To the Editor.—We appreciate the three articles in the April 2004 issue of Anesthesiology regarding GW280430A.1–3 Although remarkable advances in developing intermediate and short-acting muscle relaxants were realized, anesthetists have not yet been provided with a substance comparable to succinylcholine in terms of its rapid onset and ultrashort-acting pharmacodynamic profile.

However, this aim should not be lost. A survey in Germany4 revealed that, despite its undesirable side effects, succinylcholine is still the most used drug for both rapid sequence inductions5 and for elective case induction.6 An overwhelming majority (76.6%) of respondents answered that they would appreciate a nondepolarizing substitute for succinylcholine if a similar pharmacodynamic profile was preserved. Assuming that this is not an isolated German viewpoint, a substance replacing succinylcholine would be highly desirable.

The developers and the researchers have a great responsibility when introducing a new drug into clinical practice, particularly in neuromuscular blocking drugs.7 Dr. Caldwell addresses this issue in his editorial when he compares the side effects of rapacuronium and GW280430A.8 Because we were involved in the clinical evaluation of rapacuronium,9–11 we would like to comment on some relevant aspects of the side effect profiles of both drugs and on the drug approval processes. First, in clinically relevant concentrations, rapacuronium potentiates bronchoconstriction most probably by destabilization of the balance between M2 and M3 muscarinic receptors.12 In contrast, GW280430A seems to release histamine3 and therefore may possibly induce bronchoconstriction. Second, although many antihistaminic drugs and prophylactic strategies are available, an effective treatment to rebalance the muscarinic effects of rapacuronium was and is still missing. Third, because rapacuronium did not release histamine,13 because different M2 versus M3 muscarinic effects of muscle relaxants were unknown at that time, and because clinical symptoms of the pulmonary side effects differed from those seen during typical bronchoconstriction,12 the clearly described dose-dependent pulmonary side effects (from 10.7% with 1.5 mg/kg rapacuronium to 18.5% with 2.5 mg/kg rapacuronium)10 may have been questioned—unfortunately until patients were badly harmed. Therefore, we agree with Dr. Caldwell that the recent experience with rapacuronium must be considered during the trials with GW280430A, e.g., by in addition investigating its effects on M2 and M3 receptors. The fiasco with rapacuronium, however, must not induce pessimism if new drugs and especially GW280430A may have the potency to improve anesthesia practice.

GW280430A was, of course, not compared with rapacuronium, but it was also not compared to succinylcholine.1–3 Regardless, the hope that GW280430A will be a substitute for succinylcholine has been advanced with this first presentation. Expectations that this new drug will approximate the rapid onset of succinylcholine may in high doses, high injection speeds, and, therefore, the risk for high incidences of side effects. The presentations1–3 primarily suggest that GW280430A may be an ultrashort-acting rather than a rapid-onset muscle relaxant.

Unfortunately, preclinical and clinical trials to approve new drugs are expensive, and, in this context, the substance to be replaced is already very cheap. Nevertheless, we (and many other anesthetists) would like to encourage the recent attempts to develop better muscle relaxants (or reversal drugs, e.g., Org 2596914) to improve safety and efficiency of neuromuscular treatment during anesthesia.

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To the Editor.—We read with interest Dr. Caldwell’s recent editorial1 accompanying the articles published on GW280430A.2–4 Although we agree with his assessment of its potential advantages, we have several concerns with respect to his comments regarding its adverse effects.

GW280430A is a potent, nondepolarizing neuromuscular blocking agent with a fast onset of effect and a duration of action similar to that of succinylcholine.5–4 When administered as a rapid intravenous bolus to volunteers in doses greater than or equal to 2.8 times the ED95, it

GW280430A: Pharmacodynamics and Potential Adverse Effects

To the Editor.—We read with interest Dr. Caldwell’s recent editorial1 accompanying the articles published on GW280430A.2–4 Although we agree with his assessment of its potential advantages, we have several concerns with respect to his comments regarding its adverse effects.

GW280430A is a potent, nondepolarizing neuromuscular blocking agent with a fast onset of effect and a duration of action similar to that of succinylcholine.5–4 When administered as a rapid intravenous bolus to volunteers in doses greater than or equal to 2.8 times the ED95, it
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may cause histamine release and subsequent hemodynamic changes, a decrease in blood pressure, and an increase in heart rate. As reported by Belmont et al., such changes were observed in 25% of volunteers receiving a dose of 2.8 times the ED$_{50}$ of mivacurium. As is typical of cardiovascular effects due to histamine release, these hemodynamic changes were all self-limited and required no pharmacologic treatment. As anticipated, observed symptoms of histamine release increased in frequency as the dose of GW280430A was increased. No attempt was made to decrease histamine release in that all doses were administered as a rapid intravenous bolus (5 s) into a rapidly flowing intravenous line. To put this further into perspective, as demonstrated in figure 1, the average maximal heart rate and blood pressure changes after administration of GW280430A are less than those observed after administration of mivacurium, when mivacurium is injected slowly (over 15 s). Doses of 2.5 times the ED$_{50}$ of GW280430A do not cause histamine release even when administered over 5 s. Furthermore, because 90% suppression of T$_{1}$ occurs within 1.3 min after administration of 0.36 mg/kg GW280430A, tracheal intubation may likely be accomplished in 60 s after administration of 3.8 times the ED$_{95}$. Decreases in MAP occurred at a dose of 3.8 times the ED$_{95}$. The average maximal decrease at this dose of GW280430A was 18%. At all doses, GW280430A was administered as a bolus (5 s) into a rapidly flowing intravenous line. In the study of mivacurium, the neuromuscular blocking agent was injected three times more slowly (15 s).

