A holistic view of anesthesia-related neurotoxicity in children

Nicola G Clausen
Tom G Hansen
Department of Anesthesia and Intensive Care, Odense University Hospital, Odense, Denmark

Introduction: Animal studies (including in nonhuman primates) have shown that most general anesthetics cause enhanced neuroapoptosis in the immature brain with subsequent long-term neurocognitive deficits later in life. Whether human neurons are equally affected is yet unknown, but a final answer to this issue is still pending. To date, most human studies within the field are of observational nature and the results are conflicting. Some studies indicate an association between exposure to anesthesia and surgery while others do not.

Objective: This review summarizes results from preclinical and observational studies. Controversies and challenges regarding the interpretation of these results are presented. Crucial aspects of neurocognitive safety during pediatric anesthesia and surgery are highlighted. International initiatives aiming to improve the safe conductance of pediatric anesthesia are introduced.

Conclusion: So far, anesthesia-related neurotoxicity in humans remains an area of concern but it cannot be completely excluded. Clinical practice should not be changed until there are definite proofs that anesthetic exposure causes neurocognitive impairment later in life. Withholding necessary and timely surgeries as a consequence of any such concerns could result in worse harm. Focus of current research should also be redirected to include other factors, than merely anesthetics and surgery, that influence the neurocognitive safety of children perioperatively.

Keywords: pediatric anesthesia, neurotoxicity, anesthesia safety, neurocognitive development

Introduction
In recent years, clinicians and parents have inquired whether anesthetic agents may be neurotoxic to the developing human brain.1–7 Animal studies (including in nonhuman primates) have shown that most general anesthetics (GA) cause enhanced neuroapoptosis with subsequent long-term neurocognitive deficits later in life.3,8–10 Some human cohort studies have indicated an association between anesthesia/surgery and adverse neurocognitive outcome, whereas other studies have not.11

Objective
This article summarizes results from preclinical and observational studies on anesthesia-related neurotoxicity. Controversies and challenges regarding the interpretation of these results are presented and aspects of neurocognitive safety during pediatric anesthesia highlighted.

Background
A variety of ion channels scattered throughout the central and peripheral nervous system are sensitive to GA. However, gamma-aminobutyric acid type A (GABA_A)
and glutamate receptors seem to play a pivotal role in facilitating the beneficial state of anesthesia.\textsuperscript{12}

Anesthetics enhance inhibitory postsynaptic ion-channel activity by increasing glycine – and GABA\textsubscript{A} – receptors sensitivity to GABA.\textsuperscript{13,14} GABA is the predominant inhibitory neurotransmitter in the mature brain,\textsuperscript{15} facilitating the influx of chloride ions through opening of GABA\textsubscript{A} channels. This leads to hypersensibilization of the postsynaptic membrane and an overall reduced activity, which is observed as anxiolysis, sedation, amnesia, and anticonvulsion clinically.\textsuperscript{16} Propofol, volatile anesthetics, barbiturates, and benzodiazepines are examples of agents with these properties. In the immature brain, GABA has depolarizing properties, which facilitates the refinement of neuronal circuits early in postnatal development by acting on cell migration, synaptogenesis, DNA synthesis, and cell proliferation.\textsuperscript{17} This excitatory/inhibitory switch depends on the developmental upregulation of the potassium-chloride-cotransporter isoform 2 (KCC2) concordant with the downregulation of the potassium-chloride-cotransporter isoform 1 (NKCC1), which facilitates the net extrusion of intracellular anions in the immature neuron. While studies exposing newborn rats to intravenous anesthetics did not show any influence on the expression on KCC2,\textsuperscript{18} caspase-3 activity was increased in brains of rats receiving sevoflurane without pretreatment with an NKCC1-blocker.\textsuperscript{19} The latter indicates cellular apoptosis as a response to sevoflurane exposure, mediated by GABAergic activation of NKCC1.

Anesthetics inhibit excitatory synaptic channel activity mediated by nicotinic acetylcholine, serotonin, and NMDA (N-methyl-D-aspartate)-sensitive glutamate receptors.\textsuperscript{20,21} Ketamine and nitrous oxide are examples of NMDA-receptor antagonizing drugs.

