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THE EFFECT OF OBESITY ON DOSE OF DEXMEDETOMIDINE WHEN ADMINISTERED WITH FENTANYL DURING POSTOPERATIVE MECHANICAL VENTILATION - RETROSPECTIVE

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Abstract: We carried out a retrospective investigation on the effect of obesity on dexmedetomidine (DEX) requirements when administered with fentanyl (FEN) during mechanical ventilation after major surgeries. After Institutional Review Board approval, 14 obese patients with a body mass index (BMI) ≥ 30 kg/m² and the same number of non-obese patients with similar backgrounds to the obese patients were selected from medical records. Doses of DEX in the first 48 h or until the end of sedation or extubation were calculated for comparison. In addition to comparison of dosing between the groups, associations between total body weight (TBW), BMI, and lean body mass (LBM) values and doses of DEX (mcg/h), between BMI and various indices (i.e., amount per TBW per hour and amount per LBM per hour) of DEX doses, and between above indices of DEX and FEN doses were also examined. There were no significant differences in DEX dose indices between the groups. However, DEX requirements (mcg/h) were significantly increased with TBW (kg) ($r = 0.51, P = 0.003$), BMI ($r = 0.49, P = 0.006$) and LBM (kg) ($r = 0.42, P = 0.02$), which might have enhanced the DEX metabolism with physiological changes with obesity. These findings will be beneficial for future clinical pharmacological analysis of DEX.

Key words: dexmedetomidine, pharmacokinetics, postoperative, mechanical ventilation, obesity

INTRODUCTION

Dexmedetomidine (DEX) is a highly selective Alpha 2-adrenoceptor agonist with sedative and analgesic effects, which is administered intravenously in the perioperative period. It is recommended that DEX is initiated with a loading dose of 6 mcg/kg/h over 10 to 15 minutes and titrated over a dose range of 0.2 to 0.7 mcg/kg/h based on the total body weight (TBW) of patients. However, in many anesthetic agents, obese patients do not require as large a dose as that calculated by TBW1-3) and some types of body weight indices such as lean body mass (LBM) are often used as dosing scalars to calculate administration doses for such patients4). Until now, sophisticated pharmacokinetic analysis of DEX and its necessary amount in obese patients when administered with fentanyl (FEN) during postoperative mechanical ventilation have not been well investigated.
In this study, as a preliminary investigation for future pharmacokinetic analyses of DEX, the effect of obesity on DEX requirements when administered with FEN was studied retrospectively.

**MATERIALS AND METHODS**

The study obtained hospital Institutional Review Board approval and was registered as a UMIN-CTR clinical trial (UMIN000012466). Patients with American Society of Anesthesiologists (ASA) physical status 1 to 3, aged 18–80 years old, had a body mass index (BMI) ≥ 30 kg/m² and received DEX and FEN during mechanical ventilation after elective major surgeries between January 2011 and July 2013, were selected as an obese group. The same number of patients, whose background including age-group and sex were matched to those in the obese group and whose BMI was < 30 kg/m², were randomly selected as a non-obese group. The exclusion criteria included patients with history of neuropsychiatric disorder, brain surgery, psychotropic drug usage, preoperative disturbance of consciousness, or hepatectomy, patients with brain injuries that were later identified, and patients with prolonged intra- or postoperative shock and did not respond to treatment.

Total doses of DEX and FEN between the initiation and the time point when sedation was terminated or the patient’s trachea was extubated within 48 h, or during the first 48 h when sedation was prolonged further, were extracted from medical records. DEX and FEN were titrated according to protocols mainly by nursing staff as below:

**Protocol**: The initial and maximum doses of DEX were decided by attending anesthesiologists or intensivists. Nursing staff then titrated DEX dose to maintain patients in the target sedation range; Richmond Agitation and Sedation Scale ^5^ 0 to -1 (i.e., alert and calm to drowsy) and -2 to -3 (i.e., light to moderate sedation) during daytime and nighttime, respectively. FEN was also titrated to the Numeric Rating Scale for pain less than or equal to 3 (0 as no pain and 10 as most severe pain).

Indices of DEX and FEN doses, including; 1) amount per hour (mcg/h), 2) amount per TBW per hour (mcg/kg/h), and 3) amount per LBM ^6^ per hour (mcg/LBM/h) were analyzed.

