Assessment of a medetomidin/propofol intravenous anaesthesia (TIVA) for clinical anaesthesia in equidae

Bettschart-Wolfensberger, Regula; Freeman, S; Bettschart, R W; Fürst, Anton; Clarke, K W

Abstract: This study investigated the clinical use of a medetomidine/propofol TIVA technique in horses. Twenty-seven equidae of mixed breed, age [mean ± SD (range)] 2.9 ± 2.74 (0.11–11) years, weight (n = 15) 237 ± 130.22 (64–470) kg were anaesthetized for the following procedures: intra-abdominal (3); castration (15); joint flush (4); screw removal (1); desmotomy (3) and dermatoma removal (1). Horses were sedated with 7 µg kg−1 medetomidine IV and 10 minutes later anaesthesia was induced with 2 mg kg−1 propofol IV. Incremental doses of medetomidine were given prior to induction if the horse lifted its head when approached. Following induction, trachea were intubated and oxygen administered at FIO2 > 90%. Anaesthesia was maintained with 3.5 µg kg−1 h−1 medetomidine IV and propofol infused to effect (initial dose 0.1 mg kg−1 min−1) 1. Heart rate, respiratory rate, propofol dose and arterial blood pressure were recorded every five minutes and arterial blood gases every 15 minutes. Quality of sedation, anaesthetic induction and recovery were evaluated by the first author according to previously described scores (Bettschart-Wolfensberger et al. 1998). Data were analysed in a preliminary manner by descriptive statistical methods. Twelve horses required additional medetomidine for adequate sedation prior to induction (2 µg kg, n = 11; 6 µg kg, n = 1). Induction to anaesthesia was variable; excellent in 3, good in 18, fair in 4 and poor (unsatisfactory) in 2 horses. In an additional horse, induction to anaesthesia was very poor and transfer from the induction area to theatre was not possible, due to limb movement, even after several IV doses of propofol (total dose 2 mg kg−1) and thiopentone (total dose 2.5 mg kg−1). The use of propofol as an induction agent was stopped after this incident. During anaesthesia, range of mean values (absolute range) were, for respiratory rate 4.8–12.4 (4–32) min−1, heart rate 36.3–46.8 (15–58) min−1, mean arterial blood pressure 95.6–111 (84–128) mm Hg, propofol infusion 0.063–0.18 mg kg−1 min−1, PaO2 23.2–31.8 (11.7–59.7) kPa, PaCO2 6.1–7 (5.2–8.5) kPa. Anaesthetic quality was excellent and spontaneous movement occurred in six horses but was slow and easily controlled with increments of propofol. Anaesthetic duration was 57.9 ± 38.16 (15–175) minutes. Recovery was completed unaided within 24.3 ± 10.66 (10–44.8) minutes. It was uneventful in all horses. Medetomidine/propofol TIVA at the dose rates studied produced good quality anaesthesia with cardiopulmonary depression within ranges commonly recorded during inhalation anaesthesia in horses. However the quality of the anaesthetic induction was variable, and appeared inferior to that achieved with induction techniques which included ketamine.
Assessment of a medetomidine/propofol total intravenous anaesthesia (TIVA) for clinical anaesthesia in equidae

Regula Bettschart-Wolfensberger, Sarah Freeman, R. W. Bettschart, A. Fürst and Kathleen W. Clarke

Veterinär-Chirurgische Klinik der Universität Zürich, Schweiz

Summary

This study investigated the clinical use of a medetomidine/propofol TIVA technique in horses. Twenty seven equidae of mixed breed, age [mean (SD (range)] 2.9 ± 2.74 (0.11–11) years, weight 237 ± 130.22 (64–470) kg (in 12 horses, anaesthetised in the field weight was not scaled but judged by the first author) were anaesthetised. The following surgeries were performed: intraabdominal (3), castration (15), joint flush (4), screw removal (1), desmotomy (3), dermotoma removal (1).

In the field 6 wild horses were premedicated with 0.03 mg/kg acepromazine IM. All the horses were sedated with 7 mcg/kg medetomidine IV, anaesthesia was induced 10 minutes later with 2 mg/kg propofol IV. If the patient lifted its head when being approached for propofol administration, sedation was deepened with incremental bolii of medetomidine. After anaesthesia induction trachea was intubated and oxygen administered at FIO2 >90% in 15 horses. Anaesthesia was maintained with 3.5mcg/kg/h medetomidine IV and propofol infused to effect (initial dose 0.1mg/kg/min). Heart rate, respiratory rate, propofol dose and arterial blood pressure were recorded every five minutes and arterial blood gases every 15 minutes. Quality of sedation, anaesthetic induction and recovery were graded. Arterial bloodpressures and arterial blood gas values were analysed in a preliminary manner by descriptive statistical methods. Heart rates and respiratory rates were analysed using ANOVA for repeated measures. To detect differences to preanaesthetic values Dunnett’s post test was performed. Eleven horses required additional 2 mcg/kg and one horse 6 mcg/kg medetomidine for adequate sedation prior to induction. Induction of anaesthesia was variable: excellent in 3, good in 18, fair in 4 and poor (unsatisfactory) in 2 horses. In a 28th horse anaesthesia induction was very poor and transfer from induction area to theatre was not possible even after application of several boluses of propofol (total dose 2 mg/kg) and thiopentone (total dose 2.5 mg/kg) as whenever lifted paddeling of the limbs occurred. The use of propofol as an induction agent was stopped after this incidence. After sedation three horses showed second degree AV blocks two of which were treated with atropine. Compared to presedation values respiratory rate as well as heart rate were significantly reduced throughout anaesthesia. During anaesthesia, range of mean values (absolute range) were, for respiratory rate 12.4–17.7 (4–32) breaths/min, heart rate 36.3–46.8 (15–58) bpm, mean arterial blood pressure 95.6–111 (84–128) mmHg, propofol infusion 0.063–0.18mg/kg/min, pO2 23.2–31.8 (11.7–59.7) kPa, pCO2 6.1–7 (5.2–8.5) kPa. Anaesthetic quality was excellent: spontaneous movement occurred in 6 horses, but was slow and easily controlled with increments of propofol (0.05–0.1 mg/kg). Anaesthetic duration was 57.9 (38.16 (15–175) min. Recovery was completed unaided within 24.3 (10.66 (10–44.8) min. It was uneventful in all horses. Medetomidine-propofol TIVA at dose rates administered in the current study produced good quality anaesthesia with cardiopulmonary depression within ranges commonly recorded during inhalation anaesthesia in horses. However anaesthetic induction was variable, and appeared to be inferior to that achieved with induction regimes including ketamine.

