Table of Contents
January, 2013

Cardiovascular Anesthesiology

關於藥物治療在病人血液保護方面的當前狀況
(鄧利兵譯 薛張綱校)
Special Article: Current Status of Pharmacologic Therapies in Patient Blood Management
   o Lawrence Tim Goodnough and
   o Aryeh Shander

綜述：現代澱粉在外科手術中應用的安全性
(鄭華容譯 陳傑校)
Review Article: Safety of Modern Starches Used During Surgery
   o Philippe Van Der Linden,
   o Michael James,
   o Michael Mythen,
   o and Richard B. Weiskopf

Ambulatory Anesthesiology

地塞米松預防術後噁心嘔吐：一項隨機對照試驗的最新 Meta 分析
(王曉莉譯 馬皓琳 李士通校)
Dexamethasone to Prevent Postoperative Nausea and Vomiting: An Updated Meta-Analysis of Randomized Controlled Trials
   o Gildasio S. De Oliveira, Jr.,
   o Lucas J. Santana Castro-Alves,
   o Shireen Ahmad,
   o Mark C. Kendall,
   o and Robert J. McCarthy
Opioid-Sparing Effect of Preemptive Bolus Low-Dose Ketamine for Moderate Sedation in Opioid Abusers Undergoing Extracorporeal Shock Wave Lithotripsy: A Randomized Clinical Trial

- Babak Gharaei,
- Alireza Jafari,
- Homayoun Aghamohammadi,
- Mohammadreza Kamranmanesh,
- Mahtab Poorzamani,
- Hedayatollah Elyassi,
- Baharak Rostamian,
- and Alireza Salimi


Myocardial Accumulation of Bupivacaine and Ropivacaine Is Associated with Reversible Effects on Mitochondria and Reduced Myocardial Function

- Nicole Hiller,
- Peter Mirtschink,
- Christine Merkel,
- Lilla Knels,
- Reinhard Oertel,
- Torsten Christ,
- Andreas Deussen,
- Thea Koch,
- and Sebastian N. Stehr

Technology, Computing, and Simulation

Beat-to-Beat Tracking of Systolic Blood Pressure Using Noninvasive Pulse Transit Time During Anesthesia Induction in Hypertensive Patients

Sung-Hoon Kim, Jun-Gol Song, Ji-Hyun Park, Jung-Won Kim, Yong-Seok Park, and Gyu-Sam Hwang


The Impact of Multilumen Infusion Devices on the Occurrence of Known Physical Drug Incompatibility: A Controlled In Vitro Study

Aurélie Foinard, Bertrand Décaudin, Christine Barthélémy, Bertrand Debaene, and Pascal Odou


Special Article: Curriculum and Cases for Pain Medicine Crisis Resource Management Education

Gary J. Brenner, Jordan L. Nemack, and Daniel Raemer


Patient Safety
心臟電生理學實驗中心對多名患者氣道損傷的研究
(張怡譯 馬皓琳 李士通校)

Airway Trauma in a High Patient Volume Academic Cardiac Electrophysiology Laboratory Center
  o Zhe Yan,
  o Jonathan W. Tanner,
  o David Lin,
  o Ara A. Chalian,
  o Joseph S. Savino,
  o Lee A. Fleisher,
  o and Renyu Liu


評估琥珀膽鹼引發惡性高熱的風險
(韓敘譯 薛張綱校)

Estimate of the Relative Risk of Succinylcholine for Triggering Malignant Hyperthermia
  o Franklin Dexter,
  o Richard H. Epstein,
  o Ruth E. Wachtel,
  o and Henry Rosenberg


Obstetric Anesthesiology

比較鞘內注射30mg利多卡因和45mg利多卡因試驗劑量對產科人群影響的一項前瞻性隨機實驗
(馬霄雯譯 陳傑校)

A Prospective Randomized Trial of Lidocaine 30 mg Versus 45 mg for Epidural Test Dose for Intrathecal Injection in the Obstetric Population
  o Stephen Pratt,
  o Philip Hess,
  o and Anasuya Vasudevan


間歇性硬膜外注射與連續性硬膜外輸注用於分娩鎮痛的比較：一項系統回顧和薈萃分析
(許辛譯 馬皓琳 李士通校)
Intermittent Epidural Bolus Compared with Continuous Epidural Infusions for Labor Analgesia: A Systematic Review and Meta-Analysis

- Ronald B. George,
- Terrence K. Allen,
- and Ashraf S. Habib


使用 Episure™自動檢測™注射器與傳統的玻璃注射器實施硬膜外技術相關的學習曲線：經驗豐富的麻醉醫師對產科病人實施的一個非盲，隨機，對照，交叉試驗。

(賀盼譯 薛張綱校)

The Learning Curve Associated with the Epidural Technique Using the Episure™ AutoDetect™ Versus Conventional Glass Syringe: An Open-Label, Randomized, Controlled, Crossover Trial of Experienced Anesthesiologists in Obstetric Patients

- Jean M. Carabuena,
- Aya M. Mitani,
- Xiaoxia Liu,
- Bhavani S. Kodali,
- and Lawrence C. Tsen


一項關於處理產後出血的新鮮冰凍血漿與紅細胞比率的觀察研究

(胡曉清譯 薛張綱校)

An Observational Study of the Fresh Frozen Plasma: Red Blood Cell Ratio in Postpartum Hemorrhage

- Pierre Pasquier,
- Etienne Gayat,
- Thibaut Rackelboom,
- Julien La Rosa,
- Abeer Tashkandi,
- Antoine Tesniere,
- Julie Ravinet,
- Jean-Louis Vincent,
- Vassilis Tsatsaris,
- Yves Ozier,
- François Goffinet,
- and Alexandre Mignon

Transport Decreases the Quality of Cardiopulmonary Resuscitation During Simulated Maternal Cardiac Arrest

Steven S. Lipman, Jocelyn Y. Wong, Julie Arafeh, Sheila E. Cohen, and Brendan Carvalho


Pediatric Anesthesiology

First Evidence of a Polygenic Susceptibility to Pain in a Pediatric Cohort

Chantal Mamie, Michela C. Rebsamen, Michael A. Morris, and Alfredo Morabia


The Effect of Passive Leg Elevation and/or Trendelenburg Position on the Cross-Sectional Area of the Internal Jugular Vein in Infants and Young Children Undergoing Surgery for Congenital Heart Disease

Won Ho Kim, Jong Hwan Lee, Sangmin M. Lee, Chung Su Kim, Ryunga Kang, Chan Seon Yoo, and Hyun Sung Cho

Pediatric Neuroscience

Early Developmental Exposure to Volatile Anesthetics Causes Behavioral Defects in *Caenorhabditis elegans*

- Katherine R. Gentry,
- Louise M. Steele,
- Margaret M. Sedensky,
- and Philip G. Morgan


Neuroscience in Anesthesiology and Perioperative Medicine

Repolarization Abnormalities in Patients with Subarachnoid and Intracerebral Hemorrhage: Predisposing Factors and Association with Outcome

- Eija Junttila,
- Maarika Vaara,
- Juha Koskenkari,
- Pasi Ohtonen,
- Ari Karttunen,
- Pekka Raatikainen,
- and Tero Ala-Kokko


Technical Communication: Validation of a Stand-Alone Near-Infrared Spectroscopy System for Monitoring Cerebral Autoregulation During Cardiac Surgery

- Masahiro Ono,
- Yueying Zheng,
- Brijen Joshi,
o Jeffrey C. Sigl,
o and Charles W. Hogue


Analgesia

Pain Medicine

鞘內注射巴氯芬對複雜性區域疼痛綜合征中不同性質疼痛的功效
(唐 瑋 譯 馬皓琳 李士通校)

Efficacy of Intrathecal Baclofen on Different Pain Qualities in Complex Regional Pain Syndrome

o Anton A. van der Plas,
o Monique A. van Rijn,
o Johan Marinus,
o Hein Putter,
o and Jacobus J. van Hilten


Pain and Analgesic Mechanisms

Pan-caspase 抑制劑可減少患坐骨神經慢性擠壓傷的大鼠的心肌細胞凋亡和神經病理性疼痛
(周玲譯 薛張綱校)

A Pan-Caspase Inhibitor Reduces Myocyte Apoptosis and Neuropathic Pain in Rats with Chronic Constriction Injury of the Sciatic Nerve

o Georg Gradl,
o Philipp Herlyn,
o Burkhard Finke,
o Philip Gierer,
o Andreas Wree,
o Martin Witt,
o Thomas Mittlmeier,
o and Brigitte Vollmar
Persistent Hyperalgesia in the Cisplatin-Treated Mouse as Defined by Threshold Measures, the Conditioned Place Preference Paradigm, and Changes in Dorsal Root Ganglia Activated Transcription Factor 3: The Effects of Gabapentin, Ketorolac, and Etanercept

Hue Jung Park, Jennifer A. Stokes, Elaine Pirie, James Skahen, Yuri Shtaerman, and Tony L. Yaksh

The Involvement of Potassium Channels in the Peripheral Antiedematogenic Effect of Intrathecally Injected Morphine in Rats

Vanessa R.S. Foletto, Maria A. Martins, and Carlos R. Tonussi

The Effects of Electroacupuncture on the Extracellular Signal-Regulated Kinase 1/2/P2X3 Signal Pathway in the Spinal Cord of Rats with Chronic Constriction Injury

Jianbo Yu, Cong Zhao, and Xiaoqin Luo
High-Versus Low-Stimulation Current Threshold for Axillary Plexus Blocks: A Prospective Randomized Triple-Blinded Noninferiority Trial in 205 Patients

Timon Vassiliou, Hans-Helge Müller, Angela Ellert, Pascal Wallot, Kuo-Min Kwee, Michaela Beyerle, Leopold Eberhart, Hinnerk Wulf, and Thorsten Steinfeldt


Current Status of Pharmacologic Therapies in Patient Blood Management

Lawrence Tim Goodnough, MD* and Aryeh Shander, MD†

From the *Pathology Department, Stanford University, Stanford, California; and †Department of Anesthesiology, Critical Care Medicine, Pain Management, and Hyperbaric Medicine, Englewood Hospital and Medical Center, Englewood, New Jersey.

Patient blood management incorporates patient-centered, evidence-based medical and surgical approaches to improve patient outcomes by relying on the patient’s own (autologous) blood rather than allogeneic blood. Particular attention is paid to preemptive measures such as anemia management. The emphasis on the approaches being “patient-centered” is to distinguish them from previous approaches in transfusion medicine, which have been “product-centered” and focused on blood risks, costs, and inventory concerns rather than on patient outcomes. Patient blood management structures its goals by avoiding blood transfusion with effective use of alternatives to allogeneic blood transfusion. These alternatives include autologous blood procurement, preoperative autologous blood donation, acute normovolemic hemodilution, and intra/postoperative red blood cell (RBC) salvage and reinfusion. Reviewed here are the available
pharmacologic tools for anemia and blood management: erythropoiesis-stimulating agents (ESAs), iron therapy, hemostatic agents, and potentially, artificial oxygen carriers.