In Reply—I am grateful to Drs. Geldner and Blobner and Dr. Lien et al. for their responses to my editorial. I agree with Drs. Geldner and Blobner that succinylcholine still has a significant clinical role, and that anesthesia providers would like to see it replaced by a nondepolarizing drug with a similar time course of action. I agree also that we should not be pessimistic about prospects for new drugs, but history and pulmonary disease. Bronchospasm or any other difficulty with ventilation was not encountered over the course of the volunteer trial. This problem has been encountered rarely in the thousands of patients who have received mivacurium, which has a greater propensity to release histamine than GW280430A.

The bronchospasm noted after administration of rapacuronium is likely not due to histamine release. Bronchospasm has occurred in patients receiving rapacuronium with no evidence of histamine release. The bronchospasm after administration of rapacuronium is likely caused by its antagonism of the muscarinic M2 receptor. Non-depolarizing neuromuscular blocking agents can interact with two of the three muscarinic receptors that exist in the airways (M1, M2, and M3). Their antagonism of the M3 receptor causes bronchodiilation by inhibiting vagally induced bronchoconstriction. Antagonism of the M2 receptors, which are located presynaptically at postganglionic parasympathetic nerve endings, results in an increased release of acetylcholine that subsequently binds to M3 receptors, causing bronchoconstriction. The affinity of rapacuronium for the M2 receptor is 15 times it affinity for the M3 receptor.

As shown in experiments in cats, GW280430A is a very weak inhibitor of muscarinic receptors in general, with nearly the same safety ratio for this side effect as mivacurium. In cats, the muscarinic blocking dose (ED$_{50}$) of GW280430A is more than 25 times its ED$_{95}$ for neuromuscular block. A closer look at the data in the study of Heerdt et al. in dogs shows a complete lack of effect of GW280430A on airway pressures in the dog, even at doses of 50 times the ED$_{95}$. Nevertheless, GW280430A will have to be further tested for its relative affinity for the muscarinic receptors of the airways, as will all other nondepolarizing relaxants that may be introduced into clinical practice. Based on the data published to date, there is no reason to anticipate that GW280430A may even rarely cause life-threatening bronchospasm.

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I did not intend to suggest that GW280403A produced bronchospasm; there is no evidence for that. Rather, my intent was to emphasize that any adverse effect observed during clinical trials may assume far greater significance when the drug goes into general use. The experience with rapacuronium was a dramatic illustration of this point. In the case of GW280403A, the adverse effects raising concern are its propensity to release histamine and to produce hypotension.2

I differ with Dr. Lien et al. in the interpretation of the significance of the adverse hemodynamic effects of GW280403A. For example, take their statement “As is typical of cardiovascular effects due to histamine release, these hemodynamic changes were all self-limited and required no pharmacologic treatment.” I do not believe that the word typical can be used to characterize histamine-related effects. These effects can vary from trivial, localized, cutaneous flushing to life-threatening cardiovascular collapse. The fact that in a small sample of healthy young volunteers, who can increase heat rate to compensate for histamine-mediated vasodilatation, a blood pressure decrease of nearly 40% was observed is a matter for concern.2 The obvious question is how severely might this degree of histamine release manifest in patients with much less cardiovascuar reserve.

In addition, the fact that the blood pressure changes required no pharmacologic treatment in this limited subject population cannot be used to predict safety in the general population of patients. The subjects studied were all healthy young volunteers in whom a period of hypotension might be tolerable. Allowing the persistence of hypotension without treatment would not be an option for many patients undergoing anesthesia.

Will the doses of GW280403A causing histamine release and hypotension overlap with the doses that might be used in clinical practice? Dr. Lien et al. suggest not by speculating that GW280403A in dose of 0.36 mg/kg (2 × ED50) will provide good intubating conditions in 60 s. Because the onset time of 0.36 mg/kg GW280403A ranged up to 2 min in this small, healthy, and homogeneous group of subjects, it is difficult to believe that intubation within 60 s could be reliably achieved in the general patient population with this dose. It is likely that larger doses will be required to facilitate rapid tracheal intubation. As I described in my editorial, there is ample evidence that clinicians tend to increase doses of neuromuscular blocking drugs to achieve better and faster intubating conditions.3,5 Therefore, it is feasible that clinicians might use GW280403A in the dose range 3–4 × ED50, and the work by Dr. Lien et al. shows that at these doses, histamine release and hypotension can occur.2

Finally, it is not justified to infer the cardiovascular safety of GW280403A by using 15-yrs of historic comparisons with mivacurium.6 There is no way to ensure the comparability of the study procedures or subjects. The weakness of this comparison is further compounded by the use of “average maximal changes” as the variables for comparison. This averaging masks larger changes in individual subjects, and it is these that have the most clinical significance.

In summary, I do not believe that bronchospasm is a significant clinical issue with GW280403A, but histamine release and hypotension most assuredly are.

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In Reply—The three reports by our group in the April (2004) issue of ANESTHESIOLOGY1,3,5 on the new ultrashort-acting nondepolarizing relaxant GW280403A (AV430A) have evoked invited commentary by Caldwell7 and observations by Geldner and Blobner. Both anticipation and caution were expressed by these commentators regarding the possible future clinical performance of AV430A.

To add to the discussion, I would offer a rather optimistic viewpoint based on additional data in preparation for publication. I am certainly biased because of my closeness to the development of AV430A. Nevertheless, I do believe that future results will strengthen the candidacy of AV430A as a replacement for succinylcholine or as an excellent alternative.

AV430A is a representative of the new class of nondepolarizing relaxants, which we have called asymmetric-mixed-onium chlorofluoromar-rates.3 These compounds are inactivated by two entirely chemical (nonenzymatic) mechanisms both in vitro and in vivo: cysteine ad-duction and pH-sensitive hydrolysis.1,5,6 The cysteine addition reaction is notably manipulatable.5,6 In AV430A, both reactions are fast (combined T1/2β approximately 1–2 min) such that the duration of block is ultrashort, similar in duration to or even shorter than succinylcholine in cats, dogs, and three different primate species.1,5

Dr. Savarese has been a consultant to Glaxo Wellcome. Research Triangle Park, North Carolina, and is one of the inventors and patent holders of GW280403A and related compounds on behalf of Cornell University, New York, New York.

