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# Amnion Cells Engineering: A New Perspective in Fetal Membrane Healing after Intrauterine Surgery?

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## **Key Words**

Fetoscopy · Fetal membranes · Premature rupture of membranes · Amnion cell · Tissue engineering

#### **Abstract**

In this study we aimed to set up an in vitro culture of the rabbit amnion in order to support in vivo fetal membrane healing capacity following fetoscopy. Fetal membranes were collected from a mid-gestational rabbit, and cultured on collagen support material for 14 days. 34 rabbits at 22–23 days gestational age (GA) underwent fetoscopy. The entry site was randomly allocated to 4 closure technique study groups: group I, human amnion membrane (n = 23); group II, collagen foil (n = 16); group III, collagen

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plug (n = 19), and group IV, collagen plug with cultured amnion cells (n = 19). In all groups membrane access sites were additionally sealed with fibrin sealant, and the myometrium was closed with sutures. Fetal survival, amnion membrane integrity, and the presence of amniotic fluid were evaluated at 30 days GA. Cultures showed good survival in the collagen support material. Increased cellularity, survival and proliferations were observed. The amnion at the access site resealed in 58-64% of cases in groups II–IV, but none of the tested techniques was significantly better than the other. Histological examination indirectly revealed the anatomic repair of the membranes, since no entrapment of the membranes could be demonstrated in the myometrial wound.

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# Introduction

Although endoscopic intrauterine surgery is gaining more and more acceptance for the treatment of different congenital malformations, iatrogenic preterm premature

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Table 1. Fetal outcome parameters (fetal survival, amniotic integrity, and amniotic fluid presence) at 30 days of gestational age

Treatment group	Survival		Amniotic integrity		Amniotic fluid presence	
	n	%	n	%	n	%
Negative control group (n = 121)	90/121	74.7	79/90	87.8	83/90	92.2
Positive control group $(n = 23)$	17/23	73.9	9/17	52.9**	13/17	76.5
Human amnion membrane (group I; $n = 23$ )	19/23	82.6	5/19	26.3***	9/19	47.4*
Collagen foil (group II; n = 16)	14/16	87.5	9/14	64.3**	11/14	78.6
Collagen plug (group III; n = 19)	17/19	89.5	10/17	58.8**	11/17	64.7**
Collagen plug with amniotic cells (group IV; n = 19)	16/19	84.2	10/16	62.5**	10/16	62.5**

rupture of the membranes still remains its main limitation, although membrane trauma appears to be more minor than with the open approach [1]. Therefore, the investigation of an effective membrane closure technique is mandatory for the further development of intrauterine surgery, especially with a view to endoscopic surgery. Hence, different closure techniques have been proposed [2]. Nevertheless, only in the last years have these techniques been evaluated in various experimental models [3–8]. The pregnant rabbit appears still to be the most valuable experimental model toward a better understanding and evaluation of membrane closure techniques and healing [4, 5, 9, 10]. With this model we recently demonstrated the best functional and anatomic membrane sealing after multilayer closure of uterine trauma following fetoscopy [11]. In the present study we suggest a further development of membrane closure techniques of fetoscopic access sites, proposing the combination of this highly promising multilayer closure with the in vitro culture of amnion cells on collagen support material, which was used also as a plug, in the same animal model.