**Opioid-Sparing Effect of Preemptive Bolus Low-Dose Ketamine for Moderate Sedation in Opioid Abusers Undergoing Extracorporeal Shock Wave Lithotripsy: A Randomized Clinical Trial**

Babak Gharaei, MD*, Alireza Jafari, MD*, Homayoun Aghamohammadi, MD*, Mohamadreza Kamranmanesh, MD*, Mahtab Poorzamani, MD*, Hedayatollah Elyassi, MD†, Baharak Rostamian, MD* and Alireza Salimi, MD‡

From the *Anesthesiology Research Center, Labbafinejad Hospital, Shahid Beheshti University of Medical Sciences; †Anesthesiology Research Center, Taleghani Hospital; and ‡Anesthesiology Research Center, Loghman Hospital, Tehran, Iran

Anesth Analg January 2013 116:75-80

**BACKGROUND:** Ketamine has been used as part of a multimodal analgesia regime in opioid abusers undergoing general anesthesia. We studied the opioid-sparing effect of a very low-dose bolus of ketamine as part of moderate sedation for opioid abuse patients undergoing extracorporeal shock wave lithotripsy.

**METHODS:** In this randomized, placebo-controlled clinical trial, 190 opioid abusers were enrolled. They were stratified into 2 blocks based on their daily opioid consumption. Both blocks were then randomized to receive 0.1 mg/kg IV ketamine (group K) or placebo (group P). Lithotripsy was performed under moderate sedation with intermittent bolus doses of remifentanil (0.2 µg/kg) to alleviate pain. The total remifentanil dose (primary outcome) and respiratory adverse events (secondary outcome) were compared in the 2 groups.
RESULTS: Remifentanil administration in the group with low-opioid consumers was 1.6 ± 0.4 µg/kg (group P) compared with 1.0 ± 0.2 µg/kg in group K (confidence interval [CI] of difference 95%, 0.4–0.7; P < 0.001). Patients who had high-opioid consumption received 2.0 ± 0.5 µg/kg (group P) vs 1.5 ± 0.3 µg/kg (group K) remifentanil (CI of difference 95%, 0.40–0.75; P < 0.001). Ready to discharge time was statistically longer in high-consumption opioid abusers who received placebo compared with group K (55 ± 13 minutes vs 44 ± 8 minutes, CI of difference 95%, 6–15; P < 0.001). The incidences of bradypnea, apnea, nausea, vomiting, and hemodynamic changes were not statistically different between the ketamine and placebo groups.

CONCLUSION: Preemptive low-dose ketamine (0.1 mg/kg) as a bolus has opioid-sparing effects in opioid abusers undergoing moderate sedation.

多腔輸液設備對於已知藥物物理不相溶現象的影響：一項控制性體外研究
The impact of multilumen infusion devices on the occurrence of known physical drug incompatibility: a controlled in vitro study.

Aurélie Foinard, MSc*, Bertrand Décaudin, PhD*†, Christine Barthélémy, PhD*, Bertrand Debaene, PhD† and Pascal Odou, PhD*†
From the *Department of Biopharmacy, Galenic and Hospital Pharmacy, Université Lille Nord de France, Lille; †Department of Pharmacy, Lille University Hospital, Lille; and ‡Anesthesia and Intensive Care Department, University Hospital, Poitiers, France.

Background: Drug incompatibility is a problem, especially when managing patients in intensive care units. We designed the present study to assess the impact of multilumen infusion access devices on the occurrence of known physical drug incompatibility through a controlled in vitro study.

Methods: Three infusion devices connected to a single-lumen catheter were studied: a standard set with 2-port manifold and 1-m extension set and 2 multilumen infusion access
devices: a 3-lumen extension set and a 9-lumen extension set (Edelvaiss-Multiline™; Doran International, Toussieu, France). For the 9-lumen extension set, 3 infusion access combinations were studied. Furosemide, midazolam, and saline were infused simultaneously through 3 infusion devices. Three concentrations of furosemide were tested. The infusion rate of saline (carrier) was initially set at 100 mL/h and stepwise decreased by 10 mL/h until precipitate formation. Physical incompatibility was assessed by 2 tests: visual inspection and the subvisible particle count test according to the European Pharmacopeia. The lowest saline infusion rate to prevent visible precipitate and attain an acceptable particle count (i.e., to pass "the 2 tests") was reported for each infusion set.

**RESULTS:** The standard set revealed visible precipitate even at the highest saline flow rate (100 mL/h). The 3-lumen device prevented drug precipitation using the 2 lowest furosemide concentrations with a saline infusion rate that decreased with furosemide concentration. The 9-lumen infusion access device prevented drug precipitation whatever the furosemide concentration for 2 access combinations using saline infusion rates of between 20 and 60 mL/h but not for a third access combination, despite saline infusion rates equal to 100 mL/h.

**CONCLUSIONS:** Infusion device characteristics appear to have an impact on the physical compatibility of the 2 drugs. Under specified conditions, the 9-lumen infusion access device prevents physical furosemide-midazolam incompatibility.

**Estimate of the relative risk of succinylcholine for triggering malignant hyperthermia.**

Franklin Dexter, MD, PhD*, Richard H. Epstein, MD, CPHIMS†, Ruth E. Wachtel, PhD, MBA‡ and Henry Rosenberg, MD§

From the *Division of Management Consulting, Department of Anesthesia, University of Iowa, Iowa City, IA; †Department of Anesthesiology, Jefferson Medical College, Philadelphia, PA; ‡Department of Anesthesia, University of Iowa, Iowa City, IA; and §Malignant Hyperthermia Association of the United States; Department of Medical Education and Clinical Research, Saint Barnabas Medical Center, Livingston, NJ.

Anesth Analg January 2013 116:118-122

**評估琥珀膽鹼引發惡性高熱的風險**

**背景**: 惡性高熱 MH 應用丹曲林和吸入麻醉藥治療，麻醉中出現的惡性高熱，應用丹曲林只需 1 天 1 個劑量即可起效；不用吸入麻醉劑，通過已註冊資料和麻醉期間使用司可林的次數，估計丹曲林的需要量，以此判斷司可林引起 MH 的幾率。

**方法**: 司可林引發 MH 的原因有兩個，來自 MHAUS 的 284 名患者分別被分為 2 組，一組接受司可林的 MH 患者，另一種不給予司可林的 MH 患者，麻醉劑與琥珀膽鹼的比例的估計，使用麻醉資訊管理系統的資料來自一個典型的北美醫院由三級手術室、產科手術中心, 和內鏡和放射室。

**結果**: 用司可林引起 MH 的相對危險度比沒用司可林組高 19.6（低 95% 可信區間大於 16.1）。揮發性麻醉藥的相對風險為 9.1（大於 7.5）。兩個相對風險超過 1.0(P < 0.0001)。在北美，使用司可林的患者超過一半出現 MH，而沒使用司可林的患者出現 MH 的不足一半。與揮發性麻醉藥相比，琥珀膽鹼的發生率在醫院使用分別為 5.8% 和 11.6%。
BACKGROUND: Facilities with volatile anesthetic agents stock dantrolene for the treatment of malignant hyperthermia (MH). The availability of dantrolene at these facilities satisfies cost-utility norms even for sites with as few as 1 anesthetic per workday, based on the overall incidence of MH per anesthetic. We considered the stocking of dantrolene at facilities with succinylcholine alone (i.e., where volatile anesthetics are not available), by using registry data and estimates of the frequency of administration of succinylcholine during anesthesia. We determine the magnitude of the relative risk of the administration of succinylcholine for triggering MH.

METHODS: The relative risk of triggering MH by succinylcholine versus volatile agents was calculated using data from 2 sources. The ratio of the number of cases of MH among patients receiving succinylcholine to number among patients not receiving succinylcholine was estimated from the previously published cohort of 284 cases of MH from the North American MH Registry of the MH Association of the United States (MHAUS). The percentage of anesthetics with succinylcholine was estimated using anesthesia information management system data from a typical North American hospital comprising tertiary operating rooms, obstetrics unit, ambulatory surgical center, and endoscopy and radiological suites.

RESULTS: The relative risk of MH with versus without succinylcholine was 19.6 (lower 95% confidence limit > 16.1). Limiting to cases with volatile anesthetics, the relative risk was 9.1 (>7.5). Both relative risks exceed 1.0 (P < 0.0001). Because more than half of the reported cases of MH included the use of succinylcholine, the relative risk exceeded 1.0 provided fewer than half of anesthetics in North America included the use of succinylcholine. The incidences of succinylcholine use at the hospital were 5.8% and 11.6% for all anesthetics and for anesthetics with volatile agents, respectively.

CONCLUSIONS: Our results provide no insight into the triggering mechanism for MH (i.e., succinylcholine could in isolation have an extremely low incidence of inducing MH, yet markedly increase the risk when administered in combination with volatile anesthetics). Until more epidemiologic data are collected and analyzed, having dantrolene available, where succinylcholine may be used, is reasonable, and this practice should be maintained.
背景：在2相初步研究中，我們已經觀察到使用Episure™ 自動檢測™（彈簧）注射器能成功地確定硬膜外腔。在這項研究中，我們評估經驗豐富的麻醉師使用彈簧式注射器對建立成功的硬膜外分娩鎮痛（主要結果），放置硬膜外導管的時間，以及學習曲線（累計匯總分析，即CUSUM）的影響。

方法：14位麻醉科主治醫生隨機使用帶弹簧式注射器或傳統的玻璃注射器每人實施50個連續硬膜外麻醉技術。10位參加者使用這兩種注射器每人完成另外一個50個連續硬膜外麻醉技術。

結果：一共實施了1200個硬膜外技術。使用彈簧式注射器與使用傳統的玻璃注射器實施硬膜外麻醉相比在獲得鎮痛成功方面兩者成功率無顯著差異，（絕對差異為1.0%，95%可信區間，CI：-8.9%至10.8%），前者硬膜外導管放置平均時間更短（比為0.92 95% CI:0.89-0.96），P = 0.003）兩者有類似累積曲線相。麻醉科醫生鎮痛成功更常見（絕對差為34.6%，95%CI，14.9%-54.3%，P <0.001），首選連續生理鹽水阻力消失法與使用間歇空氣法相比鎮痛成功更常見（絕對差為 33.8% 95%CI，12.6%-55.0%，P <0.001）。我們也觀察到首選使用彈簧式注射器組所用的平均時間更短（比為0.65 95% CI，0.62 -0.67, P= 0.02）。

結論：在建立成功的硬膜外分娩鎮痛方面，經驗豐富的產科麻醉醫師使用彈簧式注射器與使用常規的玻璃注射器相比有類似的整體比率，硬膜外導管插入時間更短，有類似的累積曲線，尤其是當麻醉師第一次隨機使用新型注射器時。參加的麻醉師使用新型注射器時偏愛連續生理鹽水阻力消失法時鎮痛成功率更高。

（習盼譯 薛張綱校）

BACKGROUND: The Episure™ AutoDetect™ (spring-loaded) syringe has been observed to successfully identify the epidural space in 2 pilot studies. In this study we evaluated the impact of the spring-loaded syringe on the establishment of successful epidural labor analgesia (primary outcome), elapsed time for catheter placement, and learning curve (cumulative summary analysis, i.e., Cusum) of experienced anesthesiologists.

