

Pain and Its Effects in the Human Neonate and Fetus

Author: Anand, K J S; Hickey, P R

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Full Text: The evaluation of pain in the human fetus and neonate is difficult because pain is generally defined as a subjective phenomenon.¹ Early studies of neurologic development concluded that neonatal responses to painful stimuli were decorticate in nature and that perception or localization of pain was not present.² Furthermore, because neonates may not have memories of painful experiences, they were not thought capable of interpreting pain in a manner similar to that of adults.³⁵ On a theoretical basis, it was also argued that a high threshold of painful stimuli may be adaptive in protecting infants from pain during birth.⁶ These traditional views have led to a widespread belief in the medical community that the human neonate or fetus may not be capable of perceiving pain.^{7,8} Strictly speaking, nociceptive activity, rather than pain, should be discussed with regard to the neonate, because pain is a sensation with strong emotional associations. The focus on pain perception in neonates and confusion over its differentiation from nociceptive activity and the accompanying physiologic responses have obscured the mounting evidence that nociception is important in the biology of the neonate. This is true regardless of any philosophical view on consciousness and "pain perception" in newborns. In the literature, terms relating to pain and nociception are used interchangeably; in this review, no further distinction between the two will generally be made. One result of the pervasive view of neonatal pain is that newborns are frequently not given analgesic or anesthetic agents during invasive procedures, including surgery.⁹⁻¹⁹ Despite recommendations to the contrary in textbooks on pediatric anesthesiology, the clinical practice of inducing minimal or no anesthesia in newborns, particularly if they are premature, is widespread.⁹⁻¹⁹ Unfortunately, recommendations on neonatal anesthesia are made without reference to recent data about the development of perceptual mechanisms of pain and the physiologic responses to nociceptive activity in preterm and full-term neonates. Even Robinson and Gregory's landmark paper demonstrating the safety of narcotic anesthesia in preterm neonates cites "philosophic objections" rather than any physiologic rationale as a basis for using this technique.²⁰ Although methodologic and other issues related to the study of pain in neonates have been discussed,^{21,23} the body of scientific evidence regarding the mechanisms and effects of nociceptive activity in newborn infants has not been addressed directly.

ANATOMICAL AND FUNCTIONAL REQUIREMENTS FOR PAIN PERCEPTION

The neural pathways for pain may be traced from sensory receptors in the skin to sensory areas in the cerebral cortex of newborn infants. The density of nociceptive nerve endings in the skin of newborns is similar to or greater than that in adult skin.²⁴ Cutaneous sensory receptors appear in the perioral area of the human fetus in the 7th week of gestation; they spread to the rest of the face, the palms of the hands, and the soles of the feet by the 11th week, to the trunk and proximal parts of arms and legs by the 15th week, and to all cutaneous and mucous surfaces by the 20th week.²⁵⁻²⁶ The spread of cutaneous receptors is preceded by the development of synapses between sensory fibers and interneurons in the dorsal horn of the spinal cord, which first appear during the sixth week of gestation.²⁷⁻²⁸ Recent studies using electron microscopy and immunocytochemical methods show that the development of various types of cells in the dorsal horn (along with their laminar arrangement, synaptic interconnections, and specific neurotransmitter vesicles) begins before 13 to 14 weeks of gestation and is completed by 30 weeks.²⁹ Lack of myelination has been proposed as an index of the lack of maturity in the neonatal nervous system³⁰ and is used frequently to support the argument that premature or full-term neonates are not capable of pain perception.⁹⁻¹⁹ However, even in the peripheral nerves of adults, nociceptive impulses are carried through unmyelinated (C-polymodal) and thinly myelinated (A-delta) fibers.³¹ Incomplete myelination merely implies a slower conduction velocity in the nerves

or central nerve tracts of neonates, which is offset completely by the shorter interneuron and neuromuscular distances traveled by the impulse.³² Moreover, quantitative neuroanatomical data have shown that nociceptive nerve tracts in the spinal cord and central nervous system undergo complete myelination during the second and third trimesters of gestation. Pain pathways to the brain stem and thalamus are completely myelinated by 30 weeks; whereas the thalamocortical pain fibers in the posterior limb of the internal capsule and corona radiata are myelinated by 37 weeks.³³ Development of the fetal neocortex begins at eight weeks of gestation, and by 20 weeks each cortex has a full complement of 109 neurons.³⁴ The dendritic processes of cortical neurons undergo profuse arborization and develop synaptic targets for the incoming thalamocortical fibers and intracortical connections.^{35,36} The timing of the thalamocortical connection is of critical importance for cortical perception, since most sensory pathways to the neocortex have synapses in the thalamus. Studies of primate and human fetuses have shown that different neurons in the thalamus produce axons that arrived in the cerebrum before mid-gestation. These fibers then "wait" just below the neocortex until migration and dendritic arborization of cortical neurons are complete and finally establish synaptic connections between 20 and 24 weeks of gestation (Fig. 1).³⁶⁻³⁸

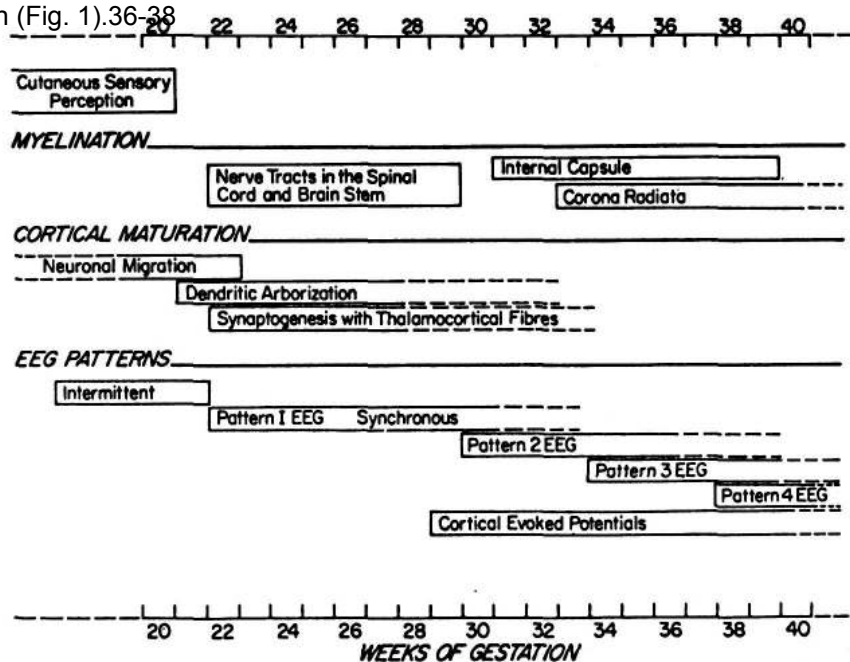


Figure 1. Schematic Diagram of the Development of Cutaneous Sensory Perception,²⁵ Myelination of the Pain Pathways,³² Maturation of the Fetal Neocortex,³³⁻³⁷ and Electroencephalographic Patterns³⁸⁻⁴⁰ in the Human Fetus and Neonate.