In the immature brain, exposure to nonphysiologic stressors, eg, drugs, hypoxia, ischemia, and hypoglycemia, at the time of peak synaptogenesis leads to neurodegeneration.\textsuperscript{22} In mice, this period occurs in the early postnatal period, but this period may continue from midgestation to young childhood in humans.\textsuperscript{23} Apoptosis of neurons is part of normal development. Anesthetics have been shown to enhance this process by mechanisms not yet fully understood – but most likely involved are the mitochondria-dependent (intrinsic) and death-receptor mediated (extrinsic) caspase pathways.\textsuperscript{24,25} Neurodevelopment seems to be highly dependent on external stimuli and neuronal trafficking. Hence, exaggerated apoptosis is believed to follow after interference with interneuronal signaling pathways and/or an imbalance between inhibitory and excitatory stimuli.\textsuperscript{26}

**Animal studies**

A growing number of animal studies has demonstrated increased neuronal apoptosis following exposure to GA\textsuperscript{8,9,26–29} In one of the landmark studies, Ikonomidou\textsuperscript{8} exposed rats to NMDA antagonists on day 7 postnatally (PD7). Neurons showed signs of excessive apoptosis, preferentially in the frontal and parietal cortex as well as in the thalamus.\textsuperscript{30} Additional studies have shown that the extent of apoptosis seems to vary between brain areas, suggesting a regional difference in susceptibility to neurotoxins.\textsuperscript{31,32} Cellular structures other than neurons seem to be affected; exposure to volatile anesthetics has been shown to result in altered dendritic spine architecture.\textsuperscript{33,34} Neonatal rhesus macaques exposed to isoflurane showed extensive apoptosis of oligodendrocytes compared to astrocytes, microglia, and interstitial neurons.\textsuperscript{35} How these histopathologic changes relate to neurocognition remains to be resolved, since there is yet no evidence of a causative link. Jevtovic-Todorovic et al\textsuperscript{8} demonstrated impaired learning in rats in the Morris water maze following exposure to midazolam, isoflurane, and nitrous oxide in combination. Most disturbingly, this impaired learning persisted into adulthood. Similar results have been obtained in various animal species, including rhesus monkeys.\textsuperscript{36} In a recent study, rhesus monkeys of both sexes were subject to sevoflurane anesthesia for 4 hours on postnatal days 6–10, and again 14 and 28 days later.\textsuperscript{17} At the age of 6 months, exposed and nonexposed monkeys were tested for their emotional reactivity toward intrusion of a human (human intruder paradigm). The frequency of anxiety-related behavior was higher in exposed than unexposed monkeys, which the authors speculate might reflect long-term effects of anesthesia. In contrast, another study on cynomolgus monkeys, exposing 6-day-old male animals to a similar sevoflurane anesthesia, did not affect their behavior tested by the “holding cage method” when they were tested at 3 and 7 months. Nor were the animals affected in learning or memory.\textsuperscript{38} Although the studies investigated two different subspecies of monkeys, it is not apparent as to why their results point in opposite directions. Overall, it is unknown how any of these findings correlate to the human pediatric population.

**Observational human studies**

So far, a number of observational studies have been published. Some of these studies argue against any association between early exposure to anesthesia and surgery and negative neurocognitive outcome.

In a Dutch study, academic performance and cognition was assessed in 1,143 twin pairs identified in the Young Netherlands’ Twin Register.\textsuperscript{39} Information on exposure to
unspecified surgery and anesthesia was collected by mailed surveys to the parents. Overall, lower equal standardized educational attainment scores and more cognitive problems/inattention as rated by teachers were found among exposed than unexposed twins. Interestingly, the 71 monozygotic twin pairs discordant for exposure showed no difference in performance between the exposed and the unexposed twin. Two Danish nationwide cohort studies comprising the complete birth cohort from 1986 to 1990 assessed, during adolescence, the academic performance of children who underwent surgery for pyloric stenosis repair before 3 months of age and inguinal hernia repair before 1 year of age, respectively. In both studies, the outcome of the exposed children was compared to that of a 5% randomly selected group of unexposed children within the same cohorts. The average mean test score did not differ between children exposed to pyloric stenosis repair and nonexposed controls (mean difference: -0.01; 95% confidence interval [CI]: 0.09–0.08 lower). The same tendency was seen for children undergoing inguinal hernia repair: their estimated mean of test scores was 0.04 below that of the control group (95% CI: 0.01–0.09). However, in both studies, rates of nonattainment were slightly higher among exposed versus nonexposed individuals: after hernia repair the odds ratio for not obtaining test scores was 1.18 (95% CI: 1.04–1.35); in the pyloric stenosis repair group odds ratio was 1.37 (95% CI: 1.11–1.68). Academic performance in children who had spinal anesthesia for inguinal hernia repair, circumcision, and pyloric stenosis repair was compared to nonexposed controls matched by grade, sex, year of testing, and socioeconomic status. On elementary school level, exposure to spinal anesthesia and surgery did not increase the odds for having very poor academic achievement.

