Perioperative Care of a Child With Cri Du Chat Syndrome

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Abstract

Cri du chat syndrome (CdCS) is a chromosomal disorder resulting from a deletion in the short arm of chromosome 5. Anatomical abnormalities of the larynx result in a distinctive high-pitched, cat-like cry for which the disorder is named. Typical findings of the syndrome involve the upper airway, cardiovascular, and central nervous system (CNS). Of particular concern during anesthetic care is the potential for airway abnormalities leading to difficulties with endotracheal intubation as well as the presence of congenital heart disease (CHD). We present a 15-month-old child with CdCS who required anesthetic care during direct laryngoscopy and supraglottoplasty. The perioperative concerns of such patients are discussed, and previous reports of anesthetic care reviewed.

Keywords: Cri du chat syndrome; Cat cry syndrome; 5p minus syndrome; Anesthesia; Perioperative care

Introduction

Cri du chat syndrome (CdCS), also known as 5p minus syndrome or cat cry syndrome, is a chromosomal disorder that was first described by Lejeune et al in 1963 [1]. A total or partial deletion of the short arm of chromosome 5 (5p-) results in an anatomically abnormal larynx causing the distinctive high-pitched, cat-like cry for which the disorder is named [2]. Additional phenotypic characteristics include microcephaly, facial dysmorphism (epicanthus, wide nasal bridge, short philtrum, flattened maxilla, micrognathia, hypertelorism), hypotonia, intellectual disability with developmental delay, and congenital heart disease (CHD) [2, 3]. The severity of its clinical manifestations and phenotypic expression varies based on the magnitude of the chromosomal deletion [3-6].

With an incidence of 1:15,000 - 50,000 live births, CdCS is one of the more common human chromosomal disorders [3]. Given the multi-system involvement of the disorder, anesthetic care may be required during various surgical procedures. We present a 15-month-old child with CdCS who required anesthetic care during direct laryngoscopy and supraglottoplasty. The perioperative concerns of such patients are discussed, and previous reports of anesthetic care reviewed.

Case Report

Preparation of this case report followed the guidelines of the Institutional Review Board (IRB) of Nationwide Children’s Hospital (Columbus, Ohio). The patient was a 15-month-old male toddler with CdCS presenting for direct laryngoscopy and supraglottoplasty. The patient was a former 27-week, preterm infant with worsening stridor after a history of bronchiolitis 1 month ago. A previous sleep study was significant for an apnea-hypopnea index of 17 and an oxygen saturation nadir of 76%. The patient was receiving home oxygen at night via a nasal cannula at 0.1 L/min. A prior echocardiogram revealed mild pulmonary stenosis. There were no recent acute illnesses or changes in physical status other than a mild increased work of breathing. Previous surgery included circumcision and left eye frontalis sling at 12 months of age to treat congenital ptosis. At that time, airway management including bag-valve-mask ventilation and endotracheal intubation were uneventful. The patient’s trachea was intubated with a 3.5-mm cuffed endotracheal tube (ETT) with a Miller 1 blade and a Cormack-Lehane grade 1 view. Preoperative home medications included budesonide (0.5 mg/2 mL) aerosol twice a day and as needed albuterol/ipratropium for wheezing. Vital signs were normal for age. The craniofacial structures were dysmorphic with abnormal facial features and micrognathia. Cardiovascular examination revealed a 2 - 3/6 systolic murmur. The breath sounds were coarse with transmitted upper airway noise. The patient was held nil per os for 6 h and transferred to the operating room where routine American Society of Anesthesiologists’ monitors were placed. Anesthesia was induced by the inhalation of incremental concentrations of sevoflurane in 50% nitrous oxide/oxygen and a peripheral intravenous cannula was placed. The larynx was sprayed with 1% lidocaine. Direct laryngoscopy and bronchoscopy were performed during deep inhalational anesthesia with sevoflurane while main-
taining spontaneous ventilation. Anesthesia was supplemented with incremental doses of intravenous dexametomidine (3 μg, a total of five doses administered at 10 - 15 min intervals). Following diagnostic airway examination, a supraglottoplasty was performed while continuing sevoflurane anesthesia via insufflation and spontaneous ventilation. The surgical procedure lasted approximately 45 - 60 min. Blood loss was minimal and total fluids included 100 mL of lactated Ringer’s solution. The patient was admitted to the pediatric intensive care unit following the procedure. There was slight worsening of the stridor following the surgical procedure, but this improved overnight. Four doses of dexamethasone were administered postoperatively. On postoperative day 1, the patient was discharged to the inpatient ward. The mild inspiratory stridor and subcostal retractions continued to improve. There were some feeding concerns with a question of aspiration which required a feeding evaluation with occupational therapy and an intervention with thickening of the feedings. The remainder of the postoperative course was uneventful. The patient was discharged home on postoperative day 6 with improved air exchange and decreased stridor.