Functional maturity of the cerebral cortex is suggested by fetal and neonatal electroencephalographic patterns, studies of cerebral metabolism, and the behavioral development of neonates. First, intermittent electroencephalographic bursts in both cerebral hemispheres are first seen at 20 weeks' gestation; they become sustained at 22 weeks and bilaterally synchronous at 26 to 27 weeks.³⁹ By 30 weeks, the distinction between wakefulness and sleep can be made on the basis of electroencephalographic patterns.³⁹⁻⁴⁰ Cortical components of visual and auditory evoked potentials have been recorded in preterm babies (born earlier than 30 weeks of gestation),⁴⁰⁻⁴¹ whereas olfactory and tactile stimuli may also cause detectable changes in electroencephalograms of neonates.⁴⁰⁻⁴² Second, in vivo measurements of cerebral glucose utilization have shown that maximal metabolic activity is located in sensory areas of the brain in neonates (the sensorimotor cortex, thalamus, and midbrain-brain-stem regions), further suggesting the functional maturity of these regions.⁴³ Third, several forms of behavior imply cortical function during fetal life. Well-defined periods of quiet sleep, active sleep, and wakefulness occur in utero beginning at 28 weeks of gestation.⁴⁴ In addition to the specific behavioral responses to pain described below, preterm and full-term babies have various cognitive,

coordinative, and associative capabilities in response to visual and auditory stimuli, leaving no doubt about the presence of cortical function.⁴⁵ Several lines of evidence suggest that the complete nervous system is active during prenatal development and that detrimental or developmental changes in any part would affect the entire system.^{25-26,42,46} In studies in animals, Ralston found that somatosensory neurons of the neocortex respond to peripheral noxious stimuli and proposed that "it does not appear necessary to postulate a subcortical mechanism for appreciation of pain [in the fetus or neonate]."⁴⁷ Thus, human newborns do have the anatomical and functional components required for the perception of painful stimuli. Since these stimuli may undergo selective transmissions, inhibition, or modulation by various neurotransmitters, the neurochemical mechanisms associated with pain pathways in the fetus and newborn are considered below.

NEUROCHEMICAL SYSTEMS ASSOCIATED WITH PAIN PERCEPTION

The Tachykinin System

Various putative neurotransmitters called the tachykinins (substance P, neurokinin A, neuromedin K, and so forth) have been identified in the central nervous system, but only substance P has been investigated thoroughly and shown to have a role in the transmission and control of pain impulses.⁴⁸⁻⁵⁶ Neural elements containing substance P and its receptors appear in the dorsal-root ganglia and dorsal horns of the spinal cords at 12 to 16 weeks of gestation.⁵⁷ A high density of substance P fibers and cells has been observed in multiple areas of the fetal brain stem associated with the pathways for pain perception and control and the visceral reactions to pain.⁵⁸⁻⁶³ Substance P fibers and cells have also been found in the hypothalamus, mamillary bodies, thalamus, and cerebral cortex of human fetuses early in their development.⁵⁸ Many studies have found higher densities of substance P and its receptors in neonates than in adults of the same species, although the importance of this finding is unclear.^{61,64-68}

The Endogenous Opioid System

With the demonstration of the existence of stereospecific opiate receptors⁶⁹⁻⁷⁰ and their endogenous ligands,⁷¹ the control of pain was suggested as a primary role for the endogenous opioid system.⁷² Both the enkephalinergic and the endorphinergic systems may modulate pain transmission at spinal and supraspinal levels.^{56,73} In the human fetus, however, there are no data on the ontogeny and distribution of specific cells, fibers, and receptors (μ -, δ -, and κ -opiate receptors) that are thought to mediate the antinociceptive effects of exogenous and endogenous opioids.⁷⁴ However, functionally mature endorphinergic cells in fetal pituitary glands have been observed at 15 weeks of gestation and possibly earlier.⁷⁴⁻⁷⁶ Beta-endorphin and betalipotropin were found to be secreted from fetal pituitary cells at 20 weeks in response to in vitro stimulation by corticotropin-releasing factor.⁷⁷ In addition, more production of beta-endorphin may occur in fetal and neonatal pituitary glands than in adult glands.^{78,79} Endogenous opioids are released in the human fetus at birth and in response to fetal and neonatal distress.⁸⁰ Umbilical-cord plasma levels of beta-endorphin and beta-lipotropin from healthy full-term neonates delivered vaginally or by caesarean section have been shown to be three to five times higher than plasma levels in resting adults.⁷⁸⁻⁸¹ Neonates delivered vaginally by breech presentation or vacuum extraction had further increases in beta-endorphin levels, indicating beta-endorphin secretion in response to stress at birth.⁸² Plasma beta-endorphin concentrations correlated negatively with umbilical-artery pH and partial pressure of oxygen and positively with base deficit and partial pressure of carbon dioxide suggesting that birth asphyxia may be a potent stimulus to the release of endogenous opioids.^{81,83-87} Cerebrospinal fluid levels of beta-endorphin were also increased markedly in newborns with apnea of prematurity,⁸⁸⁻⁹⁰ infections, or hypoxemia.^{83,91,92} These elevated values may have been caused by the "stress" of illness,⁹³ the pain associated with these clinical conditions, or the invasive procedures required for their treatment. However, these high levels of beta-endorphin are unlikely to decrease anesthetic or analgesic requirements,⁹⁴ because the cerebrospinal fluid levels of beta-endorphin required to produce analgesia in human adults have been found to be 10,000 times higher than the highest recorded levels in neonates.⁹⁵ The high levels of beta-endorphin and beta-lipotropin in cord plasma decreased substantially by 24 hours after birth.^{87,96} and reached adult levels by five days, whereas the levels in the cerebrospinal fluid fell to adult values in 24 hours.^{87,97,98} In newborn infants of women addicted to narcotics, massive increases in plasma concentrations of beta-endorphin, beta-lipotropin, and met-enkephalin occurred within 24 hours, with

some values reaching 1000 times those in resting adults. Markedly increased levels persisted for up to 40 days after birth.⁸⁷ However, these neonates were considered to be clinically normal, and no behavioral effects were observed (probably because of the development of prenatal opiate tolerance).

PHYSIOLOGIC CHANGES ASSOCIATED WITH PAIN

Cardiorespiratory Changes

Changes in cardiovascular variables, transcutaneous partial pressure of oxygen, and palmar sweating have been observed in neonates undergoing painful clinical procedures. In preterm and full-term neonates undergoing circumcision^{99,100} or heel lancing,¹⁰¹⁻¹⁰³ marked increases in the heart rate and blood pressure occurred during and after the procedure. The magnitude of changes in the heart rate was related to the intensity and duration of the stimulus¹⁰⁴ and to the individual temperaments of the babies.¹⁰⁵ The administration of local anesthesia to full-term neonates undergoing circumcision prevented the changes in heart rate and blood pressure,⁹⁹⁻¹⁰⁰⁻¹⁰⁶ whereas giving a "pacifier" to preterm neonates during heel-stick procedures did not alter their cardiovascular or respiratory responses to pain.¹⁰¹ Further studies in newborn and older infants showed that noxious stimuli were associated with an increase in heart rate, whereas non-noxious stimuli (which elicited the attention or orientation of infants) caused a decrease in heart rate.^{22,107,108} Large fluctuations in transcutaneous partial pressure of oxygen above and below an arbitrary "safe" range of 50 to 100 mm Hg have been observed during various surgical procedures in neonates.¹⁰⁹⁻¹¹¹ Marked decreases in transcutaneous partial pressure of oxygen also occurred during circumcision,^{106,112} but such changes were prevented in neonates given local analgesic agents,^{100,106,112} Tracheal intubation in awake preterm and full-term neonates caused a significant decrease in transcutaneous partial pressure of oxygen, together with increases in arterial blood pressure¹¹³⁻¹¹⁵ and intracranial pressure.¹¹⁶ The increases in intracranial pressure with intubation were abolished in preterm neonates who were anesthetized.¹¹⁷ In addition, infants' cardiovascular responses to tracheal suctioning were abolished by opiate-induced analgesia.¹¹⁸ Palmar sweating has also been validated as a physiologic measure of the emotional state in full-term babies and has been closely related to their state of arousal and crying activity.¹¹⁹ Substantial changes in palmar sweating were observed in neonates undergoing heel-sticks for blood sampling, and subsequently, a mechanical method of heel lancing proved to be less painful than manual methods, on the basis of the amount of palmar sweating.¹²⁰

