Fertility Preservation Service

The Royal Children’s Hospital
fertility preservation principles
of care and guidance

For health professionals to use when helping newly diagnosed patients and families make choices about fertility preservation
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fertility preservation principles
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Amended draft for discussion December 2019 (3rd edition)
1st edition 2014
2nd edition 2016
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Introduction letter

Dear Colleagues,

Families report that information about fertility is one of their highest unmet needs at the time of cancer diagnosis. Suboptimal discussions and/or those occurring ‘too late’ (after onset of treatment) lead to regret. It is an international standard of care to inform families about the risks of infertility due to gonadotoxic treatment in a timely manner. The benchmark of care is to provide clear and consistent information about the impact of treatment on fertility where there is curative intent in line with the Australasian Oncofertility Charter.

Fertility preservation (FP) is not considered standard practice in children. FP is now approved as a novel technology at The Royal Children’s Hospital (RCH), with research governance for data collection and clinical ethics approval for individual cases (including for all pre-pubertal patients). This provides a governance framework within which clinicians can practice safely.

This guidance aims to provide clear and consistent information and resources for healthcare professionals for use in the introductory discussion of fertility for families of children and adolescents receiving medical therapy or having medical conditions that can affect fertility. It is hoped that these resources may reduce disparities in clinical practice and assist all families to receive good quality care irrespective of gender, culture, education and socioeconomic status. Where there is discord around these decisions, clinical ethics pathways can be activated by any team member so that consensus decisions can be made.

Further information and clinical pathways and forms can be found on http://www.rch.org.au/fertility/health-prof/. These pathways are under continuous renewal, informed by research as well as consumer and clinician voices. The Fertility Preservation Steering Committee is happy to provide information and advice to clinicians at the point of clinical care.

The oncofertility coordinator Rafael Serrano Real can be contacted via Tel: (03) 9345 5896 Ext: 55896 Pager 7047, and for non-urgent matters emailed on fertility@rch.org.au.

1. Talking to families about fertility risks

The following points will help prepare you for a discussion.

1.1 Prepare

We have included talking points below, a series of tables on risk of infertility based on treatment, potential recommendations according to risk of infertility, and fertility preservation options and their pros and cons. We have also included clinical ethics guidance, and a list of forms/hand-outs (clinical and research) which are also readily available on the RCH intranet. This information is a summary. It is not perfect and we endeavour to update it annually. We are aiming to provide this information in a more streamlined electronic format soon, integrated into the electronic medical record.

1.2 Who should be there?

Health providers can struggle to provide all of the critical information families require during fertility consultations, or may not consider it within their scope of practice. At the RCH, there are many providers who now feel confident in the quality of their fertility consultations and are happy to play a leading role. If you are uncomfortable or unsure about discussion of this topic, you may find it helpful to contact the Oncofertility Coordinator or identify another individual who can lead the discussion in your place, until you feel ready. It is helpful to have another team member present for the discussion if possible.

Give some thought to the age and gender of the young person, and whether it is appropriate for them to attend the discussion or receive age appropriate explanation and discussion separately. Cognitive capacity, well-being, emotional distress and pubertal status are important factors to consider, as well as the cultural background of the family. It is important not to exclude families from fertility discussions based on their background, age and gender of the child or the risk to fertility as families cope and adjust in different ways.

Young persons should, if possible, be key participants in the discussion, and supported through decision making about fertility in a way that is comfortable to them. There is a heavy reliance on parents during this time who are surrogate decision makers for young children. Be sure to ask mature adolescents and/or families who they would like involved in this conversation. Pubertal males may feel most comfortable discussing fertility matters and sperm preservation privately with a parent or a health professional. They may prefer a male staff member.

1.3 Timing and documentation

There is so much information to give to families at this distressing and overwhelming time. If you are discussing the details of the cancer diagnosis at the same time, consider taking a break and having a separate discussion. Some families have reported that they prefer the discussion to happen in a different space so that they can pay more attention to the issue of fertility after receiving the diagnosis. It is vital to provide written resources to families and to document your discussion in the notes. There are specialised fertility templates in the EMR for documentation and referral, designed to assist medical decision-making about fertility preservation.


1.4 Discussion points

1.4.1 Risk:

1. Discuss young person’s level of risk based upon your assessment of diagnosis and treatment plan.
2. Discuss flexibility of time for initiating treatment.
3. Explain what the level of risk means if no action is taken.
4. Explain that the experimental procedures are undertaken as a novel technology rather than standard practice.