# **Materials and Methods**

34 time-dated pregnant New Zealand rabbits were housed 2 days prior to surgery. The housing was quiet, at normal room temperature and normal daylight. At 22-23 days gestational age (GA) the first operation was performed. After induction of anesthesia with 4-5 mg/kg propofol i.v. (10 mg/ml, Propofol<sup>®</sup> 1% Fresenius Emulsion i.v., Fresenius Kabi Deutschland GmbH, Bad Homburg, Germany) and antibiotic prophylaxis with trimethoprim sulfadoxine 0.5 ml/kg s.c. (Borgal-Lösung® 7.5%, Hoechst Roussel Vet Vertriebs GmbH, Unterschleissheim, Germany), the animals were intubated and mechanically ventilated (Hallowell EMC Model AWS® Veterinary Anesthesia Workstation, Voelker GmbH, Kaltenkirchen, Germany). To achieve the desired blood propofol concentration during surgery, propofol (20 mg/ml) was given by a computer-programmed syringe driver (1.5 mg/kg/min i.v., Perfusor® ED 1-300, Braun GmbH, Melsungen, Germany). Additional fentanyl 0.01 mg/kg i.v. (0.5 mg/ml, Fentanyl® 0.5 mg Curamed Injektionslösung, Dr. W. Schwabe GmbH & Co., Abt. Curamed Pharma GmbH, Karlsruhe, Germany) and metamizol 50 mg/kg 1:3 dilute with NaCl i.v. (Metalgin® Injektionslösung, Hexal AG, Holzkirchen, Germany) were given as an analgesic component. Maternal heart rate and oxygen saturation were monitored with a pulse oximeter (Ohmeda, Helsinborg, Sweden) and the end-tidal CO<sub>2</sub> concentration was displayed by a capnometer. The animals were positioned in the supine position and the abdomen was shaved, disinfected with povidone iodine and draped in a sterile way. Interventions on the uterus and membranes were done with an operation microscope (Carl Zeiss, Oberkochen, Germany, magnification  $\times 5-25$ ) and micro-instruments. Fetoscopy was done with a short, 1.2-mm, 10,000-pixel, 0° fiber endoscope (Karl Storz, Tuttlingen, Germany) housed within a 14-gauge needle.

A midline abdominal incision was made to expose the pregnant uterus. First the gestational sacs were counted and numbered. Up to a maximum of one in two amniotic sacs were randomly assigned to 4 study groups, according to the closure technique that was to be tested (n = 77), excluding the gestational sacs above the cervix. The others served as positive (n = 23), as well as negative controls (n = 23) 121; table 1).

Under microscope vision and using micro-scissors a 2- to 3-mm myometrial incision was performed as already described [8] in order to separate the chorion and allow the amniotic membrane to bulge through the chorionic incision after gentle uterine pressure. The amniotic sac was then entered under microscopic control with the 14-gauge needle. Once in the amniotic cavity, the fetoscope was introduced into the needle to explore the amniotic cavity. During fetoscopy the fetal body, extremities, face, and tail, as well as the placenta or the umbilical cord were identified. The fetoscopy was limited to a maximum inspection time of 5 min with the use of up to 5 ml Ringer lactate amniodistention infusion at body tempera-

<sup>\*</sup> Versus negative controls: p < 0.001 ( $\chi^2$  Fisher's exact two-tailed test). \*\* Versus negative controls: p < 0.05 ( $\chi^2$  Fisher's exact two-tailed test). \*\*\* Versus negative controls and groups II–IV: p < 0.005 ( $\chi^2$  Fisher's exact two-tailed test).

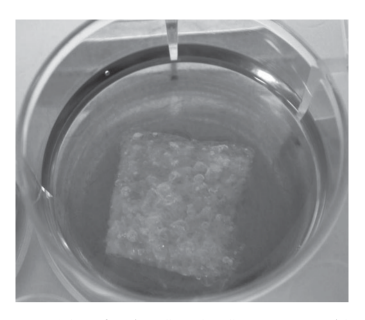
After withdrawal of the fetoscope, the access site was closed according to the study group (table 1). In study group I (n = 23) the fetoscopic access site was covered with human amnion membrane (Tissue Bank of the German Institute for Cell and Tissue Replacement, Leipzig, Germany) and 0.3 ml fibrin sealant was applied on the membrane defect area (Tissucol Duo S 2 ml Immuno<sup>®</sup>, Baxter Deutschland GmbH, Unterschleissheim, Germany). In group II (n = 16) the amnion membrane was covered with collagen foil (TissuFoil E<sup>®</sup>, Baxter Deutschland GmbH) and fibrin sealant. In group III (n = 19) the entry site was plugged with a collagen plug (Tissu-Fleece®, Baxter Deutschland GmbH) and fibrin sealant [11]. Prior to surgery, sterile dry collagen was precut and sized to the needle's lumen, with a maximal plug length of about 1 cm. To position the plug we made a purpose-designed pusher of a few millimeters shorter than the length of the needle. After withdrawal of the fetoscope from the needle, a plug was loaded into the lumen. The plug and the custom-made pusher were inserted one after the other into the needle, advancing the plug exactly to the tip of the needle point. Once in place, the plug inflates by absorbing surrounding fluid, and locks the access site. In group IV (n = 19) the above-mentioned collagen plug was covered with in vitro cultured allogeneic amnion cells, and positioned into the amnion and chorion membrane defect, as well as covered with fibrin sealant, in the same way as in group III.