Janmahasatian’s equation for LBM ^6^:

**Male**:

\[ \frac{(9.27 \times 10^3 \times \text{TBW})}{(6.68 \times 10^3 + 216 \times \text{BMI})} \]

**Female**:

\[ \frac{(9.27 \times 10^3 \times \text{TBW})}{(8.78 \times 10^3 + 244 \times \text{BMI})} \]

To investigate the confounding effects, the relationships between age, infusion duration and doses of DEX and FEN were also evaluated.

**Statistical Analysis**

Student t tests, Welch t tests, or Fisher’s exact tests were used to compare the data between the groups. Based on the data from the two groups, correlation analysis was used to examine associa-

| Table 1 |
|---------|
|         | Non-obese | Obese | *P* values |
| Age (yr) | 55.5 (9.5) | 55.6 (11.7) | 0.972 |
| Male / Female (n) | 10 / 4 | 10 / 4 | 1.000 |
| ASA I / II / III (n) | 1 / 2 / 11 | 0 / 4 / 9 | 0.510 |
| Height (cm) | 164.4 (9.1) | 165.4 (10.5) | 0.799 |
| Body weight (kg) | 62.3 (11.8) | 96.0 (24.2) | < 0.001* |
| BMI (kg/m²) | 22.9 (3.4) | 34.8 (5.8) | < 0.001* |
| Procedures Cardiovascular / Others (n) | 10 / 4 | 9 / 5 | 1.000 |
| DEX infusion duration (h) | 20.7 (13.0) | 27.0 (16.8) | 0.274 |
| FEN infusion duration (h) | 20.8 (12.3) | 27.3 (16.7) | 0.249 |
| DEX dose (mcg/h) | 27.8 (8.4) | 34.3 (17.3) | 0.227 |
| FEN dose (mcg/h) | 44.8 (8.9) | 60.4 (20.9) | 0.020* |
| DEX dose (mcg/kg/h) | 0.46 (0.14) | 0.36 (0.15) | 0.077 |
| FEN dose (mcg/kg/h) | 0.74 (0.22) | 0.65 (0.24) | 0.336 |
| DEX dose (mcg/LBM/h) | 0.57 (0.17) | 0.55 (0.23) | 0.760 |
| FEN dose (mcg/LBM/h) | 0.93 (0.24) | 0.99 (0.35) | 0.575 |

Data were presented as mean (standard deviation). *, *P* < 0.05. ASA : American Society of Anesthesiologists physical status. DEX : dexmedetomidine, FEN : fentanyl, LBM : Janmahasatian’s equation for Lean Body Mass
tions between values of TBW and doses of DEX and FEN, between BMI and doses of DEX and FEN, between values of LBM and doses of DEX and FEN, between BMI and the two indices (2 and 3) of DEX and FEN doses, and between the three indices of DEX and FEN doses. Pearson’s correlation coefficients were calculated. $P < 0.05$ was considered statistically significant. R (version 2.13.0, the R foundation for Statistical Computing) was used for statistical analysis.

RESULTS

The demographic data, indices of DEX and FEN doses in obese and non-obese groups are shown in Table 1. Both groups comprised of 14 patients. There were significant differences in TBW and BMI between the groups. There was no difference in infusion durations of DEX and FEN between the groups. There were no significant differences in all DEX dose indices between the groups. The FEN dose (mcg/h) was significantly larger in the obese group than in the non-obese group despite there being no differences in the other two dose indices of FEN.

Fig. 1 shows the correlation diagram between TBW and DEX amount per hour (A) and between TBW and FEN amount per hour (B). All patients were analyzed together. TBW, total body weight (kg).

![Correlation diagram between TBW and DEX amount per hour (A) and between TBW and FEN amount per hour (B). All patients were analyzed together. TBW, total body weight (kg).](image)

Fig. 2(A) and (B) show significant correlations between BMI and DEX and FEN amounts per hour ($r = 0.49, P = 0.006$ and $r = 0.434, P = 0.021$, respectively). Fig. 3(A) and (B) show significant correlations between LBM and DEX amounts per hour ($r = 0.42, P = 0.02$). However, there were no significant correlations between BMI and DEX and FEN amounts per TBW per hour (Fig. 4A and B), and between BMI and DEX and FEN amounts per LBM per hour (Fig. 4C and D). As mentioned in the methods section,
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Fig. 1 to 4 are based on the data from the two groups. Fig. 5(A), (B) and (C) show the correlation between each of the three dose indices of DEX and that of FEN. No significant correlations were observed in the control group, obese group and the combined data from the two groups.