1.4.2 Options:

1. Introduce potential options available for preserving fertility based on gender, pubertal stage, cancer, treatment plan and other relevant factors.
2. Clearly describe what is experimental and what is considered standard, and pros and cons.
3. Describe future options available if no action is taken.
4. Consider referral for fertility preservation (FP) according to medical/surgical risks, age and interest of patient (if mature) and family.

Discussions regarding experimental procedures should be measured and made with consideration: the final decision for experimental procedures will rest with the clinical teams and family, and depend on comorbidities. The surgical teams will ultimately decide upon the decision to proceed to surgery.

1.5 Research

RCH is undertaking one of the largest studies on the safety and efficacy of paediatric fertility preservation (HREC 33064). We recommend that all families be provided with information on this and invited to participate whether they decide to pursue fertility preservation or not, allowing outcomes to be compared between groups. In this research, parents can give permission for researchers to use information from the medical record. Optional consents include permission for data linkage (with the register of births, IVF centres), and permission for contact for future research.

1.6 What to remember:

1. All patients and families have the right to know if their fertility is at risk, even if:
   a. There is no time to do anything about it.
   b. There is nothing that can be done.
   c. Prognosis is poor.
   d. We do not think it is a good idea or necessary to preserve fertility.
2. When it is judged to be medically and ethically safe (by all relevant teams), with our information and support, the decision then rests with the patient and family.
3. Document discussions in the medical record.
4. Provide written resources to families.
5. Provide information sheet about the fertility audit (HREC 33064).
2. RCH fertility preservation principles for biological males

2.1 Purpose
To ensure that consistent discussion of infertility risk occurs with patients and their families having gonadotoxic treatment with curative intent, including the discussion of relevant preservation options for those having options available.

2.2 Principles
The RCH Children’s Cancer Centre, RCH Endocrinology, RCH Immunology, RCH Nephrology, RCH Rheumatology, RCH Surgery, RWH or Monash Andrology and RWH or Monash Reproductive Biology Units will work collaboratively to:
1. Provide education and consultation to all male patients and/or families where there is curative intent, about potential fertility impairment as a result of treatment in a clear and consistent manner with use of appropriate resource tools.
2. Discuss the pros and cons of fertility preservation (FP) options with these families including transparent discussion of standard versus experimental options.
3. Infertility risk can be based on risk stratification tables, but all situations are to be considered individually.
4. The medical teams must judge if FP is medically safe, in which case, the decision to proceed is value driven, made by patient/family in consultation with treating team.
5. Involve clinical ethics as appropriate.
6. Provide age appropriate discussions with patients when a FP procedure is being considered.
7. Document fertility discussions in the medical record.
8. Discuss participation in HREC 33064 prospective fertility audit for all patients having a fertility discussion whether they have a FP procedure or not.
9. Provide feedback of results, and appropriate follow up over the treatment journey and during survivorship.

2.3 Fertility preservation team can include:
1. Oncofertility coordinator.
2. Oncology or other treating team.
3. Endocrine consultant/fellow.
4. Surgical team.
5. Reproductive Biology Unit/Andrology liaison.
6. Oncology liaison.
7. Clinical ethics.
8. Lines coordinator.

2.4 Eligibility for the discussion
All new male patients having gonadotoxic treatment with curative intent should have a discussion about the impact of cancer treatment on fertility.

2.5 Initial discussion
It is recommended that the initial fertility discussion be undertaken by the treating clinician (e.g. oncology, renal, rheumatology, immunology) and include:
1. Assessment and discussion of risk of infertility attributable to planned chemotherapy, radiation or surgical procedure even if the risk is zero.
2. Discussion of possible fertility preservation (FP) options as appropriate.
3. Address any issues regarding sexual health in a confidential manner.
4. FP consultation can be formally documented in EMR using a template or just use your own notes.
5. Provide written resources to patient/family depending on age and type of FP procedure.
6. Invitation and consent for participation in the FP audit (which also includes consent for linkage to RWH FP database, consent to linkage with Register of Births, consent for contact for future research).

2.6 When to consider referral in biological males

2.6.1 Sperm collection
All post-pubertal male patients at any risk of infertility (can be referred directly to Andrology for sperm collection).

2.6.2 Testicular tissue cryopreservation:
1. Pre-pubertal patients at moderate to high risk of infertility.
2. Post-pubertal patients with inability to ejaculate or poor quality of sperm.
3. At request of patient/family, (note: investigational procedures are not usually undertaken if risk of infertility is low).
4. At discretion of treating team.

