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Erring on the Side of Life: Children with Rare Trisomy Conditions, Medical Interventions and Quality of Life

Deborah A. Bruns*†

Abstract

The prevailing viewpoint on children with rare trisomy conditions such as trisomy 18 (t18) and trisomy 13 (t13) is almost uniformly negative. Yet, case studies offer information about long-term survivors. What is missing in the discussion is an unbiased examination of surviving children within the context of necessary, rather than “aggressive”, medical interventions and overall quality of life. A move beyond palliative or comfort care must be an option for this population. There must be a move toward valuation of life and corresponding provision of treatment and examination of developmental gains rather than limited intervention or palliative care for infants with lethal fetal abnormalities. This article presents a call to examine the individual child rather than decision making by diagnosis framed by recommendations from the Convention on the Rights of the Child (CRC) and Convention on the Rights of Persons with Disabilities (CRPD) Medical professionals and parents must work together to ensure medical needs are met and a positive quality of life can be achieved.

Keywords
Trisomy 18; Trisomy 13; Decision-making; Medical interventions; Quality of life

The prevailing viewpoint on children with rare trisomy conditions such as trisomy 18 (t18) and trisomy 13 (t13) is almost uniformly negative. Population based studies describe low survival rates for infants commonly labeled as possessing “lethal fetal anomalies” or who are considered “incompatible with life” [1-6]. Yet, case studies are available highlighting long-term survival [7] as well as parents sharing their experiences online via blogs and Facebook. The paradox is clear. Studies describing large number of children are bleak while individual cases are more positive. The majority of medical professionals look to the former while parents the latter. This often results in virtual absence of cognitive function, e.g. trisomy 18...the potential for cognitive development – and therefore the achievement of human and social goods, e.g. relationships with others.
– are virtually absent” (p. 255). It appears that the authors have not reviewed the case studies or experienced contact with living infants and children with t18 or t13. As this author can attest, cognition may be significantly impaired but interactions and relationships with parents, siblings and significant others (e.g., grandparents, teachers) are evidenced in a variety of ways including eye contact, smiling, laughing, reaching and vocalizing by children with these conditions.

Everett and Albersheim [27] describe Baby Smith, an infant with t18. The authors describe a framework to “…increase transparency, dialogue, understanding, and trust, which, in turn, may achieve greater consensus” (p. 55) for medical care decision making. The medical team refused to intubate Baby Smith and did not aggressively treat his unspecified cardiac condition. The infant passed away on the 82nd day of life. What is most disheartening in this case study is the lack of value placed on the infant’s life and his parents’ wishes for more aggressive treatment. This is also evidenced in the care decisions presented by Bruns and Crosier [28]. The infant, Simon, passed away on the 88th day of life. As the case study indicates, quality of life is only partially dependent on length and medical needs. The infant encountered many difficulties during his brief life but all who came into contact with him were positively affected including medical professionals such as neonatal nurses and a pediatric cardiologist. Their views on medical treatments changed as a result of caring for Simon.

Interestingly, a study by Carey [29] of neonatologists’ views on resuscitation of newborns with t18 states “It is critical that the emphasis should remain on fostering the best interest of the infant at every branch of the treatment decision tree” (p. 1109). Yet, this positive outlook is tempered by the authors’ recommendation that “infants with lethal congenital anomalies and profound neurologic impairment continue to have immense inherent worth as human beings, and these infants are without doubt as deserving of love, care, and dignity as any child”. However, we contend that having intensive care measures such as intubation and corrective surgery available as potential options for infants with a confirmed lethal trisomy gives the impression to parents that these are reasonable interventions to consider…” (p. 1108).

It is puzzling how both viewpoints can co-exist in terms of identifying treatment options and reinforcing the notion of “lethal congenital anomalies”. The authors offer some positive views on decision making but reinforce the negative stance of many in the palliative care field [9,12,15,16,27].