Keywords:
anaesthesia, horse, medetomidine, propofol, total intravenous anaesthesia

Anwendung von Medetomidin-Propofol Total Intravenöser Anästhesie (TIVA) in der Pferdepraxis

In der vorliegenden Studie wurde bei 16 Pferden in einer Klinik und bei 12 Pferden unter Feldbedingungen die Anwendung von Medetomidin und Propofol zur Allgemeinanästhesie getestet. Sie waren 0,11–11 Jahre alt (2,9 ± 2,74; Mittelwert ± Standardabweichung), wogen 64–470 (237 ± 130,22) kg und waren von unterschiedlicher Rasse. Folgende Operationen wurden durchgeführt: Entfernung von Knochenimplantaten (1), Desmotomie (3), Gelenkspülung (4), Kastration (15), Dermatom Entfernung (1), Kastration intraabdominaler Hoden (3).

Unter Feldbedingungen wurden die Pferde mit 0,03 mg/kg Acepromazin i.m. prämediziert. Alle Pferde erhielten 7 µg/kg Medetomidin i.v. zur Sedation. Nach 10 Minuten wurde die Anästhesie mit 2 mg/kg Propofol i.v. eingeleitet. Die Pferde atmeten spontan, in der Klinik wurden sie intubiert und atmeten > 90% O₂. Die Anästhesie wurde aufrechterhalten durch eine konstante Infusion von Medetomidin (3,5 µg/kg/h) und Propofol infundiert nach Wirkung (initial Dosis 0,1 mg/kg/min). Arterielle Blutdruck- und arterielle Blutzuckergefährlich wurden durch deskriptive Statistik ausgewertet. Herz- und Atemfrequenzen wurden mittels Varianzanalyse für wiederholte Messungen untersucht und Unterschiede zu pränästhesischen Werten mittels Dunnet’s Test ermittelt.


Schlüsselwörter:
Anästhesie, Pferd, Medetomidin, Propofol, total intravenöse Anästhesie
Introduction

Anaesthesia related fatality rate is higher in horses than in any other domestic species (Johnston et al., 1995). One of the few facts known about this phenomenon is that the length of anaesthesia significantly influences the outcome (Johnston et al., 1995). An increase in risk with anaesthesia duration was noted, probably related to the cardiopulmonary depressant effects of halothane and the fact that horses are more prone to develop problems due to cardiopulmonary depression than small animals or humans. Total intravenous anaesthesia (TIVA) is less cardiopulmonary depressant than clinically used inhalational anaesthetic regimes (Luna et al., 1996). Thus it would seem to be wise to anaesthetise horses for long anaesthesias using TIVA combinations. However, most TIVA in horses include ketamine, but with infusions of 2 hours or longer, ketamine’s major metabolite norketamine accumulates (Taylor and Luna, 1995; Nolan et al., 1996). Due to its hallucinatory properties (Chang and Glazko, 1974), norketamine may lead to signs of apparent ketamine overdose during the horses recovery (Wolfensberger, 1993; Taylor and Luna, 1995; Bettschart-Wolfensberger et al., 1996; Nolan et al., 1996). Therefore TIVA involving ketamine is not ideal for longer anaesthesias.

To create a TIVA regime suitable for long anaesthesias in horses, a combination of medetomidine and propofol, has been elaborated (Bettschart-Wolfensberger et al., 2001a). Compared to other work that explored the use of propofol in this species in combination with other sedatives (Nolan and Hall, 1985; Taylor, 1989a; Hartsfield et al., 1994; Matthews et al., 1994; 1997; Mama et al., 1998; Taylor et al., 1997), medetomidine induced major dose reductions of propofol. The resulting cardiovascular depression was remarkably low (Bettschart-Wolfensberger et al., 2001b) and was comparable to other TIVA regimes including ketamine (Greene et al., 1986; Young et al., 1993; Bettschart-Wolfensberger et al., 1996; Luna et al., 1996), but less than with clinically used inhalational anaesthetic regimes (Luna et al., 1996). Medetomidine-propofol TIVA has been tested in experimental animals in anaesthesias as long as 4 hours (Bettschart-Wolfensberger et al., 2001a-b). Anaesthesia was easy to maintain and control. Even after the long duration of 4 hours of anaesthesia, recoveries were quick and of good quality. Under clinical conditions other factors than length of anaesthesia influence the outcome as for example type of surgery (Young and Taylor, 1993), nature (Whitehair et al., 1993), clinical condition and body weight of the horse (Johnston et al., 1995a). Thus a definitive testing of any new drug combination is only possible under clinical circumstances.