Discussion

CdCS commonly involves multiple body systems, including the upper airway (micrognathia, laryngeal and epiglottic abnormalities), cardiovascular (CHD), and central nervous system (CNS) (hypotonia) [2, 3]. Morbidity and mortality are greatest during the first year of life, with 75% of deaths occurring in the first month and about 90% occurring within the first year due to pneumonia, aspiration, and congenital heart defects [3]. After one year of age, patients have a high survival rate and life expectancy, with some individuals living into their 50’s and one into their 70’s [3, 7]. Reports show that 47-75% of patients with CdCS will undergo a surgical procedure or diagnostic testing/imaging requiring immobilization, sedation or general anesthesia [2]. The majority of these procedures are performed in the first few years of life [5].

Anesthetic care begins with a thorough preoperative evaluation and identification of end-organ and comorbid involvement. Of particular importance during anesthesia care is the potential for difficulties with airway management due to micrognathia and upper airway abnormalities. As evident in our patient, the characteristic cry is related to the anatomical abnormalities of the larynx. The larynx, epiglottis, and glottis structures are frequently malformed. The larynx is commonly hypoplastic, narrow, and diamond-shaped [8]. The epiglottis may be small, floppy, and/or curved [8]. These anatomical abnormalities of the upper airway combined with micrognathia may lead to problems with airway management and endotracheal intubation as noted by previous reports [2, 9-12]. Although problems with airway management were not encountered with our patient, previous reports describe difficulties involving bag-valve-mask ventilation, upper airway obstruction during the induction of anesthesia, subglottic narrowing with the need to use a smaller than age-appropriate endotracheal tube, limited visualization of the glottic structures requiring video laryngoscopy, laryngeal mask airway placement, or tracheostomy [2, 10-12].

Given these concerns, the appropriate equipment for dealing with the difficult airway including an indirect videolaryngoscope should be readily available prior to anesthetic induction or airway management [13, 14]. If there are concerns regarding airway management, general anesthesia can be induced by the incremental inhalation of sevoflurane with the maintenance of spontaneous ventilation until the airway is secured or adequate bag-valve-mask ventilation is demonstrated.

Abnormal laryngeal and epiglottic anatomy in combination with neuromuscular hypotonia, particularly of the pharyngeal muscles, may lead to intermittent, partial soft tissue airway obstruction during sleep or the administration of sedative, opioid or general anesthetic agents. As demonstrated by polysomnography, our patient was diagnosed with obstructive sleep apnea (OSA). Previous reports have described postoperative respiratory events following anesthetic care in patients with CdCS [2, 12, 15]. Anatomical issues impacting upper airway control and patency may be magnified by the residual effects of anesthetic agents or the ongoing use of opioid analgesics. Furthermore, hypotonia is a frequent comorbid finding in CdCS and may further impact upper airway control [5, 16]. Given these concerns, the use of short acting anesthetic agents should be considered. The risk of perioperative respiratory failure may be increased by pre-existing respiratory dysfunction from hypotonia, poor cough effort, chronic aspiration or recurrent pneumonia. Given the potential for postoperative events related to airway patency and respiratory function, continuous postoperative monitoring of respiratory function is suggested.