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Dr. Savarese has been a consultant to Glaxo Wellcome, Research Triangle Park, North Carolina, and is one of the inventors and patent holders of GW280403A and related compounds on behalf of Cornell University, New York, New York.
clinical practice with rapacuronium than foreseen before its release.
This is the crux of the problem: Minimal effects of AV430A on the
airway must be shown to provide convincing data regarding the safety
of AV430A. Caldwell & Geldner and Blobner both allude to this.

Following is a discussion of some of the data regarding AV430A that
are already published, which suggest that AV430A should be safe as far
as airway effects are concerned. Minimal airway effects have already
been found in dogs in doses up to 50 × ED95. The nonvagolytic
properties of AV430A found in cats mean that it is unlikely to have any
blocking effects on M2 or M3 receptors in the human airway. This
indicates minimal possibility of bronchospasm on this basis, in contrast
with the severe responses noted after rapacuronium. Jooste et al. have
shown that mivacurium has minimal interactions with M2 and M3
receptors where the actions of rapacuronium are clearly demonstra-
bable. Because mivacurium, which is a predecessor of AV430A, has
approximately the same large safety ratio (> 25) as does AV430A
regarding lack of interaction with muscarinic receptors in cats, the
imbalanced blocking property on muscarinic receptors in the human
airway, which is the likely mechanism of rapacuronium-induced bron-
chospasm, is a most unlikely clinical scenario in the case of AV430A.

What about histamine release? Both Caldwell and Geldner and
Blobner express concern about this, but in the rhesus monkey, the
safety ratio of AV430A for this side effect is approximately four times
greater than that of mivacurium. This results in the following com-
parison in humans: mivacurium, at 2.5–5 × ED95 (0.20–0.25 mg/kg),
gives causes greater symptomatology of histamine release when injected
over 15 s11 than does AV430A when injected over only 5 s at 3–4 × ED95 (0.5 mg/kg). Since the original study in human volunteers by
Belmont et al. in 1998, AV430A has been reformulated. This reformu-
lation has further improved the safety ratio for histamine release to
approximately 4 × ED95 in humans. As a reminder, this ‘safety ratio’
is defined as the dose required to cause an average decrease in blood
pressure of 10%, divided by the ED95 for neuromuscular blockade: ED
Hist/ED95.1

Both Caldwell and Geldner and Blobner caution that anesthesiolo-
gists might increase dosage to cause faster onset and thereby increase
the possibility of histamine release by AV430A. We have already com-
pared the onset and duration of AV430A in the human larynx and the
thumb vs atriceis succinylcholine, rapacuronium, rocuronium, mivacu-
rium, and cisatracurium. The data are not yet published but suggest
that the onset of AV430A is as fast as that of succinylcholine and faster
than those of the others. So why administer AV430A at a dosage higher
than 0.4–0.5 mg/kg? Onset of block after AV430A does not get any
faster at dosages higher than this and is probably circulation limited, as
pointed out by Caldwell. The onset is fast in all species—dogs, cats,
monkeys, and humans. Consequently, Geldner and Blobner, who believe
that ‘GW280430A may be an ultrashort-acting rather than a
rapid-onset muscle relaxant’ are most likely incorrect. AV430A is
clearly ultrashort and very rapid in onset in all studies.1–3 figure 1
shows a mechanomyograph recording of the response to AV430A (0.4
mg/kg) in the adductor pollicis in a healthy human volunteer during
nitrous oxide–oxygen–fentanyl–propofol anesthesia. The pattern of
block seen in figure 1 was noted in every volunteer subject of the more
than 100 humans treated so far. The dose of 0.4 mg/kg is approxi-
mately 3 × ED95.7

Doses up to 5 × ED95 caused minimal side effects. A 5-s bolus
dose of AV430A (0.4 mg/kg) was given at the arrow. Two control train-of-
four (TOF) responses are followed by the elicited twitch in the thumb
at 0.15 Hz. Twitch is abolished within approximately 80 s. Recovery
begins at approximately +6 min (time scale at top). The time scale
changes at and after this point. At +8 min, TOF shows appearance
of T1 and T2 is at approximately 25% of control. At +10 min, TOF is 45%
and T1 is 75% of control. At +12 min, TOF is 85% and T1 is 95% of
control. At +13.5 min, TOF is 95% of control (control TOF is at the far
left). The time from 5% T1 to a TOF of 90% is 6 min. Heart rate (upper
record) and blood pressure (lower record) show no change.

Time and experience will tell, as clinical studies in patients are
undertaken, whether early data from volunteers accurately predict the
performance of AV430A in practice. The dosage recommended to
achieve certain clinical endpoints, e.g., intubation of the trachea within
60 s, among others, must be defined by these future studies.

In volunteer studies thus far, at dosages as high as 0.8–0.9 mg/kg
(approximately 6 × ED95 or 2 × intubating dosage) where AV430A is
given as a rapid (5 s) bolus, the manifestations of histamine release
after AV430A are rather mild (facial flushing and brief decrease in
blood pressure, not requiring treatment).7 There has been no bron-
chospasm. This suggests that clinicians could give these very large
dosages safely, particularly by injecting them a little more slowly (such
as over 15 s). The side effect of histamine release as caused by AV430A,
because it is four times weaker than it is in mivacurium,7 may consti-
tute a minor concern in future clinical practice. Time again will tell.
Only after a couple of years of experience in thousands of administra-
tions will the pattern be fully defined.

Nevertheless, we can be reassured by this data. If, as Kopman et al.12
have suggested, a dose of 2.0 or 2.5 × ED95 of AV430A is enough for
good to excellent intubating conditions within 60 s (this dose would
be in the range 0.3–0.5 mg/kg), there may very well be minimal side
effects.