The present study wanted to test clinically medetomidine-propofol anaesthesia during all types of surgeries in horses of different breed, weight, character and age.

Materials and methods

Animals

The previously elaborated TIVA regime with medetomidine and propofol was tested in a total of 27 equidae. In a 28th horse anaesthesia was only induced, but then the horse was allowed to recover without surgery because of occurring problems. 15 of the animals that underwent surgery were surgical patients and were anaesthetised inside a hospital henceforth termed “under clinical conditions”. Another 12 animals were anaesthetised under field conditions. These were all stallions that had been raised in the field and had not been handled at all before the day of the surgery. Details of the animals and the surgeries performed are listed in table 1.

Protocol

The patients, that were anaesthetised under clinical conditions, were starved for 8–12 hours, if circumstances allowed this. A clinical examination was performed and if any abnormalities were detected, blood testing was done. Resting heart and respiratory rates were noted. Before the anaesthesia a catheter (Secalon®, Ohmeda, Hatfield, U.K.) was placed in the jugular vein and 8 horses were treated with penicillin (Peni G, Streuli AG, Switzerland) and gentamycin (Gentamycin Streuli, 10%, Streuli AG, Switzerland) i.v.. No analgesics were administered. The stallions that were castrated under field conditions were starved for 8–12 hours. To avoid injuries of people involved, 6 of the stallions were premedicated with 0.03 mg/kg acepromazine (ACP®, C-Vet, Leyland, U.K.) i.m. and left unattended for 30 minutes. A clinical examination followed then, resting heart and respiratory rate were noted and a catheter (Secalon®, Ohmeda, Hatfield, U.K.) was placed in the jugular vein. In both groups the animals were sedated by slow i.v. injection (over 2 minutes) of 7 µg/kg of medetomidine. Then they were given ten minutes to achieve maximal sedation. If sedation was considered unsatisfactory (i.e. the animal still reacting with a lift of the head or some steps backwards when being approached),

Tab. 1: Signalement of 27 equidae anaesthetised with medetomidine-propofol and type of surgery

| Signalement und Art des chirurgischen Eingriffs von 27 Equiden anästhesiert mit Medetomidin-Propofol |
|-------------------------------------------------|-------------------------------------------------|
| **Anaesthesias under clinical conditions** | **Anaesthesias under field conditions** |
| age, mean ± SD (range) | 3.5 ± 3.29 (0.1–11) years | 1.8 ± 0.41 (1–2) years |
| weight mean ± SD (range) | 240.0 ± 1.07 (64–470) kg | 231.6 ± 135.12 (120–450) kg; the weight of these stallions was not scaled but only judged by an experienced anaesthetist |
| breed | 6 Warm Blood horses, 4 Thoroughbred, 1 donkey, 2 Shetland ponies, 2 ponies | 7 Warm Blood horses, 5 ponies |
| type of surgery | joint flush (3), intraabdominal (3), screw removal (1), desmotomy (1), wound revision (3), dermatora removal (1), castration (3) | all castrations |
another 2–5 μg/kg of medetomidine were administered and anaesthesia induction delayed until satisfactory sedation was present. Just before induction heart and respiratory rate were noted again.

Anaesthesia was induced by i.v. injection of 2 mg/kg of propofol in all animals. Under clinical conditions patients were intubated just after anaesthesia induction, and were allowed to breath >90 % oxygen spontaneously. Anaesthesia was maintained with an infusion of propofol started at an infusion rate of 0.1 mg/kg/min and adjusted to the patients needs, using a syringe pump (P 4000, IVAC, Basingstoke, U. K.). Infusion rates of propofol were noted every 5 minutes. The clinical patients further got a continuous infusion of medetomidine at a dose rate of 3.5 μg/kg/h of medetomidine throughout the whole anaesthesia. Ringers lactate at a dose rate of 5 ml/kg/h was infused to clinical patients as well. 10 minutes after the end of the propofol infusion clinical patients were given 2 μg/kg of medetomidine to calm recovery.

Under field conditions bolus of propofol (0.085–0.105 mg/kg) were administered by hand injection every minute, according to the patients need and every five minutes the amount of propofol given was noted.

**Measurements**

After anaesthesia induction phases of apnoea were recorded and the position of the horses body during surgery was noted. Heart rate and respiratory rate were recorded every 5 minutes in all animals. In the field, heart rate was counted by palpating the pulse and respiratory rate by counting chest wall movements. Under clinical circumstance the patients were connected to an ECG (Kontron, Watford, U. K.) and intra-arterial blood pressure was measured continuously via a catheter in either the facial or the transverse facial artery. Arterial blood gas analysis was performed every 15 minutes. Arterial blood pressures and arterial blood gases were analysed in a preliminary manner by descriptive statistical methods. Heart rates and respiratory rates were analysed using ANOVA for repeated measures. To detect differences to preanaesthetic values Dunnett’s post test was performed. If horses moved or any other side effects occurred, it was noted.

Surgeons were asked to judge relaxation and surgery conditions at the end of the procedure as good, adequate or unsatisfactory.

**Induction and recovery**

During induction one person stayed with the animal holding its head, to give a little support. We recorded during induction: Time from the end of propofol injection to recumbency and any striking events concerning induction. Quality of induction was graded based on a scale of 1 (worst) to 5 (best) represented in table 2.