### Amnion Cell Isolation

Amnion cells were isolated under sterile conditions 3 weeks prior to surgery from amnion membrane biopsies of mid-gestational pregnant rabbit. The fetal amnion membrane was harvested under aseptic conditions after euthanasia. The membrane was minced into small pieces. These pieces of fetal membrane were then immersed in 0.25% trypsin-EDTA (Biochrom AG, Berlin, Germany) in PBS-Dulbecco (Biochrom AG) for 20 min at 37°C with agitation. Subsequently, the cells were isolated by digestion with 0.02% clostridial collagenase (Roche AG, Mannheim, Germany) dissolved in Dulbecco's modified Eagle's medium (Gibco, Paisley, UK) supplemented with 10% fetal calf serum (Gibco), 500 U penicillin/500 ml, as well as 50 mg streptomycin/500 ml (Biochrom AG, Berlin, Germany). The cells were cultured at 37°C, and 5% CO<sub>2</sub> in keratinocyte medium (Keratinocyte-SFM® & supplements, bovine pituitary extract, h-EGF recombinant, Gibco).

7–10 days prior to surgery the isolated amnion cells were 90% confluent and were removed by 0.25% trypsin-EDTA treatment. Cells were counted (using a Neubauer-cell counting chamber) and vitality was determined by trypan blue staining. Afterwards, the cells were resuspended, and 1  $\times$  10<sup>6</sup> cells in 1 ml keratinocyte medium were injected into each collagen plug using a 1-ml syringe and a 22-gauge needle. Cells were injected to infiltrate different injection sites and to guarantee a regular spread of cells. Drained-off cell suspension was seeded on the top of the collagen sponge. After 20 min of incubation, the dish was filled with 3–4 ml keratinocyte medium so that the top of the sponge was covered with medium. Then the collagen plugs, loaded with the cells, were incubated at 37°C and 5% CO<sub>2</sub> up to the time of implantation (fig. 1).

In all study groups the myometrial layers were closed microsurgically with a nylon 6/0 suture (Prolene®, Ethicon GmbH, Norderstedt, Germany) [11]. 23 sacs were left unclosed (positive controls), and the 121 unmanipulated sacs served as negative controls.

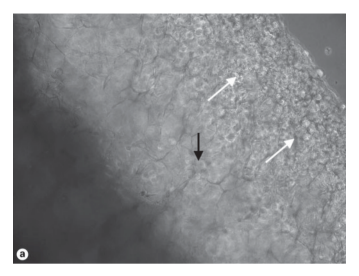


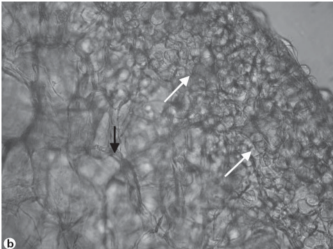
**Fig. 1.** Culture of amnion cells on the collagen support material (TissuFleece E<sup>®</sup>) at 37°C, and 5% CO<sub>2</sub> in keratinocyte medium.

After repositioning the uterus, the abdomen was closed in layers using polyglactin 3/0 (Vicryl<sup>®</sup>, Ethicon GmbH) for the fascia and intracutaneous nylon 2/0 suture (Ethilon<sup>®</sup>, Ethicon GmbH) for the skin. Postoperative uterine relaxation was achieved by administering 4.5 mg/kg medroxyprogesterone-acetate i.m. (Depo-Clinovir<sup>®</sup> 150 mg, Pharmacia & Upjohn GmbH, Erlangen, Germany). The animals were housed for the following 8 days under the same conditions as prior to surgery.