The chemical pathways of inactivation of AV430A are, in my opin-
ion, its most promising feature. The chemical breakdown will ensure
no prolonged neuromuscular blockade. Problems with atypical
pseudocholinesterase will not be at issue. Cysteine, given intrave-
nously as a “reversal drug” or “rescue agent,” will rapidly inactivate
AV430A. Complete recovery from 100% twitch inhibition can be in-
duced in monkeys with exogenous cysteine within 1–2 min.13 In
the future, anesthesiologists may have the choice of spontaneous recovery (12–14 min) from AV430A via endogenous cysteine or induced recov-
ery by giving additional cysteine, e.g., in case of an airway emergency.

fig. 1. A recording of evoked twitch (at 0.15 Hz) and train-of-four (2 Hz for 2 s every 10 s) of the adductor pollicis in an anesthetized human volunteer. AV430A (0.4 mg/kg) was given intravenously at the arrow. Time scale (minutes) at top. Note change of time scale and change of evoked response from single twitch to train-of-four at minute 7. Onset of block is 80 s. Recovery to train-of-four of 95% is 13.5 min.
Cysteine, given within 2–3 min after injection of AV430A, should abolish complete paralysis within 1–2 min. The latter treatment with cysteine may shorten the total duration of action in humans to an estimated 5 min to return of full neuromuscular function such as cough, normal vital capacity, and head lift.

**References**


To the Editor:—I read with interest the article “Lumbar Plexus in Children” by Lukas Kirchmair et al. published in the August 2004 issue of *Anesthesiology*. The authors have nicely shown that sonography of the lumbar plexus in children is feasible, they have clarified the anatomical understanding, and they have given additional information on the depth of the lumbar plexus in pediatric patients.

In addition, they applied their technique to five children scheduled to undergo inguinal hernia repair and concluded that all lumbar plexus blocks provided effective anesthesia and analgesia of the inguinal region during surgery and for postoperative pain relief. Although I am unable to check the clinical effects in these five individual patients, I have considerable doubt that lumbar plexus block is a suitable anesthetic technique for inguinal hernia repair and would not recommend it for the following reasons:

First, the iliohypogastric nerve arises from the roots T12 and L1, and the ilioinguinal nerve arises from L1 and L2, respectively. Therefore, a posterior lumbar plexus block with an injection at the L4–L5 level in combination with a sciatic nerve block, incisions coming high up, close to the inguinal ligament, are often troublesome. The ilioinguinal and genitofemoral nerves are often not sufficiently blocked.

In summary, lumbar plexus block can be used in children. However, its use should be restricted to indications where this technique is clearly effective, e.g., for lower limb procedures involving parts of the hip or extensive knee surgery, and where no other less invasive alternative exists.

**Reference**


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In the current cases, the local anesthetic solution must have reached the cranial parts of the lumbar plexus because sufficient anesthesia and analgesia of the inguinal region was observed in all five patients. Recent investigations of ultrasound-guided posterior lumbar plexus block in pediatric patients revealed a greater extent of anesthesia and analgesia compared with adults.

It was not the aim of our study to investigate posterior lumbar plexus block for inguinal hernia repair in children, and we agree with Dr. Jörh that it should not be the first choice for this surgical procedure. However, in our opinion, it might be a useful alternative and still represents a peripheral nerve block.

Dr. Jörh also had concerns about the strategy of applying small volumes in the current setting. Nevertheless, one of the proven benefits of ultrasound-guided techniques is the decreased need of local anesthetics compared with traditional approaches.

To the Editor—Over the years, Olney et al.1 have enlightened the scientific community with their research on the N-methyl-D-aspartate receptor and its role in human disease. Recently, Dr. Olney’s group has examined the effects of commonly used anesthetics and anticonvulsants on the developing brain. In several high-profile scientific journals, they reported that the administration of these drugs, including ketamine, ethanol, phencyclidine, nitrous oxide, isoflurane, propofol, barbiturates, diazepam, and other anticonvulsants all increase apoptotic neurodegeneration in developing rat brain.2–5 We and others have replicated some of the studies with ketamine, and there is no doubt regarding the scientific validity of their findings.6–8

The direct applicability of these experimental findings to the clinical practice of pediatric anesthesia and critical care, however, should be questioned. Likewise, the implication that all types of anesthetic and sedative agents may have similar potentially deleterious effects on neuronal development in neonates is premature and inappropriate. Multiple lines of evidence cast doubt on the clinical relevance of these experimental paradigms, as recently reviewed in our Special Article published in the August 2004 issue of Anesthesiology.9 Dr. Olney et al.9 eloquently provided a counterpoint editorial in the same issue of Anesthesiology, which mainly questioned our assertion that repeated ketamine administration, particularly in the absence of surgical induction of pain or stress, might have similar potentially deleterious effects on the developing nervous system. Nevertheless, one of the proven benefits of ultrasound-guided techniques is the decreased need of local anesthetics compared with traditional approaches.

Recent data from the Neurotoxicology division at the National Center for Toxicological Research (Jefferson, Arkansas) further confirm the experiments by Olney et al.9 because of the lack of a robust pattern of neuroapoptosis and excitation,7,8 which we and others have replicated some of the studies with ketamine, and there is no doubt regarding the scientific validity of their findings.6–8

In their editorial, Olney et al.10 disagree with our concern that neurodegenerative changes after repeated high doses of ketamine may result from hypoxic/ischemic damage because of the hemodynamic effects of high-dose ketamine. ‘It is excitotoxic, and ultrastructurally does not resemble apoptosis,’ they state. The neuronal cell death that occurs soon after exposure to hypoxia/ischemia does result from excitotoxicity, but rat brains were not sampled at this time period in the studies by Olney et al.2,4 A voluminous literature shows a delayed neuronal cell death after hypoxia/ischemia that results from neuronal apoptosis in the immature rat brain and also occurs at the same time periods (4–24 h) as those sampled by Olney et al.10–13 Excitotoxicity cannot account for hypoxia-induced injury in immature neurons but may explain the effects of hypoxia on more mature neurons,14 and even the excitotoxic damage of immature neurons leads to apoptosis at 6–24 h at the time periods studied by Olney et al.2,4

In their editorial, Olney et al.10 state that arterial blood gases showed oxygen saturations of 97–99% (contained in the legend for their fig. 2). To our knowledge, the complete data from arterial blood gases obtained at 0, 15, 30, 60, 120, 180, or 240 min in neonatal rats given high-dose ketamine have not been reported in the literature and cannot guarantee the absence of significant hypoxia occurring between these sampling times. In contrast, pediatric anesthesiologists routinely and continuously monitor oxygenation (oximetry), ventilation (capnography), and other hemodynamic variables during neonatal anesthesia and respond to these monitors within seconds after noting a clinically important change.

