Introduction

Oncological treatment is becoming increasingly effective and cancer patients often have a good long-term prognosis, allowing them to lead normal lives. This includes fulfilling their desire to have children [1]. As chemotherapy – depending on the therapy regime – can be gonadotoxic, a possible irreversible decrease or complete loss of ovarian reserve may result. Cryopreservation of ovarian tissue with subsequent orthotopic re-transplantation after recurrence-free remission...
is a promising method to preserve fertility in affected women [2, 3]. To date, more than 25 cases have been published worldwide reporting on live births after orthotopic re-transplantation of cryopreserved ovarian tissue. In Germany the first live birth after re-transplantation of cryo-preserved ovarian tissue occurred in 2011 [4].

A 25-year-old patient developed a Hodgkin lymphoma in 2003. As the patient was from Dresden, she underwent laparoscopic removal of ovarian tissue in the Gynaecological Clinic of Dresden University Hospital to preserve her fertility. Cryopreservation was done in Bonn, as cryopreservation of ovarian tissue is not done routinely in Dresden. For this, the tissue was transported in a suitable transport container to the Gynaecological Clinic of Bonn University Hospital where it was cryopreserved and the tissue samples were stored.

Re-transplantation was done in Erlangen in view of the acknowledged expertise of the Gynaecological Clinic of Erlangen University Hospital in thawing and re-transplanting ovarian tissue. After 5 years with no recurrence of cancer and in view of the patient’s continued wish to have a child, the thawed ovarian tissue was re-transplanted in the Gynaecological Clinic of Erlangen University Hospital in 2010, with laparoscopic re-transplantation of the tissue into a peritoneal pocket of the ovarian fossa in the right pelvic wall. Three months after re-transplantation the patient reported her first spontaneous menstrual bleeding, and low-dose stimulation therapy with follicle-stimulating hormone (FSH) was started. After confirmation of a dominant follicle in the transplant area, ovulation was induced with human chorionic gonadotropin (hCG), and timed intercourse was recommended. A pregnancy ensued. The pregnancy was completely uneventful, and a normal boy was delivered in the Gynaecological University Clinic in Dresden on October 10, 2011 [4].

We describe here in detail the method of tissue transportation from Dresden to Bonn, the preparation and cryopreservation of the ovarian tissue together with the method used for thawing and re-transplantation.

**Transport and Preparation of Ovarian Tissue for Freezing**

Except where otherwise stated, all chemicals were obtained from Sigma (Sigma Chemical Co., St. Louis, MO, USA). Laparoscopic extraction of ovarian tissue for cryopreservation was done on August 22, 2005 in the Gynaecological Clinic of Dresden University Hospital. After extraction, the tissue was sent to the Gynaecological Clinic of Bonn University Hospital, a member of the FertiPROTEKT network. Within 22 hours, the extracted ovarian tissue fragments were sent from Dresden to Bonn in a Brahma I solution (Cryo Bio System, Paris, France) in a special transport flask cooled to 4°C for the transportation of ovarian tissue. The company Cryo Bio System no longer produces the Brahma I solution. We now use a solution produced by the Gynaecological Clinic of Cologne University Hospital consisting of Leibovitz’s L-15 medium with 5% SSS (Serum Substitute Supplement, Irvine Scientific, Santa Ana, CA, USA) as an alternative to the Brahma I solution for the transport and preparation of ovarian tissue. The next day, at the Gynaecological Clinic of Bonn University Hospital the ovarian tissue fragments were transferred to a Brahma II medium (Cryo Bio System, Paris, France) and the cortex was partially separated from the stroma using tweezers and a no. 22 scalpel. The Brahma II medium is currently also no longer produced by Cryo Bio System. Instead of the Brahma II solution, clinics can make up their own solution, e.g. using Leibovitz’s L-15 medium with 5% SSS. The ovarian tissue fragments were divided into 5–8 strips with a length each of 2–3 × 1.0–1.2 mm and transferred to 2 ml cryo tubes (Nunc, Roskilde, Denmark) for freezing.
Freezing of Ovarian Tissue

To freeze the ovarian tissue, 2 ml standard cryo tubes (Nunc, Roskilde, Denmark) were filled with 1.8 ml freezing solution, consisting of L-15 medium (Leibovitz) with L-glutamine + 1.5 M (11.7%) DMSO + 10% SSS. The ovarian strips of tissue were transferred to the cryo tubes, placed in ice water for 30 min (Fig. 3a) and subsequently frozen using the IceCube 14S freezer (SyLab, Neupokkersdorf, Austria) (Fig. 3b). The freezing protocol was as follows: (1) starting temperature 4°C; (2) cooling from 4 to −6°C at a speed of −1°C/min; (3) auto-seeding at −6°C; (4) cooling from −6 to −34°C at a speed of −0.3°C/min, and finally, immersion of the cryo tubes in liquid nitrogen (Fig. 3c). The tubes were then transferred to the cryobank and stored (Fig. 3d).