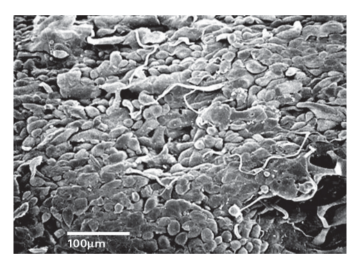
At 30 days GA the pregnant rabbits were sacrificed with a pentobarbital injection, 60 mg/kg (Narcoren®, Rhone-Merieux, Laupheim, Germany) to undergo a second-look laparotomy. A myometrial incision was made at the antimesenterial side, starting over the first cornual gestational sac [4]. The incision was extended over the full length of the horn, both over the treated as well as over the control sacs. Under microscopic control the myometrium was gently dissected with microsurgical instruments. The main outcome measurements were the integrity of amniotic membranes and the presence or absence of amniotic fluid. In case of an intact amniotic membrane, integrity was verified by an intra-amniotic injection of 5-10 ml saline solution dyed with methylene blue. The injection was given through the membrane, opposite to the fetoscopic access site with a 30-gauge needle. This way any leak eventually missed before could be visualized. The fetus was then delivered and it was noted whether the fetus was alive or not. The living fetuses were weighed and dissected to assess the wet fetal lung weight and to calculate the fetal lung to body weight ratio. Macerated stillborn fetuses were noted as non-surviving and not further included in statistical processing.

Statistics were done using the  $\chi^2$  Fisher's exact two-tailed test for nominal variables and ANOVA for continuous variables using the SPSS 12.0 for Windows software package (SPSS Inc., Chicago, Ill., USA).





**Fig. 2.** Vital amnion cells adherent (white arrow) on the collagen sponge matrix (black arrow; TissuFleece<sup>®</sup>). Original magnifications:  $\mathbf{a} \times 10$ ;  $\mathbf{b} \times 40$ .



**Fig. 3.** 10 days after seeding the isolated amnion cells, an almost confluent layer of amnion cells firmly attached to the collagen support material (TissuFleece  $E^{\otimes}$ ) can be seen. SEM. Original magnification  $\times$  240. Bar = 100  $\mu$ m.

## **Results**

Amnion cell cultures showed good survival in the collagen support material (fig. 2). Increased cellularity, survival and proliferations were observed (fig. 3). No adverse effects on animal pregnancy and the fetus could be documented after the use of in vitro cultured amnion cells in the collagen plugs (group IV).

Fetal survival was similar in the study groups except in the positive and negative control group (table 1).

The integrity of the amniotic membranes of unmanipulated sacs near term was 87.8%. After fetoscopy without any attempt to close the membranes (positive control group) the amniotic integrity rate 7 days later was 52.9%. The use of human amnion membrane combined with fibrin sealant and myometrial suture (group I) did not improve the amniotic integrity rate compared to that of the positive control group at second-look operations. In group II (Collagen foil®, Tissucol® and myometrial suture) the integrity of fetal membranes was 64.3%, and 58.8% in group III (Collagen plug, Tissucol® and myometrial suture). The use of the collagen plug in combination with in vitro cultured amniotic cells improved the integrity of fetal membranes to 62.5% (group IV).

The presence of amniotic fluid differs according to the results of amnion membrane integrity. In absolute numbers, collagen foil showed the best result with 78.6%, while covering the access site with human amnion membrane resulted in only 47.4% amniotic integrity.

Fetal lung/body weight ratio of unmanipulated sacs with intact membranes and treated gestational sacs (positive control and study groups: n = 83) was compared with intact and ruptured membranes. Fetal lung and body weights of fetuses from gestational sacs with unmanipulated membranes and treated groups with intact membranes were similar, and higher than in fetuses from treated gestational sacs with ruptured membranes. The fetal

**Table 2.** Comparison of fetal lung/body weight ratio according to integrity of amnion (mean  $\pm$  SD)

	Negative control	Treatment groups			
	group (n = 79)	sacs with intact membranes $(n = 43)$	sacs with ruptured membranes (n = 40)		
Body weight, g Lung weight, g Fetal lung/body weight ratio	$31.35 \pm 11.731$ $1.29 \pm 0.510$ $0.0412 \pm 0.00651$	$31.64 \pm 9.887$ $1.25 \pm 0.447$ $0.0394 \pm 0.00667$	$29.47 \pm 11.987$ $1.08 \pm 0.423*$ $0.0369 \pm 0.00710*$		

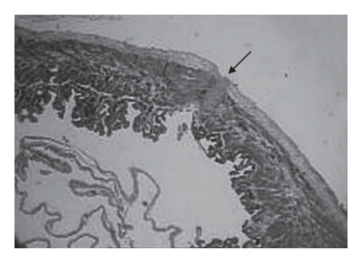
Stillborn fetuses were not included.

lung/body weight ratio between the treated groups with ruptured membranes and the negative control group with amnion integrity were statistically different (p < 0.05, table 2).

