



Bountra, K., & Gill, H. (2020). The Physiological Effect of 1MAC Sevoflurane with or without 50% Xenon in Immature Rats. *Paediatric Anaesthesia*. <https://doi.org/10.1111/pan.13933>

Peer reviewed version

Link to published version (if available):  
[10.1111/pan.13933](https://doi.org/10.1111/pan.13933)

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# Pediatric Anesthesia

## The Physiological Effect of 1MAC Sevoflurane with or without 50% Xenon in Immature Rats

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| Journal:                      | <i>Pediatric Anesthesia</i>  |
| Manuscript ID                 | PAN-2020-0200.R1   |
| Wiley - Manuscript type:      | Short Report   |
| Date Submitted by the Author: | n/a  |
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| Key Words:                    | General anesthesia, pharmacodynamics < Drugs, Neurodevelopment, inhaled agents < Drugs, sevoflurane < inhaled agents < Drugs   |
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3 i. Title: The Physiological Effect of 1MAC Sevoflurane with or without 50% Xenon in  
4 Immature Rats  
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8 ii. Running title: MAC Sevoflurane +/- Xenon in Immature Rats  
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44 Keywords: Xenon, anesthesia, postnatal development, pharmacokinetics  
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49 Funding Statement: This work was funded by The Academy of Medical Sciences Starter  
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52 Grant for Clinical Lecturers (AMS-SGCL13-Gill). HG was funded by NIHR during this work  
53  
54 (CL2014-25-003).  
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## Background and Methods

Anesthetic-induced neurodegeneration in rats occurs across the first three postnatal weeks<sup>1</sup> (human equivalent: fetus to pre-schooler), during which the central nervous system, respiratory control and pharmacodynamics undergo huge changes. In immature rats, hypercarbia alone (mimicking isoflurane anesthesia-induced hypercarbia) induced neurodegeneration<sup>2</sup> and Xenon reduces anesthetic-induced neurodegeneration<sup>3</sup>.

To facilitate comparison of equipotent anesthetic mixtures of sevoflurane and Xenon, we aimed to measure the mean concentration of sevoflurane +/- 50% Xenon preventing purposeful movement in 50% of immature rats (MACsevo and MACsevoXe) on postnatal days (P)5, 10 and 15. 50% Xenon was used as it is clinically useful; offering potential neuroprotection and delivery with up to 50% oxygen.

These experiments were carried out and approved under Home Office License and with University ethical approval. For the measurement of MACsevo and MACsevoXe, three, mixed-sex, rats of equal age were exposed to: 3.0% sevoflurane or 2.5% sevoflurane in 50% Xenon, respectively (estimated, a-priori, as MAC). Gasses were delivered using calibrated, low flow, rotameters and monitoring ensured that CO<sub>2</sub> rebreathing was limited to 2%. After 30 minutes, a 50mm bulldog clamp was applied to the mid-portion of the tails until purposeful movement was seen (or 30secs maximum). According with the up-and-down method, this response rate dictated the concentration the subsequent three animals were exposed to (i.e. where 2 or 3 animals moved, the concentration was increased by 0.2% or 0.4%, respectively, and vice versa). In this way, nine doses of sevoflurane +/- Xenon were administered to each age group and doses 2-9 (dictated by rat responses) were used to calculate a mean: MAC (SD).

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3 In further experiments, groups of six rats on P5, 10 and 15, were randomised to two-hour  
4 exposures to: MACsevo, MACsevoXe, sevoflurane alone in equal concentration to  
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6 MACsevoXe (sevoControl) or 30% oxygen alone (Control). Respiratory rate was monitored  
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8 throughout. Immediate decapitation allowed collection of mixed arterio-venous blood for  
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10 analysis. Unless severe circulatory failure is present, the mean difference between venous  
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12 and arterial pH and partial pressure of CO<sub>2</sub> (PCO<sub>2</sub>) is negligible<sup>4</sup>. We defined acidosis as  
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14 pH<7.35 and normal lactate as <3.0mmol/L.  
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19 An independent samples T-test was used to compare MACsevo with MACsevoXe at each  
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21 age. ANOVA with Tamhane's post-hoc test (equal variance not assumed) was used to  
22  
23 compare differences in MAC across the three ages and differences in physiology between  
24  
25 treatment groups.  
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## 30 Results

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34 MACsevo (SD) was higher on P10: 3.83% (0.42) than P5: 3.24% (0.27) ( $p<0.001$ ), and P15:  
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36 3.28% (0.30) ( $p<0.001$ ) and higher than MACsevoXe (SD) on P5, P10 and P15: 2.26% (0.22)  
37  
38 ( $p<0.001$ ), 2.39% (0.26) ( $p<0.001$ ) and 2.37% (0.22) ( $p<0.001$ ) with no age-difference. Of  
39  
40 note, five pups on P5 died (all exposed to the highest concentrations used). They were  
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42 allocated as *no response to tail clamping* for calculations (excluding them changed MAC  
43  
44 slightly, but not the statistical comparisons).  
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49 The physiology results are presented in the Figure. Blood was successfully analysed from  
50  
51 65/72 animals (lactate in 57). In the Control group pH was normal at all ages. 53/54 animals  
52  
53 exposed to sevoflurane +/- Xenon survived but were acidotic. A dose effect, where  
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55 sevoflurane alone lowered RR, was seen at all ages. However, this effect was only seen in  
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57 PCO<sub>2</sub> and pH on P5 and P10. Differences between the MAC groups were only seen on P10.  
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3 Only animals in the MACsevoXe groups had raised lactate (three on P5 with pH 6.72-6.82,  
4 lactate 8.22-10.03mmol/L, one on P10 with pH 7.05, lactate 3.02mmol/L). An additional  
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8 group was therefore exposed on P5 and showed similar results.  
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## 10 11 Conclusions