Thawing of Ovarian Tissue

Thawing of the ovarian tissue was done at the Gynaecological Clinic of Erlangen University Hospital. The cryo tubes with the fragments of ovarian tissue were left at room temperature for 30 s and subsequently placed in 37°C warm water for 1 min until the ice had completely melted. After defrosting, the pieces of ovarian tissue were removed from the cryo tubes and placed in 5 ml thawing solution (0.75 M sucrose solution in PBS with 10% human albumin solution [5% HAS]) for 15 min at room temperature. The tissue fragments were then placed in 0.5 M sucrose solution for 15 min, then in 0.25 M sucrose solution for 15 min, and finally transferred to a PBS solution (with 10% HAS). Directly afterwards, the tissue was transplanted using the technique previously described [5].
A number of different procedures are successfully used world-wide for the cryopreservation of ovarian tissue. Births have only ensued when slow freezing procedures were used. Cryoprotectants include ethylene glycol, propylene glycol and DMSO. The procedure described here used only DMSO as cryoprotectant and resulted in the birth of the first child to be born in Germany after re-transplantation of cryopreserved ovarian tissue. The procedure described is also suitable for freezing ovarian tissue. The Gynaecological Clinic of Cologne University Hospital currently uses a modified freezing protocol, in which ethylene glycol is used for cryopreservation in addition to DMSO [5–10]. The new freezing solution consists of L-15 medium (Leibovitz) with L-glutamine + 6% DMSO + 6% ethylene glycol + 0.15 M sucrose + 10% SSS (serum substitute supplement).

In cryopreservation, the time-point at which crystal formation is initiated during the freezing process is particularly important, because if crystal formation is not initiated, so-called spontaneous crystal formation will occur at around −18 °C. This generates exothermic crystallisation heat, which has an indirectly injurious effect on biological structures. Up until the moment of seeding, the rapidity of the freezing process is not critical. After seeding, the cooling speed must not exceed −0.3 °C/min.

In January 2005, an auto-seeding block was designed and produced at the Gynaecological University Clinic in Bonn. The prototype of this block is shown in Fig. 3b. The ovarian tissue obtained in the case described here was frozen, and seeding was done using this prototype. The basic principle of the auto-seeding block is as follows. To initiate seeding, liquid nitrogen is conveyed through a metal tube at a certain time-point during the freezing process (at −6 °C in our protocol). Seeding is initiated by thermal transfer created by the mechanical contact of the cryo tube with the nitrogen-filled metal tube. The special feature of this auto-seeding block is that contact between the cryo tube and the seeding tube occurs passively, without any oscillation, only through gravitational force. The cryo tube with the cryo solution is pressed against the seeding tube by its own weight.

The Cologne Study Group recommends the following alternative thawing protocol with continuous addition of a thawing medium. The cryo tube is first left to stand at room temperature for 30 s and then immersed for a period of 60 to 75 s in a water bath with boiling water (100 °C). The precise immersion time of the cryo tube in boiling water is controlled visually by monitoring the ice: the cryo tube is removed from the water bath as soon as the ice at the tip is thinner than −1 mm. The final temperature of the freezing solution after removal of the cryo tube from the 100 °C water bath should be between 4 and 10 °C.

If thawing is done in boiling water, there is always a risk of overheating the ovarian tissue fragments. To prevent this we recommend the following. First a compact sponge (Telaprep No. 1, Paul Hartmann AG, Heidenheim, Germany) is placed at the bottom of the 5 ml cryo tube. The ovarian tissue fragment is then placed on the sponge. This prevents the tissue from coming into direct contact with the walls of the 5 ml cryo tube and ensures rapid but “delicate” warming of the ovarian tissue fragments.

After initial thawing, the strips of ovarian tissue are transferred within 5–7 s to a 100 ml sample container (Sarstedt, Nürnberg, Germany) with 10 ml solution (0.5 M sucrose + 10% SSS + L-15 medium) to remove the cryoprotectants.

Removal of cryoprotectants is carried out following the same principles used for ethylene glycol saturation (cf. Figure 1 in Isa-chenko et al., 2006 [7]). The container is constantly agitated by a shaker (200 osc/min at room temperature for 15 min). Multiple stage rehydration of the strips of ovarian tissue is done at room temperature for 30 min. Drop rehydration is done using 50 ml culture medium (L-15 medium + 10% SSS) in 50 ml tubes (Greiner Bio-One GmbH, Frickenhausen, Germany). The method requires the slow addition (drop by drop) of a culture medium to the sucrose solution. The final sucrose concentration is 0.083 M, which practically corresponds to cellular isotonic concentrations. The strips of ovarian tissue are subsequently washed for 10 min in culture medium.

It is important to maintain constantly low temperatures during transportation and preparation of the strips of ovarian tissue to preserve tissue vitality (Fig. 1 and 2). Below, we briefly describe the results of our own tests, carried out in the Gynaecological Clinic of Cologne University. The overall aim of these studies was to investigate the effect of low positive temperatures on the viability of human ovarian tissue and to prove or refute any benefit arising from the use of low positive temperatures.