Hormonal and Metabolic Changes

Hormonal and metabolic changes have been measured primarily in neonates undergoing surgery, although there are limited data on the neonatal responses to venipuncture and other minor procedures. Plasma renin activity increased significantly five minutes after venipuncture in full-term neonates and returned to basal levels 60 minutes thereafter; no changes occurred in the plasma levels of Cortisol, epinephrine, or norepinephrine after venipuncture.¹²¹ In preterm neonates receiving ventilation therapy, chest physiotherapy and endotracheal suctioning produced significant increases in plasma epinephrine and norepinephrine; this response was decreased in sedated infants.¹²² In neonates undergoing circumcision without anesthesia, plasma Cortisol levels increased markedly during and after the procedure.^{123,124} Similar changes in Cortisol levels were not inhibited in a small number of neonates given a local anesthetic,¹²⁵ but the efficacy of the nerve block was questionable in these cases. Further detailed hormonal studies¹²⁶ in preterm and full-term neonates who underwent surgery under minimal anesthesia documented a marked release of catecholamines,¹²⁷ growth hormone,¹²⁸ glucagon,¹²⁷ Cortisol, aldosterone, and other corticosteroids,^{129,130} as well as suppression of insulin secretion.¹³¹ These responses resulted in the breakdown of carbohydrate and fat stores,^{127,132,133} leading to severe and prolonged hyperglycemia and marked increases in blood lactate, pyruvate, total ketone bodies, and nonesterified fatty acids. Increased protein breakdown was documented during and after surgery by changes in plasma amino acids, elevated nitrogen excretion, and increased 3-methylhistidine:creatinine ratios in the urine (Anand KJS, Aynsley-Green A: unpublished data). Marked differences also occurred between the stress responses of premature and full-term neonates (Anand KJS, Aynsley-Green A: unpublished data) and between the responses of neonates undergoing different degrees of surgical stress.¹³⁴ Possibly because of the lack of deep anesthesia, neonatal stress responses were found to be three to five times greater than those in adults,

although the duration was shorter.¹²⁶ These stress responses could be inhibited by potent anesthetics, as demonstrated by randomized, controlled trials of halothane and fentanyl. These trials showed that endocrine and metabolic stress responses were decreased by halothane anesthesia in full-term neonates¹³⁵ and abolished by low-dose fentanyl anesthesia in preterm neonates.¹³⁶ The stress response of neonates undergoing cardiac surgery were also decreased in randomized trials of high-dose fentanyl and sufentanil anesthesia.^{126,137,138} These results indicated that the nociceptive stimuli during surgery performed with minimal anesthesia were responsible for the massive stress responses of neonates. Neonates who were given potent anesthetics in these randomized trials were more clinically stable during surgery and had fewer postoperative complications as compared with neonates under minimal anesthesia.^{126,129} There is preliminary evidence that the pathologic stress responses of neonates under light anesthesia during major cardiac surgery may be associated with an increased postoperative morbidity and mortality (Anand KJS, Hickey PR: unpublished data). Changes in plasma stress hormones (e.g., Cortisol) can also be correlated with the behavioral states of newborn infants,^{124,139,140} which are important in the postulation of overt subjective distress in neonates responding to pain.

BEHAVIORAL CHANGES ASSOCIATED WITH PAIN

Simple Motor Responses

Early studies of the motor responses of newborn infants to pinpricks reported that the babies responded with a "diffuse body movement" rather than a purposeful withdrawal of the limb,² whereas other studies found reflex withdrawal to be the most common response.¹⁴¹⁻¹⁴³ More recently, the motor responses of 124 healthy full-term neonates to a pinprick in the leg were reported to be flexion and adduction of the upper and lower limbs associated with grimacing, crying, or both, and these responses were subsequently quantified.^{144,145} Similar responses have also been documented in very premature neonates, and in a recent study, Fitzgerald et al. found that premature neonates (<30 weeks) not only had lower thresholds for a flexor response but also had increased sensitization after repeated stimulation.¹⁴⁶

Facial Expressions

Distinct facial expressions are associated with pleasure, pain, sadness, and surprise in infants.¹⁴⁷ These expressions, especially those associated with pain, have been objectively classified and validated in a study of infants being immunized.^{102,148} With use of another method of objectively classifying facial expressions of neonates, different responses were observed with different techniques of heel lancing and with different behavioral states¹⁴⁹ (and Grunau RVE, Craig KD: unpublished data). These findings suggest that the neonatal response to pain is complex and may be altered by the behavioral state and other factors at the time of the stimulus.¹⁵⁰

Crying

Crying is the primary method of communication in newborn infants and is also elicited by stimuli other than pain.¹⁵¹ Several studies have classified infant crying according to the type of distress indicated and its spectrographic properties.^{152,154} These studies have shown that cries due to pain, hunger, or fear can be distinguished reliably by the subjective evaluation of trained observers and by spectrographic analysis.¹⁵⁵⁻¹⁶⁰ This has allowed the cry response to be used as a measure of pain in numerous recent studies.^{22,99,100,102,106,152} The pain cry has specific behavioral characteristics and spectrographic properties in healthy full-term neonates.¹⁶¹⁻¹⁶⁴ Pain cries of preterm neonates and neonates with neurologic impairment, hyperbilirubinemia, or meningitis are considerably different, thereby indicating altered cortical function in these babies.¹⁶⁵⁻¹⁶⁸ Changes in the patterns of neonatal cries have been correlated with the intensity of pain experienced during circumcision and were accurately differentiated by adult listeners.¹⁶⁹ In other studies of the cry response to painful procedures, neonates were found to be more sensitive to pain than older infants (those 3 to 12 months old) but had similar latency periods between exposure to a painful stimulus and crying or another motor response.^{99-100,103,152,170} This supports the contention that slower conduction speed in the nerves of neonates is offset by the smaller interneuron distances traveled by the impulse.

Complex Behavioral Responses

Alterations in complex behavior and sleep-wake cycles have been studied mainly in newborn infants undergoing circumcision without anesthesia. Emde and coworkers observed that painful procedures were followed by prolonged periods of non-rapid-eye-movement sleep in newborns and confirmed these observations in a controlled study of neonates undergoing circumcision without anesthesia.¹⁷¹

Similar observations have been made in adults with prolonged stress. Other subsequent studies have found increased wakefulness and irritability for an hour after circumcision, an altered arousal level in circumcised male infants as compared with female and uncircumcised male infants, and an altered sleep-wake state in neonates undergoing heelstick procedures.^{103,172,173} In a double-blind, randomized controlled study using the Brazelton Neonatal Behavioral Assessment Scale, 90 percent of neonates had changed behavioral states for more than 22 hours after circumcision, whereas only 16 percent of the uncircumcised infants did.¹⁷⁴ It was therefore proposed that such painful procedures may have prolonged effects on the neurologic and psychosocial development of neonates.¹⁷⁵ A similar randomized study showed the absence of these behavioral changes in neonates given local anesthetics for circumcision.¹⁷⁶ For two days after circumcision, neonates who had received anesthetics were more attentive to various stimuli and had greater orientation, better motor responses, decreased irritability, and a greater ability to quiet themselves when disturbed. A recent controlled study showed that intervention designed to decrease the amount of sensory input and the intensity of stressful stimuli during intensive care of preterm neonates was associated with improved clinical and developmental outcomes.¹⁷⁷ Because of their social validity and communicational specificity, the behavioral responses observed suggest that the neonatal response to pain is not just a reflex response.¹⁷⁸⁻¹⁸⁰