12  
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14 MACsevo was higher on P10 than P5 and P15, in keeping with another rat study<sup>5</sup> and with  
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16 humans (peaking during infancy). 50% Xenon had a significant sevoflurane-sparing effect.  
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18  
19 Respiratory depression and acidosis were severe, but comparable between MACsevo and  
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21  
22 MACsevoXe, potentially enabling robust comparison of any neuroprotective effect in future.  
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25 In the younger animals, respiratory depression was marked, and co-administration of Xenon  
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27 and sevoflurane appeared to increase lactate. On P15 there is evidence that the rats were  
28  
29 better able to compensate for respiratory depression. The physiological derangements  
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31 shown in our study highlight the need for large animal models for optimal translation to the  
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34 clinic.  
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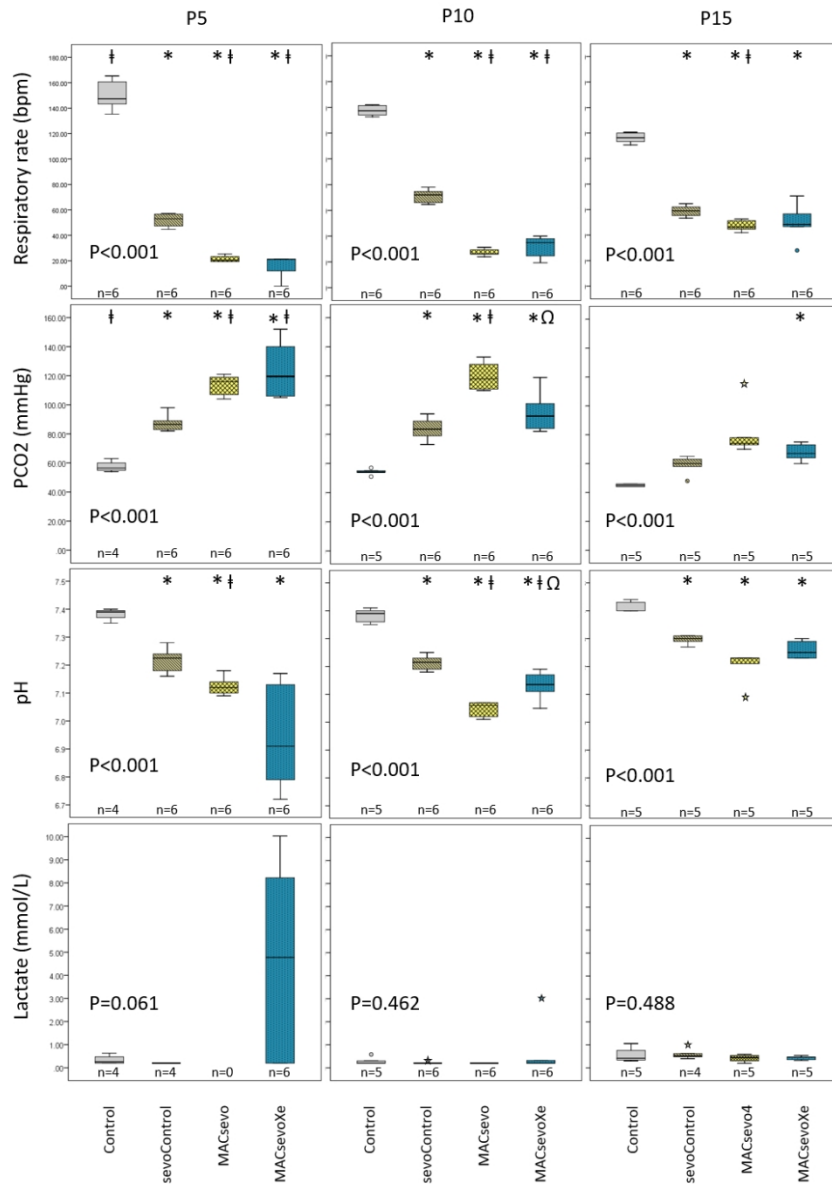
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18 Figure legend:  
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20 Median (IQR and range) of the average respiratory rates, and the partial pressure of CO<sub>2</sub>  
21 (PCO<sub>2</sub>), pH and lactate concentration in mixed arterio-venous blood taken from immature  
22 rats at three ages (postnatal days (P)5, 10 and 15), during and immediately after two-hour  
23 exposures to: 30% oxygen alone (Control), sevoControl (sevoflurane alone in equal  
24 concentration to that in MACsevoXe), MACsevo, or MACsevoXe.  
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32 The differences between the exposure groups at each age were tested using one-way  
33 analysis of variance (P value shown in each panel and significant differences from Control  
34 (\*), sevoControl (‡), or MACsevo (Ω) are shown (Tamhane's post hoc test). An open circle or  
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star represents individual outliers and extreme outliers, respectively.



Median (IQR and range) of the average respiratory rates, and the partial pressure of CO<sub>2</sub> (PCO<sub>2</sub>), pH and lactate concentration in mixed arterio-venous blood taken from immature rats at three ages (postnatal days (P)5, 10 and 15), during and immediately after two-hour exposures to: 30% oxygen alone (Control), sevoControl (sevoflurane alone in equal concentration to that in MACsevoXe), MACsevo, or MACsevoXe. The differences between the exposure groups at each age were tested using one-way analysis of variance (P value shown in each panel and significant differences from Control (\*), sevoControl (†), or MACsevo (Ω) are shown (Tamhane's post hoc test). An open circle or star represents individual outliers and extreme outliers, respectively.

178x250mm (144 x 144 DPI)