The literature reinforces the statistic of only 10% of affected infants reaching their first birthday. This percentage is often cited to parents at the time of prenatal diagnosis [2,6,13]. Yet, the 10% amount does not grant the authors cited here exclusive emphasis on the 90% of non-survivors. Investigations into differences between long-term survivors and non-survivors are absent. There is a dearth of information concerning characteristics of infants who die prior to one month, for example, compared with those living between one and three years. Analyses of these data are especially needed in the face of recommendations for comfort care at birth and palliative care after the immediate newborn period.

Koogler et al. [30] point out, parents should be presented with all possible outcomes to arrive at informed decisions on their infant’s behalf rather than being told their infant has a lethal anomaly and should not receive care. In order for guidelines for decision making to be developed, there is a necessity to gather more extensive data (population-based and directly from parents and caregivers), build consensus and, only then, offer recommendations. In fact, Janvier et al. [31] describe positive perspectives from parents coupled with the need to learn more about their experiences raising children with rare trisomy conditions. Without these voices, there will be a continued emphasis on comfort care and non-treatment.

Offering a Positive Viewpoint and Associated Recommendations

In this author’s experience, infants with t18 and t13 can and do flourish after cardiac repair and similar types of aggressive medical intervention [21,32]. Along with this data, there is a concomitant need to examine the needs of individual children rather than over-reliance on statistical information for decision making. Data collected for the Tracking Rare Incidence Syndromes (TRIS) project includes multiple data points on medical needs and services and developmental outcomes. Two children in the TRIS database are described below:

Annabel was born in March 2005 at 40 weeks gestation via planned c-section. She weighed 2781 grams. Medical issues at birth included respiratory difficulties, heart murmur, ASD, VSD, feeding difficulties and jaundice. She was diagnosed with trisomy 18, 11 days after birth. Currently, Annabel’s most pressing issues are kidney and liver functioning as well as related to feeding including PICC and central line concerns. Annabel had ureteral reimplantation surgery at 19 months. She currently receives medical care in the areas of cardiology, dermatology, gastroenterology, genetics, ophthalmology, pulmonology, and urology. Annabel imitates simple gestures. Annabel associates names of objects with their representation. She also displays preferences for familiar adults.

Arianna was born in June 2006 at 39 weeks gestation via vaginal birth. She weighed 2781 grams. Medical issues at birth included respiratory difficulties, ASD, PDA, VSD, feeding difficulties, and jaundice. She was diagnosed at three days. Arianna required oxygen until the spring of 2012. She is fed via a G-tube. She is also diagnosed with hyperopia and conductive hearing loss. At seven months of age, Arianna had cranial stenososis surgery. She also had her tonsils and adenoids removed at 23 months. Arianna indicates preferences for familiar adults and uses her hands to explore objects. She also interacts during social games.

It is important to note that both children were diagnosed postnatally. “Soft markers” were not identified for Annabel but were present for Arianna. Her family did not consider termination due to their religious beliefs. There is also much additional data on both children describing educational and therapy services and use of assistive items such as a gait trainer and augmentative communication devices.

Most current literature does not provide this type of in-depth discussion of long-term survivors and generally reinforces the findings on early death or negative outcomes. Yet, this information is needed to expand the current knowledge base for professionals and provide parents with more positive, yet realistic (e.g., surgeries, chronic conditions), outcomes. Annabel and Arianna’s families do not focus on what their daughters cannot do or compare her to their typically developing siblings. They celebrate their children for their personality, temperament and abilities rather than their diagnosis.

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Lantos and Meadow [33] agree that decisions should be made based on an individual infant’s response to treatment. Derrington and Dworetz [34] also emphasize an individualized approach coupled with understanding the values of each infant’s family. Information from medical professionals is used to arrive at decisions regarding medical interventions. Best interests of child were discussed with information from physicians about possible outcomes. It is unfortunate that the recommendations of Derrington and Dworetz [34] and Lantos and Meadow [33] are not followed as evidenced by Thiele’s [35] description of the circumstances she faced with her son with t18. What makes this case particularly discouraging was the absence of compassion after prenatal diagnosis of the condition. The author’s account illustrates the medical profession’s bias toward a paradigm of medical futility and placing minimal value on a life due to the child’s rare trisomy diagnosis. With the continued rise of prenatal testing, the end point of this perspective becomes increasingly disheartening.

