Fertility Preservation: Successful Transplantation of Cryopreserved Ovarian Tissue in a Young Patient Previously Treated for Hodgkin’s Disease

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LEARNING OBJECTIVES

After completing this course, the reader will be able to:

1. Discuss recent advances in the use of cryopreserved ovarian tissue to restore fertility.
2. Explain the main aspects of the procedure for transplantation of cryopreserved ovarian tissue.
3. Discuss the options to preserve fertility of young patients with a high risk for premature ovarian failure after cancer therapy.

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ABSTRACT

Cryopreservation of ovarian tissue is now offered as an experimental procedure to preserve the fertility of young patients with a high risk for premature ovarian failure resulting from cancer therapy. This is the only available option to preserve the fertility of prepubertal patients treated with gonadotoxic chemotherapy. At present, thousands of patients all over the world have undergone this procedure with the hope of later restoring their fertility. Although the efficiency of the transplantation of cryopreserved ovarian tissue to restore ovarian function has been established, reports of pregnancy are still very scarce. Here, we describe the second published full-term spontaneous pregnancy after an orthotopic and heterotopic transplantation of cryopreserved ovarian tissue in a 31-year-old woman previously treated by conditioning therapy for bone marrow transplantation for Hodgkin’s disease. This birth gives compelling evidence for the graft origin of the gamete and confirms the efficacy of ovarian tissue transplantation in restoring human natural fertility after oncological treatment. This case report stresses the importance of proposing the ovarian tissue cryopreservation procedure to all young patients who require potentially sterilizing treatment, with all alternative options to preserve fertility being duly taken into consideration.

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INTRODUCTION
As a result of improvements in oncological treatments, most young cancer patients achieve prolonged survival [1, 2]. For these young patients, ensuring their reproductive capacity after the oncological treatments has become a main concern, as it is directly related to their quality of life. However, premature ovarian failure is one of the most common long-term adverse effects affecting premenopausal patients treated with alkylating agents and/or radiation therapy [2–4]. The risk for premature ovarian failure depends mainly on the age of the patient, the type and dose of chemotherapy, and the irradiation field. Moreover, the resumption of cyclic menses after oncological treatment does not guarantee normal fertility. In a recent review, the chances of spontaneous pregnancy were estimated to be 28% and only 5% in women <20 years and >25 years of age, respectively, at the time of cancer treatment [5]. Conditioning therapy for bone marrow transplantation (BMT) is considered the most gonadotoxic treatment, inducing premature ovarian failure in >80% of cases, even when performed during childhood [6].

In recent years, the cryopreservation of ovarian tissue prior to gonadotoxic treatments has offered new hope for these young patients and is considered as an investigational procedure with the aim of restoring fertility [7–10]. Restoration of ovarian function after cryopreserved ovarian tissue transplantation has been described previously [11, 12]. However, data concerning fertility restoration after this procedure are still very limited. The first spontaneous full-term pregnancy after orthotopic cryopreserved ovarian tissue transplantation was reported by Donnez et al. [13] in 2004. However, the gamete’s origin has been questioned in this case, and indeed spontaneous ovulation from the remaining in situ ovary could not be excluded [14]. Following cryopreserved ovarian tissue transplantation, another full-term pregnancy was reported after an in vitro fertilization (IVF) procedure and embryo transfer [15].

We report here the second spontaneous full-term pregnancy after orthotopic and heterotopic transplantation of cryopreserved ovarian tissue in a patient previously treated by BMT. The discussion addresses the indications, advantages, and limitations of this procedure considering its recent success and the other available options.

CASE PRESENTATION
At age 24, the patient received intensive chemotherapy before autologous BMT for Hodgkin’s disease, which consisted of six courses of ABVD (doxorubicin, 50 mg/m²; bleomycin, 20 mg; vincristine, 12 mg/m²; dacarbazine, 750 mg/m² per course), three courses of EVA (etoposide, 300 mg/m²; vinblastine, 6 mg/m²; doxorubicin, 50 mg/m² per course), and a CBV conditioning regimen (cyclophosphamide, 6,400 mg/m²; carmustine, 450 mg/m²; etoposide, 1,500 mg/m²). Unilateral ovariectomy was performed after the first ABVD course, allowing the cryopreservation of 40 pieces of ovarian cortex, as previously described [16]. After BMT, the patient was considered to be disease free but never recovered spontaneous menstrual cycles. Four years later, she expressed a wish to conceive. However, amenorrhea and hormonal blood tests confirmed premature ovarian failure (follicle-stimulating hormone [FSH] of 87 mIU/ml and inhibin B <15 ng/ml). Biopsy of the remaining ovary revealed the absence of follicles.

