
| Yes                                                  | 45          | 26 (19–38)       | 0.02             | 82 (61–93)            | 0.9     | 90 (67–100) | 0.8 |
| No                                                   | 55          | 23 (17–30)       |                  | 82 (64–93)            |         | 90 (71–100) |     |
| Use of public housing                                |             |                  |                  |                       |         |           |     |
| Yes                                                  | 9           | 29 (22–41)       | 0.04             | 85 (57–93)            | 0.7     | 74 (47–100) | 0.4 |
| No                                                   | 91          | 23 (18–31)       |                  | 82 (64–93)            |         | 90 (71–100) |     |
| Receive supplemental security income (SSI)           |             |                  |                  |                       |         |           |     |
| Yes                                                  | 23          | 23 (17–31)       | 0.8              | 86 (69–100)           | 0.1     | 95 (71–100) | 0.1 |
| No                                                   | 77          | 24 (18–32)       |                  | 82 (61–93)            |         | 90 (71–95) |     |
| Use a community based clinic for primary care         |             |                  |                  |                       |         |           |     |
| Yes                                                  | 12          | 23 (19–41)       | 0.5              | 73 (57–96)            | 0.7     | 81 (67–100) | 0.4 |
| No                                                   | 88          | 23 (18–31)       |                  | 83 (63–93)            |         | 90 (71–100) |     |
| Financial Burden since infant discharge               |             |                  |                  |                       |         |           |     |
| Participant employed                                 |             |                  |                  |                       |         |           |     |
| Yes                                                  | 68          | 23 (18–31)       | 0.7              | 82 (64–93)            | 0.9     | 90 (71–100) | 0.4 |
| No                                                   | 32          | 24 (18–35)       |                  | 85 (61–93)            |         | 90 (67–100) |     |
| Partner employed                                     |             |                  |                  |                       |         |           |     |
| Yes                                                  | 88          | 21 (17–30)       | 0.2              | 85 (64–96)            | 0.5     | 90 (71–100) | 0.5 |
| No                                                   | 12          | 27 (22–28)       |                  | 83.5 (64–89)          |         | 90 (67–95) |     |
| Any member has taken ≥ 3 h taken off from work without pay weekly |             |                  |                  |                       |         |           |     |
| Yes                                                  | 51          | 27 (18–40)       | <0.005           | 71 (53–89)            | <0.005  | 81 (52–95) | <0.005 |
| No                                                   | 49          | 21 (17–29)       |                  | 86 (71–96)            |         | 90 (80–100) |     |
| Receives employer-based compensation for time off    |             |                  |                  |                       |         |           |     |
| Yes                                                  | 31          | 22 (18–31)       | 0.6              | 86 (75–96)            | 0.02    | 90 (76–100) | 0.2 |
| No                                                   | 69          | 24 (17–33)       |                  | 79 (57–93)            |         | 90 (69–100) |     |
| No compensation (from any source) for time off       |             |                  |                  |                       |         |           |     |
| No compensation                                      | 17          | 29 (18–41)       | 0.1              | 65 (50–89)            | 0.01    | 85 (47–100) | 0.2 |
| Compensation                                         | 83          | 23 (17–30)       |                  | 85 (68–89)            |         | 90 (76–100) |     |
| Unexpected costs incurred                            |             |                  |                  |                       |         |           |     |
| Yes                                                  | 41          | 29 (19–39)       | <0.01            | 75 (57–93)            | <0.01   | 81 (59–95) | <0.01 |
| No                                                   | 59          | 21 (17–29)       |                  | 86 (69–96)            |         | 90 (76–100) |     |
| Increased bills                                      |             |                  |                  |                       |         |           |     |
| Yes                                                  | 19          | 30 (23–44)       | <0.02            | 71 (53–86)            | <0.02   | 76 (47–100) | <0.02 |
| No                                                   | 81          | 22 (17–30)       |                  | 86 (68–95)            |         | 90 (76–100) |     |
| Increased out-of-pocket expenses                      |             |                  |                  |                       |         |           |     |
| Yes                                                  | 13          | 33 (23–44)       | <0.04            | 75 (61–93)            | <0.04   | 76 (43–100) | <0.04 |
| No                                                   | 87          | 23 (18–30)       |                  | 82 (64–93)            |         | 90 (71–100) |     |
expenses, bills, debt and financial worry) and social isolation were associated with both greater family and parental impact.