MEMORY OF PAIN IN NEONATES The persistence of specific behavioral changes after circumcision in neonates implies the presence of memory. In the short term, these behavioral changes may disrupt the adaptation of newborn infants to their postnatal environment,¹⁷⁴⁻¹⁷⁶ the development of parent-infant bonding, and feeding schedules.¹⁸¹⁻¹⁸² In the long term, painful experiences in neonates could possibly lead to psychological sequelae,²² since several workers have shown that newborns may have a much greater capacity for memory than was previously thought.¹⁸³⁻¹⁸⁶ Pain itself cannot be remembered, even by adults¹⁸⁷; only the experiences associated with pain can be recalled. However, the question of memory is important, since it has been argued that memory traces are necessary for the "maturation" of pain perception,³ and a painful experience may not be deemed important if it is not remembered. Long-term memory requires the functional integrity of the limbic system and diencephalon (specifically, the hippocampus, amygdala, anterior and mediodorsal thalamic nuclei, and mamillary nuclei)¹⁸⁸; these structures are well developed and functioning during the newborn period.⁴² Furthermore, the cellular, synaptic, and molecular changes required for memory and learning depend on brain plasticity, which is known to be highest during the late prenatal and neonatal periods.^{189,190} Apart from excellent studies in animals demonstrating the long-term effects of sensory experiences in neonatal period,¹⁹¹ evidence for memories of pain in human infants must, by necessity, be anecdotal.^{178,192,193} Early painful experiences may be stored in the phylogenically old "procedural memory," which is not accessible to conscious recall.¹⁸²⁻¹⁸³⁻¹⁹⁴ Although Janov¹⁹⁵ and Holden¹⁹⁶ have collected clinical data that they claim indicate that adult neuroses or psychosomatic illnesses may have their origins in painful memories acquired during infancy or even neonatal life, their findings have not been substantiated or widely accepted by other workers.

CONCLUSIONS Numerous lines of evidence suggest that even in the human fetus, pain pathways as well as cortical and subcortical centers necessary for pain perception are well developed late in gestation, and the neurochemical systems now known to be associated with pain transmission and modulation are intact and functional. Physiologic responses to painful stimuli have been well documented in neonates of various gestational ages and are reflected in hormonal, metabolic, and cardiorespiratory changes similar to but greater than those observed in adult subjects. Other responses in newborn infants are suggestive of integrated emotional and behavioral responses to pain and are retained in memory long enough to modify subsequent behavior patterns. None of the data cited herein tell us whether neonatal nociceptive activity and associated responses are experienced subjectively by the neonate as pain similar to that experienced by older children and adults. However, the evidence does show that marked nociceptive activity clearly constitutes a physiologic and perhaps even a psychological form of stress in premature or full-term neonates. Attenuation of the deleterious effects of pathologic neonatal stress responses by the use of various anesthetic techniques has

now been demonstrated. Recent editorials addressing these issues have promulgated a wide range of opinions, without reviewing all the available evidence. The evidence summarized in this paper provides a physiologic rationale for evaluating the risks of sedation, analgesia, local anesthesia, or general anesthesia during invasive procedures in neonates and young infants. Like persons caring for patients of other ages, those caring for neonates must evaluate the risks and benefits of using analgesic and anesthetic techniques in individual patients. However, in decisions about the use of these techniques, current knowledge suggests that humane considerations should apply as forcefully to the care of neonates and young, nonverbal infants as they do to children and adults in similar painful and stressful situations. References

- REFERENCE NOTES
1. Merskey H, Albe-Fessard DG, Bonica JJ, et al. Pain terms: a list with definitions and notes on usage: recommended by the IASP Subcommittee on Taxonomy. *Pain* 1979; 6:249-52.
 2. McGraw MD. The neuromuscular maturation of the human infant. New York: Columbia University Press, 1943.
 3. Merskey H. On the development of pain. *Headache* 1970; 10:116-23.
 4. Levy DM. The infant's earliest memory of inoculation: a contribution to public health procedures. *J Gen Psychol* 1960; 96:3-46.
 5. Harris FC, Lahey BB. A method for combining occurrence and nonoccurrence interobserver agreement scores. *J Appl Behav Anal* 1978; 11:523-7.
 6. Bondy AS. Infancy. In: Gabel S, Erickson MT, eds. *Child development and developmental disabilities*. Boston: Little, Brown, 1980:3-19.
 7. Eland JM, Anderson JE. The experience of pain in children. In: Jacox AK, ed. *Pain: a source book for nurses and other health professionals*. Boston: Little, Brown, 1977:453-73.
 8. Wallerstein E. Circumcision: the uniquely American medical enigma. *Urol Clin N Am* 1985; 12:123-32.
 9. Anand KJS, Aynsley-Green A. Metabolic and endocrine effects of surgical ligation of patent ductus arteriosus in the human preterm neonate: Are there implications for further improvement of postoperative outcome? *Mod Probl Paediatr* 1985; 23:143-57.
 10. Lippmann N, Nelson RJ, Emmanouilides GC, Diskin J, Thibeault DW. Ligation of patent ductus arteriosus in premature infants. *Br J Anaesth* 1976; 48:365-9.
 11. Shaw EA. Neonatal anaesthesia. *Hosp Update* 1982; 8:423-34.
 12. Katz J. The question of circumcision. *Int Surg* 1977; 62:490-2.
 13. Swafford LI, Allan D. Pain relief in the pediatric patient. *Med Clin North Am* 1968; 52:131-6.
 14. Rees GJ. Anesthesia in the newborn. *Br Med J* 1950; 2:1419-22.
 15. Betts EK, Downes JJ. Anesthetic considerations in newborn surgery. *Semin Anesth* 1984; 3:59-74.
 16. Inkster JS. Paediatric anaesthesia and intensive care. *Int Anesthesiol Clin* 1978; 16:58-91.
 17. Norman EA. Pulse oximetry during repair of congenital diaphragmatic hernia. *Br J Anaesth* 1986; 58:934-5.
 18. Hatch DJ. Analgesia in the neonate. *Br Med J* 1987; 294:920.
 19. Shearer MH. Surgery on the paralysed, unanesthetized newborn. *Birth* 1986; 13:79.
 20. Robinson S, Gregory GA. Fentanyl-air-oxygen anesthesia for ligation of patent ductus arteriosus in preterm infants. *Anesth Analg* 1981; 60:331-4.
 21. Weiss C. Does circumcision of the newborn require an anesthetic? *Clin Pediatr (Phila)* 1968; 7:128-9.
 22. Owens ME. Pain in infancy: conceptual and methodological issues. *Pain* 1984; 20:213-30.
 23. Richards T. Can a fetus feel pain? *Br Med J* 1985; 291:1220-1.
 24. Gleiss J, Stuttgen G. Morphologic and functional development of the skin. In: Stave U, ed. *Physiology of the perinatal period*. Vol. 2. New York: Appleton-Century-Crofts, 1970:889-906.
 25. Humphrey T. Some correlations between the appearance of human fetal reflexes and the development of the nervous system. *Prog Brain Res* 1964; 4:93-135.
 26. Valman HB, Pearson JF. What the fetus feels. *Br Med J* 1980; 280:223-4.
 27. Okado N. Onset of synapse formation in the human spinal cord. *J Comp Neurol* 1981; 201:211-9.
 28. Wozniak W, O'Tahilly R, Olszewska B. The fine structure of the spinal cord in human embryos and early fetuses. *J Hirnforsch* 1980; 21:101-24.
 29. Rizvi T, Wadhwa S, Bijilani V. Development of spinal substrate for nociception. *Pain [Suppl]* 1987; 4:195.
 30. Tilney F, Rosett J. The value of brain lipoids as an index of brain development. *Bull Neurol Inst NY* 1931; 1:28-71.
 31. Schulte FJ. Neurophysiological aspects of brain development. *Mead Johnson Symp Perinat Dev Med* 1975; 6:32-47.
 32. Idem. Gestation, wachstum und hirnentwicklung. In: Linneweh F, ed. *Fortschritte der Paedologie*. Vol. 2. Berlin: Springer-Verlag, 1968:46-64.
 33. Gilles FJ, Shankle W., Dooling EC. Myelinated tracts: growth pattern. In: Gilles FH, Leviton A., Dooling EC, eds. *The developing human brain: growth and epidemiologic neuropathology*. Boston: John Wright, 1983:117-83.
 34. Marin-Padilla M. Structural organization of the human