For recovery the animals were left unattended in a smoothly padded recovery box (clinical cases) or in the field. Clinical patients were given 2 mg/kg of medetomidine 10 minutes after the end of the propofol infusion. When they started swallowing the endotracheal tube was removed, and the time noted. Quality of recovery was scored based on a scale of 1 (worst) to 5 (best) as described in table 2. Additionally during recovery the time of first movement, the time of first head lift, duration of sternal recumbency, time to gain standing position and the number of attempts to stand up were noted.

**Results**

**Anaesthesia induction**

A foal of 6 weeks, that was suffering from polyarthritis and a temperature of 39.6 °C was given only 3 μg/kg of medetomidine, which induced sedation that seemed adequate for anaesthesia induction. In all 12 of the stallions in the field an additional bolus of 2 μg/kg and in one a bolus of 6 μg/kg of medetomidine was necessary, to achieve sedation that, subjectively assessed by the anaesthetist, seemed to be adequate for anaesthesia induction. Overall induction was excellent (5) in 3, good (4) in 18, fair (3) in 4, poor (2) in 2 and very poor (1) in one animal. One horse that showed a poor induction was a 2 year old stallion weighing 170 kg that had been very difficult to handle. It showed paddling of the limbs and spasms of the whole body up to one minute. The second one was a 6 year old cryptorchid thoroughbred stallion weighing 470 kg that was anaesthetised under clinical conditions. Both animals had seemed adequately sedated before anaesthesia induction. The horse that showed a very poor induction was an 11 year old Warm Blood of 330 kg, that required the removal of screws from the radius. It had shown deep sedation, but its heart rate had dropped from 48 before medetomidine administration to

**Fig. 2: Induction and recovery scores.**

<table>
<thead>
<tr>
<th>Score</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>very poor, unpredictable fall of the horse, the horse being injured</td>
</tr>
<tr>
<td>2</td>
<td>poor, attaining recumbency unpredictably, but no injuries</td>
</tr>
<tr>
<td>3</td>
<td>fair, horse slowly attains sternal or lateral recumbency, marked paddling of limbs or shaking of head</td>
</tr>
<tr>
<td>4</td>
<td>good, horse slowly attains recumbency, only slight paddling of limbs or shaking of the head</td>
</tr>
<tr>
<td>5</td>
<td>excellent, recumbency achieved slowly and smoothly, no paddling or head shaking</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Score</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>very poor, bad recovery with high risk of injury</td>
</tr>
<tr>
<td>2</td>
<td>poor, more than one attempt to stand, horse excited</td>
</tr>
<tr>
<td>3</td>
<td>good, but more than one attempt to stand, horse stays calm</td>
</tr>
<tr>
<td>4</td>
<td>good, 1 attempt to stand, some ataxia</td>
</tr>
<tr>
<td>5</td>
<td>excellent, 1 attempt to stand, minimal or no ataxia</td>
</tr>
</tbody>
</table>
15 beats/min. Thus a bolus of 1 mg atropine was given i.v. 5 minutes later its heart rate was regular with a frequency of 32 beats/min, sedation seemed to be adequate and anaesthesia was induced. The horse fell over backwards 45 seconds after the propofol and started paddling extensively and trying to lift its head. By sitting on its head it was maintained on the floor and the immediate administration of propofol to effect was started. During the next 2–3 minutes the horse was administered 66 ml of propofol (1%), the equivalent of its induction dose. As paddling became more severe, thiopentone was administered, until after a dose of 1 gr the horse became calm. It was intubated, supplementation of oxygen (15 l/min) started immediately and then it was connected to a hoist. After the thiopentone injection about 2–3 minutes had passed. Respiration seemed to be adequate, mucous membranes were pink and pulse regular and strong. The horse was lifted to transport it into surgery but within less than 30 seconds, it struggled so hard, that it had to be lowered again. It needed another 1.5 gr of thiopentone to prevent it from moving. As the anaesthetist judged the horse to be uncontrollable it was transported back to the recovery box, where it lay calm on its side for 42 minutes. Then it got up excellently. The data from this horse are excluded from the other paragraphs.

Cardiopulmonary effects

Arrhythmias

Three horses showed II degree AV blocks just prior to anaesthesia induction. One horse was treated with atropine before anaesthesia induction (see above). In another horse heart rate before anaesthesia induction had fallen from 48 to 28 beats/min and was not treated, but atropine was administered 15 min after anaesthesia induction as heart rate had dropped to 15. The horse was 370 kg and the initial dose of atropine was 1 mg i.v. followed by another 2 mg i.v. after 5 minutes as heart rate was still 15. Within another 2 minutes AV blocks disappeared, heart rate rose to 52 beats/min and remained there until the end of the surgery 65 minutes after anaesthesia induction. In the third horse, that showed II degree AV blocks just prior to anaesthesia induction these blocks disappeared spontaneously 40 minutes after anaesthesia induction.

Fig. 2: Mean heart rates (± SD) of 12 horses and ponies anaesthetised in the field; time points at which heart rates of all animals (n=15) are available.

Mittlere Herzfrequenz (± SD) von 12 Pferden und Ponies, anaesthesiert unter Feldbedingungen, vor der Applikation jeglicher Medikamente (pre), vor der Anästhesieeinleitung (bef) und alle 5 Minuten unter Medetomidin-Propofol Anästhesie; Zeitpunkte zu welchen Herzfrequenzen aller Tiere gemessen wurden

Arterial blood pressures

The application of an intra-arterial catheter proved to be difficult in some horses. In 5 horses the anaesthetist was not able to successfully place an intra-arterial catheter. Arterial blood pressures were very stable in individual animals over time and did not show variations greater than ± 11 mmHg. Occurring variations coincided with presence or absence of surgical stimuli. Mean arterial blood pressure (MAP) ranged from 95.6–111 with a range from 84–128.