On the other hand, some observational studies do suggest adverse neurocognitive outcome following anesthesia and surgical exposure.

In Olmstead County, Minnesota, all children born from 1976 to 1982 were included in a retrospective study investigating the association between general anesthesia for all types of surgeries before the age of 4 and learning disability (LD). Multiple exposures were found to be a significant risk factor for LD, the incidence among exposed individuals at age 19 years being 35.1% (95% CI: 26.2%–42.9%) compared to 20.0% (95% CI: 18.8%–21.3%) for children not exposed at all. Using the same birth cohort, the same group compared the need for individual educational programs and the results in tests of cognition and achievement between children exposed to any kind of surgery before the age of 2 and unexposed controls. Controls were matched for maternal level of education, birth weight, gestational age, and sex, all factors knowingly associated with LD and age.

Again, multiple exposures increased the risk of LD (hazard ratio: 2.16; 95% CI: 1.35–3.46) as well as the need for individual educational programs (hazard ratio: 4.76; 95% CI: 2.48–9.12).44 In a third study within the same birth cohort, the authors investigated the association between exposure to surgery and GA and diagnosis of attention deficit hyperactivity disorder (ADHD): multiple exposures were associated with an increased risk of being diagnosed with ADHD (hazard ratio: 1.95; 95% CI: 1.03–3.71).45

Enrollees in the New York State Medicaid program born from 1999 to 2001 were included in a retrospective cohort analysis. Compared to controls, frequency-matched in age (in months) but not in other parameters, children undergoing GA for inguinal hernia repair before the age of 3 were more than twice as likely to be diagnosed with a developmental or behavioral disorder (hazard ratio: 2.3; 95% CI: 1.3–4.1).46 Controls were randomly selected among children in the same birth cohort and may have been exposed to anesthesia and any other surgery but inguinal hernia repair. In a retrospective twin-sibling study based on children born from 1999 to 2005 and enrolled in the same Medicaid program, exposure to GA for any kind of surgery before the age of 3 increased the risk of behavioral disorders by 1.6 (95% CI: 1.4–1.8). Interestingly, the risk increased from 1.1 (95% CI: 0.8–1.4) for one exposure to 2.9 (94% CI: 2.5, 3.1) for two, and 4.0 (95% CI: 3.5, 4.5) for three or more exposures.47

The Western Australian Pregnancy Cohort (the Raine Study) contains information on 2,868 subjects born between 1989 and 1992.48 Until birth, demographic and medical data were collected on pregnant women who had good English language skills, had planned to deliver in hospital, and expected to stay in Western Australia for the decade to come. Postnatal data were based on parent reporting, such as information on exposures and nonexposures to anesthesia and surgery. Based on this information, children exposed to GA for all types of surgery before the age of 3 were tested neuropsychologically at the age of 10, and their results were compared to outcome in unexposed children within the same cohort. Exposure was associated with an increased risk of poor performance in language (risk ratio: 1.87; 95% CI: 1.12–2.64) and cognition (risk ratio: 1.69; 95% CI: 1.3–2.53). This association persisted with a single exposure to anesthesia.48

In Iowa, Block et al49 compared composite scores in the Iowa test of basic skills and education between the general population and 185 children previously exposed to anesthesia and surgery for circumcision, pyloric stenosis,
or inguinal hernia repair without orchidopexy before the age of 1. Exposed children were identified from the department of anesthesia’s billing records, and data were based on medical records and retrieved after written consent from parents. Within the cohort, a subgroup of 75 children at high risk for cognitive dysfunction (e.g., due to central nervous system disorders) were recorded separately. Compared to the general Iowa population, exposed children had very low achievement test scores (below the fifth percentile), both overall and within the “high-risk cohort”.49