cerebral cortex prior to the appearance of the cortical plate. *Anat Embryol (Berl)* 1983; 168:21-40. 35. Molliver ME, Kostovic I, Van der Loos H. The development of synapses in cerebral cortex of the human fetus. *Brain Res* 1973; 50:403-7. 36. Rakic P, Goldman-Rakic PS. Development and modifiability of the cerebral cortex: early developmental effects: cell lineages, acquisition of neuronal positions, and areal and laminar development. *Neurosci Res Prog Bull* 1982; 20:433-51. 37. Kostovic I, Rakic P. Development of prestriate visual projections in the monkey and human fetal cerebrum revealed by transient cholinesterase staining. *J Neurosci* 1984; 4:25-42. 38. Kostovic I, Goldman-Rakic PS. Transient cholinesterase staining in the mediodorsal nucleus of the thalamus and its connections in the developing human and monkey brain. *J Comp Neurol* 1983; 219:431-47. 39. Spehlmann R. In: *EEG primer*. New York: Elsevier/North-Holland, 1981:159-65. 40. Torres F, Anderson C. The normal EEG of the human newborn. *J Clin Neurophysiol* 1985; 2:89-103. 41. Henderson-Smart DJ, Pettigrew AG, Campbell DJ. Clinical apnea and brain-stem neural function in preterm infants. *N Engl J Med* 1983; 308:353-7. 42. Prechtl HFR, ed. *Continuity of neural functions from prenatal to postnatal life*. Oxford: Blackwell, 1984. 43. Chugani HT, Phelps M. Maturation changes in cerebral function in infants determined by ¹⁸F-FDG positron emission tomography. *Science* 1986; 231:840-3. 44. Arduini D, Rizzo G, Giorlandino C, Valensise H, Dell'acqua S, Romanini C. The development of fetal behavioural states: A longitudinal study. *Prenat Diagn* 1986; 6:117-24. 45. Sammons WAH. Premature behavior and the neonatal intensive care unit environment. In: Cloherty JP, Stark AR, eds. *Manual of neonatal care*. Boston: Little, Brown, 1980:359-63. 46. Flower MJ. Neuromaturation of the human fetus. *J Med Philos* 1985; 10:237-51. 47. Ralston HJ. Synaptic organization of spinothalamic projections to the thalamus, with special reference to pain. *Adv Pain Res Ther* 1984; 6:183-95. 48. Nawa H, Hirose T, Takashima H, Inayama S, Nakanishi S. Nucleotide sequences of cloned cDNAs for two types of bovine brain substance P precursor. *Nature* 1983; 306:32-6. 49. Watson SP, Sandberg BEB, Hanley MR, Iversen LL. Tissue selectivity of substance P alkyl esters: suggesting multiple receptors. *Eur J Pharmacol* 1983; 87:77-84. 50. Mantyh PW, Maggio JE, Hunt SP. The autoradiographic distribution of kassinin and substance K binding sites is different from the distribution of substance P binding sites in rat brain. *Eur J Pharmacol* 1984; 102:361-4. 51. Valentino KL, Tatemoto K, Hunter J, Barchas JD. Distribution of neuropeptide K-immunoreactivity in the rat central nervous system. *Peptides* 1986; 7:1043-59. 52. Pernow B. Substance P. *Pharmacol Rev* 1983; 35:85-141. 53. Otsuka M, Konishi S. Substance P-the first peptide neurotransmitter? *Trends Neurosci* 1983; 6:317-20. 54. Henry JL. Relation of substance P to pain transmission: neurophysiological evidence. In: Porter R, O'Connor M, eds. *Substance P in the nervous system*, Ciba Foundation Symposium 91. Loudon: Pitman, 1982:206-24. 55. Pearson J, Brandeis L, Cuello AC. Depletion of substance P-containing axons in substantia gelatinosa of patients with diminished pain sensitivity. *Nature* 1982; 295:61-3. 56. Jessel T, Iversen LL. Opiate analgesics inhibit substance P release from rat trigeminal nucleus. *Nature* 1977; 268:549-51. 57. Charnay Y, Paulin C, Chayvialle J-A, Dubois PM. Distribution of substance P-like immunoreactivity in the spinal cord and dorsal root ganglia of the human foetus and infant. *Neuroscience* 1983; 10:41-55. 58. Paulin C, Charnay Y, Dubois PM, Chayvialle J-A, Localisation de substance P dans le systeme nerveux du foetus humain: resultats preliminaires. *C R Acad Sci Paris [SERIES D]* 1980; 291:257-60. 59. Pickel VM, Sumal KK, Reis DJ, Miller RJ, Hervonen A. Immunocytochemical localization of enkephalin and substance P in the dorsal tegmental nuclei in the human fetal brain. *J Comp Neurol* 1980; 193:805-14. 60. Roizen MF, Newfield P, Eger EI II, Hosobuchi Y, Adams JE, Lamb S. Reduced anesthetic requirement after electrical stimulation of periaqueductal gray matter. *Anesthesiology* 1985; 62:120-3. 61. Del Fiacco M, Dessi ML, Leranti MC. Topographical localization of substance P in the human post-mortem brainstem: an immunohistochemical study in the newborn and adult tissue. *Neuroscience* 1984; 12:591-611. 62. Nomura H, Shiosaka S, Inagaki S, et al. Distribution of substance P-like immunoreactivity in the lower brainstem of the human fetus: an immunohistochemical study. *Brain Res* 1982; 252:315-25. 63. Helke CA, Charlton CG, Keeler JR. Bulbosplinal substance P and sympathetic regulation of the cardiovascular system: a review. *Peptides* 1985; 6:Suppl 2:69-74. 64. Inagaki S, Sakanaka M, Shiosaka S, et al. Ontogeny of substance P-containing neuron system of the