In the one horse, in which heart rate 15 minutes after induction had dropped to 15 beats/min, for the first 20 minutes pulse was hardly palpable and mucous membranes looked very pale with no assessable capillary refill time (CRT). In this horse, for the first 25 minutes no intraarterial catheter could be placed and thus no pressure measurements were obtained. After 30 minutes (i.e. after the atropine) mucous membranes looked pink and CRT was 2 secs. MAP was 84 mmHg and remained like this until the end of the anaesthesia. This horse underwent emergency surgery, because it had suffered from severe injuries on its legs including loss of an unknown amount of blood. PCV before anaesthesia induction had been 24% and plasma protein contents 68 g/L, otherwise clinical examination had revealed no other problems.
Respiratory rate and arterial blood gases
Mean respiratory rates of clinical cases at time points at which data of all animals were available (during 45 minutes) are shown in figure 3. All animals showed a slowing of the respiratory rate after sedation with medetomidine. After anaesthesia induction four horses showed an increase of respiratory rate to presedation values (± 2 breaths/min). In the first minute after becoming recumbent all horses showed pauses in respiration of 20–60 secs, but phases of apnoea no longer occurred.
During anaesthesia respiratory rate was stable in all animals. Careful observation detected that some changes of tidal volume or respiratory rate occurred with surgical stimulation or changes in anaesthetic depth.

From anaesthesia induction onwards until the recovery phase, 4 animals, 3 of the horses in the field and one under clinical conditions showed irregular breathing patterns with respiratory breaks of up to 20 seconds, occurring every 1–3 minutes. Time course of mean arterial blood gas values are given in figure 4 and 5. Mean $p_{o_2}$ values ranged from to 23.2–31.8 kPa and mean $p_{aCO_2}$ values ranged from 6.1 to 7.0 kPa. Lowest and highest registered $p_{o_2}$ values were 11.7 and 59.7 kPa respectively. Lowest and highest registered $p_{aCO_2}$ values were 5.2 and 8.5 kPa respectively.

Other than cardiopulmonary effects
All animals appeared very lightly anaesthetised, showing brisk palpebral reflexes and nystagmus of various degrees. Observers not involved in the experiments, were all amazed, that the animals did not move.
During anaesthesia a total of six animals (2 in the field and 4 in clinics) showed movements in response to surgical stimuli. Once this was only a shivering in the neck region, twice a slight nodding with the head and 3 times a movement with the foot. These movements were always slow in nature and easily controlled by bolus injections of 0.05–0.1 mg/kg propofol. In horses that had an intra-arterial catheter in place a rise in mean arterial blood pressure of 10–15 mmHg preceded these movements.

Urination during or just after anaesthesia occurred in all animals. In clinical patients with estimated anaesthesia duration of >60 minutes, from the fourth clinical case onwards, urinary catheterisation was performed after anaesthesia induction.

**Fig. 3**: Mean respiratory rates (± SD) in 15 clinical cases anaesthetised with medetomidine-propofol, previously (pre) to any drug administration, before (bef) anaesthesia induction and every five minutes during anaesthesia. Time in mins after anaesthesia induction, n decreasing with time.

**Fig. 4**: Mean $p_{aCO_2}$ in 11 horses anaesthetised under clinical conditions using medetomidine-propofol.

**Fig. 5**: Mean $p_{o_2}$ in 11 horses anaesthetised under clinical conditions using medetomidine-propofol. Spontaneous ventilation, inspired $O_2$ concentration >90%.

**Anaesthesia duration, propofol infusion rates and surgical conditions**
In the clinical cases mean anaesthesia time was 79.0 ± 36.04 (50–175) mins and in the field 30.0 ± 11.40 (20–50) mins. During all surgeries, especially the castrations, surgeons were satisfied with the good relaxation of the animals. The obvious lack of violent „purposeful movements“ and the ease of controllability is another noted advantage of this protocol. Propofol infusion rates of the two groups are presented separately; as under field conditions the horses weights were only judged and they were also premedicated with acepromazine and not infused with medetomidine. Mean propofol infusion rates in the clinical patients are represented in figure 6. They ranged from 0.085–0.105 mg/kg/min with an individual range of 0.05–0.18 mg/kg/min. In all but three animals, the individual infusion rate could be maintained within ± 0.015 mg/kg/min from the beginning of the anaesthesia onwards. In two animals, that suffered from fever (foal with polyartheritis) or had suffered from acute blood loss (injured horse, in which blood pressure and heart rate problems occurred initially) the initially chosen dose rate of 0.1 mg/kg/min was reduced immediately to 0.06 mg/kg/min, as their cardiovascular function...
and the reduction of reflexes suggested a too deep level of anaesthesia. One pony that was anaesthetised for castration and had been very difficult to handle before anaesthesia, had to be bolused with propofol several times within the first 10 minutes reaching an average dose rate of 0.18 mg/kg/min of propofol. In this pony dose rate could be reduced with time reaching levels of 0.13 mg/kg/min 50 minutes after anaesthesia induction.

In the horses and ponies anaesthetised under field conditions mean propofol infusion rates reached from 0.063–0.18 mg/kg/min, according to the judged weight.

**Discussion**

The aim of the present study was to test the practical use of a new TIVA regime for equidae (Bettschart-Wolfensberger et al., 2001a+b). After a total of 27 anaesthetics the clinical testing of this regime was terminated, because of a relatively high incidence of unsatisfactory anaesthesia inductions, and one totally unacceptable induction phase in a horse, that almost resulted in injuries of people involved.