**Adjusted associations of potentially modifiable characteristics with outcomes**

Table 3 shows associations of potentially modifiable characteristics with the total IOF Scale scores, adjusting for potential confounders. Taking time off from work without pay, increased bills, financial worry, an unsafe home environment, and social isolation were all associated with higher total IOF scores, indicating greater impact. Similarly, Table 4 shows adjusted associations of potentially modifiable characteristics with impact on Parent Emotion and Time Limitation scores. Taking time off from work without pay and social isolation were associated with greater impact on parent time limitation, as was an unsafe home environment. In contrast, enrollment in early intervention and Medicaid programs were associated with higher parent time limitation scores, indicating less parental impact.

**Discussion**

In this study, we described the impact of preterm birth on parents and families in the first 2 years after neonatal discharge. Our results support our conceptual model, which posits modifiable factors that are associated with the impact of preterm birth on parents and families, independent of infant health and socio-demographic characteristics. We identified several potentially modifiable factors that were associated with both higher and lower impact. In particular, social isolation, financial burdens such as taking unpaid time off from work, increased bills and financial worry, and an unsafe home environment were all associated with higher impact on at least one of our main outcomes. In contrast, enrollment in early intervention and Medicaid and use of public housing were associated with less parent impact.

Predisposing characteristics such as infant co-morbidities affected both impact on family scores and parental scores. Infant development affected parental scores for increased anxiety and emotion. Our findings were consistent with previous studies that the impact was greater among families whose preterm children demonstrated either a functional handicap or low
Financial burden since infant discharge

- Family member has taken ≥ 3 h/week off from work without pay
  4.7 (0.8, 8.7) 0.02
- Family member has received compensation by employer for time off
  1.3 (−2.9, 5.6) 0.5
- No compensation (from any source) for time off
  4.8 (−0.3, 9.9) 0.06

Financial burden

- Unexpected costs
  1.5 (−2.8, 5.7) 0.7
- Increased bills
  6.1 (0.4, 11.9) 0.04
- Increased out-of-pocket expenses
  −1.5 (−7.4, 5.5) 0.6
- Financial worry
  4 (0.2, 7.7) 0.04
- Collections discussed prior to discharge
  2.5 (−2.5, 7) 0.4
- Enrollment in medi-caid
  1.3 (−3.5, 6) 0.6

Health related social problems

- Unsafe home environment
  6.3 (0.2, 12.3) 0.04
- Social isolation
  8.5 (4, 12.9) 0.0003

Models adjusted for: infant birthweight (per 500 g increase), race, family annual household income, infant co-morbidities during hospitalization in NICU (including of the following: growth restriction, surfactant deficiency, necrotizing enterocolitis, intraventricular hemorrhage grade 3 or 4, patent ductus arteriosus, retinopathy of prematurity)

*Estimates represent differences in points on Impact on Family Scale (range of possible scores, 0 to 56). Higher scores represent higher impact

Use of one or more public assistance programs: Supplemental nutritional assistance program for Women, Infants and Children, Transitional Aid to Families with Dependent Children, Low Income Home Energy Assistance Program

Unsafe home environment constitutes water leaks, pests, or no heat

Social isolation defined by positive response to query about feelings of isolation

We also found that the use of prescription medications and durable medical equipment affected both parental impact scores and impact on family scores, which was consistent with other publications [20, 60]. Specifically, the use of medications and medical equipment may contribute to the substantial out-of-pocket expenditures that families may incur [61, 62]. As pressure mounts to reduce hospital length of stay and readmission rates, and as we move more complex care into the community, high out-of-pocket costs is an important factor that can contribute to parental and family strain.

Several studies have shown that preterm birth and an infant’s hospitalization can adversely affect the finances of families after the birth of a preterm or VLBW infant [20, 60–63]. However, little is known about a more modifiable determinant such as the specifics of financial burden faced by families, or about the impact of financial burden on parent quality of life. In our study, we found that a lack of compensation for time off work was associated with both family and parent-time impact scores. Also, increased bills due to hospitalization and increased financial worry were associated with greater impact. Complementing our findings, 2 studies have reported the out-of-pocket costs incurred by families of preterm infants for outpatient services, medications, as well as indirect costs like lost productivity are significant especially during the first year after discharge [60, 64]. Specifically, Hodek et al. cited that co-payments for outpatient ancillary services and medications increased parental out-of-pocket expenses. Moreover, lost wages for missing work days may increase income losses [60]. Overall, by highlighting the specific aspects of financial burden most closely associated with parent and family impact, such as lack of compensation and increased bills, our results may inform targeted financial support programs for families of preterm infants after discharge. Moreover, our findings support that while annual income was not associated with impact on family, parental perspective on financial burden was, which should also be considered when caring for these families.