In a recently conducted matched-control study, Backeljauw et al50 assessed academic achievements in the Oral and Written Language scales (OWLS) and Wechsler Performance IQ Intelligence Scale for Children. A total of 53 children with an existing MRI (magnetic resonance imaging) scan of the cerebrum and previous exposure to general anesthesia for all kinds of surgery before the age of 4 were compared to 53 controls identified in the same cross-sectional MRI database. Controls were chosen if they were found neurologically healthy on examination and had no history of neurological or psychiatric illness, head trauma, previous or current LD, or prematurity. Furthermore, they were matched on age, sex, socioeconomic status, and left- or right-handedness. Besides other surgical procedures, all exposed children underwent at least one ear-nose-throat intervention. Whereas mean cognitive test scores were found to be within population norms, the exposed group presented significantly lower scores for performance IQ and OWLS listening comprehension. Furthermore, there was a decrease in gray matter in distinct cortical areas that have previously been associated with impaired cognition and language skills. Due to the retrospective study design, no conclusions about causality can be made. Notably, the frequencies of both the Wechsler performance IQ and OWLS listening comprehension scores for exposed children follow a Gaussian distribution, as opposed to unexposed controls. Although exposed children are matched according to relevant parameters, this could be explained by systematic differences between the study groups, confounding the outcome under investigation. Furthermore, any association between exposure and outcome might be overestimated due to the high occurrence of ear-nose-throat procedures in the exposed group, since this group of patients often presents with impaired language skills and cognition problems.

Table 1 gives an overview of selected observational studies.51-54

### Ongoing clinical studies

Three clinical studies are currently in progress: the Pediatric Anesthesia and Neurodevelopment Assessment Multicenter study (http://www.kidspandastudy.org/index.html), the Mayo Safety for Kids study,55 and the General Anaesthesia compared to Spinal Anaesthesia (GAS) study.56,57 While these studies have the advantage of being prospective and based on cohorts selected for the purpose, they are time-consuming and expensive. Table 2 summarizes the studies’ objectives, study populations, and outcomes.

### Interpretation of current knowledge: challenges and pitfalls

The discrepancy between results in preclinical and observational studies can, to some extent, be explained by many of the impediments of the study designs employed and will now be considered.

Results from animal studies cannot by default be transferred to a human population for the following, but assumingly not exhaustive, reasons:

1. The course of an anesthetic is different in animals compared to a human setting since vital signs and end tidal levels of inhalational gases are rarely monitored, neither are changes in blood glucose, acid–base status, body temperature, and partial pressures of oxygen and carbon dioxide. Furthermore, airways are often not secured and unsupported, spontaneous respiration is maintained, especially in small animals using high concentrations of oxygen. These variables may each and/or in combination influence perfusion and oxygenation of the cerebrum and hence have an impact on neuronal cell function. As a consequence, the isolated effect of the anesthetics cannot be demonstrated.

2. Doses of anesthetics administered and durations of exposures are not analogous to those usually used in clinical pediatric practice; for instance, Paule et al56 found impaired learning in rhesus monkeys after 24 hours of anesthesia induced with up to 50 mg/kg of ketamine. Furthermore, the routes of administration often vary due to the reduced size of the animals, rendering pharmacokinetics and pharmacodynamics of anesthetics unpredictable.

3. Great effort has been put into translating developmental stages of the animal central nervous system into the human corollary (http://www.translatingtime.net), creating a theoretical model based on mathematical algorithms.58 However, these do not account for inter- and intraindividual variations.
Table 1  Retrospective epidemiological studies – an overview stratified according to positive findings and negative findings