rat: immunohistochemical analysis. *Neuroscience* 1982; 7:251-77, 1097-126. 65. Quirion R, Dam T-V. Ontogeny of substance P receptor binding sites in rat brain. *J Neurosci* 1986; 6:2187-99. 66. Jonsson G, Hallman H. Substance P counteracts neurotoxin damage on norepinephrine neurons in rat brain during ontogeny. *Science* 1982; 215:75-7. 67. Idem. Effect of substance P on neonatally axotomized noradrenaline neurons in rat brain. *Med Biol* 1983; 61:179-85. 68. Narumi S, Fujita T. Stimulatory effects of substance P and nerve growth factor (NGF) on neurite outgrowth in embryonic chick dorsal root ganglia. *Neuropharmacology* 1978; 17:73-6. 69. Pert CB, Snyder SH. Opiate receptor: demonstration in nervous tissue. *Science* 1973; 179:1011-4. 70. Terenius L. Stereospecific interaction between narcotic analgesics and a synaptic plasma membrane fraction of rat cerebral cortex. *Acta Pharmacol Toxicol (Copenh)* 1973; 32:317-20. 71. Hughes J. Isolation of an endogenous compound from the brain with pharmacological properties similar to morphine. *Brain Res* 1975; 88:295-308. 72. Jacob JJC, Ramabadran K. Role of opiate receptors and endogenous ligands in nociception. In: Williams NE, Wilson H, eds. *Pain and its management*. Oxford: Pergamon Press, 1983:13-32. 73. Hosobuchi Y, Li CH. The analgesic activity of human beta-endorphin in man. *Commun Psychopharmacol* 1978; 2:33-7. 74. Paterson D J, Robson LE, Kosterlitz HW. Classification of opioid receptors. *Br Med Bull* 1983; 39:31-6. 75. Begeot M, Dubois MP, Dubois PM. Immunologic localization of α and β -endorphins and β -lipotropin in corticotropic cells of the normal and anencephalic fetal pituitaries. *Cell Tissue Res* 1978; 193:413-22. 76. Li JY, Dubois MP, Dubois PM. Ultrastructural localization of immunoreactive corticotropin, β -lipotropin, α - and β -endorphin in cells of the human fetal anterior pituitary. *Cell Tissue Res* 1979; 204:37-51. 77. Gibbs DM, Stewart RD, Liu JH, Vale W, Rivier J, Yen SSC. Effects of synthetic corticotropin-releasing factor and dopamine on the release of immunoreactive β -endorphin/ β -lipotropin and α -melanocyte-stimulating hormone from human fetal pituitaries in vitro. *J Clin Endocrinol Metab* 1982; 55:1149-52. 78. Csontos K, Rust M, Höllt V, Mahr W, Kromer W, Teschemacher HJ. Elevated plasma β -endorphin levels in pregnant women and their neonates. *Life Sci* 1979; 25:835-44. 79. Vuolteenaho O, Leppaluoto J, Hoyhtya M, Hirvonen J. β -endorphin-like peptides in autopsy pituitaries from adults, neonates and fetuses. *Acta Endocrinol (Copenh)* 1983; 102:27-34. 80. Gautray JP, Jolivet A, Vielh JP, Guillemin R. Presence of immunoassayable β -endorphin in human amniotic fluid: elevation in cases of fetal distress. *Am J Obstet Gynecol* 1977; 129:211-2. 81. Wardlaw SL, Stark RI, Baxi L, Frantz AG. Plasma β -endorphin and β -lipotropin in the human fetus at delivery: correlation with arterial pH and pO_2 . *J Clin Endocrinol Metab* 1979; 49:888-91. 82. Puolakka J, Kauppila A, Leppaluoto J, Vuolteenaho O. Elevated beta-endorphin immunoreactivity in umbilical cord blood after complicated delivery. *Acta Obstet Gynecol Scand* 1982; 61:513-4. 83. Shaaban MM, Hung TT, Hoffman DI, Lobo RA, Goebelsmann U. β -endorphin and β -lipotropin concentrations in umbilical cord blood. *Am J Obstet Gynecol* 1982; 144:560-9. 84. Browning AJF, Butt WR, Lynch SS, Shakespear RA, Crawford JS. Maternal and cord plasma concentrations of β -lipotropin, β -endorphin and λ -lipotropin at delivery: effect of analgesia. *Br J Obstet Gynaecol* 1983; 90:1152-6. 85. Pohjavuori M, Rovamo L, Laatikainen T. Plasma immunoreactive β -endorphin and Cortisol in the newborn infant after elective caesarean section and after spontaneous labour. *Eur J Obstet Gynecol Reprod Biol* 1985; 19:67-74. 86. Pohjavuori M, Rovamo L, Laatikainen T, Kariniemi V, Pettersson J. Stress of delivery and plasma endorphins and catecholamines in the newborn infant. *Biol Res Pregnancy Perinatol* 1986; 7:1-5. 87. Panerai AE, Martini A, Di Giulio AM, et al. Plasma β -endorphin, β -lipotropin, and met-enkephalin concentrations during pregnancy in normal and drug-addicted women and their newborn. *J Clin Endocrinol Metab* 1983; 57:537-43. 88. MacDonald MG, Moss IR, Kefale GG, Ginzburg HM, Fink RJ, Chin L. Effect of naltrexone on apnea of prematurity and on plasma beta-endorphin-like immunoreactivity. *Dev Pharmacol Ther* 1986; 9:301-9. 89. Orłowski JP. Cerebrospinal fluid endorphins and the infant apnea syndrome. *Pediatrics* 1986; 78:233-7. 90. Sankaran K, Hindmarsh KW, Watson VG. Plasma beta-endorphin concentration in infants with apneic spells. *Am J Perinatol* 1984; 1:331-4. 91. Hindmarsh KW, Sankaran K, Watson VG. Plasma beta-endorphin concentrations in neonates associated with acute stress. *Dev Pharmacol Ther* 1984; 7:198-204. 92. Sankaran K, Hindmarsh KW, Watson VG. Hypoxic-ischemic encephalopathy and plasma β -endorphin. *Dev Pharmacol*

Ther 1984; 7:377-83. 93. Hindmarsh KW, Sankaran K. Endorphins and the neonate. *Can Med Assoc J* 1985; 132:331-4. 94. Lerman J, Robinson S, Willis MM, Gregory GA. Anesthetic requirements for halothane in young children 0-1 month and 1-6 months of age. *Anesthesiology* 1983; 59:421-4. 95. Foley KM, Kourides IA, Inturrisi CE, et al. β -endorphin: analgesic and hormonal effects in humans. *Proc Natl Acad Sci USA* 1979; 76:5377-81. 96. Facchinetti F, Bagnoli F, Bracci R, Genazzani AR. Plasma opioids in the first hours of life. *Pediatr Res* 1982; 16:95-8. 97. Moss IR, Conner H, Yee WFH, Iorio P, Scarpelli EM. Human β -endorphin-like immunoreactivity in the perinatal/neonatal period. *J Pediatr* 1982; 101:443-6. 98. Burnard ED, Todd DA, John E, Hindmarsh KW. Beta-endorphin levels in newborn cerebrospinal fluid. *Aust Paediatr J* 1982; 18:258-63. 99. Williamson PS, Williamson ML. Physiologic stress reduction by a local anesthetic during newborn circumcision. *Pediatrics* 1983; 71:36-40. 100. Holve RL, Bromberger BJ, Groverman HD, Klauber MR, Dixon SD, Snyder JM. Regional anesthesia during newborn circumcision: effect on infant pain response. *Clin Pediatr (Phila)* 1983; 22:813-8. 101. Owens ME, Todt EH. Pain in infancy: neonatal reaction to a heel lance. *Pain* 1984; 20:77-86. 102. Johnson CC, Strada ME. Acute pain response in infants: a multidimensional description. *Pain* 1986; 24:373-82. 103. Field T, Goldson E. Pacifying effects of nonnutritive sucking on term and preterm neonates during heelstick procedures. *Pediatrics* 1984; 74:1012-5. 104. Clifton RK, Graham FK, Hatton HM. Newborn heart-rate response and response habituation as a function of stimulus duration. *J Exp Child Psychol* 1968; 6:265-78. 105. Kagan J. Heart rate and heart rate variability as signs of a temperamental dimension in infants. In: Izard CE, ed. *Measuring emotions in infants and children*. Cambridge: Cambridge University Press, 1982; 38-66. 106. Maxwell LG, Yaster M, Wetzel RC. Penile nerve block reduces the physiologic stress of newborn circumcision. *Anesthesiology* 1986; 65:A432. abstract. 107. Berg KM, Berg WK, Graham FK. Infant heart rate response as a function of stimulus and state. *Psychophysiology* 1971; 8:30-44. 108. Campos JJ. Heart rate: a sensitive tool for the study of emotional development in the infant. In: Lipsitt LD, ed. *Developmental psychobiology*. Hillsdale, J.J.: Lawrence Erlbaum Associates, 1976:1-31. 109. Welle P, Hayden W, Millter T. Continuous measurement of transcutaneous oxygen tension of neonates under general anesthesia. *J Pediatr Surg* 1980; 15:257-60. 110. Venus B, Patel KC, Pratap KS, Konchigeri H, Vidyasagar D. Transcutaneous P02 monitoring during pediatric surgery. *Crit Care Med* 1981; 9:714-6. 111. Messner JT, Loux PC, Grossman LB. Intraoperative transcutaneous p02 monitoring in infants. *Anesthesiology* 1979; 51:S319. abstract. 112. Rawlings DJ, Miller PA, Engel RR. The effect of circumcision on transcutaneous p02 in term infants. *Am J Dis Child* 1980; 134:676-8. 113. Kelly MA, Finer NN. Nasotracheal intubation in the neonate: physiologic responses and effects of atropine and pancuronium. *J Pediatr* 1984; 105:303-9. 114. Marshall TA, Deeder R, Pai S, Berkowitz GP, Austin TL. Physiologic changes associated with endotracheal intubation in preterm infants. *Crit Care Med* 1984; 12:501-3. 115. Gibbons PA, Swedlow DB. Changes in oxygen saturation during elective tracheal intubation in infants. *Anesth Analg* 1986; 65:S58. abstract. 116. Raju TNK, Vidyasagar D, Torres C, Grundy D, Bennett EJ. Intracranial pressure during intubation and anesthesia in infants. *J Pediatr* 1980; 96:860-2. 117. Friesen RH, Honda AT, Thieme RE. Changes in anterior fontanel pressure in preterm neonates during tracheal intubation. *Anesth Aialg* 1987; 66:874-8. 118. Hickey PR, Hansen DD, Wessel DL, Lang P, Jonas RA, Elixson EM. Blunting of stress responses in the pulmonary circulation of infants by fentanyl. *Anesth Analg* 1985; 64:1137-42. 119. Harpin VA, Rutter N. Development of emotional sweating in the newborn infant. *Arch Dis Child* 1982; 57:691-5. 120. Idem. Making heel pricks less painful. *Arch Dis Child* 1983; 58:226-8. 121. Fiselier T, Monnens L, Moerman E, Van Munster P, Jansen M, Peer P. Influence of the stress of venepuncture on basal levels of plasma renin activity in infants and children. *Int J Pediatr Nephrol* 1983; 4:181-5. 122. Greisen G, Frederiksen PS, Hertel J, Christensen NJ. Catecholamine response to chest physiotherapy and endotracheal suctioning in preterm infants. *Acta Paediatr Scand* 1985; 74:525-9. 123. Talbert LM, Kraybill EN, Potter HD. Adrenal cortical response to circumcision in the neonate. *Obstet Gynecol* 1976; 48:208-10. 124. Gunnar MR, Fisch RO, Korsvik S, Donhowe JM. The effects of circumcision on serum Cortisol and behavior. *Psychoneuroendocrinology* 1981; 6:269-75. 125. Williamson PS, Evans ND. Neonatal Cortisol response to circumcision with anesthesia. *Clin*