Another modifiable determinant are health related social problems. These are economic and social problems that can affect health such as food insecurity and substandard housing [45]. Prior studies have demonstrated the impact of substandard housing on child health such as increased infectious disease and injury [65]. However, we found that an unsafe home environment was associated with adverse parent-time impact and impact on the family. Timely receipt of public housing has been associated with improved health in other medical conditions [66]. Addressing housing concerns for families of preterm infants through existing public housing programs is a feasible approach to reducing the parental and family impact.

Another health related social problem that was associated with greater parental and family impact was social isolation. Other studies that have examined families of premature infants have found that “alienation” [67] and social isolation may have profound impact on parental emotion [68]. Jackson et al. described the paradigm of the process of acclimatization of caring for a premature
### Table 4 Adjusted associations of potentially modifiable characteristics with impact on parent emotion and time limitation scores (n = 196)\(^a\)

| Characteristics | Total (n = 196)\(^b\) | Parent emotion points | p-value |
|-----------------|------------------------|------------------------|---------|
| Use of community based resources | | | |
| Enrollment in early intervention | | 4.2 (−12, 20.5) | 0.6 |
| Use of social services\(^c\) | | −2.3 (−14, 9.5) | 0.7 |
| Use of public housing | | 7.9 (−13, 29) | 0.5 |
| Receive supplemental security income | | 0.5 (−12, 13.3) | 0.9 |
| Use of a community based clinic for primary care | | 1.9 (−12.3, 16) | 0.8 |
| Financial burden since infant discharge | | | |
| Family member has taken ≥ 3 h/week off from work without pay | | −12.4 (−20, −4) | 0.004 |
| Financial burden | | | |
| Unexpected costs | | −2.7 (−11, 6) | 0.6 |
| Increased bills | | −5 (−17, 7) | 0.4 |
| Increased out-of-pocket expenses | | 2.5 (−9, 14) | 0.4 |
| Financial worry | | −4.7 (−13, 3) | 0.3 |
| Collections discussed prior to discharge | | 1.1 (−9, 12) | 0.8 |
| Enrollment in medi-caid | | 4.9 (−5, 15) | 0.4 |
| Health related social problems | | | |
| Unsafe home environment\(^d\) | | −1.2 (−14, 12) | 0.8 |
| Social isolation\(^e\) | | −11.2 (−21, −1) | 0.03 |

### Table 4 Adjusted associations of potentially modifiable characteristics with impact on parent emotion and time limitation scores (n = 196)\(^{\text{Continued}}\)

| Characteristics | Total (n = 196)\(^b\) | Parent limitation on time score points (95% Confidence Interval) | p-value |
|-----------------|------------------------|-----------------------------------------------------------------|---------|
| Use of community based resources | | | |
| Enrollment in early intervention | | 20.2 (5.6, 34) | 0.007 |
| Use of social services\(^c\) | | 1.9 (−8.7, 12.5) | 0.7 |
| Use of public housing | | 3.5 (−15, 22) | 0.7 |
| Receive supplemental security income | | 0.1 (−15, 22) | 0.7 |
| Use of a community based clinic for primary care | | 5.5 (−7, 18) | 0.4 |
| Financial burden since infant discharge | | | |
| Family member has taken ≥ 3 h/week off from work without pay | | −8.8 (−16.9, −1) | 0.03 |
| Family member has received compensation by employer for time off | | 5.4 (−3.5, 15) | 0.2 |

---

infant as alienation, responsibility, confidence and familiarity, and that alienation may be protracted in this population [67]. Intervention strategies that have improved parental emotions often include education-behavioral models. For example, the Creating Opportunities for Parent Empowerment (COPE) program created by Melnyk, et al. was associated with reduced parental impact during transition home from the NICU [69, 70]. It is possible that programs like that one benefit families by reducing social isolation in months to years after discharge.