METHODS: Fourteen attending and fellow anesthesiologists were randomized to perform 50 consecutive epidural technique attempts using a spring-loaded or conventional glass syringe. Ten participants completed an additional 50 attempts with the alternate syringe in a crossover design.

RESULTS: A total of 1200 epidural placement attempts were performed. Use of the spring-loaded syringe was associated with a nonsignificant difference of estimated success rate in obtaining analgesia success (absolute difference of 1.0% 95% confidence interval, CI: -8.9% to 10.8%), shorter elapsed mean time to epidural catheter placement (ratio of 0.92 95% CI, 0.89-0.96); P = 0.003) and similar Cusum curves when compared with a conventional glass syringe. Analgesia success was more common with attending versus fellow anesthesiologists (absolute difference of 34.6% 95% CI, 14.9% to 54.3%; P < 0.001), and when the initial preferred technique was loss-of-resistance to continuous saline versus intermittent air (absolute difference of 33.8% 95% CI, 12.6% to 55.0%; P < 0.001). Shorter elapsed mean times were also observed in the group exposed to the spring-loaded syringe first (ratio of 0.65 95% CI, 0.62-0.67; P = 0.02).

CONCLUSIONS: When used by experienced obstetric anesthesiologists, the spring-loaded syringe was associated with a similar overall rate for establishing successful epidural labor analgesia, a shorter elapsed time to epidural catheter insertion, particularly when the anesthesiologist was randomized to use the novel syringe first, and a similar Cusum curve when compared with a conventional glass syringe. Attending versus fellow anesthesiologists and an
An observational study of the fresh frozen plasma: red blood cell ratio in postpartum hemorrhage.

Pierre Pasquier, MD*, Etienne Gayat, MD, PhD†, Thibaut Rackelboom, MD‡, Julien La Rosa, MD‡, Abeer Tashkandi, MD†, Antoine Tesniere, MD, PhD‡, Julie Ravinet, MD§, Jean-Louis Vincent, MD, PhD∥, Vassilis Tsatsaris, MD, PhD§, Yves Ozier, MD, PhD*, François Goffinet, MD, PhD§ and Alexandre Mignon, MD, PhD‡

From the *Département d’Anesthésie-Réanimation, Hôpital d’Instruction des Armées Bégin, Saint-Mandé; †Département d’Anesthésie-Réanimation, Hôpital Lariboisière, Assistance Publique—Hôpitaux de Paris, Université Paris Diderot, Paris; ‡Département d’Anesthésie-Réanimation and §Maternité Port-Royal, Hôpital Cochin, Assistance Publique—Hôpitaux de Paris, Université Paris Descartes, Paris, France; and ‖Department of Intensive Care, Erasme Hospital, Université Libre de Bruxelles, Brussels, Belgium.

Anesth Analg January 2013 116:155-161

BACKGROUND: Postpartum hemorrhage is the leading cause of maternal death worldwide. Recent data from trauma patients and patients with hemorrhagic shock have suggested that an increased fresh frozen plasma:red blood cell (FFP:RBC) ratio may be of benefit in massive bleeding. We addressed this issue in cases of severe postpartum hemorrhage.
METHODS: We reviewed data from all patients diagnosed with severe postpartum hemorrhage during a 4-year period (2006-2009). Patients who were treated with sulprostone and required transfusion within 6 hours of delivery were included in the study and were divided into 2 groups according to their response to sulprostone: bleeding controlled with sulprostone alone (sulprostone group) and bleeding requiring an additional advanced interventional procedure including arterial angiographic embolization and/or surgical procedures (arterial ligation, B-Lynch suture, or hysterectomy; intervention group). The requirement or no requirement for advanced procedures constituted the primary end point of the study. Propensity scoring was used to assess the effect of a high FFP:RBC ratio on bleeding control.

RESULTS: patients were transfused with both RBCs and FFP. The FFP:RBC ratio increased over the study period (P < 0.001), from 1:1.8 at the start to 1:1.1 at the end of the study period. After propensity score modeling (inverse probability of treatment weighting), a high FFP:RBC ratio was associated with lower odds for advanced interventional procedures (odds ratio [95% confidence interval], 1.25 [1.07-1.47]; P = 0.008). There were no deaths, severe organ dysfunction, or other complications as a consequence of severe postpartum hemorrhage.

CONCLUSIONS: In this retrospective study, a higher FFP:RBC ratio was associated with a lower requirement for advanced interventional procedures in the setting of postpartum hemorrhage. The benefits of transfusion using a higher FFP:RBC ratio should be confirmed by randomized-controlled trials.

Repolarization abnormalities in patients with subarachnoid and intracerebral hemorrhage: predisposing factors and association with outcome
Eija Juntila, MD＊, Maarika Vaara, MD＊, Juha Koskenkari, MD, PhD＊, Pasi Ohtonen, MSc†, Ari Karttunen, MD, PhD‡, Pekka Raatikainen, MD, PhD§ and Tero Ala-Kokko, MD, PhD＊ From the ＊Department of Anesthesiology, Division of Intensive Care, Oulu University Hospital; †Departments of Anesthesiology and Surgery, Oulu University Hospital; ‡Department of Radiology, Oulu University Hospital; §Department of Internal Medicine, Oulu University Hospital, Oulu, Finland; and ＊Heart Center Co, Tampere University Hospital, Tampere, Finland. Anesth Analg January 2013 116:190-197

背景：顱內病變患者中心電圖異常十分常見。在本研究中, 我們評估了複極異常的誘發因素，例如，校正的 QT (QTc) 間期延長，心電圖缺血樣改變和複極末形態異常，並且考察了需要重症監護的蛛網膜下腔出血和腦出血患者中這些異常指標的預示價值。

方法：本研究在重症監護室進行，是一個前瞻性、觀察性的研究。入院時記錄患者的臨床特徵，意識水準，早期的頭顱 CT 結果。研究分為三階段，各兩天。每個階段均記錄12導聯心電圖、經胸心超、電解質和肌鈣蛋白I等指標，以及血管活性藥物和靜脈藥物輸注速度。複極異常比如 QTc 間期延長（毫秒），心電圖缺血樣改變，複極末形態異常（存在/缺失）等被評估和分析。格拉斯哥預後評分評估一年功能恢復。

結果：本研究歷時兩年，共108名患者參加研究。在兩種出血類型中不同的複極異常均頻繁發生。QTc 間期延長易發生於女性 (β, 24.5; P = 0.010) 和使用異丙酚者 (β, 30.5; P = 0.001)。心電圖缺血樣改變易發生於男性 (比值比 [OR], 5.9; P = 0.003)，複極末形態異常易
BACKGROUND: Electrocardiographic (ECG) abnormalities are frequent in patients with intracranial insult. In this study, we evaluated the factors predisposing to the repolarization abnormalities, i.e., prolonged corrected QT (QTc) interval, ischemic-like ECG changes and morphologic end-repolarization abnormalities, and examined the prognostic value of these abnormalities in patients with subarachnoid and intracerebral hemorrhages requiring intensive care.

METHODS: This was a prospective, observational clinical study in a university-level intensive care unit. Clinical characteristics, the level of consciousness, and findings in primary head computed tomography were recorded on admission. The study period was divided into three 2-day sections. In each section, a 12-lead ECG, transthoracic echocardiography, the results of standard blood electrolytes and cardiac troponin I, as well as the rate of vasoactive and sedative drug infusions were recorded. Repolarization abnormalities such as prolongation of the QTc interval (millisecond), ischemic-like ECG changes, and morphologic end-repolarization abnormalities (present/absent) were evaluated and analyzed. The 1-year functional outcome was determined using the Glasgow Outcome Score.

RESULTS: During the 2-year study period, 108 patients were included in the study. Different repolarization abnormalities were frequent in both types of hemorrhage. Prolongation of the QTc interval was predisposed by female gender (β, 24.5; P = 0.010) and the use of propofol (β, 30.5; P = 0.001). The predisposing factor for ischemic-like ECG changes were male gender (odds ratio [OR], 5.9; P = 0.003) and for morphological end-repolarization abnormalities aneurysmatic bleeding (OR, 13.0; P = 0.002). Ischemic-like ECG changes were common, in 87/108 patients during the study period, and were associated with a poorer 1-year functional outcome (OR, 4.7; lower 95% confidence interval, 1.5; P = 0.010).

CONCLUSIONS: Each repolarization abnormality has characteristic predisposing factors. Ischemic-like ECG changes are common and are associated with a poorer 1-year functional outcome.

pan-caspase 抑制劑可減少患坐骨神經慢性擠壓傷的大鼠的心肌細胞凋亡和神經病理性疼痛

A pan-caspase inhibitor reduces myocyte apoptosis and neuropathic pain in rats with chronic constriction injury of the sciatic nerve.

Georg Gradl, MD*, Philipp Herlyn, MD†, Burkhard Finke, MD‡, Philip Gierer, MD*, Andreas Wree, MD‡, Martin Witt, MD§, Thomas Mittlmeier, MD* and Brigitte Vollmar, MD†

From the Departments of *Traumatology and Reconstructive Surgery and †Anatomy, ‡Institute for Experimental Surgery, and §Electron Microscopy Centre, Medical Faculty, University of Rostock, Rostock, Germany.

