3-20-2020

Pediatric resident knowledge, experience, comfort, and perceived competency in providing sibling psychosocial support.

David Buchbinder
Sonam Sidhu
Melissa A. Alderfer
Anne Lown
Russ C. Kolarik

See next page for additional authors

Follow this and additional works at: https://jdc.jefferson.edu/pedsfp

Part of the Pediatrics Commons

Let us know how access to this document benefits you

This Article is brought to you for free and open access by the Jefferson Digital Commons. The Jefferson Digital Commons is a service of Thomas Jefferson University's Center for Teaching and Learning (CTL). The Commons is a showcase for Jefferson books and journals, peer-reviewed scholarly publications, unique historical collections from the University archives, and teaching tools. The Jefferson Digital Commons allows researchers and interested readers anywhere in the world to learn about and keep up to date with Jefferson scholarship. This article has been accepted for inclusion in Department of Pediatrics Faculty Papers by an authorized administrator of the Jefferson Digital Commons. For more information, please contact: JeffersonDigitalCommons@jefferson.edu.
Authors
David Buchbinder, Sonam Sidhu, Melissa A. Alderfer, Anne Lown, Russ C. Kolarik, and Tommy Wang
To the Editor

There is a growing number of children with life-threatening chronic health conditions in the United States.1-2 Care must encompass consideration of psychosocial support for family members.3 There is a variety of unmet psychosocial support needs among siblings of children with chronic conditions.4-6 Despite a focus on family-centered care,7-8 it is unclear whether pediatric training programs prepare physicians to address the psychosocial needs of their patients’ siblings.9-10 To address this gap in our knowledge, we sought to determine the extent of pediatric residents’ training, knowledge, experience, comfort, and perceived competence in areas relating to the psychosocial support of siblings. We also aimed to learn from residents their views regarding psychosocial support for siblings, their desire for training, and the content areas in which they are interested.

From June to December 2017, residents were approached for participation during a monthly residency meeting. Completion of the survey was voluntary. Residents from a single center were asked to provide their year of training if they had been involved in the care of a child whose sibling may have benefited from evaluation and referral for support services. Only 23% endorsed ever evaluating or referring a sibling. Approximately half (55%), endorsed any personal experiences with a sibling of a child with a life-threatening illness.

Less than half of residents endorsed at least some training, experience, knowledge, competency, and comfort with respect to counseling parents on topics (e.g., emotional, behavioral, quality of life impact on siblings, identifying signs that a sibling has difficulty adjusting) pertinent to the psychosocial care of siblings including ways to talk with siblings about illness, treatment, and family impact as well as talking to siblings when a brother or sister is near death. Less than one-quarter of residents endorsed at least some training, experience, knowledge, competency, and comfort with screening to identify at-risk siblings as well as providing a referral to psychosocial support services.
The majority (>90%) of residents agreed that primary care providers, specialty care providers, and oncologists have an important role in the provision of psychosocial support for siblings. Across the ten topic areas presented to participants, the top three areas that participants indicated were most important to include in training curricula were (in descending order of importance): 1) counseling parents about siblings’ needs; 2) the types of challenges siblings face when a brother or sister has a serious illness, and 3) screening siblings to determine if they are at risk for problems.

Few pediatric residents endorsed adequate training, experience, knowledge, competence and comfort relating to the counseling of parents and siblings directly on topics pertinent to the psychosocial care of siblings. Moreover, counseling parents and siblings directly was a topic of importance for future training needs. Counseling parents of children with a chronic health condition should include a focus on tending to the siblings’ needs. For example, pediatric trainees must be able to educate parents to be cognizant of “red flags” that would alert them to a sibling that is having a difficult time adjusting to having a brother or sister with a chronic health condition. Pediatricians must be able to counsel siblings in ways that facilitate their involvement in the family with greater understanding about the care needs of their ill brother or sister. We must be able to create opportunities to engage siblings through various means such as providing siblings with developmentally appropriate education regarding their brother or sister’s disease and its treatment.

We found that most pediatric residents lacked training, experience, knowledge, competence and comfort in counseling parents about ways to talk with siblings when their brother or sister was near death as well as communicating with siblings directly when their brother or sister was near death. The death of a child is one of the most challenging experiences a pediatric resident will face during their training. Pediatric residents must be able to provide support to soon-to-be bereaved siblings ensuring open and honest communication with these siblings in a developmentally appropriate context. In the context of bereavement, there is a greater focus on palliative care in pediatrics; however, residents are still ill-equipped to address the needs of bereaved or soon-to-be bereaved siblings.

Most pediatric resident’s lack training, experience, knowledge, competency, and comfort with respect to conducting a screening assessment to identify at-risk siblings. Moreover, screening siblings to determine if they are at risk for problems was also noted to be a topic of importance for future training. Barriers exist in the context of conducting screening assessments of siblings, including a lack of familiarity with screening tools as well as a lack of time and resources required to conduct assessments. There is now an increasing number of instruments that may be utilized in the identification of at-risk siblings of children with chronic health conditions. We must ensure that residents can identify appropriate screening tools as well as utilize, interpret, and apply the results of these screening assessments in their clinical practice.

Although the psychosocial support of siblings has been under the purview of child life specialists and social workers or other healthcare professionals, pediatric residents perceive that they have an essential role to play in the provision of sibling psychosocial support. Perceptions of pediatric residents regarding their role in support of siblings are positive. Despite this, residents face challenges in fulfilling this role, including a lack of training, time, and comfort in the provision of such care. Strategies must be put into place to change our current training system to ensure that these efforts can be supported.

We found that pediatric residents, in general, lacked training, experience, knowledge, competence and comfort in addressing topics that are vital to the provision of sibling psychosocial support. Findings from this study should inform the development of curricula that will be able to ease the burden of providing psychosocial care to siblings. It is critical that we meet the needs of the siblings’ in order to ensure the delivery of family-centered comprehensive care to patients and their families.

Conflicts of Interest
The authors declare that they have no conflicts of interest.

References
5. Patterson P, McDonald FE, White KJ, Wallczaek A, Butow PN. Levels of unmet needs and distress amongst adolescents and young adults (AYAs) impacted by familial cancer. Psychooncology. 2017;26(9):1285-1292.