In terms of investigation of potential confounding factors, there were no significant correlations between age, infusion duration, and indices of DEX and FEN amount (data not shown).

DISCUSSION

From the clinical viewpoint, whether the drug doses for obese patients can be calculated simply on per kg basis (i.e., double dose for double body weight) is of paramount importance. With several intravenous anesthetic drugs, such dosing regimen can cause overdose in obese patients. For example, propofol dose calculated not with TBW but with LBM is enough for anesthetic induction in high BMI patients. FEN dose can also be reduced compared to that calculated with TBW. Because DEX overdosing could cause side effects including hypotension, bradycardia, or prolonged sedation due to drug accumulation, discovering an appropriate dose, schedule for administration, and dosing scalar are essential through investigating pharmacokinetics and pharmacodynamics of DEX in obese patients. As a preliminary analysis for a sophisticated pharmacological analysis, we performed a retrospective descriptive study.

The metabolism of DEX is dependent upon hepatic blood perfusion because of the high hepatic extraction ratio as with propofol. Enlarged liver mass caused by obesity results in increased hepatic blood flow and more hepatic extraction, which might enhance the increased administered DEX dose associated with the increase of BMI (and its constitute TBW) in this study. Peeters et al reported that hepatic blood flow was a more predictive indicator than cardiac output for propofol clearance,
suggesting that the results can most likely be extrapolated to other highly extracted drugs like DEX, which support our prediction. Meanwhile, it is estimated that the necessary dose of drugs with high extraction ratio does not increase linearly with body weight as observed in many pharmacokinetic studies of propofol and FEN where drug clearances do not increase in direct proportion to weight. Also in our study, when administered with FEN, a non-linear requirement (i.e., double body weight does not need double dose) of DEX with regards to TBW and LBM was observed. Meanwhile, the increase in DEX amount per hour with BMI is considered to be a ‘spurious correlation’ because simply TBW and LBM increased with obesity (data not shown). In addition, in the BMI range in our study, the necessary dose of DEX per TBW tended to decrease with BMI increase although it was not statistically significant and the necessary dose of DEX per LBM did not change with BMI, which suggests that LBM can be one of the better options as a dosing scalar for ‘per kg’ calculations of DEX than TBW.

There have been some published reports on pharmacokinetics of DEX in which body size covariates were searched. Dyck et al built a pharmacokinetic model of DEX which included height as a covariate for central clearance in non-obese (mean body weight of 82 kg) adult volunteers. Valitalo et al. performed a population pharmacokinetic analysis of DEX in critically ill patients in the intensive care unit (ICU) and found that TBW strongly correlates to clearance, and their final model included a TBW covariate on clearance. Because the formula structure that they used was based on the allometric scaling technique which is frequently used for pharmacokinetic analysis of anesthetics in obese patients, the increase in clearance with body weight was at an exponential rate. However, the number of obese patients included in the study was not specified in the literature. Talke et al. also built a pharmacokinetic model of DEX in young and non-obese patients who underwent hypophysecto-

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**Fig. 3.** Correlation diagram between LBM and DEX amount per hour (A) and between LBM and FEN amount per hour (B). All patients were analyzed together. LBM, Jannmahasatian’s equation for Lean Body Mass (kg).
my and found body weight or height not to be a co-
variate for clearances or distribution volumes. In a
pharmacokinetic model of DEX reported by Lin et
al.\textsuperscript{14)}, height was included as a covariate in central
clearance but the study population did not cover
obese patients. Another pharmacokinetic model
built by Iirola et al.\textsuperscript{15) showed that DEX systemic
clearance significantly decreased with decrease in
normalized cardiac output (cardiac output / cardiac
output at baseline) which potentially enhances the
relationship between body weight and clearance.
Lee et al.\textsuperscript{16) investigated DEX pharmacokinetics in
young, healthy, non-obese adult volunteers and
found that body weight or height were not covariates
for clearances or distribution volumes.