Another modifiable determinant is enrollment in a community-based developmental program like “Early Intervention (EI),” was associated with less impact on parental limitation of time. A recent meta-analysis suggests that community-based developmental programs had beneficial pooled effects on maternal anxiety, depressive symptoms, and self-efficacy [71]. Moreover, other studies have suggested that these programs can also empower families because of the collaborative process that EI offers; in turn, they have a deeper understanding of their child’s developmental needs [72]. Similarly to early intervention, we found that receipt of Medicaid was associated with lower impact scores on limitations on time. Other studies have demonstrated families who were registered with Medicaid showed improved “parent role confidence” and “parent-baby
interaction" than those with private insurance [69]. While this result was unexpected, it has been speculated that parents with a higher socioeconomic status and private insurance may have higher expectations for themselves [69] and therefore may perceive an increased parental impact on their time versus those who utilize Medicaid. Overall, our results suggest that greater participation in public assistance programs may lessen familial and parental burden for this patient population.

A strength of our study was our high response rate (82%). Characteristics of the infants were very similar to other follow-up programs [25]. However, we studied families of infants presenting for follow-up care rather than the underlying population of families of infants receiving neonatal intensive care, potentially limiting generalizability. Moreover, we did sample both mothers and fathers, which may affect how some of our results are interpreted and parent gender may influence some of the measures including financial burden and social isolation [73]. Also, our study was cross-sectional making us unable to establish causation. While we adjusted for a number of potential confounders, like all observational studies, ours is subject to residual confounding. While we did not have a full term cohort control in comparison [25], nor data on those not enrolled, our aim was to elicit the experience of parents and families of preterm infants, as well as relationships with modifiable characteristics specific to this population.

Conclusions
In summary, we identified several predictors of increased family and parent impact in families of preterm infants. Of particular interest were the potentially modifiable factors including social isolation and financial burden, which were associated with greater impact, and use of community-based developmental services, public housing, and Medicaid which were associated with less impact.

Our results suggest that interventions to target these factors, for example social and financial support programs, and efforts to increase enrollment in community-based developmental services and public health insurance programs, might lessen the impact of preterm birth on parents and families.

Acknowledgments
This study was supported by the Richardson Fund, Department of Neonatology, Beth Israel Deaconess Medical Center, Boston MA; the Program for Patient Safety and Quality (PPSQ) at Boston Children’s Hospital, Boston, MA and the Medical Staff Organization at Boston Children’s Hospital, Boston, MA. We would also like to recognize Jane Stewert, MD, the director of the Infant Follow Up Program (IFUP) at Boston Children’s Hospital, Larry Rhein, MD, the director of the Center for Healthy Infant Lung Development (CHILD) program at Boston Children’s Hospital and Janet Soul, MD, the director of the Fetal Neurology Program at Boston Children’s Hospital who facilitated our study. We would also like thank Brooke Corder, the social worker at the IFUP who assisted with patient recruitment. We would also like to acknowledge Aaron Picklins who developed the web-interface for the study questionnaire and Drs. Dionne Graham and Sheree Schrager who assisted with statistical analysis.

Funding
This study was also supported by the Richardson Fund with the Department of Neonatology, Beth Israel Deaconess Medical Center, Boston MA; the Program for Patient Safety and Quality (PPSQ) at Boston Children’s Hospital, Boston, MA and the Medical Staff Organization at Boston Children’s Hospital, Boston, MA.

Availability of data and materials
The datasets generated during and/or analyzed during the current study are not publicly available due to patient information, but are available from the corresponding author on reasonable request.

Authors’ contributions
Dr. Lakshmanan and Dr. Belfort made substantial contributions to designing the study, analyzing the data, and interpreting the results. Dr. Lakshmanan wrote the first draft of the manuscript. Ms. Agni collected the data and contributed to data analysis and drafting of the manuscript. Drs. Lieu, Fleegler and McCormick made substantial contributions to designing the study, interpreting the results, and critically revising the manuscript. Drs. Kipe and Friedlich also assisted in interpreting the results and revising the manuscript. All authors read and approved the final version of the manuscript.

Consent for publication
Not applicable. There is no identifiable data in this study.

Competing interests
The authors declare that they have no competing interests.

Ethics approval and consent
The Boston Children's Hospital and Children's Hospital Los Angeles human subjects committees approved the study protocol and participation consent was obtained from all subjects.

Financial disclosure
Dr. Lakshmanan is supported by National Center for Advancing Translational Science (NCATS) of the U.S. National Institutes of Health (KL2TR001854). Dr. Belfort was supported by National Institutes of Health K23DK83817. The remaining authors have indicated they have no financial relationships relevant to this article to disclose.