Similarly, our concern for nutritional deprivation was also dismissed by Olney et al.10 because of the lack of a robust pattern of neuroapoptosis in the controls from their experiments, stating that “all animals are sacrificed 4 to 8 h later for histologic evaluation of the brains.” Their experiments, however, showed that rat pups were killed at 4, 8, 12, 16, 24, or 48 h after ketamine and that apoptosis was most prominent in the brains sampled at 12, 16, and 24 h.7 If newborn rat pups were exposed to nutritional deprivation for these periods of time, increasing patterns of neuronal apoptosis would be expected.16–17

Last, we must restate our concern that neonatal rats or humans exposed to anesthetic agents in the presence versus the absence of surgically induced pain or stress would manifest drastically different effects and that the observations that the neuronal effects of all anes-

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Of Mice and Men: Should We Extrapolate Rodent Experimental Data to the Care of Human Neonates?

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at issue in the study of Newburger and de Ferranti et al.25 WISC-III = Wechsler Intelligence Scale for Children–III.

Wechsler Preschool and Primary Scale of Intelligence score measures at 4 and 8 yr after surgical repair of complex congenital heart defects. Population norms are 100 ± 10. There were no significant differences between the study groups and the population norms. Data compiled from Bellinger et al.21,22 and de Ferranti et al.25

Fig. 1. Wechsler Preschool and Primary Scale of Intelligence scores at 4 and 8 yr after surgical repair of complex congenital heart defects. Population norms are 100 ± 10. There were no significant differences between the study groups and the population norms. Data compiled from Bellinger et al.21,22 and de Ferranti et al.25

Wechsler Preschool and Primary Scale of Intelligence

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In Reply—In their letter to the editor, Soriano et al. state that in our recent counterpoint editorial,1 we “mainly questioned [their] assertion that repeated large doses of ketamine were responsible for mediating the neurodegenerative changes noted in neonatal rat brains.” This is not what we said, nor do we know that repeated large doses of ketamine can cause extensive neuroapoptosis. What we mainly questioned was whether repeated large doses are necessary, or whether neuroapoptosis can be triggered by a single low dose of ketamine. We then presented evidence that a single subcutaneous dose of ketamine (20, 30, or 40 mg/kg)—one that does not fully immobilize, anesthetize, or abolish pain responses in infant mice—does trigger a significant increase in the rate of neuroapoptosis. The scientifically appropriate response would be for Soriano et al. to administer these single subanesthetic doses of ketamine to infant mice and, using the same methods we used, to see whether they could reproduce our findings. Instead, Soriano et al. imply that it requires “repeated large doses” to trigger neuroapoptosis and argue that ketamine is safe for pediatric anesthesia because such large “doses and durations . . . are never used in pediatric anesthesia.”

To bolster their claim that a single dose is ineffective, they cite a recent report of Scallet et al.2 in which a single dose of ketamine at 20 mg/kg did not trigger apoptosis in infant rats, although repeated 20-mg/kg doses did. Because, in our single-dose experiments, 20 mg/kg was the threshold dose for triggering apoptosis, it is not surprising or very meaningful that one laboratory would report a barely significant effect and another would report a barely insignificant effect at this dose. What is surprising is that instead of directly acknowledging and discussing the implications of our finding that a single dose of ketamine at 20, 30, or 40 mg/kg does trigger apoptosis in a dose-dependent manner, Soriano et al. continue to promote their original position that ketamine is safe because, in their hands, a single dose as high as 75 mg/kg does not trigger neuroapoptosis. It is difficult to reconcile this position with their introductory statement that “there is no doubt regarding the scientific validity” of our findings.

Soriano et al. suggest that we should have measured blood ketamine concentrations in our mouse experiments. However, Soriano et al. did not measure ketamine blood concentrations in their rodent experiments, and anesthesiologists do not routinely measure, much less rely on, ketamine blood concentrations to determine depth of anesthesia. We reported that a single dose of ketamine, in the range of 20–40 mg/kg, that does not fully immobilize, anesthetize, or render an infant mouse insentient to pain does trigger neuroapoptosis in the infant mouse brain. This is a message that is not difficult to understand. Presumably, we can all agree that regardless of ketamine blood concentrations, it would be unacceptable to perform surgery on an infant mouse, or infant human, whose depth of anesthesia is such that the infant is squirming around, failinf the extremities, and responding to skin pinch by vigorous antalgic movements.

Soriano et al. continue to argue that the neuroapoptosis response to anesthetic drugs is due to hypoxia/ischemia. How is this possible in light of our demonstration1 that arterial oxygen saturation remains in the 97–99% range over a 4-h period after a dose of ketamine that triggers neuroapoptosis within this same time interval? Soriano et al. postulate that the oxygen saturation fleetingy decreased to brain-damaging levels during intervals between our sampling time points but abruptly resumed normal levels at each time point (15, 30, 60, 120, 180, 240 min) just before we drew blood. We doubt that the readership of Anesthesiology will be persuaded by this argument, especially because we are talking about a subanesthetic dose of ketamine, a drug that reputedly, even at anesthetic doses, does not compromise cardio-respiratory function.