In general, pharmacokinetic simulations can be
performed to predict plasma DEX concentrations
using drug administration data and pharmacokinetic
model parameters indicated in literatures as in those
previously mentioned. However, it is strictly incor-
rect to extrapolate those results into an obese popu-
lation because commonly the results from pharma-
cokinetic simulations for patients from special
populations using pharmacokinetic models which are
not built using data from such populations can be in-
accurate. Therefore, pharmacokinetic analysis is
required in obese population. Our real clinical data
might support such study.

Besides the sedative effect, DEX activates Al-
pha-2 adrenergic receptors in the spinal cord and in-
hibits pain transmission, which causes an analgesic
effect. According to the predicted plasma DEX
concentration resulting from the usual clinical dose,
the strong analgesic effect by a single agent cannot
be proposed\textsuperscript{17). However, a synergic effect of DEX
with opioids is promising\textsuperscript{18) and clinically the con-
sumption of opioids is reported to be reduced when
administered together with DEX\textsuperscript{19,20). Our study
design is not enough to discuss the synergic effect of
the two drugs because there was no group to which
a single agent or a fixed dose was administered.

In order to examine if the ratio of the doses of
Fig. 5. Plots of the three indices of DEX and FEN doses in terms of the control group (open circle), obese group (closed circle), and combined.
DEX and FEN changes with obesity, we performed the exploratory analysis as illustrated in Figure 5 although no significant correlations between DEX and FEN consumption were observed in the control, obese and combined groups. However, these results may be interpreted as that the real time drug titration by nursing staff were modestly well performed in which sedation and analgesia were assessed independently. Strong significant correlations between two drug dose indices would rather imply the monotonic increase in two drug doses even though clinically there should be various requirements of the ratio of the doses of DEX and FEN. At least, during the observation periods after the initial recovery of consciousness, no patients had RASS score of 5 (i.e., Unarousable) for two consecutive observation points, which represents that the oversedation was potentially avoided in all patients (data not shown).

There are several limitations in this study. First, we had small number of patients because it has not been long since DEX infusion lasting more than 24 h after surgery in the ICU was approved (2010) and we have relatively few obese patients in Japan. Second, patients’ conditions were various and co-administered cathecolamine dose differed. Cardiac output affects the metabolism of DEX as mentioned earlier but it was not investigated. Third, the inclusion/exclusion criteria did not include administration of lidocaine21,22, which is considered to have a potential analgesic effect. Fourth, the initial and maximum doses of DEX were decided at the discretion of attending anesthesiologists or intensivists who were not blinded to weight of patients, which might have reflected to drug dose. These limitations should be considered when interpreting the results.

In conclusion, DEX requirements were increased with obesity when administered with FEN during mechanical ventilation after surgeries but not as much as those calculated by per kg basis. These findings will be beneficial for future clinical pharmacological analysis of DEX.

DISCLOSURES

Shinju Obara and Masahiro Murakawa received a speaker’s honorarium from Maruishi Pharmaceutical CO., Ltd. (Osaka, Japan) which sells dexmedetomidine in Japan for delivering a lecture and for presiding at a meeting, respectively; the contents of the present study were not included in the lecture. All other authors have no Conflict of Interest to declare.