Author details
1 Center for Fetal and Neonatal Medicine, USC Division of Neonatal Medicine, Children’s Hospital Los Angeles; Keck School of Medicine, University of Southern California, 4650 Sunset Boulevard, MS #31 CA 90027 Los Angeles, USA. 2 Newborn and Infant Critical Care Unit, Children’s Hospital Los Angeles, 4650 Sunset Boulevard, MS #31 CA 90027 Los Angeles, USA. 3 Department of Preventive Medicine, Keck School of Medicine, University of Southern California, Los Angeles, CA, USA. 4 Center for Health Policy and Economics, University of Southern California, Los Angeles, CA, USA. 5 Drexel School of Medicine, Philadelphia, PA, USA. 6 Division of Research, Kaiser Permanente, Oakland, CA, USA. 7 Division of Emergency Medicine, Boston Children’s Hospital, Boston, MA, USA. 8 Saban Research Institute, Children’s Hospital Los Angeles, Los Angeles, CA, USA. 9 Division of Newborn Medicine, Boston Children’s Hospital, Boston,
References

1. Hamilton BE, Hoyert DL, Martin JA, Strobino DM, Guyer R. Annual summary of vital statistics: 2010–2011. Pediatrics. 2013;131:S48–58.

2. Wang CJ, McGlynn EA, Brook RH, et al. Quality-of-care indicators for the neurodevelopmental follow-up of very low birth weight children: results of an expert panel process. Pediatrics. 2006;117:2086–92.

3. Newborn AaOPCoFta. Hospital discharge of the high-risk neonate. Pediatrics. 2008;122:1119–26.

4. Garfield L, Holditch-Davis D, Carter CS, et al. Risk factors for postpartum depressive symptoms in low-income women with very low-birth-weight infants. Adv Neonatal Care. 2015;15:53–74.

5. Howe TH, Sheu CF, Wang TN, Hsu YW. Parenting stress in families with very low birth weight preterm infants in early infancy. Res Dev Disabil. 2014;35:1748–56.

6. Thygesen SK, Olsen M, Ostergaard JR, Sorensen HT. Respiratory distress syndrome in moderately late and late preterm infants and risk of cerebral palsy: a population-based cohort study. BMJ Open. 2016;6011643.

7. Vohr BR, Wright LL, Dusick AM, et al. Neurodevelopmental and functional outcomes of extremely low birth weight infants in the National Institute of Child Health and Human Development Neonatal Research Network, 1993–1994. Pediatrics. 2000;105:1216–26.

8. Hack M, Taylor HG, Drotar D, et al. Chronic conditions, functional limitations, and special health care needs of school-aged children born with extremely low-birth-weight in the 1990s. JAMA. 2005;294:318–25.

9. Saigal S, Doyle LW. An overview of mortality and sequelae of preterm birth and special health care needs of school-aged children born with extremely low birth weight children. Pediatrics. 2009;124:717–27.

10. Stein RE, Jessop DJ. Measurement of neurobehavioral outcomes in very preterm and/or very low birth weight infants on the family is long lasting. A matched control study. Health Serv Res. 2017; 15:38.

11. Aday LA, Andersen R. A framework for the study of access to medical care. Health Serv Res. 1974;9:208–20.

12. Raina P, O’Donnell M, Schweller H, et al. Caregiving process and caregiver burden: conceptual models to guide research and practice. BMC Pediatr. 2004;4:1.

13. Drotar D, Hack M, Taylor G, Schluchter M, Andreasi L, Klein N. The impact of extremely low birth weight on the families of school-aged children. Pediatrics. 2006;117:2006–13.

14. Stein RE, Jessop DJ. Uncertainty and its relation to the psychological and social correlates of chronic illness in children. Soc Sci Med. 1985;20:993–9.

15. Stein RE, Jessop DJ. What diagnosis does not tell: the case for a noncategorical approach to chronic illness in childhood. Soc Sci Med. 1989;29:769–78.

16. Stein RE, Jessop DJ. Measuring health variables among Hispanic and non-Hispanic children with chronic conditions. Public Health Rep. 1989;104:377–84.

17. Stein RE, Jessop DJ. The impact on family scale revised: further psychometric data. J Dev Behav Pediatr. 2003;249–16.

18. Landgraf JM, Vogel I, Oostenbrink R, van Baar ME, Raat H. Parent-reported health outcomes in infants/toddlers: measurement properties and clinical validity of the ITQOL-SF47. Qual Life Res. 2013;22:635–46.