細胞凋亡被認為可能是疼痛和運動障礙的一個啓動子。我們的這項研究是要證明在此動物模型中，細胞凋亡是否會引起神經病理性疼痛。

方法：為了弄清這個問題，我們對患有坐骨神經的慢性擠壓傷而產生神經病理性疼痛的大鼠進行治療，並觀察其結果。動物接受 pan-caspase 抑制劑 ZVAD（OMe）-fmk（N = 5）或相等劑量的賦形藥（n = 6）。假手術組作為對照組（n = 6）。

結果：在神經結紮後第 4 天，各實驗組沒有任何灌注異常或肌肉組織炎症的跡象。然而，賦形藥組出現明顯的細胞凋亡，而這在用 ZVA-D 治療的動物中是幾乎完全阻斷的。與假手術對照組的結果相比，用 ZVA-D 治療的動物對由熱、冷和機械性刺激引起的疼痛明顯減少。

結論：從本次神經病理性疼痛的實驗模型發現，細胞凋亡可能有助於產生溫度和機械性異常痛覺。神經病理性疼痛的症狀發展並不依賴於微循環紊亂或肌肉組織的炎症。

細胞外信號調節激酶 1/2/P2X3 信號轉導通路在慢性坐骨神經損傷的大鼠脊髓的電針影響。

The Effects of Electroacupuncture on the Extracellular Signal-Regulated Kinase 1/2/P2X3 Signal Pathway in the Spinal Cord of Rats with Chronic Constriction Injury.
Jianbo Yu, PhD*, Cong Zhao, MD*† and Xiaqin Luo, MD‡
From the *Department of Anesthesiology, Tianjin Nankai Hospital, Tianjin Medical University, Tianjin; †Department of Anesthesiology, Chengdu Fifth People’s Hospital, Sichuan Province; and ‡Department of Pathology, Xiangyang First People’s Hospital, Hubei Province, China.
Anesth Analg January 2013 116:239-246
背景：作为传统的治疗疼痛的方法，电针（EA）被广泛用于临床，但其镇痛疗效尚未明确。在本研究中，我们探讨EA对慢性疼痛的疗效以及患有慢性压迫性脊髓损伤的大鼠P2X3受体的表达。

方法：本研究分2部分。第1部分，将SD大鼠分成6组（n=10）：假-CCI，CCI，LEA；CCI+2Hz（电针穴位），HEA；CCI+15Hz（电针刺激穴），NA-LEA（CCI+2Hz电针非穴位）以及NA-LEA（CCI+15Hz电针非穴位）。4至9天CCI后电针治疗隔天一次，伤害性刺激由电子触觉测量仪和热板装置执行。印迹检查测量脊髓内蛋白和P2X3受体的mRNA以及即时聚合酶链反应。第2部分，大鼠被分成5组（n=10）：假-CCI，CCI，EA（CCI+EA电针穴位），NA-EA（CCI+EA非穴位）以及U0126（CCI+鞘内注射U0126）。EA治疗和第一部分相似，给予U0126组大鼠鞘内注射5ugU0126和5%二甲基亚砜。十微升作为其他四组的载体在CCI治疗后4至9天每两天给一次。使用印迹法检查细胞外信号调节激酶1/2（ERK1/2）和其磷酸化。

结果：电针治疗具有显著的镇痛效果，减少了CCI引起的脊髓P2X3受体蛋白和mRNA表达增加。此外，相比15Hz，2Hz电针具有较好的止痛效果，同时15Hz电针穴位的取穴比2Hz电针治疗大鼠脊髓P2X3受体在蛋白和mRNA水平的降低。无论是EA在穴位或鞘内注射U0126释然异常性疼痛和痛觉过敏和降低P2X3受体的表达和ERK1/2磷酸化的脊髓。

结论：本研究显示，EA能部分缓解神经性疼痛的行，並通過ERK1/2信號通路在脊髓P2X3受体的表达减少。相比高频EA，低频EA对神经病理性疼痛有较好的止痛效果。

（杨琰译 薛张纲校）

BACKGROUND: Electroacupuncture (EA), as a traditional clinical method, is widely accepted in pain clinics, but the analgesic effect of EA has not been fully demonstrated. In the present study, we investigated the effect of EA on chronic pain and expression of P2X3 receptors in the spinal cord of rats with chronic constriction injury (CCI).

METHODS: The study was conducted in 2 parts. In part 1, Sprague Dawley rats were divided into 6 groups (n = 10): sham-CCI, CCI, LEA; CCI + 2 Hz EA at acupoints), HEA; CCI + 15 Hz EA at acupoints), NA-LEA (CCI + 2 Hz EA at nonacupoints), and NA-HEA (CCI + 15 Hz EA at nonacupoints). EA treatment was performed once a day on days 4 to 9 after CCI. Nociception was assessed using von Frey filaments and a hotplate apparatus. The protein and the messenger RNA (mRNA) levels of P2X3 receptors in the spinal cord were assayed by Western blotting and real-time polymerase chain reaction, respectively. In part 2, rats were divided into 5 groups (n = 10): sham-CCI, CCI, EA (CCI + EA at acupoints), NA-EA (CCI + EA at nonacupoints), and U0126 (CCI + intrathecal injection of U0126). EA treatment was conducted similar to part 1. Rats were given 5 µg U0126 in the U0126 group and 5% dimethyl sulfoxide intrathecally. Ten microliters was used as a vehicle for the other 4 groups twice a day on days 4 to 9 after CCI. Extracellular signal-regulated kinase 1/2 (ERK1/2) and ERK1/2 phosphorylation in the spinal cord were also assayed by Western blotting.

RESULTS: EA treatment exhibited significant antinociceptive effects and reduced the CCI-induced increase of both protein and mRNA expression of P2X3 receptors in the spinal cord. Furthermore, 2 Hz EA had a better analgesic effect than 15 Hz EA, and the protein and mRNA level of P2X3 receptor in spinal cord were lower in rats treated with 2 Hz EA at acupoints than 15 Hz EA at acupoints. Either EA at acupoints or intrathecal injection of U0126 relieved allodynia and hyperalgesia and reduced the expression of P2X3 receptors and ERK1/2 phosphorylation in the spinal cord.
CONCLUSIONS: The data demonstrated that EA alleviates neuropathic pain behavior, at least in part, by reducing P2X3 receptor expression in spinal cord via the ERK1/2 signaling pathway. Low frequency EA has a better analgesic effect than high frequency HEA on neuropathic pain.

Review Article: Safety of Modern Starches Used During Surgery
Philippe Van Der Linden, MD, PhD*, Michael James, MB ChB, PhD, FRCA, FCA(SA)‡, Michael Mythen, MD FRCA‡§ || and Richard B. Weiskopf, MD ||
From the *Service D’Anesthesie–Réanimation, CHU Brugmann, Bruxelles, Belgium; †Department of Anaesthesia, University of Cape Town, Cape Town, South Africa; ‡University College London; §University College London Hospitals NHS Foundation Trust/University College London and Royal Free London NHS Foundation Trust, Research Support Centre; || Department of Health, Enhanced Recovery Partnership, London, United Kingdom; and || Department of Anesthesia and Perioperative Care, University of California, San Francisco, San Francisco, California.
Anesth Analg January 2013 116:35-48

Various hydroxyethyl starch (HES) preparations have been used for decades to augment blood volume. There has been concern recently regarding possible adverse outcomes when using HES in the intensive care setting, especially in patients with septic shock. However, the pharmacokinetic and pharmacodynamic properties of HES preparations depend on their chemical composition and source material. Thus, different clinical conditions could result in differing effectiveness and safety for these preparations. Consequently, we assessed the safety of tetrastarches when used during surgery, using a formal search, that yielded 59 primary full publications of studies that met a priori inclusion criteria and randomly allocated 4529 patients with 2139 patients treated with tetrastarch compared with 2390 patients treated with a comparator. There were no indications that the use of tetrastarches during surgery induces adverse renal effects as assessed by change or absolute concentrations of serum creatinine or need for renal replacement therapy (39 trials, 3389 patients), increased blood loss (38 trials, 3280 patients), allogeneic erythrocyte transfusion (20 trials, 2151 patients; odds ratio for HES transfusion 0.73 [95% confidence interval = 0.61–0.87], P = 0.0005), or increased mortality (odds ratio for HES mortality = 0.51 [0.24–1.05], P = 0.079).
布比卡因和羅呱卡因的心肌蓄積與線粒體逆轉效應和心功能降低相關

Myocardial Accumulation of Bupivacaine and Ropivacaine Is Associated with Reversible Effects on Mitochondria and Reduced Myocardial Function

Nicole Hiller, MSc*, Peter Mirtschink, MD‡, Christine Merkel, MD*, Lilla Knels, MD‡, Reinhard Oertel, PhD§, Torsten Christ, MD ||, Andreas Deussen, MD||, Thea Koch, MD* and Sebastian N. Stehr, MD, DESA*

From the *Department of Anesthesiology and Intensive Care Medicine, ‡Institute of Anatomy, §Institute of Clinical Pharmacology, || Department of Pharmacology and Toxicology, and ¶Institute of Physiology, University Hospital Dresden, Dresden, Germany; and †Institute of Molecular Health Science, ETH Zurich, Switzerland.


背景：局麻藥引起心臟毒性的機理至今仍未完全清楚。此項研究分析了能引起臨床毒性

的局麻藥濃度是否會影響心肌線粒體結構和氧耗。

方法：攜離出豚鼠心臟建立 Langendorff 離體心臟模型，將其暴露於布比卡因 (3.0 和 7.5 

μg/mL) 和羅呱卡因 (3.6 和 9.0 μg/mL) 10min。記錄心率，收縮壓，左室內壓變化速率，

心電圖和冠脈流量。給藥後即刻或 20min 洗脫期、60min 洗脫期後測量組織的局麻藥濃

度。另外，使用電子顯微鏡觀察心肌線粒體並對其結構損傷進行評分。將心肌細胞放入布

比卡因液進行培養，測定氧耗率、細胞外酸化情況和 PGC-1α mRNA 相對量（一種細胞能

量代謝的調節因數）的表達。

結果：布比卡因和羅呱卡因引起 PR 間期及 QRS 可逆性延長，左室內壓及其變化率降

低。心肌組織中局麻藥濃度是動脈濃度的 3 倍。局麻藥應用後線粒體表現出顯著的濃度依

賴性形態腫脹。對於羅呱卡因和布比卡因，這些變化分別在進行 20min、60min 洗脫期後

被逆轉。布比卡因在乳鼠心肌細胞培養中，降低線粒體氧耗，增加 PGC-1α 的表達，而脂

肪酸代謝不受影響。

結論：布比卡因和羅呱卡因可在心肌組織中蓄積。當達到負性肌力作用的濃度時，局麻

藥可逆性引起線粒體腫脹。布比卡因降低細胞代謝，然而此效應可被脂肪酸逆轉。與線粒

體的相互作用可能是導致局麻藥負性肌力作用的原因。

（諸琳婕 譯 陳傑 校）

BACKGROUND: Mechanisms of local anesthetic cardiac toxicity are still not completely
understood. In this study, we analyzed whether concentrations of local anesthetics found in
clinical toxicity affect myocardial mitochondrial structure and oxygen consumption.

METHODS: Guinea pig isolated heart Langendorff preparations were exposed to bupivacaine
(3.0 and 7.5 μg/mL) and ropivacaine (3.6 and 9.0 μg/mL) for 10 minutes. Heart rate, systolic
blood pressure, the first derivative of left ventricular pressure (+dP/dt), electrocardiogram, and
coronary flow were recorded. The local anesthetic tissue concentration was measured either
immediately after local anesthetic exposure, or after 20- and 60-minute washout periods. In
addition, electron microscopy of myocardial mitochondria was performed using a scoring system
for structural damage of mitochondria. Cardiomyocyte cell culture was incubated with
bupivacaine, and oxygen consumption ratio, extracellular acidification, and relative amounts of
PGC-1α mRNA, a regulator of cellular energy metabolism, were determined.