The histological examination of the access sites indirectly revealed the anatomic repair of the membranes as no entrapment of fetal membranes could be confirmed in the myometrial wound (fig. 4).

#### **Discussion**

After the first successful intrauterine intervention in the pregnant guinea pig animal model [12], intensive development of pioneering methods for prenatal diagnosis [13], as well as pre-, intra-, and postoperative management to maximize feto-maternal survival, only recently began in the last decades of the 20th century [14]. Today, high-resolution ultrasound, magnetic resonance imaging (MRI), and other diagnostic procedures allow an earlier and more precise intrauterine diagnosis of congenital anomalies [15], and even their intrauterine treatment has become reality. Therefore, prenatal treatment can nowadays be considered as a treatment option not only for life-threatening malformations but also for non-lifethreatening anomalies like spina bifida [16–18]. Moreover, recent advances in endoscopic interventions have allowed a safer intrauterine approach to the fetus, thereby increasing the number of centers performing such interventions, although the risk of preterm premature rupture of the membranes and preterm labor exists [1]. Although the prospects for fetal treatment of non-lifethreatening disorders such as craniofacial malformations are still unclear, their intrauterine treatment will depend upon whether and when the known complications of



**Fig. 4.** Van Gieson staining of the access site of treatment group IV. Evidence of scar tissue in the myometrial wound (arrow), but no entrapment of the fetal membranes, representative of all amnion sacs in which fetoscopy was performed, followed by myometrial suture. Lower left site: Presence of the membranes at the central part of the amniotic sac. Original magnification  $\times 4.5$ .

these invasive procedures (amniotic fluid leakage, membrane disruption, and oligohydramnios) can be treated satisfactorily [19, 20].

For these reasons, the mid-gestational rabbit was proposed as the first animal model to investigate fetal membrane healing after hysterotomy and fetoscopic exploration of the amniotic sac [4, 5, 8–11]. In the meantime, this model has been established as the gold standard for this kind of study, and will surely contribute substantially toward the further development of intrauterine surgery.

<sup>\*</sup> p < 0.05 as compared with negative control with intact membranes (ANOVA (LSD)).

Consequently, in the animal model mentioned, our laboratory modified the previously described multilayer surgical closure technique of the fetal membranes [11], introducing for the first time as an additional option the in vitro culture of amnion cells and their utilization on collagen support material, which could likewise be used as a plug. This combination of a collagen plug for the fetal membranes with in vitro cultured amnion cells and fibrin sealant as well as myometrium closure with sutures following fetoscopy was found to be reproducible and achieved the promising goals we expected with regard to anatomic healing of the membrane defects. In our opinion this method is presently the most encouraging one, and can surely be developed further considering the learning curve of such a new technique in our hands. Therefore, we speculate that further improvement in the in vitro culture of the amnion cells would result in better sealing rates of the fetal membranes after endoscopic fetal surgery. Additionally, as the 'pure' plug technique (group IV), this method is applicable to percutaneous interventions as well.

Nevertheless, further experimental studies are obligatory, not only in order to investigate possible postoperative complications using these collagen plugs to seal the iatrogenic rupture of membranes, but also to better understand the biology of the amnion cells and their healing process, and to further develop this technique in order to achieve safe and excellent surgical results. Only under such conditions will it be possible in the future to clinically assist the human fetus in a wider manner and, therefore, also those with non-life-threatening malformations such as the craniofacials [19, 20].

#### **Conclusions**

This animal culture system represents a new model for the study of fetal membrane repair after iatrogenic defect. Despite the good survival and localized proliferation observed in the collagen support material, no obvious statistically better closure of fetoscopy-induced permanent membrane defects was seen under the conditions described. Therefore, additional research is needed to further explore the mechanisms of in vitro culture of fetal membranes on support materials, as well as the use of other possible support materials to improve fetal membrane healing following iatrogenic trauma.

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