Although not required for the procedure performed in our patient, the associated hypotonia may also impact the choice of neuromuscular blocking agents (NMBAs). Patients with pre-existing motor weakness and hypotonia may be sensitive to the effects of non-depolarizing NMBAs and the depolarizing agent, succinylcholine, may be contraindicated. A prolonged effect can be expected even with intermediate-acting non-depolarizing agents (atracurium, rocuronium or vecuronium) [17]. The novel reversal agent, sugammadex, offers the potential to reverse even profound neuromuscular blockade in patients with neuromyopathic conditions [18]. The limited literature available has not demonstrated increased sensitivity to the effects of NMBAs in patients with CdCS, but this possible adverse effect should be considered due to the associated hypotonia in these patients.

Despite the potential lethal side effect of hyperkalemia in susceptible populations and its association with malignant hyperthermia (MH) and MH-like symptoms in patients with neuromuscular disorders, succinylcholine has been administered without adverse effects to patients with CdCS [12, 15, 19, 20]. Given the limited and anecdotal experience of these reports, no definitive conclusions can be drawn regarding the safety of succinylcholine in this patient population. Additionally, a single case report suggests the possible occurrence of MH in a patient with CdCS [21]. The clinical symptoms occurred immediately after the induction of anesthesia with a volatile agent during attempted placement of a laryngeal mask airway. Mask ventilation had become difficult, the teeth were clenched, and the mouth could not be opened. No change was noted after the administration of propofol and succinylcholine. Associated tachycardia and an increase in body temperature from 35.7 to 37.7 °C were noted. The anesthetic was abort-
ed, the clinical signs and symptoms resolved without the administration of dantrolene, and the remainder of his hospital course was uneventful. Many other reports have demonstrated the successful and uneventful use of volatile anesthetic agents thereby limiting the suggestion that there is any concern of MH-susceptibility in this patient population.

A significant percentage (15-20%) of patients with CdCS have CHD (most commonly patent ductus arteriosus, ventricular septal defect, atrial septal defect, and, tetralogy of Fallot) [5]. Although spontaneous deletions result in the majority of CdCS cases, a greater incidence and higher complexity of cardiac anomalies are noted in those with unbalanced translocations [3, 4]. Given the high association of CHD in CdCS, a preoperative echocardiogram is suggested in all patients.

Other health problems associated with CdCS include swallowing and feeding difficulties, poor weight gain and physical growth retardation, recurrent infections such as otitis and pneumonia, orthopedic deformities including scoliosis, and involvement of the CNS (hyperacusis and abnormalities involving the brainstem and cerebellar hypoplasia) [5, 22]. Seizures have been reported in up to 10% of patients [22-24]. Preoperative management includes ensuring therapeutic anticonvulsant levels prior to the surgical procedure. Routine anticonvulsant medications should be administered the morning of the procedure despite concerns of the patient’s nil per os status with subsequent intraoperative dosing as needed [25].

Joint and soft tissue contractures may lead to problems with intraoperative positioning [10, 15, 24]. Growth retardation, nutritional issues, and CNS disabilities may predispose to temperature instability with hypothermia. Thus, care must be taken to reduce the risk of intraoperative hypothermia with continuous intraoperative temperature monitoring and the use of overhead heating lights, forced airway warming, increased room temperature, and heated and humidified inspired gases.

Given the multisystem involvement of CdCS and the likelihood of the need for surgical procedures or diagnostic imaging, anesthetic care is frequently required in this patient population. The preoperative evaluation should seek to identify the associated end-organ involvement. Of particular importance are upper airway and laryngeal abnormalities with the potential for problematic airway management or difficult endotracheal intubation, associated CHD, and CNS involvement including intellectual disability, seizures, and hypotonia.

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Financial Disclosure

None to declare.

Conflict of Interest

None to declare.

Informed Consent

In accordance with the IRB guidelines of Nationwide Children’s Hospital, IRB review and written informed consent are not required.

Author Contributions

Catherine Davis prepared the initial, subsequent, and final drafts; Jonathan Grischkan did the review of final draft, perioperative care of patient; Joseph Tobias did the concept, review of all drafts, postoperative care of patient.

Data Availability

Any inquiries regarding supporting data availability of this study should be directed to the corresponding author.

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