Analysis of hypothalamic tissue function showed that cold temperatures resulted in a twofold increase in thyrotrophin-releasing hormone mRNA synthesis [11]. Moreover a good survival of trophoblastic fragments subjected to 48 h cooling at 4 °C was shown. The survival rate did not differ from that of untreated controls (98 and 98%, respectively) [12].

Previous studies have shown that storage of human ovarian tissue for up to 26 h at low positive temperatures has no inhibitory effect on follicle development during later in vitro culture. Compared to untreated controls the number of primordial follicles was significantly decreased in all treatment groups due to developments in advanced stages [13].

Wood et al. [14] investigated the impact of storing domestic cat ovaries for a longer period under hypothermic conditions (at 4 °C for 48 h) on follicle atresia and recorded a positive impact. The data, described below, support their results.

In contrast, the study by Yin et al. [15] reported that hypothemric storage of rat ovaries at 4 °C for 24 h did not have an atretic effect on ovarian function. Although the ovaries had fewer follicles, this is contrary to the results of the Cologne Study Group. We would explain this phenomenon as species-specific. The most important factor for the successful transplantation of ovarian tissue is ensuring sufficient blood supply within a very short time. This is decisive for follicular survival [16].

Given that neo-vascularisation is important for nidation and the development of ovarian tissue, the Cologne Study Group uses neo-vascularisation as the criterion for tissue viability and has verified this criterion. The formation of new vessels from already existing vessels is vitally important for many physiological processes. Dysregulation of vascularisation contributes to many diseases, from ischemia to cancer [17].

In fact, neo-vascularisation starts with the formation of very small vessels. The Cologne Study Group used desmin as a marker to demonstrate the formation of these vessels. In our experi-

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ments pieces of human ovarian tissue were divided into the following groups:

- Group 1: pieces cultivated immediately after operation,
- Group 2: pieces cooled to 5 °C for 24 h after operation and then cultivated,
- Group 3: pieces frozen immediately after operation, then thawed and cultivated,
- Group 4: pieces cooled to 5 °C for 24 h after operation, then frozen, thawed and cultivated.

Ovarian tissue fragments were cultivated for 5 days on chorioallantoic membrane (CAM). The impact of cooling on follicles and the intensity of neo-vascularisation was recorded and assessed based on the detection of desmin expression. Increased neo-vascularisation in the presence of desmin expression was demonstrated. The intensity of immunoreactivity was recorded semi-quantitatively [18]: no immunoreactivity (−), weak immunoreactivity (+), moderate immunoreactivity (++), strong immunoreactivity (+++).

Histological investigation after 5-day CAM culture of pieces of ovarian tissue showed that the primordial, primary and secondary follicles remained viable. Normal follicle proportions (92, 93, 89 and 90%) were recorded for groups 1, 2, 3 and 4, respectively (p > 0.1).

Immunoreactivity for desmin was only seen in the medullary part of the tissue fragments. Groups had the following immunoreactivity:

- Group 1 (pieces cultivated immediately after operation): weak (+),
- Group 2 (pieces cooled to 5 °C for 24 h after operation): strong (+++),
- Group 3 (pieces frozen, then thawed after operation): none (−),
- Group 4 (pieces cooled to 5 °C for 24 h after operation, then thawed): moderate (++).

The results of other experiments described elsewhere showed that 24-hour cooling to 5 °C prior to cryopreservation has a positive effect on human ovarian tissue [19].

Our article also contains data previously published elsewhere [4, 20]. In their study, Müller et al. described various aspects of the procedure used for cryobanking of ovarian tissue [4]. The publication also describes the first birth in Germany after the re-transplantation of cryopreserved ovarian tissue [4].

It was noted that oncologists should inform their young female patients with tumour disease scheduled to undergo radio- or chemotherapy more about the different possibilities of preserving fertility and offer more counselling on fertility preservation to these patients.

Dittrich et al. [20] reported on the biological part of the cryobanking process, describing the successful cryopreservation of ovarian tissue after lengthy transportation under hypothermic conditions. Moreover, they proposed taking the possibilities to preserve fertility into consideration when planning oncological treatments and offering patients counselling on fertility preservation by a member of the FertiPROTEKT network (www.fertiprotekt.de).

More than 25 births worldwide have been recorded after re-transplantation of cryopreserved ovarian tissue [21]. The aim of our paper was to describe the protocol for ovarian tissue cryopreservation. The focus was on providing detailed information which can be used directly for laboratory protocols. The birth of a child has confirmed the reliability of our protocol. However, we have not confined ourselves to describing technologies from 2005 but instead present more recent versions of these technologies. The practical application of the results of our experiments into the cooling of human ovarian tissue is described. Nowadays, the Gynaecological Clinic of Cologne University Hospital carries out cooling of human ovarian tissue for 22–23 h prior to cryopreservation as part of its standard protocol.

The reader is thus given the opportunity to decide for himself whether he wishes to utilise the technology used in 2005 or whether he prefers to opt for the technology developed over the last 6 years [2, 3, 6, 7, 19].

**Conflict of Interest**

None.

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**References**


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