<table>
<thead>
<tr>
<th>Study title</th>
<th>Data source</th>
<th>Study subjects</th>
<th>Outcome</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>The effects of exposure to GA in infancy on AP at age 12&lt;sup&gt;14&lt;/sup&gt;</td>
<td>Children born at term between 1998 and 1999 at KK Children’s and Women’s Hospital Singapore</td>
<td>Exposure group: children within the cohort who had surgery and anesthesia for any kind of surgery (n=257) Controls: Children within the cohort without any history of surgery and anesthesia</td>
<td>1. PSLE aggregate score; PSLE is administered by the Ministry of Education. 2. LD – parental reported 3. LD – formally diagnosed Information on anesthetic agents, periprocedural vital parameters and type of surgery retrieved in patient journal</td>
<td>N=100 exposed children were included in analysis and compared to n=107 controls; Adjusted difference in PSLE score between exposed and unexposed not significant. OR for exposed relative to controls of being formally diagnosed with LD 4.5% (95% CI: 1.44–14.4)</td>
</tr>
<tr>
<td>Neurosurgical conditions and procedures in infancy are associated with mortality and APs in adolescence: a nationwide cohort study&lt;sup&gt;51&lt;/sup&gt;</td>
<td>Danish birth cohort 1986–1990; retrospective birth cohort study based on Danish National CPR registry&lt;sup&gt;4&lt;/sup&gt;</td>
<td>Exposed to surgery and anesthesia for the following neurosurgical procedures: n=228 (n=130 hydrocephalus; n=43 cranioectodermal encephalocele); Controls: randomly selected, age-matched 5% sample of the same cohort, n=14,698</td>
<td>Comparison of 1. APs at ninth grade exam 2. Mortality 3. NA at exam between exposed and controls</td>
<td>Average test scores were significantly lower than those of controls in the hydrocephalus and cranioectodermal encephalocele children; Mortality was higher among exposed than controls NA at exam was significantly higher among exposed than controls</td>
</tr>
<tr>
<td>Cognition and brain structure following early childhood surgery with anesthesia&lt;sup&gt;50&lt;/sup&gt;</td>
<td>Matched-control design study; cross-sectional MRI database including 5–18-year-old healthy volunteers; anesthesia records</td>
<td>Exposed n=53 with history of anesthesia and any kind of surgery before 4 years of age and n=53 matched controls</td>
<td>OWLS and Wechsler Performance IQ Intelligence Scale for Children Findings on MRI scans conducted without anesthesia</td>
<td>Lower performance IQ was associated with decreased gray matter in cerebellum; mean cognitive test scores were comparable between exposed and unexposed but subgroup analysis showed significant lower scores among exposed for performance IQ and OWLS-listening comprehension. Single GA exposure not a risk factor for development of LD Multiple GA exposure: significant risk factor for LD</td>
</tr>
<tr>
<td>Early exposure to anesthesia and learning disabilities in a population-based birth cohort&lt;sup&gt;63&lt;/sup&gt;</td>
<td>Population-based retrospective birth cohort study in Olmstead County, Minnesota</td>
<td>593 children born during 1976–1982 and exposed to GA for any kind of surgery &lt; 4 years of age; exposed n=449 (single GA) n=100 (2x GA) n=44 (3x GA)</td>
<td>LD</td>
<td></td>
</tr>
<tr>
<td>Comparative analysis of outcome used in examining neurodevelopmental effects of early childhood anesthesia exposure&lt;sup&gt;52&lt;/sup&gt;</td>
<td>Western Australian Pregnancy Cohort; n=2,868 subjects born 1989–1992 included</td>
<td>Individuals within the cohort exposed to GA for any kind of surgery &lt; 3 years of age and with available records of complete outcome data</td>
<td>At age 10 years: 1. Neuropsychological testing. 2. ICD, ninth revision, clinical modification-coded clinical disorders 3. Academic achievement</td>
<td>Individuals within the cohort with complete outcome data n=781; Exposed to GA &lt; 3 years of age n=112 Unexposed to GA &lt; 3 years of age n=669 Exposed showed 1. Lower scores in neuropsychological testing Increased diagnosis of behavioral and language deficits, cognitive disorders</td>
</tr>
</tbody>
</table>

(Continued)
### Table 1 (Continued)