Medetomidine proved to be suitable as a preanaesthetic sedative in a practical situation. The previously tested dose of 7 µg/kg was increased, based on subjective decision by an experienced anaesthetist. This would be routine practice with other alpha2-adrenoceptor agonists under similar circumstances. None of the clinically anaesthetised horses needed further sedation but all the stallions castrated under field conditions did. These horses had not been handled at all, had been transported just before the anaesthetic and in six of them the application of an intravenous catheter had only been possible after the administration of some intramuscular acepromazine. Nevertheless the further administration of 2–6 µg/kg medetomidine resulted in good sedation with total lack of response to under clinical circumstances normal external stimuli, such as people talking or approaching and other horses walking by. Because of this deep level of sedation, achieved in all horses, the author is convinced that none of the bad anaesthesia inductions or nonideal recoveries in this study can be attributed to lack of sedation.

The incidence of nonideal or bad anaesthesia induction phases in the present study, was relatively high compared to previous propofol reports. Nolan (1989) reported one fair anaesthesia induction phase out of four ponies that were sedated with detomidine and anaesthesia induced with propofol and Nolan and Hall (1985) reported some paddling after anaesthesia induction when the horses had been sedated with acepromazine and butorphanol but only smooth induction phases after sedation with xylazine. Taylor (1989a), Aguiar et al. (1993) or Pablo et al. (1997) all used detomidine or xylazine for premedication and propofol as an induction agent and only reported good anaesthesia inductions. None of the ponies or horses that weighed < 250 kg showed any problems during anaesthesia induction but the incidence of paddling after induction increased with increasing weight of the horse. In the larger horses the injection of the total dose of propofol took up to 55 secs. When these trials were performed, propofol was only available as a 1% solution resulting in very high injection volumes. Since the completion of this current work a 2% solution has become available. The onset of unconsciousness and anaesthesia is dependent on the time which the drug takes to achieve certain levels in the brain within a certain time and thus delays onset of unconsciousness. The number of horses in which propofol was used as an induction agent is low (Nolan and Hall, 1985; Nolan, 1989; Aguiar et al., 1993; Pablo et al., 1997) and thus only a blind comparison of different sedatives followed by propofol inductions would be able to really test if anaesthesia inductions are worse with medetomidine premedication than following other sedatives.
One of the stallions anaesthetised in the field showed not only paddling of the limbs, but also spasms of the whole body immediately after anaesthesia induction as well as during recovery. In dogs different authors have reported spasms and stiffness after the use of propofol combined with different sedatives (Hall and Chambers, 1987; Watney and Pablo, 1992; Nolan et al., 1993; Kramer et al., 1995), the causes of which remained unknown. However Lagerweij et al. (1993) reported spontaneous limb movements during 2 hours of propofol infusion in dogs that did never occur if the same dogs were premedicated with medetomidine.

The horse that showed a very poor anaesthesia induction had been anaesthetised already in the same clinic 7 months before this episode. At that time anaesthesia had been induced with xylazine (1.1 mg/kg)-ketamine (2mg/kg)-diazepam (0.02 mg/kg) and maintained with halothane. On this first occasion also a bolus of 0.5 gr thiopentone had been necessary to intubate and transport the horse into theatre, but otherwise anaesthesia had been uneventful. Why the use of medetomidine-propofol resulted in an absolutely unacceptable, and even with thiopentone, hardly controllable situation remains unknown. Horses dying of hypoxaemia can react violently and show extensive movement. We were not able to take an arterial blood sample in this horse, as it struggled so hard. Nevertheless we are convinced, that no severe hypoxaemia was present as mucous membranes were always pink, pulse rate regular and strong, the horse seemed to the anaesthetist to be conscious and the paddling ceased when thiopentone was administered. The influence of atropine administered just prior to induction in this horse is another unknown factor.

Already in previous medetomidine-propofol reports it was obvious that judgement of depth of this anaesthesia differs from that during inhalation anaesthesia. In the present study the author had already been familiar with the technique and did not have problems to control depth of anaesthesia. Clinical judgement of depth of anaesthesia is one of the major “holy grails” in anaesthesia. As long as a horse doesn’t show purposeful movements during anaesthesia, the anaesthetist can not be certain, if depth of anaesthesia is adequate. The prompt recoveries noted in all animals suggest, that none of the horses were overdosed. In all animals propofol infusion was started at an infusion rate of 0.1 mg/kg/min. Only in a foal and one horse, both in poor condition, was the dose rate reduced to 0.06 mg/kg/min rapidly after anaesthesia induction. If the patient did not show any reaction on surgical stimulus the infusion rate was decreased by 0.01 mg/kg/min every 15–20 minutes. Mean infusion rates of propofol of 0.85–0.105 mg/kg/min that were necessary to maintain anaesthesia were within the ranges determined in previous studies (Bettschart-Wolfensberger et al., 2001a+b). These infusion rates were much lower than those used by other authors that combined propofol with various sedatives, analgesics or muscle relaxants (Nolan and Hall, 1985; Taylor, 1989a; Matthews et al., 1994; Taylor et al., 1997; Mama et al., 1998). Only combination of propofol with constant ketamine infusion resulted in propofol infusion rates comparable with the one described here (Flaherty et al., 1997). In this current study we maintained anaesthesia as light as possible, to reduce cardiopulmonary side effects, thus making a fatal outcome less likely. This resulted in purposeful movements in 6 horses. Contrary to TIVA based on ketamine (Wallensberger, 1992), these movements were always gentle and easy to control by slow injection of a propofol bolus of 0.05–0.1 mg/kg. Although anaesthesia was maintained at such a light level the surgeons always considered muscle relaxation good and as the occurring movements were always gentle everybody in theatre gained quickly confidence to the new anaesthetic.