RESULTS: Bupivacaine and ropivacaine induced reversible PR interval and QRS prolongation,
and left ventricular pressure and +dP/dt reduction. Myocardial tissue concentration of local
anesthetics was 3-fold the arterial concentration. Mitochondria showed a significant concentration-dependent morphological swelling after local anesthetic application. These changes were reversed by a 20-minute washout period for ropivacaine and by a 60-minute washout for bupivacaine. Bupivacaine reduced mitochondrial oxygen consumption and increased PGC-1α expression in neonatal cardiomyocyte cell cultures, whereas fatty acid metabolism remained unaffected.

CONCLUSIONS: Bupivacaine and ropivacaine accumulate in the myocardium. Reversible local anesthetic-induced mitochondrial swelling occurs at concentrations that induce a negative inotropic effect. Bupivacaine reduces cellular metabolism, whereas this reduction is reversible by fatty acids. Interaction with mitochondria may contribute to the negative inotropic effect of local anesthetics.

**Special Article: Curriculum and Cases for Pain Medicine Crisis Resource Management Education**

Gary J. Brenner, MD, PhD*,†‡, Jordan L. Nemark, MD* and Daniel Raemer, PhD*†‡

From the *Department of Anesthesia, Critical Care and Pain Medicine, Massachusetts General Hospital, Boston; †Center for Medical Simulation, and ‡Harvard Medical School, Cambridge, Massachusetts.


Medical crises that may occur in the setting of a pain medicine service are rare events that require skillful action and teamwork to ensure safe patient outcome. A simulated environment is an ideal venue for both acquisition and reinforcement of this knowledge and skill set. Here, we present an educational curriculum in pain medicine crisis resource management for both pain medicine fellows and attending physicians as well as the results of a successful pilot program.
METHODS: In this prospective, randomized, double-blind trial, patients scheduled for cesarean delivery were assigned to 1 of 4 groups: lidocaine 30 mg in the spinal or epidural space, or lidocaine 45 mg by the same routes. A blinded observer assessed the degree of sensory and motor block. The ability to identify intrathecal injection of each dose was compared. Sensory block above T6 dermatome and hypotension were recorded as side effects.

RESULTS: Intrathecal administration of lidocaine 30 mg produced rapid subjective and objective signs of neuroblockade within 3 minutes (100%, 95% confidence interval CI, 85%–100% for each). Lidocaine 45 mg produced similar results. All patients in both groups described their legs as warm or heavy after 3 minutes and had a motor block by 5 minutes. On the basis of an intrathecal catheter rate of 1:380, the observed negative predictive value for intrathecal placement if the patient described no sensory changes at 3 minutes was 100% (95% CI, 99.95%–100%) for 30 mg and 100% (95% CI, 99.93%–100%) for 45 mg. We did not identify a decrease in the rate of side effects with the lower dose.

CONCLUSIONS: Our results suggest that there is unlikely to be a large difference in the ability of these doses to detect unintentional intrathecal catheter placement. While the negative predictive value for intrathecal injection is very high for both doses, the 95% CI for the sensitivity of either dose is too wide to demonstrate clinical safety to identify all intrathecal catheters. A much larger study is warranted to assess whether there is a lower sensitivity with the 30-mg dose, or a propensity toward high cephalad motor block levels with the 45-mg dose.
BACKGROUND: There is currently no evidence about the genetic bases of postoperative pain variability in children.

METHODS: We prospectively followed a cohort of 168 children after orthopedic or abdominal surgery, who were under morphine patient-controlled analgesia. The children and their parents were genotyped for 6 candidate-gene polymorphisms (single-nucleotide polymorphisms [SNPs]) implicated in nociception and opiate metabolism: ABCB1C3435T, COMTVal158Met, NTRK1His40Tyr, OPRM1118G, POMCArg236Gln, and a haplotype of CYP2D6. Postoperative pain was assessed using the Faces Pain Scale (FPS), at rest and during mobilization, 11 times during the first 24 postoperative hours.

RESULTS: At rest, and to a lesser extent, at mobilization, having at least 4 pain peaks of FPS score >6 in 24 hours was more frequent in children with ABCB1_CC than in children with ABCB1_CT and ABCB1_TT (adjusted risk ratio = 4.5; 95% confidence interval [CI], 1.5–13.4; corrected CI for multiple comparisons, 0.98–20.55) and was more frequent in children with OPRM_GA than those with OPRM_AA (adjusted risk ratio = 3.5; 95% CI, 1.1–11.2; corrected CI, 0.70–17.30). After adjusting for parental mating type and correcting for multiple comparisons, mean FPS scores across the 24 postoperative hours were higher for OPRM_GA.
than for OPRM_AA at rest (P < 0.0002), higher for NTRK1_CT or NTRK1_TT than NTRK1_CC during mobilization (P = 0.002), and lower for COMT_GG than COMT_AA and
COMT_GA, during mobilization (P = 0.005).

CONCLUSIONS: ABCB1 and OPRM genotypes are associated with clinically meaningful pain variability, whereas NTRK1 and COMT are linked to subclinical effects. This first but small cohort study provides clues to further explore the genetic foundations of pediatric pain.
functional consequences in adulthood. Clinical retrospective reviews have suggested that multiple anesthetic exposures in early childhood are associated with learning disabilities later in life as well. Despite much concern about this phenomenon, little is known about the mechanism by which anesthetics initiate neuronal cell death. Caenorhabditis elegans, a powerful genetic animal model, with precisely characterized neural development and cell death pathways, affords an excellent opportunity to study anesthetic-induced neurotoxicity. We hypothesized that exposing the nematode to volatile anesthetics early in life would induce neuron cell death, producing a behavioral defect that would be manifested in adulthood.

METHODS: After synchronization and hatching, larval worms were exposed to volatile anesthetics at their 95% effective concentration for 4 hours. On day 4 of life, exposed and control worms were tested for their ability to sense and move to an attractant (i.e., to chemotax). We determined the rate of successful chemotaxis using a standardized chemotaxis index.

RESULTS: Wild-type nematodes demonstrated striking deficits in chemotaxis indices after exposure to isoflurane (ISO) or sevoflurane (SEVO) in the first larval stage (chemotaxis index: untreated, 85 ± 2; ISO, 52 ± 2; SEVO, 47 ± 2; P < 0.05 for both exposures). The mitochondrial mutant gas-1 had a heightened effect from the anesthetic exposure (chemotaxis index: untreated, 71 ± 2; ISO, 29 ± 12; SEVO, 24 ± 13; P < 0.05 for both exposures). In contrast, animals unable to undergo apoptosis because of a mutation in the pathway that mediates programmed cell death (ced-3) retained their ability to sense and move toward an attractant (chemotaxis index: untreated, 76 ± 10; ISO, 73 ± 9; SEVO, 76 ± 10). Furthermore, we discovered that the window of greatest susceptibility to anesthetic neurotoxicity in nematodes occurs in the first larval stage after hatching (L1). This coincides with a period of neurogenesis in this model. All values are means ± SD.

CONCLUSION: These data indicate that anesthetics affect neurobehavior in nematodes, extending the range of phyla in which early exposure to volatile anesthetics has been shown to cause functional neurological deficits. This implies that anesthetic-induced neurotoxicity occurs via an ancient underlying mechanism. C elegans is a tractable model organism with which to survey an entire genome for molecules that mediate the toxic effects of volatile anesthetics on the developing nervous system.

技術交流：驗證單機近紅外光譜系統在心臟手術中監測腦血流自動調節的作用
Technical Communication: Validation of a Stand-Alone Near-Infrared Spectroscopy System for Monitoring Cerebral Autoregulation During Cardiac Surgery
Masahiro Ono, MD, PhD*, Yueying Zheng, MD†, Brijen Joshi, MD‡, Jeffrey C. Sigl, PhD§ and Charles W. Hogue, MD ||
From the *Division of Cardiac Surgery, Department of Surgery, The Johns Hopkins University School of Medicine, Baltimore, Maryland; †Department of Anesthesiology, The First Affiliated Hospital, School of Medicine, Zhejiang University, Hangzhou, China; ‡Department of Internal Medicine, Sinai Medical Center, The Johns Hopkins University School of Medicine, Baltimore
Anesth Analg January 2013 116:198-204

背景:在體外迴圈(CPB)期間基於腦血流(CBF)自動調節功能監測的動脈血壓(ABP)個體化控制，可以提供一個比現行使用的監護標準更為有效的預防腦血流灌注不足的方法。經顱多普勒(TCD)可即時監測血流自動調節功能。之前研究證實近紅外光譜(NIRS)衍生出的局部腦氧飽和度(rScO2)監測可以在臨床上替代腦血流(CBF)自動調節功能的監測。本研
BACKGROUND: Individualizing arterial blood pressure (ABP) targets during cardiopulmonary bypass (CPB) based on cerebral blood flow (CBF) autoregulation monitoring may provide a more effective means for preventing cerebral hypoperfusion than the current standard of care. Autoregulation can be monitored in real time with transcranial Doppler (TCD). We have previously demonstrated that near-infrared spectroscopy (NIRS)–derived regional cerebral oxygen saturation (rScO2) provides a clinically suitable surrogate of CBF for autoregulation monitoring. The purpose of this study was to determine the accuracy of a stand-alone “plug-and-play” investigational system for autoregulation monitoring that uses a commercially available NIRS monitor with TCD methods.

METHODS: TCD monitoring of middle cerebral artery CBF velocity and NIRS monitoring were performed in 70 patients during CPB. Indices of autoregulation were computed by both a personal computer–based system and an investigational prototype NIRS-based monitor. A moving linear correlation coefficient between slow waves of ABP and CBF velocity (mean velocity index [Mx]) and between ABP and rScO2 (cerebral oximetry index [COx]) were calculated. When CBF is autoregulated, there is no correlation between CBF and ABP; when CBF is dysregulated, Mx and COx approach 1 (i.e., CBF and ABP are correlated). Linear regression and bias analysis were performed between time-averaged values of Mx and COx derived from the personal computer–based system and from COx measured with the prototype monitor. Values for Mx and COx were categorized in 5 mm Hg bins of ABP for each patient.
The lower limit of CBF autoregulation was defined as the ABP where Mx incrementally increased to ≥0.4.

RESULTS: There was correlation and good agreement between COx derived from the prototype monitor and Mx ($r = 0.510$; 95% confidence interval, 0.414–0.595; $P < 0.001$; bias, –0.07 ± 0.19). The correlation and bias between the personal computer–based COx and the COx from the prototype NIRS monitor were $r = 0.957$ (95% confidence interval, 0.945–0.966; $P < 0.001$ and 0.06 ± 0.06, respectively). The average ABP at the lower limit of autoregulation was 63 ± 11 mm Hg (95% prediction interval, 52–74 mm Hg). Although the mean ABP at the COx-determined lower limit of autoregulation determined with the prototype monitor was statistically different from that determined by Mx (59 ± 9 mm Hg; 95% prediction interval, 50–68 mm Hg; $P = 0.026$), the difference was not likely clinically meaningful.