<table>
<thead>
<tr>
<th>Study title</th>
<th>Data source</th>
<th>Study subjects</th>
<th>Outcome</th>
<th>Results</th>
</tr>
</thead>
</table>
| Cognitive and behavioral outcomes after early exposure to anesthesia and surgery       | Matched cohort study based on birth cohort in Rochester, Minnesota       | n=350 children exposed to any kind of surgery <2 years of age compared to n=700 controls matched for age and known risk factors for LD | Learning disabilities  
Need for IEP  
Results of TCA | Multiple exposures <2 years of age  
1. Increased risk of LD, but not of behavioral problems;  
2. Increased risk of need for IEP  
Incidence of ADHD among nonexposed approximately 7%;  
Increased risk of being diagnosed with ADHD after exposure with ≥2× GA |
| ADHD after early exposure to procedures requiring GA | Retrospective birth cohort study, Rochester, Minnesota | Within the birth cohort of n=5,357 children, n=341 had a diagnosis of ADHD; exposed to surgery and GA prior to 2 years of age, n=350 | Association between ADHD and exposure to surgery and GA <2 years of age | Questionnaires n=243 returned;  
Adjusted OR for presence of clinically deviant outcome when operated  
1. <6 months: 1.38 (95% CI: 0.59–3.22)  
2. 6–12 months: 1.19 (95% CI: 0.45–3.18)  
and – between 12 and 24 months: 1.20 (95% CI: 0.45–3.20) (reference: operated at age >24 months) |
| Behavior and development in children and age at the time of first anesthetic exposure | Children undergoing urological surgery in the years 1987, 1991, 1993, and 1995 at the University Medical Center Utrecht  
Retrospective cohort study | n=314 children operated for 1. Ureteropelvic junction obstruction  
2. Obstructive megaureter  
3. Posterior urethral valves between 0 and 6 years of age;  
A properly powered cohort study would require at least 2,268 children | 120-item parental child behavior checklist/4–18 – a set of measures for assessing children from parent, teacher, and self-report | Increased risk of diagnosis of behavioral development disorders among exposed children  
Compared to a random sample from the same birth cohort, n=5,050 children  
60% increase in BDP among exposed |
| A retrospective cohort study of the association of anesthesia for hernia repair surgery with behavioral and developmental disorders in young children | New York State Medicaid program;  
Retrospective cohort analysis | n=383 children within the birth cohort from 1999 to 2001 and having surgery for inguinal hernia repair <3 years of age | Diagnostic codes for  
1. Delayed development or behavioral disorder  
2. Mental retardation  
3. Autism  
4. Language/speech problems  
Diagnosis of BDP compared to 10,980 unexposed twin siblings in the cohort | Children undergoing anesthesia and surgery during infancy had very low achievement test scores (below the fifth percentile), both in overall sample and the subgroup of 58 patients without CNS problems/potential risk factors |
| Early childhood exposure to anesthesia and risk of developmental and behavioral disorders in a sibling birth cohort | New York State Medicaid program;  
Retrospective twin-sibling study | Twin siblings born during 1999–2005; n=68 exposed to any kind of surgery <3 years of age |  | |
| Are anesthesia and surgery during infancy associated with altered AP during childhood? | Department of Anesthesia Billing Records searched for patients between 7 and 17.9 years on January 28, 2008, who had been operated on for: 1) circumcision;  
2) pyloromyotomy; and 3) inguinal hernia repair and orchiopexy | Of n=623 eligible individuals, n=185 could be included in data analysis;  
Composite scores were available for n=133 individuals; among these n=58 were identified as not having any risk factors for poor AP | Scores on Iowa tests of basic skills and education – (Iowa tests):  
standardized tests assessing basic, general intellectual skills and abilities in verbal, mathematical, and other areas – composite score in second to fourth grade, corresponding to 7–10 years of age  
Results of neuropsychological tests at age 10 years; exposed individuals compared to unexposed (n=2,287) within the same cohort | Individuals exposed to GA <3 years of age performed poorer in areas of receptive, expressive, and total language |
| Long-term differences in language and cognitive function after childhood exposure to anesthesia | Western Australian Pregnancy Cohort containing 2,868 subjects from 1989 to 1992 | Within the cohort, n=321 individuals were exposed to GA for any kind of surgery <3 years of age |  | |
### Negative findings

#### Risk of autistic disorder after exposure to GA and surgery


#### Anesthesia and cognitive performance in children: no evidence for a causal relationship

1,143 monozygotic twin pairs born during 1986–1995; exposed to any surgery and anesthesia from <3 years of age to 3–12 years of age.

#### AP in adolescence after inguinal hernia repair in infancy: a nationwide cohort study

2,689 children in the Danish birth cohort from 1986 to 1990 having surgery for inguinal hernia repair <1 year of age.

#### Cognitive outcome after spinal anesthesia and surgery during infancy

n=365 children born between January 1, 1989 and August 31, 2003 with gestational age 28 weeks or more and before 5 years of age having spinal anesthesia for

1. Pyloromyotomy
2. Inguinal hernia repair
3. Circumcision

#### Educational outcome in adolescence following pyloric stenosis repair before 3 months of age: a nationwide cohort study

779 infants in the Danish birth cohort from 1986 to 1990 having surgery for pyloric stenosis repair <3 months of age.

#### Academic results

1. New Standards Reference Examination achievement tests in reading, writing, mathematics (assessed in fourth, eighth, tenth grades) or
2. New England Common Assessment Program examination in grades three to eight and eleven and
3. Need for individual educational plan

#### Comparison

Compared to 14,665 controls randomly selected in the birth cohort, exposed children showed

1. No significant difference in AP
2. Higher risk of NA among exposed

#### Duration of surgery in spinal anesthesia did not relate to very poor academic achievement in elementary school

Duration of surgery in spinal anesthesia did not relate to poor academic outcome; Exposure to spinal anesthesia did not relate to very poor academic achievement in elementary school.