Hypoxaemia has been the biggest problem with medetomidine-propofol TIVA in the previous studies (Bettschart-Wolfensberger et al., 2001a+b). Thus clinical patients were allowed to breathe >90% oxygen from the start of anaesthesia. Mean $pO_2$ values, although lower than theoretically possible with pure oxygen ventilation, were within ranges that are common in clinically anaesthetised horses breathing spontaneously pure oxygen. It has been recognised a long time ago, that in recumbent horses often huge alveolar-arterial oxygen differences occur (Hall et al., 1968), mainly attributable to ventilation-perfusion disturbances. The minimal $pO_2$ value of 11.7 kPa registered in a horse on its back 60 minutes after anaesthesia induction cannot be interpreted as an unacceptable respiratory depression due to medetomidine-propofol. In a clinical survey in horses that were anaesthetised with halothane in oxygen and breathing spontaneously, 5% of the horses on their back developed hypoxaemia (defined as $pO_2$<60 mmHg=8 kPa) from the beginning of anaesthesia and 35% by its end, 101 minutes after anaesthesia induction (Day et al., 1995). We still do lack an explanation for the relatively large number of severe hypoxaeias, that occurred in the previous studies with medetomidine-propofol (Bettschart-Wolfensberger et al., 2001a+b). The present results suggest, that medetomidine-propofol, administered at adequate dose rates does not induce greater hypoxaemia than do other general anaesthetics in horses. $PCO_2$ values were within normal limits for horses under general anaesthesia with a relatively low maximal value of 8.5 kPa registered in a horse in lateral recumbency, 75 minutes after anaesthesia induction.

Bradyarrhythmias after the use of medetomidine are common in dogs (Vainio et al., 1986/87; Lombard et al., 1989; Nilsfors et al., 1989; Kramer et al., 1996). Their effects, other than a reduction in cardiac output and the potential danger during a

| Tab. 3: Recovery times (in minutes from end of infusion) and scores in 14 equidae anaesthetised under clinical conditions (Clin) or in the field (Field) following medetomidine-propofol infusion anaesthesia. |
|---|---|---|
| **time of extubation (mins)** | (Clin, n=14)) | (Field, n=12) |
| 1. movement (mins) | 13.5 ± 7.39 | 12.6 ± 6.66 |
| into sternal recumbency (mins) | 18.2 ± 9.24 | 18.1 ± 10.0/3 |
| duration of sternal recumbency (mins) | 8.5 ± 8.06 | 2.0 ± 1.66 |
| standing (mins) | 27.1 ± 9.96 | 20.2 ± 11.19 |
| nr. of attempts to stand | 1.5 ± 0.94 | 1.2 ± 0.39 |
| recovery score (1–5; 1 = worst, 5 = best) | 4.6 ± 0.76 | 4.2 ± 0.75 |
general anaesthetic, are a matter of discussion. Anticholinergics can prevent these bradycardias, but their use has been questioned, as it may result in very high blood pressures (Bergström, 1988; Alibhai et al., 1996). In horses a reduction of heart rate after the use of any \( \alpha \)-adrenoceptor agonist is common (England and Clarke, 1996). Pretreatment with anticholinergics is not recommended as anticholinergics also induce disturbances of vision and depression of gastrointestinal motility (Hall and Clarke, 1983). Nevertheless in two horses heart rates had dropped to 15 beats/min atropine was administered to maintain cardiac output. Compared to atropine doses of 0.005–0.01 mg/kg used by other authors (Short et al., 1986; Gasthuys et al., 1990) the doses of 0.003 and 0.008 mg/kg we used were rather low but resulted in heart rates within normal ranges within 5 minutes.

In dogs heart rate tends to rise during propofol infusion following medetomidine premedication (Lagerweij et al., 1993; Hammond and England, 1994; Hall et al., 1997). A rise in heart rate occurred in the clinical cases in this study but not in the horses in the field, probably because the observation period had been too short.

Two horses showed sinus arrhythmias during the propofol infusion, a fact that had not been observed during the anaesthesia of the ponies in the previous medetomidine-propofol studies (Bettschart-Wolfensberger et al., 2001a+b), significance of which is unknown.

After anaesthesia induction the placement of an arterial catheter was not always easy probably due to peripheral vasoconstriction induced by medetomidine (DOcherty and Mc-Grath, 1980). Such problems, but with the catheterisation of veins have been reported in cats after the premedication with medetomidine (Becker and Oechtering, 1996). Peripheral vasoconstriction was also noted by other authors. Lagerweij et al. (1993) reported frequently cadaveric appearance of the oral mucous membranes and paleness of the conjunctival membranes after medetomidine premedication and explained this with hypoperfusion of some vascular beds due to increased vascular resistance from selective vasoconstriction. We noted alarmingly pale mucous membranes in one horse only, the one in which heart rate fell to 15 beats/min. Circulation was failing probably due to the reduction of heart rate caused by medetomidine but also due to a fall in blood pressure caused by propofol. In this horse the amount of blood loss and the general status before the anaesthesia had probably been underestimated. In the horses in which arterial blood pressure was measured, it was very stable and within ranges commonly accepted as being adequate during horse general anaesthesia. The clinical significance of an adequate blood pressure should not be overinterpreted during medetomidine-propofol anaesthesia. Data from previous studies in ponies (Bettschart-Wolfensberger et al., 2001a+b) and from dog studies suggest that arterial blood pressure is mainly dependent on the proportion of propofol plasma concentration to medetomidine plasma concentration (Vainio, 1991; Lagerweij et al., 1993; Hall et al., 1997). Thus a horse can show a „good“ blood pressure even with a low cardiac output, if peripheral vasoconstriction is present.