A first orthotopic (peritoneal and ovarian sites) and heterotopic (s.c. abdominal site) transplantation of cryopreserved ovarian tissue was then performed by a two-step laparoscopy (day 0, November 2004) [16]. Before the procedure, three fragments of ovarian tissue were thawed and histologically examined for tumor cells and follicular density (12 follicles/mm²) [16].

After the transplantation procedure, ovarian function was continuously monitored. Basal FSH and inhibin B (measured between day 3 and day 5 of each menstrual cycle) were used to evaluate ovarian reserve. Estradiol was regularly measured during each cycle and ovulation was determined based on the luteinizing hormone surge. Progesterone was measured during each luteal phase. Follicular development was assessed at each transplanted site by ultrasonography [16].

The ovarian tissue transplantation allowed recovery of ovarian function, and a spontaneous clinical pregnancy was obtained following the sixth cycle. Unfortunately, the patient miscarried at 7 weeks because of aneuploidy (day 310) [16].

One year after the first ovarian tissue transplantation, a progressive increase in FSH values and a decrease in inhibin B levels, both ovarian reserve markers, suggested progressive deterioration of the graft (Fig. 1A). Ovarian stimulation was attempted without success, after 21 days of contraceptive pill administration, reducing the basal FSH value.

To recover a normal hormonal status and to increase, as much as possible, the chance of pregnancy, a second orthotopic and heterotopic transplantation was performed in May 2006. Four pieces of ovarian cortex were thawed as previously described [16] (Fig. 2A). By a two-step laparoscopy, two ovarian fragments were sutured on the remaining ovary (Fig. 2B). The other two fragments were divided and placed s.c. using the left abdominal incision made for the trocar (Fig. 2C). No intra- or postoperative complications were observed.

During the first 3 months, her basal FSH levels re-
mained high (>20 mIU/ml), although the patient again ovulated (Fig. 1B). Follicular development was observed at both the ovarian and right s.c. sites (first s.c. transplanted site, Fig. 2D). No follicular development was observed at the left s.c. site (second s.c. transplanted site). From the third cycle, basal FSH levels returned to normal premenopausal values (<10 mIU/ml). During the fifth spontaneous cycle following the second transplantation procedure, two follicles of 15 mm diameter each were observed at the ovarian site at the time of ovulation. A positive human chorionic gonadotropin level was detected at day 14 of the luteal phase and a clinical pregnancy was later confirmed by vaginal ultrasound. The pregnancy evolved normally without any complication and the patient gave birth to a healthy girl at 41 weeks’ gestation.

DISCUSSION
Considering the impact of infertility on long-term quality of life, all young patients who potentially face premature ovarian failure should receive accurate information concerning the available options to preserve their fertility. A previous report concerning young patients affected by breast cancer showed that receiving fertility-related information was rated as significantly more important than receiving menopause-related information at the time of diagnosis [17]. In the same way, the majority of pediatric oncological patients and their parents are concerned by the fertility impact of...
treatment, but only around one third of them are satisfied with the amount of information received [18]. The oncologist’s initiative in starting a discussion, supported by a multidisciplinary team, is a determinant of improving access to information and proposing the optimal options to preserve fertility.

The first option is to reduce the gonadotoxicity of the cancer treatment with the intention of preserving ovarian function and future natural fertility. This includes ovarian transposition, in the case of pelvic irradiation, or adaptation of the chemotherapy regimen (Fig. 3). Based on promising results from animal studies, concomitant administration of a gonadotropin-releasing hormone (GnRH) agonist during treatment with alkylating agents has also been proposed to reduce the gonadotoxicity of the drugs [19, 20]. The efficiency of GnRH agonists in preserving human fertility was suggested by a preliminary study [21]. However, no prospective, randomized trial confirming these results has yet been reported. Thus, at present, concomitant administration of a GnRH agonist should only be proposed as part of a clinical trial.

The second option is to cryopreserve gametes in order to later restore fertility (Fig. 3). Around 15 oocytes can be harvested after FSH ovarian stimulation of young cancer patients [22]. In the case of hormone-sensitive tumors such as breast cancer, alternatives to the gonadotropin ovarian stimulation protocol can be proposed [3]. When a partner is present, oocytes can be fertilized and the resulting embryos cryopreserved. Cryopreservation of embryos is a well-established procedure routinely performed in IVF clinics. In the absence of a partner, mature unfertilized oocytes can be cryopreserved, but the clinical pregnancy rate for thawed oocytes is still low (2%) [23]. The main limitation of the oocyte or embryo cryopreservation procedure in cancer patients is the 2- to 4-week delay needed to achieve ovarian stimulation and oocyte collection. Furthermore, this procedure cannot be proposed if the patient has already started chemotherapy treatment because of genetic risk and poor efficacy [24].