---

**Note:** CPR, a Danish national register, holding individual numbers assigned at birth.

**Abbreviations:** GA, general anesthesia; ADHD, attention deficit hyperactivity disorder; OR, odds ratio; EA, educational achievement in Dutch CITO elementary test; CP, cognitive problems; LD, learning disability; BDP, behavioral or developmental problems; AP, academic performance; NA, nonattainment; PSLE, primary school leaving examination; VIRS, Vermont infant’s spinal registry; IEP, individual education program; ICD, international classification of diseases; TCA, test of cognition and achievement; NHiRD, national health insurance research database; MRI, magnetic resonance imaging; OWLS, oral and written language scales; CPR, central person register; CNS, central nervous system; CITO-test, final test primary education in the Netherlands.
Table 2 Ongoing prospective trials

<table>
<thead>
<tr>
<th>Study name</th>
<th>Design</th>
<th>Cohort</th>
<th>Primary outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>GAS study</td>
<td>Multisite randomized controlled trial</td>
<td>Newborns randomized to spinal or GA for inguinal hernia repair</td>
<td>IQ score at age 2 and 5 years in WPPSI-III</td>
</tr>
<tr>
<td>MASK study</td>
<td>Cohort study, retrospective regarding exposure, prospective regarding outcome</td>
<td>Children born between 1994 and 2007 exposed to single or multiple GA before 3 years of age compared to controls from the same cohort</td>
<td>Results in single 4-hour neuropsychological test battery</td>
</tr>
<tr>
<td>PANDA study</td>
<td>Ambidirectional cohort study</td>
<td>Sibling exposed to GA before 3 years of age for inguinal hernia repair compared to nonexposed sibling</td>
<td>WASI-II scores and NEPSY II scores between 8 and 15 years of age</td>
</tr>
</tbody>
</table>

Note: A multisite randomized controlled trial comparing regional and general anesthesia for effects on neurodevelopmental outcome and apnea in infants.


4. In an animal setting, anesthesia is conducted solely for the purpose of the experiment. Hence, anesthesia is conducted without any concurrent surgical trauma or other types of insults. There are data suggesting that anesthetics under such conditions might be neuroprotective rather than neurotoxic. McAuliffe et al exposed 9-day-old mice to isoflurane, desflurane, sevoflurane, or room air for 3 hours. The next day the animals had 60 minutes of hypoxia-ischemia. Histological sections did not show any difference between the groups regarding neuronal injury. But those animals preconditioned with a volatile agent performed better in behavioral testing than animals preconditioned with room air alone. With regard to some parts of the tests, exposed animals even performed equal to the sham group not exposed at all. Similarly, both short-term structural and long-term functional neuroprotection has been demonstrated when volatile anesthetics were administered to 10-day-old mice after induction of brain ischemia. In a randomized study on piglets having cardiac pulmonary bypass surgery, postoperative neurologic outcome was improved among animals anesthetized with desflurane compared to animals allocated to the administration of fentanyl–droperidol. This exemplifies anesthetic neuroprotection in a setting, where the trauma of surgery is superimposed to a state of reduced cerebral blood flow. The mechanism behind the neuroprotective properties of volatile anesthetics is not fully understood. However, in an animal setting, volatile anesthetics have been shown to reduce cerebral blood flow less than intravenous sedatives. Since the underlying condition, the impact of surgery itself, and the potential neurotoxic effects of anesthetics are intertwined in a human setting, observational studies are prone to “confounding by indication”. This adds to the overall lack of control of confounders.

5. It is unknown how both short-term and long-term toxic damage to neurons will present themselves clinically. Any consequence is likely to depend on both the age and neurodevelopmental stage at exposure and at time of follow-up. Individual variations in neurodevelopmental progress cannot be taken into account. Based on findings in animal studies, it is assumed that brain areas responsible for learning and memory are affected intensively by anesthetics. Hence, learning difficulties, academic performance in standardized examinations, and behavioral disorders have been used as estimates of function in these areas. However, outcomes assessed in observational studies in order to investigate neurotoxicity were constructed for other purposes: school grades aim to reflect certain skills achieved through comprehension of teaching and learning contents communicated both verbally and in writing; codes of behavioral and psychiatric disorders intend to apprehend pathological conditions, which is also true for neuropsychological test batteries. It is assumed, not known, that they function as acceptable measures of clinically relevant neurotoxic effects. Moreover, many of these tests are interrelated. Increasing the number of tests used in a study increases the risk for type 1 statistical error.

