# **Bovine Embryo Development Rates are Affected when Oocytes are Matured in Different Vials Containing HEPES/Bicarbonate Buffered Medium**



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#### Introduction

Lab ware for *in vitro* produced embryos is generally made from embryo tested plastic instead of glass. The quality of the plastic is crucial for the outcome, as plastic is often toxic to the gametes. In addition, gas molecules permeate through the plastic at a rate that depends on a variety of factors, such as diffusion coefficient and thickness of the plastic, thus influencing medium pH. Furthermore, medium volume in the vial and the number of

cumulus-oocyte-complexes (COCs) are important factors contributing to pH alteration in the medium. Emission of CO<sub>2</sub> from cumulus cell metabolism contributes to a decrease in pH in low gas permeable vials, whereas a low medium volume increases pH in high gas permeable vials. HEPES buffer does maintain pH for a longer period of time, but not indefinitely when bicarbonate is also present.

## **Objectives**

To choose the optimal system for oocyte transportation during maturation (IVM). Blastocyst rates were compared after maturing different numbers of oocytes, 5, 20 and 45, in glass vs plastic vials.

After screening several plastic vials, the least toxic was chosen. Two different maturation medium volumes, 50 % and 95 %, were assessed in the two different vials.

#### Results

In Experiment 1 the highest blastocyst rates, 40 % and 43 %, were obtained in glass vials with COC numbers

In Experiment 2 the highest blastocyst rates were also obtained in the glass vials, though not significantly different. In both experiments the lowest blastocyst rates, 28 % and 29 %, were obtained in glass vials with 45 COCs in

between 5 and 20, in 50 % medium volume. The corresponding pH values in the maturation medium were 7.7 and 7.6 respectively, after 21 hours of IVM. The maturation medium pH was 7.3 at the start of IVM.

both medium volume groups. The corresponding pH value in both groups was 7.0. Furthermore, there was a clearly visible pH gradient in the glass vials with 95 % medium volume. See Fig. 2.1.









		- —		centage of inseminated oocvtes
Plastic 5	171	32 <u>+</u> 7.0	7.6	(Mean $\pm$ SD). Four replicates with a
Glass 20	245	43 <u>+</u> 6.2	7.6	total of 1488 COCs.
Plastic 20	223	30 <u>+</u> 6.0	7.6	Values in a column and within ex-
Glass 45	220	28 <u>+</u> 5.9	7.0	were different
Plastic 45	236	36 <u>+</u> 6.1	7.5	<sup>a,b</sup> : P ≤ 0.1; <sup>c,d</sup> : P ≤ 0.05; <sup>e,f</sup> : ≤ P 0.1
Table 1.1				
	Plastic 5 Glass 20 Plastic 20 Glass 45 Plastic 45	Plastic 5171Glass 20245Plastic 20223Glass 45220Plastic 45236	Plastic 5       171       32±7.0         Glass 20       245       43±6.2         Plastic 20       223       30±6.0         Glass 45       220       28±5.9         Plastic 45       236       36±6.1	Plastic 5       171       32±7.0       7.6         Glass 20       245       43±6.2       7.6         Plastic 20       223       30±6.0       7.6         Glass 45       220       28±5.9       7.0         Plastic 45       236       36±6.1       7.5

(Mean $\pm$ SD). Four replicates with a total of 1375 COCs.	
were different	
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#### **Materials and Methods**

pH indicator:	A pH indicator guide was created using color-matched photographs for glass (VWR DK, NSCAC 4015-96, 1 ml) and plastic vials (Sigma-Aldrich, CLS430661, 2 ml) con-			
	taining media at various pH levels with measurements taken using a pH Meter (Fig. 3).			
	The visual pH value estimated from vials was reconfirmed by measuring pH (Fig. 3).			
Media:	All media used were from IVF Bioscience, UK.			
Experiment 1:	Slaughterhouse oocytes were randomized into 6 maturation groups: 5, 20 and			
	45 COCs per vial in glass and plastic, respectively.			
	<ul> <li>Matured (IVM) 21 hours in BO-HEPES IVM, at 38.8 °C without CO<sub>2</sub>.</li> </ul>			
	Vials contained 50 % medium volume/vial.			
	<ul> <li>Fertilized (IVF) overnight in BO-IVF under a 5.5 % CO<sub>2</sub> in a 38.8 °C humidified</li> </ul>			
	atmosphere.			
	• Cultured (IVC) in BO-IVC under a 5.5 % CO <sub>2</sub> , 5.5 % O <sub>2</sub> , 89% N <sub>2</sub> in a 38.8 °C			
	humidified atmosphere.			
	Twenty-one hours post IVM the medium pH was assessed from Fig. 3 and			
	presumptive zygotes were vortexed to remove cumulus cells and then cultured in			
	BO-IVC for 7 days (8 days post-IVF) when blastocyst rates were assessed			
	(Table 1.1., 1.2, Fig. 1.2. and 2.2).			
Experiment 2:	Same protocol as above, but with 95 % maturation medium volume/vial.			
<b>Statistical Analysis:</b>	Statistical analysis was performed with Chi Square and levels of significance			
	at P $\leq$ 0.1 and $\leq$ 0.05.			





## Conclusion

Occytes can be successfully matured without CO<sub>2</sub> incubation during transport with the selection of the:

- Correct vial
- Correct medium volume
- Correct number of oocytes
- Correct maturation medium

Surprisingly, the acceptable pH range in the maturation medium is rather wide for subsequent good blastocyst rates. The highest rates, 40 %, 43 %, 38 %, and 35 % were obtained in pH 7.7, 7.6, 7.3 and 7.2, respectively.

# **Final Conclusion:**

The toxicity of plastic vials was demonstrated as the glass vial groups showed significantly higher blastocyst rates with more or less the same pH in both vial volume groups with the same number of oocytes, 5 and 20, during maturation.

The gas permeability in the plastic vials was clearly demonstrated as the pH was maintained in all plastic groups at 7.4 - 7.6 independently of number of oocytes and medium volume. In the glass vials with 45 oocytes in both medium volume groups the pH dropped to 7.0, impairing the blastocyst rates. Furthermore, when the medium volume was 95 % a pH gradient occurred (0.2 to 0.3 pH units), with the lowest pH in the medium surrounding the COCs at the bottom of the glass vial.

The optimal blastocyst rates were obtained when oocytes were matured in glass vials in 50 % medium volume/vial and with 5 - 20 oocytes per vial.

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