CONCLUSIONS: Monitoring CBF autoregulation with an investigational stand-alone NIRS monitor is correlated and in good agreement with TCD-based methods. The availability of such a device would allow widespread autoregulation monitoring as a means of individualizing ABP targets during CPB.
BACKGROUND: Painful neuropathy is a dose-limiting side effect in cancer chemotherapy. To characterize this phenomenon, we examined pain behavior and analgesic actions in a mouse model of cisplatin polyneuropathy.

METHODS: Male C57BL/6 mice received intraperitoneal cisplatin or saline (2.3 mg/kg/d) every other day 6 times over 2 weeks for a total dose of 13.8 mg/kg. Thermal escape latencies, mechanical allodynia using von Frey hairs, and observation of behavior/morbidity and body weights were assessed. After onset of allodynia, we examined the actions of intraperitoneal gabapentin (100 mg/kg), etanercept (20 and 40 mg/kg), ketorolac (15 mg/kg), and morphine (1, 3, and 10 mg/kg). Additionally, using the conditioned place preference (CPP) paradigm, we examined the effects of gabapentin and ketorolac on the presumed pain state initiated by cisplatin. Additionally, we examined the spinal cord and dorsal root ganglia (DRG) of cisplatin-treated mice.

RESULTS: Cisplatin, but not saline treatment, produced persistent hindpaw tactile allodynia, which persisted 46 days with no effect on thermal escape. Gabapentin and morphine, but neither etanercept nor ketorolac, produced a complete but transient (2-hour) reversal of the allodynia. Etanercept (40 mg/kg) pretreatment resulted in a delay in onset of mechanical allodynia. Using CPP, gabapentin, but not ketorolac, in cisplatin animals resulted in a significant preference for the drug-associated treatment compartment. There was no place preference in non–cisplatin-treated (nonallodynic) mice after gabapentin injection. Immunohistochemistry in cisplatin-treated mice showed no change in glial fibrillary acidic protein (astrocyte) or Iba1 (ionized calcium binding adaptor molecule 1) (microglia) activation states, but a significant increase in activated transcription factor 3 was observed in the DRG.

CONCLUSIONS: Cisplatin-treated mice display allodynia and an activation of DRG activated transcription factor 3, which is paralleled by its effects on behavior in the CPP system, wherein gabapentin, but not ketorolac, in the presence of the cisplatin polyneuropathy, is positively rewarding, confirming that this neuropathy is an aversive (painful) state that is ameliorated by gabapentin.

用於腋路臂叢阻滯的高低刺激電流閾值對比：一項對 205 名病人的前瞻隨機三盲非劣效性試驗

High- Versus Low-Stimulation Current Threshold for Axillary Plexus Blocks: A Prospective Randomized Triple-Blinded Noninferiority Trial in 205 Patients

Timon Vassiliou, MD*, Hans-Helge Müller, PhD†, Angela Ellert, MD*‡, Pascal Wallot, MD*, Kuo-Min Kwee, MD*, Michaela Beyerle, MD*§, Leopold Eberhart, MD*, Hinnerk Wulf, MD* and Thorsten Steinfeldt, MD*

From the *Department of Anesthesiology and Critical Care, University Hospital Giessen-Marburg, Philipps University Marburg, Marburg, Germany; †Institute for Medical Informatics, Biometry and Epidemiology, Ludwig-Maximilians-University, Munich, Germany; ‡Department of Anesthesiology and Intensive Care, Hospital Reinbek St. Adolf-Stift, Reinbek, Germany; and §Department of Anesthesiology and Intensive Care Medicine, Asklepios-Klinik Harburg, Hamburg, Germany.


背景：對於神經刺激器引導下的局部麻醉，麻醉醫師必須在假定低成功率（使用高電流閾值）和假定增加神經損傷風險（使用低電流閾值）這二者之間做出權衡。針對神經刺激
引導的臂叢神經阻滯，本研究假設使用 0.9 至 1.1mA 範圍內的高電流閾值在操作和起效時間方面並不比 0.3 至 0.5mA 範圍內的低電流閾值要差。

方法：205 名計劃擇期手術病人被隨機分為低刺激電流閾值組（0.3-0.5mA, n=103）或者高刺激電流閾值組（0.9-1.1mA, n=102），兩組都使用 40mL局麻藥混合液（1%丙胺卡因和 0.75%羅呱卡因各 20mL）進行腋路臂叢神經阻滯。研究主要終點是感覺完全阻滯時間。次要結果是手術準備時間（定義為從開始阻滯至感覺完全阻滯時間）和操作時間。非劣效性邊界設為 5 分鐘並且以雙邊 95%的自舉置信區間（[CIs] 100,000 次重複）均數差來評價。

結果：低刺激電流閾值組的感覺完全阻滯平均時間顯著減少（17.9 ± 12.1 分鐘對 22.8 ± 12.4 分鐘; 95% 置信區間，1.1 至 8.6; p = 0.012）。低刺激電流閾值組的手術準 備時間為 30.3 ± 13.8 分鐘而高刺激電流閾值組為 31.7 ± 12.9 分鐘（95% 置信區間，-2.7 至 5.5; p = 0.49）。高刺激電流閾值組的操作時間明顯縮短（9.5 ± 4.7 分鐘對11.9 ± 5.7 分鐘; 95% 置信區間，-4 至 1.1; p = 0.001）。

結論：主次要終點都不能確認高電流閾值技術的非劣效性。但考慮到臨床實踐，在滿足手術要求的平均花費時間方面約 8.5 分鐘的差異是可以接受的。

（孫曉瓊 譯 陳傑 校）

BACKGROUND: For nerve stimulator-guided regional anesthesia, one has to compromise between a presumed low success rate (using a high-current threshold) and a presumed increased risk of nerve damage (using a low-current threshold). We hypothesized that high-current thresholds in the range of 0.9 to 1.1 mA are not inferior with respect to the procedural and latency times compared with low threshold currents in the range of 0.3 to 0.5 mA for nerve stimulation in brachial plexus blocks.

METHODS: Two hundred five patients scheduled for elective surgery were randomized to a low (0.3–0.5 mA, n = 103) or a high (0.9–1.1 mA, n = 102) stimulation current threshold for the axillary plexus block with 40 mL local anesthetic mixture (20 mL, each of prilocaine 1% and ropivacaine 0.75%). The primary end point was the time to complete sensory block. The secondary outcome measures were the time to readiness for surgery (defined as the time from the start of block procedure to complete sensory block) and the block performance time. The noninferiority margin was set at 5 minutes and was evaluated using the two-sided 95% bootstrap-confidence intervals ([CIs] 100,000 replications) for differences in means.

RESULTS: The mean times to complete sensory block revealed a significant decrease with the low-current group (17.9 ± 12.1 (mean ± SD) versus 22.8 ± 12.4 minutes; 95% CI, 1.1 to 8.6; p = 0.012). The time to readiness for surgery was 30.3 ± 13.8 minutes in the low-current group and 31.7 ± 12.9 minutes in the high-current group (95% CI, –2.7 to 5.5; p = 0.49). The performance time was significantly shorter in the high-current threshold group (9.5 ± 4.7 versus 11.9 ± 5.7 minutes; 95% CI, –4 to 1.1; p = 0.001).

CONCLUSION: Noninferiority for the high-current threshold technique could neither be confirmed for the primary end point nor for secondary end points. However, we consider a difference in mean times of approximately 8.5 minutes to achieve readiness for surgery acceptable for clinical practice.

地塞米松預防術後噁心嘔吐：一項隨機對照試驗的最新 Meta 分析

Dexamethasone to Prevent Postoperative Nausea and Vomiting: An Updated Meta-Analysis of Randomized Controlled Trials
BACKGROUND: Dexamethasone has an established role in decreasing postoperative nausea and vomiting (PONV); however, the optimal dexamethasone dose for reducing PONV when it is used as a single or combination prophylactic strategy has not been clearly defined. In this study, we evaluated the use of 4 mg to 5 mg and 8 mg to 10 mg IV doses of dexamethasone to prevent PONV when used as a single drug or as part of a combination preventive therapy.

METHODS: A wide search was performed to identify randomized clinical trials that evaluated systemic dexamethasone as a prophylactic drug to reduce postoperative nausea and/or vomiting. The effects of dexamethasone dose were evaluated by pooling studies into 2 groups: 4 mg to 5 mg and 8 mg to 10 mg. The first group represents the suggested dexamethasone dose to prevent PONV by the Society for Ambulatory Anesthesia (SAMBA) guidelines, and the second group represents twice the dose range recommended by the guidelines. The SAMBA guidelines were developed in response to studies, which have been performed to examine different dosages of dexamethasone.

RESULTS: Sixty randomized clinical trials with 6696 subjects were included. The 4-mg to 5-mg dose dexamethasone group experienced reduced 24-hour PONV compared with control, odds ratio (OR, 0.31; 95% confidence interval [CI], 0.23–0.41), and number needed to treat (NNT, 3.7; 95% CI, 3.0–4.7). When used together with a second antiemetic, the 4-mg to 5-mg
The dexamethasone group also experienced reduced 24-hour PONV compared with control (OR, 0.50; 95% CI, 0.35–0.72; NNT, 6.6; 95% CI, 4.3–12.8). The 8-mg to 10-mg dose dexamethasone group experienced decreased 24-hour PONV compared with control (OR, 0.26; 95% CI, 0.20–0.32; NNT, 3.8; 95% CI, 3.0–4.3). Asymmetric funnel plots were observed in the 8-mg to 10-mg dose analysis, suggesting the possibility of publication bias. When used together with a second antiemetic, the 8-mg to 10-mg dose dexamethasone group experienced decreased 24-hour PONV compared with control (OR, 0.35; 95% CI, 0.22–0.53; NNT, 6.2; 95% CI, 4.5–10). In studies that provided a direct comparison between groups, there was no clinical advantage of the 8-mg to 10-mg dexamethasone dose compared with the 4-mg to 5-mg dose on the incidence of postoperative nausea and/or vomiting.

CONCLUSIONS: Our results showed that a 4-mg to 5-mg dose of dexamethasone seems to have similar clinical effects in the reduction of PONV as the 8-mg to 10-mg dose when dexamethasone was used as a single drug or as a combination therapy. These findings support the current recommendation of the SAMBA guidelines for PONV, which favors the 4-mg to 5-mg dose regimen of systemic dexamethasone.