Pediatr (Phila) 1986; 25:412-5. 126. Anand KJS. Hormonal and metabolic functions of neonates and infants undergoing surgery. *Curr Opin Cardiol* 1986; 1:681-9. 127. Anand KJS, Brown MJ, Bloom SR, Aynsley-Green A. Studies on the hormonal regulation of fuel metabolism in the human newborn infant undergoing anaesthesia and surgery. *Horm Res* 1985; 22:115-28. 128. Milne EMG, Elliott MJ, Pearson DT, Holden MP, Orskov H, Alberti KGMM. The effect on intermediary metabolism of open-heart surgery with deep hypothermia and circulatory arrest in infants of less than 10 kilograms body weight. *Perfusion* 1986; 1:29-40. 129. Obara H, Sugiyama D, Maekawa N, et al. Plasma Cortisol levels in paediatric anaesthesia. *Can Anaesth Soc J* 1984; 31:24-7. 130. Srinivasan G, Jain R, Pildes RS, Kannan CR. Glucose homeostasis during anesthesia and surgery in infants. *J Pediatr Surg* 1986; 21:718-21. 131. Anand KJS, Brown MJ, Causon RC, Christofides ND, Bloom SR, Aynsley-Green A. Can the human neonate mount an endocrine and metabolic response to surgery? *J Pediatr Surg* 1985; 20:41-8. 132. Pinter A. The metabolic effects of anesthesia and surgery in the newborn infant: changes in the blood levels of glucose, plasma free fatty acids, a amino-nitrogen, plasma amino-acid ratio and lactate in the neonate. *Z Kinderchir* 1973; 12:149-62. 133. Elphick MC, Wilkinson AW. The effects of starvation and surgical injury on the plasma levels of glucose, free fatty acids, and neutral lipids in newborn babies suffering from various congenital anomalies. *Pediatr Res* 1981; 15:313-8. 134. Anand KJS, Aynsley-Green A. Measuring the severity of surgical stress in newborn infants. *J Pediatr Surg* (in press). 135. Idem, Does the newborn infant require anesthesia during surgery? Answers from a randomised trial of halothane anesthesia. *Pain Res Clin Manage* (in press.) 136. Anand KJS, Sippell WG, Aynsley-Green A. Randomised trial of fentanyl anaesthesia in preterm neonates undergoing surgery: effects on the stress response. *Lancet* 1987; 1:243-8. 137. Anand KJS, Carr DB, Hickey PR. Randomized trial of high-dose sufentanil anesthesia in neonates undergoing cardiac surgery: hormonal and hemodynamic stress responses. *Anesthesiology* 1987; 67:A501. abstract. 138. Anand KJS, Hickey PR. Randomized trial of high-dose sufentanil anesthesia in neonates undergoing cardiac surgery: effects on the metabolic stress response. *Anesthesiology* 1987; 67:A502. abstract. 139. Anders TF, Sachar EJ, Kream J, Roffwarg HP, Hellman L. Behavioral state and plasma Cortisol response in the human newborn. *Pediatrics* 1970; 46:532-7. 140. Tennes K, Carter D. Plasma Cortisol levels and behavioral states in early infancy. *Psychosom Med* 1973; 35:121-8. 141. Lipsitt LP, Levy N. Electrocutaneous threshold in the neonate. *Child Dev* 1959; 30:547-54. 142. Dockeray FC, Rice C. Response of newborn infants to pain stimulation. *Ohio State Univ Stud Contrib Psychol* 1934; 12:82-93. 143. Sherman M, Sherman IC. Sensori-motor response in infants. *J Comp Psychol* 1925; 5:53-68. 144. Rich EC, Marshall RE, Volpe JJ. The normal neonatal response to pinprick. *Dev Med Child Neurol* 1974; 16:432-4. 145. Franck LS. A new method to quantitatively describe pain behavior in infants. *Nurs Res* 1986; 35:28-31. 146. Fitzgerald M, Shaw A, Macintosh N. The postnatal development of the cutaneous flexor reflex: a comparative study in premature infants and newborn rat pups. *Dev Med Child Neurol* (in press). 147. Ekman P, Oster H. Facial expressions of emotion. *Annu Rev Psychol* 1979; 30:527-54. 148. Izard CE, Huebner RR, Risser D, McGinnes GC, Dougherty LM. The young infant's ability to produce discrete emotional expressions. *Dev Psychol* 1980; 16:132-40. 149. Grunau RVE, Craig KD. Pain expression in neonates: facial action and cry. *Pain* 1987; 28:395-410. 150. Melzack R, Wall PD. Pain mechanisms: a new theory. *Science* 1965; 150:971-9. 151. Lester BM. A biosocial model of infant crying. In: Lipsitt L, ed. *Advances in infancy research*. New York: Ablex, 1984:167-212. 152. Levine JD, Gordon NC. Pain in prelingual children and its evaluation by pain-induced vocalisation. *Pain* 1982; 14:85-93. 153. Wasz-Hockert O, Lind J, Vuorenkoski V. The infant cry: a spectrographic and auditory analysis. *Clin Dev Med* 1968; 2:9-42. 154. Michelsson K, Raes J, Thoden C-J, Wasz-Hockert O. Sound spectrographic cry analysis in neonatal diagnostics: an evaluative study. *J Phonetics* 1982; 10:79-88. 155. Zeskind PL, Sale J, Maio ML, Huntington L, Weiseman JR. Adult perceptions of pain and hunger cries: a synchrony of arousal. *Child Dev* 1985; 56:549-54. 156. Boukydis CFZ. Perception of infant crying as an interpersonal event. In: Lester BM, Boukydis CFZ, eds. *Infant crying: theoretical and research perspectives*. New York: Plenum Press, 1985:187-215. 157. Murry T, Amundson P, Hollien H. Acoustical characteristics of infant cries: fundamental frequency. *J Child Lang* 1977; 3:321-8. 158.