REFERENCES

1. Egan TD, Huizinga B, Gupta SK, et al. Remifentanil pharmacokinetics in obese versus lean patients. Anesthesiology, 89 : 562-573, 1998.
2. Ingrande J, Brodsky JB, Lemmens HJ. Lean body weight scalar for the anesthetic induction dose of propofol in morbidly obese subjects. Anesth Analg, 113 : 57-62, 2011.
3. Shibutani K, Inchiosa MA, Jr., Sawada K, Bairamian M. Accuracy of pharmacokinetic models for predicting plasma fentanyl concentrations in lean and obese surgical patients: derivation of dosing weight (“pharmacokinetic mass”). Anesthesiology, 101 : 603-613, 2004.
4. Coetzee JF. Total intravenous anaesthesia to obese patients: largely guesswork? Eur J Anaesthesiol, 26 : 359-361, 2009.
5. Sessler CN, Gosnell MS, Grap MJ, et al. The Richmond Agitation–Sedation Scale: validity and reliability in adult intensive care unit patients. Am J Respir Crit Care Med, 166 : 1338-1344, 2002.
6. Janmahasatian S, Duffull SB, Ash S, Ward LC, Byrne NM, Green B. Quantification of lean body-weight. Clin Pharmacokinet, 44 : 1051-1065, 2005.
7. Dutta S, Lal R, Karol MD, Cohen T, Ebert T. Influence of cardiac output on dexmedetomidine pharmacokinetics. J Pharm Sci, 89 : 519–527, 2000.
8. Katayama M, Yamazumi K, Kino K, Tsuru M, Fukazawa T, Shimada H. Change of Liver Weight in the Elderly. (in Japanese) Jpn J Geriat, 27 : 584-588, 1990.
9. Peeters MY, Aarts LP, Boom FA, et al. Pilot study on the influence of liver blood flow and cardiac output on the clearance of propofol in critically ill patients. Eur J Clin Pharmacol, 64 : 329-334, 2008.
10. Dyck JB, Maze M, Haack C, Azarnoff DL, Vuorilehto L, Shafer SL. Computer-controlled infusion of intravenous dexmedetomidine hydrochloride in adult human volunteers. Anesthesiology, 78 : 821-828, 1993.
11. Valitalo PA, Ahtola-Satila T, Wighton A, Sarapohja T, Pohjanjousi P, Garratt C. Population pharmacokinetics of dexmedetomidine in critically ill patients. Clin Drug Invest, 33 : 579-587, 2013.
12. Coetzee JF. Allometric or lean body mass scaling of propofol pharmacokinetics: towards simplifying parameter sets for target-controlled infusions. Clin Pharmacokinet, 51 : 137-145, 2012.
13. Talpe P, Richardson CA, Scheiman M, Fisher DM.
Postoperative pharmacokinetics and sympatholytic effects of dexmedetomidine. Anesth Analg, 85: 1136-1142, 1997.

14. Lin L, Guo X, Zhang MZ, Qu CJ, Sun Y, Bai J. Pharmacokinetics of dexmedetomidine in Chinese post-surgical intensive care unit patients. Acta Anaesthesiol Scand, 55: 359-367, 2011.

15. Iirola T, Ihmsen H, Laitio R, et al. Population pharmacokinetics of dexmedetomidine during long-term sedation in intensive care patients. Br J Anaesth, 108: 460-468, 2012.

16. Lee S, Kim BH, Lim K, et al. Pharmacokinetics and pharmacodynamics of intravenous dexmedetomidine in healthy Korean subjects. J Clin Pharm Ther, 37: 698-703, 2012.

17. Ebert TJ, Hall JE, Barney JA, Uhrich TD, Colinco MD. The effects of increasing plasma concentrations of dexmedetomidine in humans. Anesthology, 93: 382-394, 2000.

18. Hendrickx JF, Eger EI, 2nd, Sonner JM, Shafer SL. Is synergy the rule? A review of anesthetic interactions producing hypnosis and immobility. Anesth Analg, 107: 494-506, 2008.

19. Arain SR, Ruehlow RM, Uhrich TD, Ebert TJ. The efficacy of dexmedetomidine versus morphine for postoperative analgesia after major inpatient surgery. Anesth Analg, 98: 153-158, table of contents, 2004.

20. Kim SY, Chang CH, Lee JS, Kim YJ, Kim MD, Han DW. Comparison of the efficacy of dexmedetomidine plus fentanyl patient-controlled analgesia with fentanyl patient-controlled analgesia for pain control in uterine artery embolization for symptomatic fibroid tumors or adenomyosis: a prospective, randomized study. J Vasc Interv Radiol, 24: 779-786, 2013.

21. De Oliveira GS, Jr., Fitzgerald P, Streicher LF, Marcus RJ, McCarthy RJ. Systemic lidocaine to improve postoperative quality of recovery after ambulatory laparoscopic surgery. Anesth Analg, 115: 262-267, 2012.

22. Kang H, Kim BG. Intravenous lidocaine for effective pain relief after inguinal herniorrhaphy: a prospective, randomized, double-blind, placebo-controlled study. J Int Med Res, 39: 435-445, 2011.