在高血壓病人麻醉誘導期間使用無創性脈搏傳導時間監測每搏收縮壓
Beat-to-Beat Tracking of Systolic Blood Pressure Using Noninvasive Pulse Transit Time During Anesthesia Induction in Hypertensive Patients
Sung-Hoon Kim, MD, Jun-Gol Song, MD, PhD, Ji-Hyun Park, MD, Jung-Won Kim, MD, Yong-Seok Park, MD and Gyu-Sam Hwang, MD, PhD
From the Department of Anesthesiology and Pain Medicine, Laboratory for Cardiovascular Dynamics, Asan Medical Center, University of Ulsan College of Medicine, Seoul, Korea.
Anesth Analg January 2013 116:94-100

背景：已有研究報導脈搏傳導時間（PTT）在清醒的人中，與動脈血壓（BP）有較好的相關性。我們測量了高血壓病人麻醉誘導期間，無創每搏 PTT 是否與持續有創動脈血壓監測有精確的相關性。

方法：23 名擇期行腎移植手術的高血壓病人被選入組。同步記錄橈動脈 BP、心電圖和指脈血氧體積描記。PTT 定義為從心電圖上的 R 波的波峰到光電容積描記的最大上升支的時間間隔。每搏 PTT 和 BP 之間的關係用相關性和受試者工作特徵（ROC）曲線分析來進行評價。

結果：在麻醉誘導期間，PTT 的變化與 BP 的變化是成正比的：當 BP 下降時，PTT 延長，反之亦然。PTT 的改變與收縮壓的相關性顯著優於與平均動脈壓（r = 0.81 ± 0.11 vs r = 0.72 ± 0.17; P < 0.001）和舒張壓（r = 0.81 ± 0.11 vs r = 0.52 ± 0.24; P < 0.001）的相關性。PTT 的改變與舒張壓下降的相關性比與舒張壓上升的相關性明顯（r = 0.83 ± 0.12 vs r = 0.68 ± 0.20; P = 0.001）。ROC 曲線分析表明，在麻醉誘導期間，PTT 增加 15％可以發現收縮壓下降 ≥30％，ROC 曲線下面積為 0.85。

結論：每搏 PTT 與有創收縮壓有良好的相關性，並且可以預測麻醉誘導期間的收縮壓下降。在高風險的高血壓患者不可用有創血壓時，每搏 PTT 可以作為潛在有用的收縮壓無創性監測指標。
（安光惠 譯 馬皓琳 李士通 校）

BACKGROUND: Pulse transit time (PTT) has been reported to show good agreement with arterial blood pressure (BP) in awake humans. We evaluated whether noninvasive beat-to-beat
PTT accurately correlated with invasively measured continuous arterial BP during anesthesia induction in hypertensive patients.

**METHODS:** Twenty-three hypertensive patients who were scheduled for kidney transplant were enrolled. Radial arterial BP, electrocardiogram, and finger pulse oximetric plethysmography were simultaneously recorded. PTT was measured as the time interval from the R-wave peak on the electrocardiogram to the maximal upslope of the photoplethysmogram. Relationships between beat-to-beat PTT and BP were evaluated by correlation and receiver operating characteristic (ROC) curve analysis.

**RESULTS:** During anesthesia induction, changes in PTT were directly proportional to changes in BP: when BP decreased, PTT lengthened, and vice versa. The inverse of PTT demonstrated significantly better correlation with systolic BP than with mean BP \( (r = 0.81 \pm 0.11 \text{ vs } r = 0.72 \pm 0.17; P < 0.001) \) or diastolic BP \( (r = 0.81 \pm 0.11 \text{ vs } r = 0.52 \pm 0.24; P < 0.001) \). The inverse of PTT was more highly correlated with decreasing than with increasing changes in systolic BP \( (r = 0.83 \pm 0.12 \text{ vs } r = 0.68 \pm 0.20; P = 0.001) \). The ROC curve analysis revealed that a 15% increase in PTT during anesthesia induction could detect a ≥30% decrease in systolic BP, with an area under the ROC curve of 0.85.

**CONCLUSION:** Beat-to-beat PTT was fairly well correlated with invasive systolic BP and could predict a reduction in systolic BP during anesthesia induction. Beat-to-beat PTT may show potential as a useful noninvasive index of systolic BP when invasive BP is unavailable in high-risk hypertensive patients.
BACKGROUND: Providing anesthesia and managing airways in the electrophysiology suite can be challenging because of its unique setting outside of the conventional operating room. We report our experience of several cases of reported airway trauma including tongue and pharyngeal hematoma and vocal cord paralysis in this setting.

METHODS: We analyzed all of the reported airway trauma cases between December 2009 and January 2011 in our cardiac electrophysiology laboratories and compared these cases with those without airway trauma. Data from 87 cases, including 16 cases with reported airway trauma (trauma group) and 71 cases without reported airway trauma from the same patient population pool at the same period (control group), were collected via review of medical records.

RESULTS: Airway trauma was reported for 16 patients (0.7%) in 14 months among 2434 anesthetic cases. None of these patients had life-threatening airway obstruction. The avoidance of muscle relaxants during induction in patients with a body mass index less than 30 was found to be a significant risk factor for airway trauma ($P = 0.04$; odds ratio, 10; 95% confidence interval, 1.1–482). Tongue or soft tissue bite occurred in 2 cases where soft bite block was not used during cardioversion. No statistically significant difference was found between the trauma and the control groups for preprocedure anticoagulation, anticoagulation during the procedure, or reversal of heparin at the end of the procedure.

CONCLUSIONS: The overall incidence of reported airway trauma was 0.7% in our study population. Tongue injury was the most common airway trauma. The cause seems to have been multifactorial; however, airway management without muscle relaxant emerged as a potential risk factor. Intubation with muscle relaxant is recommended, as is placing a soft bite block and ensuring no soft tissue is between the teeth before cardioversion.

間歇性硬膜外注射與連續性硬膜外輸注用於分娩鎮痛的比較：一項系統回顧和薈萃分析

**Intermittent Epidural Bolus Compared with Continuous Epidural Infusions for Labor Analgesia: A Systematic Review and Meta-Analysis**

Ronald B. George, MD, FRCPC*, Terrence K. Allen, MBBS, FRCA† and Ashraf S. Habib, MB, ChB, MSc, MHS, FRCA‡

From the *IWK Health Centre, Dalhousie University, Halifax, Nova Scotia, Canada; †Division of Women’s Anesthesia, Duke University Medical Center, Durham; and ‡Department of Anesthesiology, Duke University Medical Center, Durham, North Carolina.

Anesth Analg January 2013 116:133-144

背景：目前標準硬膜外分娩鎮痛方案為局麻藥結合阿片類藥物通過持續硬膜外輸注（CEI）。使用 CEI 局麻藥鎮痛的劑量可能較大，產生深度運動阻滯，從而潛在影響器材輔助分娩發生率。在本隨機對照試驗（RCTs）的系統性回顧中，我們對採用硬膜外鎮痛分娩的健康產婦，比較了間斷硬膜外注射（IEB）和標準 CEI 使用或不使用患者自控硬膜外鎮痛，對患者滿意度、人為麻醉干預的需要、產程和分娩方式的影響。

2 例未使用軟性咬口的心臟複律病人中發生了舌頭或軟組織咬傷。創傷組和控制組之間在操作前抗凝、操作期間抗凝及操作結束時逆轉肝素方面均未發現統計學顯著性差異。
方法：本研究為一項比較 CEI 和 IEB 分娩鎮痛的 RCTs 系統性評價。這些文章進行了有效性評估，資料由作者提取，並用比值比（ORs）、平均差（MDS）和 95%可信區間（CIs）進行總結。

結果：9 項隨機對照試驗被納入該系統性回顧。344 名受試者接受了 CEI，350 名受試者接受了 IEB 分娩鎮痛。所有 9 個研究被認爲為低偏倚風險。IEB 和 CEI 在剖腹產率（OR, 0.87; 95% CI, 0.56–1.35）、產程（MD, −17 分; 95% CI, −42 to 7）和麻醉干預的需要（OR, 0.56; 95% CI, 0.29–1.06）均無統計學差異。IEB 能輕度但統計學顯著性減少局部麻藥使用量（MD, 每小時 −1.2 mg 布比卡因; 95% CI, −2.2 到 −0.3）。產婦的滿意度得分（100-mm 視覺類比評分法）IEB（MD, 7.0 mm; 95%CI, 6.2–7.8）較高。

結論：IEB 是一個有研究前景的概念，目前證據表明 IEB 稍微降低了局部麻醉劑使用量並提高產婦的滿意度。考慮到收集的很多結果中 CIs 較多，不能得出明確結論，但 IEB 仍是一個潛在改善器械助產率及麻醉干預需要的方法。需要更多研究來使理想的 IEB 方案概念化，並研究其對分娩鎮痛和產科轉歸的影響。

（許辛譯，馬皓琳，李士通校）

BACKGROUND: The current standard labor epidural analgesic regimens consist of a local anesthetic in combination with an opioid delivered via continuous epidural infusion (CEI). With CEI local anesthetic, doses may be large with resulting profound motor blockade potentially affecting the incidence of instrumental deliveries. In this systematic review of randomized controlled trials (RCTs), we compared the effect of intermittent epidural bolus (IEB) to standard CEI dosing with or without patient-controlled epidural analgesia on patient satisfaction, the need for manual anesthesia interventions, labor progression, and mode of delivery in healthy women receiving labor epidural analgesia.

METHODS: A systematic review of RCTs that compared CEI with IEB for labor analgesia was performed. The articles were evaluated for validity, and data were extracted by the authors and summarized using odds ratios (ORs), mean differences (MDs), and 95% confidence intervals (CIs).

RESULTS: Nine RCTs were included in this systematic review. Three hundred forty-four subjects received CEI, whereas 350 subjects received IEB labor analgesia. All 9 studies were deemed to be low risk of bias. There was no statistical difference detected between IEB and CEI in the rate of cesarean delivery (OR, 0.87; 95% CI, 0.56–1.35), duration of labor (MD, −17 minutes; 95% CI, −42 to 7), or the need for anesthetic intervention (OR, 0.56; 95% CI, 0.29–1.06). IEB did result in a small but statistically significant reduction in local anesthetic usage (MD, −1.2 mg bupivacaine equivalent per hour; 95% CI, −2.2 to −0.3). Maternal satisfaction score (100-mm visual analog scale) was higher with IEB (MD, 7.0 mm; 95% CI, 6.2–7.8).

CONCLUSIONS: IEB is an appealing concept; current evidence suggests IEB slightly reduces local anesthetic usage and improves maternal satisfaction. Given the wide CIs of the pooled results for many outcomes, definite conclusions cannot be drawn for those outcomes, but there is also a potential that IEB improves instrumental delivery rate and need of anesthesia interventions. More study is required to conceptualize the ideal IEB regimen and investigate its effect on labor analgesia and obstetric outcomes.