Cryopreservation of ovarian tissue has the advantage of allowing the storage of a large number of primordial follicles, it can be performed at any time in the menstrual cycle without delay, and it is available for prepubertal patients. Options to restore fertility from cryopreserved ovarian tissue include follicular in vitro culture and ovarian tissue transplantation. However, the development of in vitro culture systems supporting primordial follicular growth in humans will probably need a few more years before being available. Ovarian tissue transplantation has been performed previously with success in humans. At present, >25 cases of heterotopic and orthotopic cryopreserved ovarian tissue transplantation have been reported. However, only three births have been reported after this procedure—one baby was conceived naturally after spontaneous ovulation [13] and the others were conceived using an IVF procedure ([15]; Anderson et al.

![Figure 3. Options to preserve or restore fertility in cancer patients.](http://theoncologist.alphamedpress.org/)

**Abbreviations**: GnRH, gonadotropin-releasing hormone; IVF, in vitro fertilization.
ovarian tissue transplantation is ischemic damage during return to ovarian failure status. One of the major issues after the first transplantation procedure suggests a progressive humans [16]. The increase in basal FSH values 1 year after months) reflects the time necessary for folliculogenesis in to restore ovarian function after the first transplantation (5 follicular growth. As previously described, the time needed evolutions is also compatible with the graft being the origin of the hormonal profile after the first and the second transplanta- tion procedure was well documented by hormonal function is observed in <10% of the patients [25]. Furthermore, premature ovarian failure before the first transplantation procedure was well documented by hormonal profiles as well as by biopsy of the remaining ovary, confirming the absence of follicles [16]. The evolution of the hormonal profile after the first and the second transplantations is also compatible with the graft being the origin of the follicular growth. As previously described, the time needed to restore ovarian function after the first transplantation (5 months) reflects the time necessary for folliculogenesis in humans [16]. The increase in basal FSH values 1 year after the first transplantation procedure suggests a progressive return to ovarian failure status. One of the major issues after ovarian tissue transplantation is ischemic damage during the revascularization process, limiting the life span of the graft. According to the literature, the life span of the graft varies from a few months to >2 years, depending mainly on both the age of the patient at the time of the cryopreservation procedure and the quantity of ovarian tissue trans- planted [12, 26, 27]. However, a correlation between these factors and the life span of the graft is not easy to establish.

Another limitation of the procedure is the possibility of reintroducing malignant cells. According to previous studies, there has been no evidence of disease invasion in the ovarian tissue harvested from patients with Hodgkin’s lymphoma [28]. However, the safety of the procedure must be evaluated for each request, considering the type of disease and the ovarian tissue analysis. Despite reassuring data regarding the absence of relapse and the good health of babies born following this procedure, all children, and patients who conceived in this manner, should have long-term follow-up, especially when the ovarian tissue was exposed to systemic chemotherapy before cryopreservation.

Finally, if all previously described options are not feasible, alternative approaches to parenting, like oocyte dona- tion, have to be considered (Fig. 3).

**CONCLUSION**

This is the second full-term spontaneous pregnancy reported after transplantation of cryopreserved ovarian tissue. This birth represents a great hope for all patients with a high risk for premature ovarian failure after oncological treatment and provides evidence of the efficacy of the procedure in restoring natural fertility. This case report also stresses the importance of proposing ovarian tissue cryopreservation to all young patients who require potentially sterilizing treatment for cancer, especially when other procedures, such as oocyte or embryo cryopreservation, are not recom- mend.

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<th>Table 1. Inclusion criteria for ovarian tissue cryopreservation and transplantation procedures</th>
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<td><strong>Ovarian tissue cryopreservation</strong></td>
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<tr>
<td>Maximum age 35 years</td>
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<td>Risk for premature ovarian failure</td>
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<td>No previous high gonadotoxic treatment</td>
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<td>Presence of the uterus</td>
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<td>No HIV or hepatitis (B,C)-positive serologies</td>
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<td>No surgical contraindication</td>
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communication ESHRE 2007). Thus, the procedure should be still considered experimental with selected inclusion criteria (Table 1).

We report here the second spontaneous pregnancy following ovarian tissue transplantation and we provide strong arguments for the graft origin of the follicular development. The patient had previously received conditioning treatment for BMT, after which spontaneous recovery of ovarian function is observed in <10% of the patients [25]. Furthermore, premature ovarian failure before the first transplantation procedure was well documented by hormonal profiles as well as by biopsy of the remaining ovary, confirming the absence of follicles [16]. The evolution of the hormonal profile after the first and the second transplantations is also compatible with the graft being the origin of the follicular growth. As previously described, the time needed to restore ovarian function after the first transplantation (5 months) reflects the time necessary for folliculogenesis in humans [16]. The increase in basal FSH values 1 year after the first transplantation procedure suggests a progressive return to ovarian failure status. One of the major issues after ovarian tissue transplantation is ischemic damage during...