Making treatment decisions in the best interests of an infant cannot be effectively done with a “doom and gloom” orientation as is often the case toward this population. How can hope be held when a condition is characterized as “…lethal condition that [is] not curable by intensive care…” [27]? How can parents make decisions without counseling that incorporates a variety of outcomes and an appreciation for quality of life [36-38]? Quality of life is not solely ours to decide. It must be informed by all possible outcomes gleaned from valid research including parent perspectives on their living as well as deceased children. In this author’s experience with the TRIS project, the latter group voices an almost uniform point of view of the value of their child’s life regardless of how short, medical complications and the like.

Merritt et al. [39] discuss decision making and the need for case-by-case review as the basis for decision-making rather than broad recommendations. Nelson et al. [40] state “although diagnoses of trisomy 13 and 18 are generally assumed to be fatal within days to weeks after birth, a small but significant subgroup of children with trisomy 13 and 18 are alive over the age of 1 year, and at least some of these children receive substantial inpatient hospital care” (p. 874) Yates et al. [41] point to a shift in interventions to address cardiac anomalies based on parent request. Yet, the impact of these studies is minimized by empirical reports emphasizing early mortality [42].

At the present time, there is no definitive consensus on medical interventions and quality of life due to the reasons explained here. A greater effort to collect longitudinal data is a starting point. The TRIS project has up to six years of data for some children and adults (project began in 2007; parents complete annual updates on medical interventions, therapy services etc) There are no similar databases. There continues to be a reliance on adherence to the “gloom and doom” data cited in the literature [1,2,9,12,24,27] rather than a more balanced representation of possible outcomes [17,19,29,36].

Conclusion
There is a need toward ethical case analysis for medical interventions and determining quality of life for children with t18 and t13 as described by Hentschel et al. [43]. The authors state “more attention should be focused on the parents’ information level and on their involvement in the decision-making process” (p. 568). This perspective, along with data indicating the success of medical interventions for this population [3,19,21,32,44,45] deserve greater study and consideration. In addition, changing thoughts on palliative care must be considered on behalf of this population [46]. As Fenton [47] explained, “How easy it is to assume we know what a good quality of life is for anyone other than ourselves. We assess the burdens of care, the impact on siblings, the impact on parental relationships, the impact on finances and the utilization of resources. We perform ethical analyses assuming we truly understand how to apply beneficence or malefice to a child with trisomy 18 or 13 or any other disorder in which there may be profound disability. As if the child can tell us what he or she is feeling. But we can do our best to assess and treat pain and discomfort. Smiles and laughter need no score pad. We know what they mean. The advice we give may often be centered around our personal notions about quality of life… advocate for a willingness to do whatever it takes, however long it takes, however many consults and team members it takes to fully inform and understand the goals, values and aspirations parents may have for their children. Once we have done that we need to honor the parents by helping them achieve those goals whether it is comfort care alone, a full court press or something in between.”

These along with directives established by the “…CRC and CRPD…as the comprehensive articulation of the rights of all individuals with disabilities including infants and young children” discussed by Brown and Guralnick [22] and coupled with Carey’s [48] recommendation for a “…balanced approach to counseling families of the newborn with trisomy 18 and 13 at the time of diagnosis and at decision points in management, that is, in the delivery room, newborn nursery and clinic” are words to guide decision making. The present review was intended to provide a counterpoint to discussions of “incompatibility with life”, “lethal diagnoses” and a general unwillingness for “aggressive” interventions. Long-term survivors with rare trisomy conditions must be further studied so that recommendations for their care can be made on an informed basis rather than from a biased perspective inclined toward comfort care or palliative care. Quality of life needs to enter the equation informed by data and parent voices. Children like Annabel and Arianna deserve no less.

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