Even if severe hypoxia/ischemia did occur, it could not account for the neuroapoptosis response to ketamine because 4–6 h after ketamine administration, an increase in apoptotic profiles is evident both as a caspase-3 activation response and as ultrastructurally confirmed apoptotic morphology. However, when one intentionally induces hypoxia/ischemia and examines the developing brain 4–6 h later, there is no increase in apoptotic profiles, either by caspase-3 activation or ultrastructural criteria. It is illogical to argue that anesthesia-induced apoptosis is caused by hypoxia/ischemia if one cannot demonstrate that intentionally induced hypoxia/ischemia reproduces the anesthesia-induced apoptosis phenomenon. What one does find in the brain 4–6 h after hypoxia/ischemia, as we have demonstrated previously,3 and also very recently,4 is excitotoxic neurodegeneration. (See Young et al.4 for a detailed presentation of evidence directly addressing and clarifying this issue.) Soriano et al. challenge our position by citing works from other laboratories that they believe contradict our observations. We have examined the cited works, some of which are in vitro studies, and find that these works either support our position or are irrelevant to the issue in contention. We stand by our own observations, which are based on a three-decade-long direct investigation of the specific issue in contention: in vitro excitotoxic versus apoptotic neurodegeneration in the developing brain.3–10

Regarding the nutritional deprivation issue, we stated1 that in a “typical” experiment, we expose infant rodents to a single dose of saline or an apoptogenic anesthetic drug and, without returning the pups to the maternal nest, kill them 4–8 h later. Because both the control and experimental pups are exposed to the same degree of maternal/nutritional deprivation during this 4-to 8-h period, nutritional deprivation cannot explain the higher rate of neuroapoptosis in the experimental pups. In an apparent effort to refute this interpretation, Soriano et al. note that in one study we killed animals not only at 4 and 8 h but also at 12, 16, 24, and 48 h, and determined, using a staining procedure that detects cumulative neuronal degeneration, that apoptosis became increasingly more prominent at 12, 16, and 24 h. We do not understand how this reference to our earlier comprehensive evaluation of the apoptotic response to large doses of MK80111 refutes our current interpretations pertaining to “typical” experiments focusing on the very early response to low subanesthetic doses of ketamine.

Soriano et al. point out that our most recent findings1 pertaining to threshold conditions for inducing developmental neuroapoptosis were conducted in mice and suggest that species differences between rats and mice and between rodents and humans may be of paramount importance. We have tested rats and mice and find no appreciable differences between these species, but we agree that differences between rodents and primates may be very important. Of course, species differences can go in either direction—humans may be less vulnerable or they may be more vulnerable than rodents.

Soriano et al. conclude that only human studies can provide the final answer. We do not contest the importance of human experiments, but such experiments will require many years to complete and, because of design limitations, may provide equivocal results that defy interpretation. Therefore, we recommend that the issue be addressed in nonhuman primate studies designed to test the sensitivity of the primate brain to anesthesia-induced developmental neuroapoptosis. If the pri-
Avoid Excessive Sedation during Cervical Injections

To the Editor.—I read with interest the article by Rathmell et al. regarding cervical transforaminal injection of steroids. An important safety issue not addressed was the need to minimize complications by avoiding excessive sedation. Hodges et al. reported two cases of nerve injury after cervical epidural steroid injections, both performed in heavily sedated patients using fluoroscopy. Excessive sedation may result in the inability of the patient to experience and report pain and paresthesias at the time of spinal cord or nerve root contact. In addition, some recommend that cervical injections should only be performed by experienced and well-trained practitioners.3

Richard B. Weiskopf, M.D., served as Handling Editor for this letter, the following letter by Wills and Martin, and the reply by Rathmell et al.

References


3. De Cordoba J, Bernat J: Cervical transforaminal blocks should not be attempted by anyone without extensive documented experience in fluoroscopically guided injections. Anesthesiology 2004; 100:1325–4

(See publication for December 20, 2004.)

To the Editor.—We thank Dr. Rathmell et al. for emphasizing the potential hazards of transforaminal injections. It is clear that these injections should be performed by individuals who are fully trained in advanced imaging and interventional techniques. Moreover, the practitioner must be capable of managing any adverse sequelae.

Although the treatment of radiculopathic pain with the local injection of corticosteroid is appreciated, we believe that the best treatment of radiculopathic pain is by the application of pulsed radiofrequency current to the involved dorsal root ganglion. In our extensive experience with pulsed radiofrequency, we have found the results of treatment to be superior to those of conventional corticosteroid injections in both effectiveness and duration.2,5

Furthermore, pulsed radiofrequency application exposes patients to less risk for the following reasons: (1) Cannula placement can be performed based solely on osseous anatomy and electrophysiologic stimulation results. Therefore, no injection of any material is requisite. (2) The patient is exposed to no systemic sequelae of medication or radiocontrast administration. (3) Unlike injection therapies, pulsed radiofrequency can be repeated as indicated, without fear of accumulating medicinal toxicity.

The only potential disadvantage to the use of pulsed radiofrequency versus injection therapy is the requirement of a larger cannula (20–22 vs. 26 gauge) that could cause more tissue trauma. Regardless, the take-home message is well elucidated by the authors. Spinal interventional techniques should only be performed by practitioners who have demonstrated expertise in neural imaging for interventional treatment modalities.

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(See publication for December 20, 2004.)
In Reply.—Critical to the safety of cervical transforaminal injection of steroids is an understanding of the anatomy of the cervical intervertebral foramina and their contents, coupled with disciplined and accurate imaging. We thank both Dr. Gajraj and Drs. Willis and Martin for emphasizing that cervical transforaminal injection should be performed by experienced and well-trained practitioners. Indeed, the radiographic anatomy of the cervical spine is difficult to master. We have all watched talented physicians-in-training get confused by small changes in alignment on the x-ray image caused by positioning or rotation of the neck and stray dangerously off-course during needle placement. Image-guided injection in the cervical spine requires advanced and extended training under the guidance of an experienced practitioner; as we have emphasized before, this is not something that can be mastered through a weekend cadaver course.1

Even with the best technique in skilled hands, minimal changes in needle direction and depth can lead the tip into contact with neural structures. After the needle is in proper position, the volume of the injectate itself can cause painful neural compression. We have emphasized the need to maintain an awake and responsive patient when performing cervical epidural steroid injection via a translaminar route as the only safe means to avoid injury,2 and we thank Dr. Gajraj for raising this point because it is equally relevant to any type of neural blockade.