Wasz-Hockert O, Partanen T, Vuorenkoski V, Valanne E, Michelsson K. Effect of training on ability to identify preverbal vocalizations. *Dev Med Child Neurol* 1964; 6:393-6. 159. Gladding ST. Effects of training versus non-training in identification of cry-signals: a longitudinal study. *Percept Mot Skills* 1979; 48:752-4. 160. Johnston CC, O'Shaughnessy D. Acoustical attributes of infant pain cries: discriminating features. *Pain* 1987; Suppl 4:233. 161. Wolff PH. The natural history of crying and other vocalizations in early infancy. In: Foss BM, ed. *Determinants of infant behaviour*. Vol. 4 London: Methuen, 1969:88-295. 162. Wasz-Hockert O, Michelsson K, Lind J. Twenty-five years of Scandinavian cry research. In: Lester BM, Boukydis CFZ, eds. *Infant crying: theoretical and research perspectives*. New York: Plenum Press, 1985; 83-104. 163. Michelsson K, Jarvenpaa A-L, Rinne A. Sound spectrographic analysis of pain cry in preterm infants. *Early Hum Dev* 1983; 8:141-9. 164. Friedman SL, Zahn-Waxler C, Radke-Yarrow M. Perceptions of cries of full-term and preterm infants. *Infant Behav Dev* 1982; 5:161-73. 165. Michelsson K, Sirvio P, Wasz-Hockert O. Pain cry in full-term asphyxiated newborn infants correlated with late findings. *Acta Paediatr Scand* 1977; 66:611-6. 166. Fisichelli VR, Coxe M, Rosenfeld L, Haber A, Davis J, Karelitz S. The phonetic content of the cries of normal infants and those with brain damage. *J Psychol* 1966; 64:119-26. 167. Wasz-Hockert O, Koivisto M, Vuorenkoski V, Partanen TJ, Lind J. Spectrographs analysis of pain cry in hyperbilirubinemia. *Biol Neonate* 1971; 17:260-71. 168. Michelsson K, Sirvio P, Wasz-Hockert O. Sound spectrographs cry analyses of infants with bacterial meningitis. *Dev Med Child Neurol* 1977; 19:309-15. 169. Porter FL, Miller RH, Marshall RE. Neonatal pain cries: effect of circumcision on acoustic features and perceived urgency. *Child Dev* 1986; 57:790-802. 170. Fisichelli VR, Karelitz S, Fisichelli RM, Cooper J. The course of induced crying activity in the first year of life. *Pediatr Res* 1974; 8:921-8. 171. Emde RN, Harmon RJ, Metcalf D, Koenig KL, Wagonfeld S. Stress and neonatal sleep. *Psychosom Med* 1971; 33:491-7. 172. Anders TF, Chalemian RJ. The effects of circumcision on sleep-wake states in human neonates. *Psychosom Med* 1974; 36:174-9. 173. Brackbill Y. Continuous stimulation and arousal level in infancy: effects of stimulus intensity and stress. *Child Dev* 1975; 46:364-9. 174. Marshall RE, Stratton WC, Moore JA, Boxerman SB. Circumcision. I. Effects upon newborn behaviour. *Infant Behav Dev* 1980; 3:1-14. 175. Richards MPM, Bernal JF, Brackbill Y. Early behavioral differences: gender or circumcision? *Dev Psychobiol* 1976; 9:89-95. 176. Dixon S, Snyder J, Holve R, Bromberger P. Behavioural effects of circumcision with and without anesthesia. *J Dev Behav Pediatr* 1984; 5:246-50. 177. Als H, Lawhon G, Brown E, et al. Individualized behavioral and environmental care for the very low birth weight preterm infant at high risk for bronchopulmonary dysplasia: neonatal intensive care unit and developmental outcome. *Pediatrics* 1986; 78:1123-32. 178. Darwin C. *The expression of the emotions in man and animals*. London: John Murry 1872:65-7. 179. Kazdin AE. Assessing the clinical or applied importance of behavior change through social validation. *Behav Modif* 1977; 1:427-52. 180. D'Apolito K. The neonate's response to pain. *Am J Matern Child Nurs* 1984; 9:256-8. 181. Marshall RE, Porter FL, Rogers AG, Moore JA, Anderson B, Boxerman SB. Circumcision II. Effects upon mother-infant interaction. *Early Hum Dev* 1982; 7:367-74. 182. Osofsky JD. Neonatal characteristics and mother-infant interaction in two observational situations. *Child Dev* 1976; 47:1138-47. 183. Lipsitt LP. The study of sensory and learning processes of the newborn. *Clin Perinatol* 1977; 4:163-86. 184. Stone LJ, Smith H, Murphy LB, eds. *The competent infant: research and commentary*. New York: Basic Books, 1973. 185. Moscovitch M. *Infant memory: its relation to normal and pathological memory in humans and other animals*. New York: Plenum Press, 1984. 186. Kolata G. Early signs of school age IQ. *Science* 1987; 236:774-5. 187. Jones E. Pain. *Int J Psychoanal* 1957; 38:255. 188. Squire LR. Mechanisms of memory. *Science* 1986; 232:1612-9. 189. Will B, Schmitt P, Dalrymple-Alford J. Brain plasticity, learning and memory: historical background and conceptual perspectives. *Adv Behav Biol* 1985; 28:1-11. 190. Bischof H-J. Influence of developmental factors on imprinting. *Adv Behav Biol* 1985; 28:51-9. 191. Fillion TJ, Blass EM. Infantile experience with suckling odors determines adult sexual behavior in male rats. *Science* 1986; 231:729-31. 192. Wachter-Shikora NL. Pain theories and their relevance to the pediatric population. *Issues Compr Pediatr Nurs* 1981; 5:321-6. 193. Dale JC. A multidimensional study of infants' responses to painful stimuli. *Pediatr Nurs*

1986; 12:27-31. 194. Reynolds OE, Hutchins HC. Reduction of central hyper-irritability following block anesthesia of peripheral nerve. Am J Physiol 1948; 152:658-62. 195. Janov A. The anatomy of mental illness. New York: Putnam's Sons, 1971. 196. Holden EM, Primal pathophysiology. J Psychosom Res 1977; 21:341-50. abstract. 197. Hatch DJ. Analgesia in the neonate. Br Med J 1987; 294:920. 198. Berry FA, Gregory GA. Do premature infants require anesthesia for surgery? Anesthesiology 1987; 67:291-3. 199. Booker PD. Postoperative analgesia for neonates? Anaesthesia 1987; 42:343-4. 200. Pain, anaesthesia and babies. Lancet 1987; 2:543-5. 201. Yaster M. Analgesia and anesthesia in neonates. J Pediatr 1987; 111:394-6.

AuthorAffiliation K.J.S. Anand, M.B.B.S., D. Phil., and P.R. Hickey, M.D. AuthorAffiliation Reprinted from the New England Journal of Medicine, 317:1321-1329 (November 19), 1987. Drs. Anand and Hickey are from the Department of Anaesthesia, Harvard Medical School, and Children's Hospital, Boston. Address reprint requests to Dr. Anand at the Department of Anaesthesia, Children's Hospital, 300 Longwood Ave., Boston, MA 02115.

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