轉運降低模擬孕婦心跳驟停時的心肺復甦品質
Transport Decreases the Quality of Cardiopulmonary Resuscitation During Simulated Maternal Cardiac Arrest
BACKGROUND: The purpose of this study was to compare cardiopulmonary resuscitation (CPR) for simulated maternal cardiac arrest rendered during transport to the operating room with that rendered while stationary in the labor room. We hypothesized that the quality of CPR would deteriorate during transport.

METHODS: Twenty-six teams composed of 2 providers (obstetricians, nurses, or anesthesiologists) were randomized to perform CPR on the Laerdal Resusci Anne SkillReporter™ mannequin during transport or while stationary. The primary outcome measure was the percentage of correctly delivered compressions, defined as compression rate $\geq 100$ beats per minute, correct sternal hand placement, compression depth $\geq 1.5$ inches (3.8 cm), and proper release. Secondary outcomes included interruptions in compressions, position of providers relative to the mannequin during the transport phase, and ventilation tidal volume.

RESULTS: The median (interquartile range) percentage of correctly rendered compressions during phase II was 32% (10%–63%) in the transport group and 93% (58%–100%) in the stationary group ($P=0.002$, 95% confidence interval of mean difference = 22%–58%). The median (interquartile range) compression rates were 124 (110–140) bpm in the transport group and 123 (115–132) bpm in the stationary group ($P = 0.531$). Interruptions in CPR were observed in 92% of transport and 7% of stationary drills ($P<0.001$, 95% confidence interval of difference = 61%–92%). During transport, 18 providers kneeled next to the mannequin, 2 straddled the mannequin, and 4 ran alongside the gurney. Median
INTERQUARTILE RANGE tidal volume was 270 (166–430) mL in the transport group and 390 (232–513) mL in the stationary group \( (P = 0.03) \).

CONCLUSIONS: Our data confirm our hypothesis and demonstrate that transport negatively affects the overall quality of resuscitation on a mannequin during simulated maternal arrest. These findings, together with previously published data on transport-related delays when moving from the labor room to the operating room further strengthen recommendations that perimortem cesarean delivery should be performed at the site of maternal cardiac arrest.

The Effect of Passive Leg Elevation and/or Trendelenburg Position on the Cross-Sectional Area of the Internal Jugular Vein in Infants and Young Children Undergoing Surgery for Congenital Heart Disease

Won Ho Kim, MD, Jong Hwan Lee, MD, PhD, Sangmin M. Lee, MD, PhD, Chung Su Kim, MD, PhD, Ryunga Kang, MD, Chan Seon Yoo, MD and Hyun Sung Cho, MD, PhD
From the Department of Anesthesiology and Pain Medicine, Samsung Medical Center, Sungkyunkwan University School of Medicine, Seoul, Republic of Korea.

BACKGROUND: In this study we evaluated the effect of passive leg elevation (LE) and Trendelenburg (T) position on the cross-sectional area (CSA) of the internal jugular vein (IJV) in infants and young children undergoing surgery for congenital heart disease. A secondary aim was to compare the CSA of the IJV between subjects with right-to-left (RL) shunt and left-to-right (LR) shunt.
METHODS: Ninety infants and small children from 10 days to 31 months old weighing from 1.5 to 9.7 kg were assigned to group RL (n = 48) or LR (n = 42). In both groups, the CSA, transverse, and vertical diameters of the IJV on both sides of the neck were measured using a 2-dimensional ultrasound transducer in the following positions: supine position, 15° of T position, supine position with 50° of LE, and 15° of Trendelenburg position with 50° of LE (TLE). A more than 25% increase in mean CSA of the IJV was considered clinically significant.

RESULTS: In group LR, T, LE, and TLE significantly increased CSA of both right (at least 12.3%, 10.3%, and 18.3%, respectively, “at least” refers to the lower 95% confidence limits) and left (at least 15.8%, 15.0%, and 18.9%, respectively) IJVs, whereas only TLE increased the CSA of both IJVs significantly in group RL (at least 8.2% and 7.7% in the right and left, respectively). The increase in the CSA of the right IJV related to T and TLE was larger in group LR than in group RL (at least 12.3% vs 1.2% for T and at least 18.3% vs 8.2% for TLE, respectively). A clinically significant increase in CSA was achieved in both right and left IJVs with TLE in group LR (mean 28.6% and 26.3%, respectively). The CSA of the right IJV was larger than that of the left IJV in most (at least 69.2%) patients.

CONCLUSIONS: Passive LE was as effective as T position to increase the CSA of the IJV, but there was no clinically significant increase in the CSA with any single maneuver. Only T position with passive LE achieved a clinically significant increase in the CSA of both IJVs in infants and young children with LR shunt, but not in the same age group with RL shunt.

鞘內注射巴氯芬對複雜性區域疼痛綜合征中不同性質疼痛的功效
Efficacy of Intrathecal Baclofen on Different Pain Qualities in Complex Regional Pain Syndrome
Anton A. van der Plas, MD*, Monique A. van Rijn, MD, PhD*, Johan Marinus, PhD*, Hein Putter, PhD† and Jacobus J. van Hilten, MD, PhD*
From the Departments of *Neurology and †Medical Statistics and Bioinformatics, Leiden University Medical Center, Leiden, the Netherlands.

背景：複雜性區域疼痛綜合征（CRPS）以使人嚴重衰弱的慢性疼痛為特點。患有 CRPS 的病人可能會經歷各種疼痛感覺，它們很可能體現了不同的病理生理機制。本研究中，我們評估了中樞 γ-氨基丁酸（B）受體刺激對患有肌張力障礙的 CRPS 患者中不同性質疼痛的獨特作用。

方法：在一個爲期一年的開放性設計中，我們每 3 個月對接受滴定劑量鞘內注射巴氯芬（ITB）治療的 42 名有肌張力障礙的 CRPS 患者評估一次 10 項疼痛性質的神經病理性疼痛量表、肌張力障礙程度和鎮痛藥物使用情況的改變。

結果：我們使用一個線性混合模型分析並以總體肌張力障礙程度和追加鎮痛藥物的使用情況作爲控制手段。在最初的 6 個月中，我們發現總體的劇烈痛、尖銳痛、鈍性痛和深部痛有顯著的改善。在這個時期之後，儘管肌張力障礙進一步改善並繼續提升 ITB 剤量但評分趨於平穩。

結論：ITB 刺激 γ-氨基丁酸（B）受體對有肌張力障礙的 CRPS 患者特定性質的疼痛表現出獨特的鎮痛效果。
**BACKGROUND:** Complex regional pain syndrome (CRPS) is characterized by severe debilitating chronic pain. Patients with CRPS may experience various pain sensations, which likely embody different pathophysiologic mechanisms. In this study, we evaluated the differential effects of central γ-aminobutyric acid (B) receptor stimulation on the different pain qualities in CRPS patients with dystonia.

**METHODS:** The 10 pain qualities of the neuropathic pain scale, dystonia severity, and changes in use of antinociceptive drugs were evaluated every 3 months for a period of 1 year in 42 CRPS patients with dystonia receiving titrated doses of intrathecal baclofen (ITB) treatment in an open design.

**RESULTS:** Using a linear mixed model analysis and controlling for global dystonia severity and the use of supplemental analgesics, we found a significant improvement in global intense pain, sharp pain, dull pain, and deep pain during the first 6 months. After this period, the scores leveled off despite further improvement of dystonia and continued ITB dose escalation.

**CONCLUSIONS:** γ-Aminobutyric acid (B) receptor stimulation by ITB exerts differential antinociceptive effects on specific pain qualities in CRPS patients with dystonia.

**BACKGROUND:** A previous study indicated that intrathecal administration of morphine reduces experimental inflammatory edema in rats by activating the nitric oxide/cyclic guanosine

---

**鉀離子通道在大鼠鞘內注射嗎啡所致抗外周水腫效應中的作用。**

*The Involvement of Potassium Channels in the Peripheral Antiedematogenic Effect of Intrathecally Injected Morphine in Rats*

Vanessa R.S. Foletto, DVetMed, MSc, Maria A. Martins, DVetMed, ScD and Carlos R. Tonussi, ScD

From the Departamento de Farmacologia, CCB, Universidade Federal de Santa Catarina, Florianópolis, Brasil.

Anesth Analg January 2013 116:232-238

**背景:** 既往研究顯示鞘內注射嗎啡可通過啓動一氧化氮/環磷酸鳥苷酸通路減少大鼠實驗性炎性水腫。這個證據支持了這樣一個假設，即在這種情況下鉀通道的開放在調解嗎啡的效應中起重要作用。

**方法:** 雄性韋氏大鼠接受鞘內注射藥物（20 微升）30 分鐘後，予以叉豆膠（150 微克）刺激爪。通過記錄爪體積的增加（以毫升為單位）來評估水腫情況，通過依藍染液滲出計算血漿滲漏。通過髓過氧化物酶分析來間接評價中性粒細胞遷移情況。通過組織學檢查來觀察炎性細胞浸潤和血管充血情況。

**結果:** 與磷酸鹽緩衝液相比較，嗎啡（37nmol）可以抑制炎性水腫，血漿漏出及血管充血，但是對髓過氧化物酶活性及中性粒細胞數量無作用。同時注射 4-氨基吡啶（10nmol）、格列本脲（5nmol）和地喹銨（10pmol）可逆轉嗎啡的作用，而尼克地爾（0.03nmol）可增強嗎啡的作用。

**結論:** 這些結果支持這樣一個假設，即鞘內注射嗎啡所致的外周抗水腫效應由鉀通道的啓動所介導。而且，阿片類藥物不影響急性中性粒細胞遷移的抑制，但影響毛細血管募集反應的減小。

（王贇 譯 馬皓琳 李士通 校）

**BACKGROUND:** A previous study indicated that intrathecal administration of morphine reduces experimental inflammatory edema in rats by activating the nitric oxide/cyclic guanosine
monophosphate pathway. This evidence supports the hypothesis that potassium channel opening may play an important role in mediating morphine’s effect under such conditions.

**METHODS:** Male Wistar rats received intrathecal injections of drugs (20 μL) 30 minutes before paw stimulation with carrageenan (150 μg). Edema was measured as paw volume increase (in milliliters), and plasma leakage was measured by Evans blue dye leakage. Neutrophil migration was evaluated indirectly by myeloperoxidase assay. The inflammatory infiltration and vascular congestion were observed by histologic examination.

**RESULTS:** Morphine (37 nmol) inhibited inflammatory edema, plasma leakage, and vascular congestion but had no effect on myeloperoxidase activity or neutrophil content compared with phosphate-buffered saline. Coinjection with 4-aminopyridine (10 nmol), glibenclamide (5 nmol), and dequalinium (10 pmol) reversed, but nicorandil (0.03 nmol) enhanced the effect of morphine.

**CONCLUSIONS:** These results support the hypothesis that the peripheral antiedematogenic effect produced by intrathecal morphine is mediated by potassium channel activation. Furthermore, this opioid effect does not involve the inhibition of acute neutrophil migration but does involve a reduction in capillary recruitment.