As for Drs. Willis and Martin’s advocating pulsed radiofrequency treatment of the dorsal root ganglion as a superior technique for treating cervical radicular pain, we point out that there is little evidence to support their assertion. Small, randomized trials of conventional radiofrequency treatment (i.e., resulting in a thermal lesion) of the dorsal root ganglion for the treatment of cervical radicular pain suggest time-limited efficacy.4,5 Results from similar trials in patients with lumbosacral radicular pain have been less promising: An initial large observational study suggested significant pain reduction,5 but a subsequent randomized trial by the same investigators showed no benefit over placebo.6 Pulsed radiofrequency treatment has evolved from the notion that the pain relief that ensues after radiofrequency treatment does not result from actual tissue destruction caused by conventional thermal lesions; rather, it is brought about by the large volume fluctuations in the area of treatment that induce long-term changes in the dorsal horn of the spinal cord.7 The appeal of pulsed radiofrequency treatment is immediately clear: a simple treatment that imparts long-term pain relief without tissue destruction. However, we do not have even a single randomized trial that compares the efficacy of pulsed radiofrequency to any type of control treatment or to conventional radiofrequency treatment. We hope that the evidence will soon appear to support the unbridled zeal of practitioners for this new treatment. We urge those like Drs. Willis and Martin who have extensive experience with these techniques to conduct the randomised trials we need to demonstrate the effectiveness (or lack thereof) of pulsed radiofrequency treatment.

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Endotracheal Tube Damage during Head and Neck Surgeries as a Result of Harmonic Scalpel® Use

To the Editor.—The Harmonic Scalpel® (Ethicon, Somerville, NJ) and laser are precise cutting and coagulating surgical devices. These devices are widely used worldwide for endoscopic and open surgical procedures. We describe our experience with the Harmonic Scalpel® during the anesthetic management of head and neck oncoursery.

A 58-yr-old male with postradiation recurrence of tongue and soft palate carcinoma was scheduled for wide excision and functional neck dissection with the Harmonic Scalpel® under general anesthesia. After induction of general anesthesia, direct laryngoscopy was performed and an 8 mm cuffed endotracheal tube (ETT) was introduced nasotracheally under vision. Maintenance of anesthesia was achieved with opioids, muscle relaxants, and positive pressure ventilation with 66% nitrous oxide, 34% oxygen, and 0.8% halothane delivered by anesthesia ventilator. The patient remained stable and was closely monitored while the surgeons proceeded with dissection using the Harmonic Scalpel®. While the soft palate lesion was being dissected, the ventilator suddenly emitted the low airway pressure alarm and stopped functioning. The surgical team also noticed bubbling of blood inside the oral cavity. Damage to the ETT was suspected and the head end of the operating table was immediately lowered to prevent aspiration of blood that had pooled in the oropharynx. The fraction of inspired oxygen was increased to 60% and manual ventilation was attempted after thorough suctioning of the oral cavity. However, this was unsuccessful due to the leak (fig. 1), and a fresh ETT of the same size was introduced with the aid of a tube exchanger. The patient was reversed at the end of surgery and the ETT was retained overnight. The postoperative period was uneventful.

The blade of the Harmonic Scalpel® vibrates at 55,000 Hz. It cuts and coagulates tissue at temperatures much lower than either lasers or traditional electrocautery. As a result, the risk of airway fires should be reduced. However, as shown here, it is clear that the Harmonic Scalpel®—like a laser—can cause accidental damage to the ETT when used in the pharynx.

The existing literature abounds with techniques to protect the air-
way and the ETT from lasers and to manage fires when they occur.\textsuperscript{1–6} Substitution of the Harmonic Scalpel\textsuperscript{®} can eliminate the risk of fire (and does not dictate the use of lower concentrations of oxygen), but ETT damage may still occur.

**Fig. 1.** Polyvinyl chloride endotracheal tube damaged by the Harmonic Scalpel\textsuperscript{®} during surgery on the soft palate.

Support was provided solely from institutional and/or departmental sources.
To the Editor.—Placement of a labor epidural or combined spinal–epidural in advanced labor is technically challenging. Regular painful contractions often make it difficult for the parturient to remain still during epidural placement, and this may increase the chance of an accidental dural tap or nerve injury. Decreasing the intensity and frequency of uterine contractions during neuraxial placement in this setting may be advantageous. Previous reports show that nitroglycerin produces rapid effective uterine relaxation.1–5 Nitroglycerin to facilitate the placement of a labor epidural has not previously been reported. This case describes the use of nitroglycerin in the setting of advanced labor to facilitate the placement of a labor epidural.

A 35-year-old, healthy, 90 kg, gravida 3, para 2 parturient admitted to labor and delivery in advanced labor requested an epidural for pain relief. She had two previous uncomplicated normal vaginal deliveries without the use of a labor epidural. A recent cervical examination showed her cervix to be dilated 8 cm, with the fetal head at +1 station. She was moving and uncooperative during contractions, which occurred every 45–60 s. She had not responded to 100 μg intravenous fentanyl given 5 min previously. With difficulty, we managed to position her in a sitting position to administer the epidural. After a sterile preparation of her back with a 10% povidone-iodine solution and 1% lidocaine skin infiltration, we attempted to insert the epidural catheter using a 17-gauge Tuohy needle. However, she kept moving and was uncooperative during and between uterine contractions. After informing the obstetrician and the patient that we were going to administer medication to help ease the painful contractions, we administered three sprays (400 μg per spray dose) of sublingual nitroglycerin (Nitrolingual® Pumpspray; First Horizon Pharmaceutical Corporation, Alpharetta, GA). This produced a temporary decrease in her uterine contractions (reduction in peak uterine pressures and an increased between-contraction interval as measured by external tocodynamometer) and resulted in some transient pain relief. It was then possible to perform the combined spinal–epidural during the interval between contractions. The patient experienced no hypotension or cardiovascular disturbances after administration of the nitroglycerin and resumed her normal uterine contraction pattern within a few minutes. The patient was delivered of a healthy baby vaginally 2 h later, with 1- and 5-min Apgar scores of 8 and 9, respectively.