19. Kuzina I, Janssen W, van Sprang NC, Carter AS, Raat H. The Effectiveness of the BITSEA as a Tool to Early Detect Psychosocial Problems in Toddlers, a Cluster Randomized Trial. PLoS One. 2015;10(e0136488).

20. Flink II, Beierens TM, Looman C, et al. Health-related quality of life of infants from ethnic minority groups: the Generation R Study. Qual Life Res. 2011;20:653–64.

21. Flink II, Prins RG, Mackenbach JJ, et al. Neighborhood ethnic diversity and behavioral and emotional problems in 3 year olds: results from the Generation R Study. PLoS One. 2013;8(e0107007).

22. Raat H, Landgraf JM, Oostenbrink R, Moll HA, Essink-Bot ML. Reliability and validity of the Infant and Toddler Quality of Life Questionnaire (ITQOL) in a large population and respiratory disease sample. Qual Life Res. 2007;16:445–60.

23. Oostenbrink R, Jongman H, Landgraf JM, Raat H, Moll HA. Functional abdominal complaints in pre-school children: parental reports of health-related quality of life. Qual Life Res. 2010;19:363–9.

24. van Baar ME, Essink-Bot ML, Derma, et al. Reliability and validity of the Health Outcomes Burn Questionnaire for infants and children in The Netherlands. Burns. 2006;32:557–67.

25. ITQOL: Infant Toddler Quality of Life Questionnaire. (Accessed December 2nd, 2016, at healthactchq.com.)

26. Meert KL, Slomine BS, Christensen JR, et al. Family Burden After Out-of-Hospital Cardiac Arrest in Children. Pediatrics. 2016;121:498–507.

27. van Zeillem L, Buysse C, Madderon M, et al. Long-term neuropsychological outcomes in children and adolescents after cardiac arrest. Intensive Care Med. 2015;41:1076–1082.

28. Rolfsjord LB, Skjeroven HO, Carlsen KH, et al. The severity of acute bronchitis in infants was associated with quality of life nine months later. Acta Paediatr. 2016;105:834–41.

29. Doty M, Rustgi SD, Schoen C, Collins SR. Maintaining health insurance during a recession: likely COBRA eligibility: an updated analysis using the Commonwealth Fund 2007 Biennial Health Insurance Survey. Issue brief. 2009;9491–12.

30. Doty MM, Collins SR, Nicholson JL, Rustgi SD. Failure to protect: why the individual insurance market is not a viable option for most U.S. families: findings from the Commonwealth Fund Biennial Health Insurance Survey. Issue brief. 2009;9521–16.

31. Doty MM, Collins SR, Rustgi SD, Nicholson JL. Out of options: why so many workers in small businesses lack affordable health insurance, and how health care reform can help. Findings from the Commonwealth Fund Biennial Health Insurance Survey, 2007. Issue brief. 2009;67:1–22.

32. Rustgi SD, Doty MM, Collins SR. Women at risk: why many women are foregoing needed health care. An analysis of the Commonwealth Fund 2007 Biennial Health Insurance Survey. Issue brief. 2009;6521–12.

33. Fleegler EW, Lieu TA, Wise PH, Muetet-Wagstaff S. Families’ health-related social problems and missed referral opportunities. Pediatrics. 2007;119:e1322–41.