6. Cohort studies based on administrative cohorts are sensitive to selection; the included individuals might not be representative of overall populations, thus weakening the generalizability of results. Similarly, losses to follow-up might under- or overestimate an association under investigation. The persons lost to death and migration might have one or more features in common that enhance or mitigate an association.

Discussion

If anesthesia-related neurotoxicity exists in humans, many additional answers are urgently needed: Who is at risk?
Is this an age-associated phenomenon? Which dosages and which agents cause greatest damage? What is the impact of surgery and diagnosis? How does any such potential damage present itself? Which study designs are most likely to answer these questions?

Observational studies are retrospective and based on data that were selected for other purposes, ie, administration. While this does make data more prone to selection bias and in some instances reduces the generalizability of results, they are still valuable and feasible. An observational study design will never demonstrate a causative association, but can rather illustrate which issues are important. This requires either a large cohort or a strong association between exposure and outcome. The inconsistent results of observational data so far suggest that an association between anesthesia and surgery and neurocognitive impairment is either minor or hidden behind confounding factors. Multiple studies have shown the underlying disease and/or surgery, prematurity, sex, and parental level of education to have higher impact on outcome than anesthesia itself. McCann and Schouten have recently reviewed the impact of blood pressure and perioperative cerebral perfusion on neurodevelopment in ex-prematures and infants. Infants have less cerebral autoregulatory reserve rendering them more vulnerable to hypotension, and prone to hypocapnia-induced cerebral ischemia – both these factors may contribute to the development of hypoxic-ischemic encephalopathy. In its mild form, hypoxic-ischemic encephalopathy is characterized by postoperative irritability, poor feeding, excessive crying or sleepiness, or even seizures in more severe cases.

For future research, it must be taken into account that many factors other than exposure to anesthetic drugs contribute to impaired neurodevelopment in young children exposed to anesthesia and surgery. Introducing “The 10N’s”, Weiss et al emphasizes ten factors of importance for the safe conductance of anesthesia: absence of pain and fear, normotension, normocardiya, normooxemia, normocarbia, normothermia, normovolemia, normonatremia, and normoglycemia. This multifactorial approach is the cornerstone of the safe anesthesia for every tot initiative (SAFETOTS). The initiative aims to increase focus on safe conductance of pediatric anesthesia and define the safe use of anesthetics in the pediatric population (http://www.safetots.org). Within this framework, two studies are currently in progress: Anaesthesia PRactice In Children Observational Trial (APRICOT) and NEonate-Children sTudy of Anaesthesia pRactice IN Europe (NECTARINE).

APRICOT, a prospective multicenter observational study, investigates the incidence of severe critical events in children undergoing anesthesia in Europe. From April 1, 2014 to December 31, 2014, participating centers in Europe registered variables concerning the pre-, peri- and post-anesthesia process. Data are currently being analyzed and results are expected in 2016 (ClinicalTrials.gov website: NCT01878760).

As an extension of the APRICOT study, NECTARINE, a prospective, observational multicenter audit, will provide information on morbidity and mortality related to neonatal anesthesia. Over a 12-week observation period, beginning on March 1, 2016, data on 5,000 patients in European centers will be registered (ClinicalTrials.gov website: NCT02350348).

According to a recent web-based survey among practicing European anesthetists, the majority consider neurotoxicity an important topic. Two-third of the anesthetists reported that they had changed their clinical practice in an attempt to reduce any potential harm. Based on current knowledge, a change in practice is unfounded and should be balanced against the risks related to withholding necessary surgery.

Summary
Histologic changes in neurons and long-term neurocognitive impairments due to exposure to anesthetics are well documented in animal studies. Results from human observational studies are less clear and, due to inconsistent study designs and varying measures of outcome, difficult to compare. Anesthesia-related neurotoxicity can neither be excluded nor verified based on these findings, and the significance of this issue for the children requiring surgery and anesthesia worldwide remains unknown at present. A change in clinical practice cannot be recommended at this point.

Results from the few ongoing randomized trials are still awaited and will add to current knowledge rather than completely resolve this complex issue. Future studies will broaden their search for factors other than anesthetics that have the potential of impairing neurodevelopment of infants.

Disclosure
The authors report no conflicts of interest in this work.

References


37. Raper J, Alvarado MC, Murphy KL, Baxter MG. Multiple anesthetic exposure in infant monkeys alters emotional reactivity to an acute stressor. Anesthesiology. Epub 2015 Sep 2.