Reducing contraction pain during placement of a labor epidural is potentially beneficial. However, the risks of uterine tocolysis must be balanced with the potential benefit of safer epidural placement and labor analgesia. Although there have been no studies demonstrating increased dural puncture or neural damage after epidural placement in an uncooperative and moving parturient, most clinical anesthesiologists believe that a relation must exist. Decreasing the intensity and frequency of uterine contractions during neuraxial placement in this setting should be potentially advantageous. Remifentanil has been described in this setting to improve analgesia and facilitate the insertion of a labor epidural.6 However, potent narcotics have potential adverse effects, in particular maternal apnea, dysphoria, and emesis. Nitroglycerin is a safe, effective uterine tocolytic commonly used in labor, with a rapid onset and brief half-life.2,7 Nitroglycerin has minimal, short-lived cardiovascular effects compared with β-adrenergic tocolytics. Although the safety of nitroglycerin during obstetric emergencies seems high, with no adverse maternal or neonatal outcomes,2 maternal hypotension and hemodynamics changes are possible, especially if high doses are given.9 No more than three metered sprays are recommended within a 15-min period.9

A number of studies and case reports describe the use of nitroglycerin in achieving rapid uterine relaxation.2,5–3 Nitroglycerin has been used as a tocolytic to reduce uterine hyperactivity,8,9 assist reduction of an inverted uterus,10 facilitate intrapartum external cephalic version,12 and manage preterm labor contractions.13 Nitroglycerin can be administered via a number of routes (intravenous, sublingual, or ointment), however, bioavailability is highly variable between subjects because of a pronounced first-pass metabolism. After sublingual administration, bioavailability is approximately 38%.14 Nitroglycerin may be useful in a setting where advanced labor and parturient movement during uterine contractions makes the placement of an epidural difficult and potentially dangerous. It exposes the mother and fetus to minimal risk and, in selected patients, may offer potential benefits justifying its use in this setting. However, physicians should remember that this is ‘‘off-label’’ use of nitroglycerin.9 Both the risks and the benefits must be considered, and the patient and her obstetrician must be consulted before uterine tocolytics are administered in this setting.

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Transpharyngeal Ultrasonography for Cannulation of the Internal Jugular Vein

To the Editor—Complications due to internal jugular vein (IJV) cannulation are infrequent and rarely life threatening. However, inadvertent carotid artery puncture can lead to serious problems in patients who have atheromas or bleeding disorders or who are undergoing full anticoagulation therapy, as for cardiopulmonary bypass. An external vascular ultrasound technique, using a vascular probe positioned on the neck, has previously been described as an aid for IJV cannulation.1 We have found useful in our practice an alternative method, using a transesophageal probe.

During the past year, we selected 50 cardiac patients for whom jugular vein cannulation could present a risk. Patients who required monitoring with a transesophageal probe and who had carotid artery disease, previous carotid artery surgery, and difficult anatomy, such as unclear landmarks or no palpable venous pulse, were selected. The mean age of the patients was 72.3 ± 8.8 yr (median, 74 yr).

Transesophageal echocardiographic monitoring is performed with use of a multiplane transducer and a Sonos 5500 imaging system (Hewlett Packard, Andover, MA). Induction of anesthesia and tracheal intubation are performed before insertion of the echo probe. A small towel is placed under the patient’s shoulders. The head is then extended and turned slightly to the side opposite the cannulation, and the patient’s arms are placed by his or her sides. The patient is positioned in a 25° head-down position. The ultrasound monitor is placed in front of the operator.

The transesophageal echo probe is inserted 12–20 cm from the teeth. The tip is directed along the pharyngeal lateral wall, which is why we call this method transpharyngeal. The probe is then rotated laterally 15–20° until the cervical vascular bundle is seen. The view is a mirror image of that obtained from conventional vascular ultrasonography (fig. 1).

A needle covered with a plastic hood is used to search the skin surface of the neck for the best site of cutaneous insertion to find the IJV, which is not pulsating and does not compress. The ultrasound probe is kept stable on a trolley. The operating field, the operator, and the devices are then prepared as usual under sterile conditions.

The IJV puncture was successful in 100% of the patients studied. No carotid punctures or other immediate complications occurred. Conventional ultrasound-guided cannulation of the IJV significantly improves the success rate, decreases the access time, and reduces the complication rate of cannulation.2–4 Meta-analyses and a systematic review of control data from literature5–7 were performed and suggested an advantage of ultrasonography in complicated cases and when access problems were anticipated.1,8 At our institution, the widespread use of transesophageal echocardiography, with its ability to provide a view of the vascular bundle of the neck, offers the anesthesiologist a simple way to aid in central venous cannulation, without any additive cost.

Potential advantages of transpharyngeal ultrasonography in comparison with conventional ultrasonography in intubated patients undergoing transesophageal echocardiographic monitoring are as follows: Direct compression on the IJV by the external probe is not needed; other professionals do not need to be involved in the procedure; the operator’s hands are free; and the ultrasound probe, still working in the patient, can be used for other purposes.

Further studies are needed to assess the reliability of this procedure for IJV cannulation, to determine its proper indications, and to compare this technique with other methods.

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Fig. 1. Transpharyngeal, short-axis view of the neck vessels. The direction of the needle is away from the carotid artery (CA), and the tip of the needle is inside the internal jugular vein (IJV).
References


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