34. Hassan A, Blood E, Piklingis A, Krull E, McNickles L, Mammon G, Woods E, Fleegler E. Improving Social Determinants of Health: Effectiveness of a web-based intervention. Am J Prev Med. 2015;49(6):822–31. doi:10.1016/j.amepre.2015.04.023.
47. Wylie SA, Hassan A, Krull EG, et al. Assessing and referring adolescents’ health-related social problems: concerns often overlooked during the medical visit. J Adolesc Health. 2013;53:265–71.
48. Wylie SA, Hassan A, Krull EG, et al. Assessing and referring adolescents’ health-related social problems: qualitative evaluation of a novel web-based approach. J Telemed Telecare. 2012;18:392–8.
49. Statistics NCoH. National Health Interview Survey. Washington DC: Services UDohAEd; ed, 1995.
50. Bureau UC. The American Housing Survey. Washington DC: Development DoHaH, ed; 1994.
51. Cress JFJ. The Philadelphia Survey of Child Care and Work. Philadelphia, PA: Temple University; 2003.
52. Weinreb L, Wehler C, Perloff J, et al. Hunger: its impact on children’s health and mental health. Pediatrics. 2002;110:e41.
53. Accountable Health Communities Model. 2016. https://innovation.cms.gov/initiatives/AHCM). Accessed 18 Dec 2016.
54. Hediger ML, Overpeck MD, Ruan WJ, Troendle JF. Birthweight and gestational age effects on motor and social development. Paediatr Perinat Epidemiol. 2002;16:33–46.
55. at http://www.nlsinfo.org/childya/nlsdocs/questionnaires/2006/Child2006quex/MotherSupplement2006_Assessments.html#MOTORANDSOCIALDEVELOPMENT).
56. Belfort MB, Santo E, McCormick MC. Using parent questionnaires to assess neurodevelopment in former preterm infants: a validation study. Paediatr Perinat Epidemiol. 2013;27:199–207.
57. Rivers A, Caron B, Hack M. Experience of families with very low birthweight children with neurologic sequelae. Clin Pediatr. 1987;26:223–30.
58. Taylor HG, Klein N, McNich NW, Hack M. Long-term family outcomes for children with very low birth weights. Arch Pediatr Adolesc Med. 2001;155:55–61.
59. Saigal S, Burrows E, Stokofl BL, Rosenbaum PL, Steiner D. Impact of extreme prematurity on families of adolescent children. J Pediatr. 2000;137:701–6.
60. Hodek JM, von der Schulenburg JM, Mittendorf T. Measuring economic consequences of preterm birth - Methodological recommendations for the evaluation of personal burden on children and their caregivers. Heal Econ Rev. 2011;1:6.
61. Petrou S. Economic consequences of preterm birth and low birthweight. BJOG. 2003;110 Suppl 20:17–23.
62. McCormick MC, Bembbaum JC, Eisenberg JM, Kustra SL, Finnegan E. Costs incurred by parents of very low birth weight infants after the initial neonatal hospitalization. Pediatrics. 1991;88:33–41.
63. Underwood MA, Danielsen B, Gilbert WM. Cost, causes and rates of rehospitalization of preterm infants. J Perinatol. 2007;27:614–9.
64. Schiffman JK, Dulchavny D, Mowitz M, Kirpalani H, Mao W, Roberts R, Nyberg A, Zupancic J. Quantifying the Economic Burden of Neonatal Illness on Families of Preterm Infants in the U.S. and Canada. San Diego, CA: Pediatric Academic Societies; 2015.
65. Krieger J, Higgins DL. Housing and health: time again for public health action. Am J Public Health. 2002;92:758–68.
66. Rodgers JT, Purnell JQ. Healthcare navigation service in 2-1-1 San Diego: guiding individuals to the care they need. Am J Prev Med. 2012;43:S450–6.
67. Jackson K, Temstedt BM, Schollin J. From alienation to familiarity: experiences of mothers and fathers of preterm infants. J Adv Nurs. 2003;43:120–9.
68. Meijers DE, Wolf MJ, Koldewijn K, van Wassenaer AG, Klok JH, van Baar AL. Parenting stress in mothers after very preterm birth and the effect of the Infant Behavioural Assessment and Intervention Program. Child Care Health Dev. 2011;37:195–202.
69. Melnyk BM, Osvalt KL, Sidora-Arcoleo K. Validation and psychometric properties of the neonatal intensive care unit parental beliefs scale. Nurs Res. 2013;62:105–15.
70. Melnyk BM, Alpert-Gillis LJ. The COPE program: a strategy to improve outcomes of critically ill young children and their parents. Pediatr Nurs. 1998;24:521–7.
71. Benzeis RM, Magill-Evans JE, Hayden KA, Ballantyne M. Key components of early intervention programs for preterm infants and their parents: a systematic review and meta-analysis. BMC Pregnancy Childbirth. 2013;13 Suppl 1:S10.
72. Pighini MJ, Goelman H, Buchanan M, Schonert-Reichl K, Brynelsen D. Learning from parents’ stories about what works in early intervention. Int J Psychol. 2014;49:263–70.
73. Arockiasamy V, Holsti L, Albersheim S. Fathers’ experiences in the neonatal intensive care unit: a search for control. Pediatrics. 2008;121:e215–22.
74. Phillips KA, Morrison KR, Anderson R, Aday LA. Understanding the context of healthcare utilization: Assessed environmental and provider-related variables in the behavioral model of utilization. Health Serv Res. 1998;33:571–96.