Urine production was not assessed objectively in the present study. Nevertheless horses were noted to urinate unusually large quantities either during anaesthesia or recovery, or immediately after recovery. The problem of very large urine production due to the use of medetomidine had been noted previously (Bettschart-Wolfensberger et al., 2001a+b). Nevertheless it was surprising that stallions undergoing castrations of only 20 minutes duration urinated during anaesthesia or right after recovery. Urinating during anaesthesia can contaminate the surgical site and an extended bladder may cause discomfort during recovery. The increased fluid loss during longer surgeries should be taken into account and replaced with i.v. fluid. In clinical patients we started, after the fourth case, to regularly induce an urinary catheter at the beginning of surgery to reduce problems. However this enormous urine production is certainly a relatively big disadvantage of the present TIVA regime. In the present study the protocol was such that none of the clinical patients received preemptive analgesia. To administer an analgesic before surgery is considered the best option to prevent „wind up“ phenomenon (Woolf, 1983). Medetomidine itself has marked analgesic properties and we wished to get an idea if the analgesia persists any longer than the sedative action. Thus we chose not to pretreat the horses with analgesics. We did not perform any sophisticated testing of analgesia or present pain after surgery, but only watched the horses right after recovery in their box as well as one and 4 hours later. All the horses started eating as soon as they were returned to their box and remained totally calm as long as they were observed. We were amazed by this fact as, at least in the four intraabdominal surgeries we were expecting some reactions. In such surgeries the routine preoperative application of flunixin meglumine is very often insufficient, many horses show some signs of colic after surgery and have to be treated with supplemental analgesia. This postoperative analgesic action of medetomidine could be a big advantage of the use of this drug in the perioperative period.

In previous studies (Bettschart-Wolfensberger et al., 2001a+b) the use of atipamezole after medetomidine propofol anaesthesia has been tested. It was concluded, that atipamezole should not be routinely used as recovery was not improved and, as its use could result in antagonism of not only the sedative but also the analgesic component of medetomidine. In foals it is very important, that they wake up as soon as possible from general anaesthesia to start sucking. During recovery an infusion therapy is not possible and the foals can become hypoglycaemic and their energy metabolism can dis regulate. This can be very harmful especially in foals that suffer from other diseases. Matthews et al. (1994) have reported the successful use of propofol in foals with very quick recoveries. Thus we decided to use medetomidine-propofol followed by atipamezole in recovery in a foal that underwent its fourth anaesthetic and was severely ill. As antibiotics, low plasma proteins, poor state of health and neonatal age certainly influence pharmacokinetics of a drug, we had no idea of the necessary dose rate of atipamezole. We have shown that the onset of its action is very quick (Bettschart-Wolfensberger et al., 1999). Thus we gave it by very slow injection and stopped as soon as we noted signs of awakening. It was impressive how within not even two minutes the foal was standing, totally alert and sucking.

In horses recovery quality is one of the most important features of an anaesthetic regime, as fatalities often happen during this period (Young and Taylor, 1993). In experimental animals, testing of recovery quality is very difficult, as it usually improves with repeated anaesthetics. The horses are not in an unfamiliar environment any more and most often no pain is present. Multiple observations on experimental animals will give results different from those of single observations on a large number of animals. As reported by other authors that used propofol in horses, overall recovery quality was good and of acceptable length. We
failed to test if fatalities during recovery could be reduced by the routine use of medetomidine-propofol TIVA, as a much larger number of clinical cases would be necessary to do so. In the present study we have shown that an infusion of medetomidine-propofol results in good quality and easy controllable anaesthesia. Cardiovascular changes were within expected ranges and hypoxaemia was no more severe than during routine general horse anaesthesia. However problems with anaesthesia induction limited the use of propofol and implicated the termination of the trial. Further investigations of alternative induction agents are necessary. The availability of a medetomidine antagonist proved to be very useful to speed recovery in one foal which was severely ill.

Literature


Assessment of a medetomidine/propofol total intravenous anaesthesia (TIVA) for clinical anaesthesia in equidae


Vainio O., L. Palmu, R. Virtanen and J. Weckslé (1986/87): Medetom-

Vainio O. (1991): Propofol infusion anaesthesia in dogs pre-medica-


Ylisela E. and O. Vainio (1989): Effects of medetomidine on the exper-


Pablo J., J. Bailey and C. Nicklin (1997): Evaluation of guaifenesin-
propofol and sevoflurane in premedicated horses. Proc. 6th Int. Conf. of Vet. Anaesth., 123.

Short C.E., N. Matthews, R. Harvey and C.L. Tyner (1986): Cardiovas-
cular and pulmonary function studies of a new sedative/analgetic (detomidine/domosedan®) for use alone in horses or as a preanesthe-


Taylor P.M. and S.P.L. Luna (1995): Total intravenous anaesthesia in ponies using detomidine, ketamine and guaifenesin: pharmacoki-


Regula Bettschart-Wolfensberger, Dr. med. vet, PhD, Dipl ECVA
Oberarzsestinin Änthesie
Veterinar-Chirurgische Klinik der Universität Zürich
Winterthurerstr. 260
CH 8057 Zürich
Tel: 0041 1 635 84 77
Fax: 0041 1 635 89 05
e-mail: bettwolf@vetchi.urzch.ch