2.6.3 Fertility consultations also occur at other times in response to:
1. Males with relapsed disease and a new gonadotoxic treatment plan.
2. Referrals from any provider (nephrology, rheumatology, immunology, other).
3. At request of patients or families.
4. Males who have completed treatment and having long term follow-up who require surveillance.

2.7 Referral for sperm collection

2.7.1 Inpatient procedures for sperm preservation:
If you have a patient who would like to bank sperm while an inpatient, important principles include:
1. Ensuring the patient has privacy during collection. It is unacceptable for others to be present in the room unless requested by the patient.
2. Sometimes patients may not be able to produce a sample, due to a range of factors including distress, immaturity, and sickness. Sometimes a produced sample contains no viable sperm due to concurrent illness. Therefore it is best to discuss fertility preservation early, as multiple collections may be required.
3. A referral to Surgery or Endocrinology can be made to explore other options, such as testicular biopsy.
2.7.1 If it is a weekday follow the instructions below for sperm collection:

1. Contact oncofertility coordinator.
2. Give the patient/designated family member the following materials:
   a. Sperm preservation instructions.
   b. Male fertility preservation handouts:
      i. Having Children After Cancer (RCH handout for teens developed by psycho-oncology group).
   c. Sperm banking consent:
      i. If using Andrology Services — Andrology Unit Request for Sperm Storage.
      ii. If using Monash Medical Centre — Semen Storage Consent.
      iii. For Andrology Services complete Andrology Request Form and fax to: (03) 8345 3990.
      iv. Call the Andrology Laboratory on (03) 8345 3992 to report that request slip and consent are on the way.
3. Give the patient a specimen cup and a paper bag.
4. Secure a private space e.g. single patient room and hang ‘Stop Sign’. Instruct patient to remove sign when finished.
5. When the stop sign is removed, check with the patient to ensure:
   a. Specimen is in the cup with a tightly closed lid, and the cup is in a bag.
   b. Sperm banking consent forms completed.
   c. Payment method determined (the first two years are funded by My Room Children’s Cancer Charity).
6. Arrange a timely delivery of the specimen to the Andrology Lab.
7. The referring team is responsible for relaying results to the family. Please seek advice from Andrology or Endocrinology if you cannot interpret the results. If the specimen is inadequate, recollection may need to be attempted, or consider referral for TTCP. Including the oncofertility coordinator can facilitate these discussions and follow-up.
8. Arrange follow-up with Endocrinology to discuss results formally after treatment.
9. Invite the patient to participate in the Fertility Preservation Research Audit. Patients can be invited whether they decide to pursue fertility preservation or not:
   a. 33064 Parent Information and Consent Form if child is <18 years.
   b. 33064 Patient Information and Consent Form if young person is ≥18 years.
10. Document notes in the EMR.

2.7.1.2 After hours sperm collection

In the case of emergency cancer treatment, when sperm must be collected and stored after hours, please contact the ON CALL Andrology scientist in charge via RCH or RWH switchboard.

2.7.2 Outpatient procedures for sperm preservation

If you have a patient who would like to bank sperm as an outpatient, please take the following steps:

1. Contact oncofertility coordinator.
2. Give the patient/designated family member the following materials:
   a. Sperm preservation instructions.
   b. Male fertility preservation handouts:
      i. Having Children After Cancer (RCH handout for teens developed by psycho-oncology group).
   c. Sperm banking consent:
      i. If using Andrology Services — Andrology Unit Request for Sperm Storage.
      ii. If using Monash Medical Centre — Semen Storage Consent.
3. Invite the patient to participate in the Fertility Preservation Research Audit. Patients can be invited whether they decide to pursue fertility preservation or not:
   a. 33064 Parent Information and Consent Form if child is <18 years.
   b. 33064 Patient Information and Consent Form if young person is ≥18 years.
4. If using Andrology Services:
   a. Complete Andrology request form and fax to (03) 8345 3990.
   b. Call the Andrology Laboratory on (03) 8345 3992 to say that request slip and consent are on the way, and the family will be making an appointment.
   c. Direct the patient/family to make an appointment directly with the Andrology Service on:
      Andrology Unit/Sperm Bank
      The Royal Women’s Hospital, Carlton Campus
      321 Cardigan Street
      Carlton Vic 3053
      Tel: (03) 8345 3992 Hours: Monday–Friday 9am–5pm
5. Please follow-up the results prior to cancer treatment. If the count is low, then organize a repeat collection or consider referral for TTCP. The oncofertility coordinator can assist with this if notified and can provide a summary of fertility care letter.
6. Arrange follow-up with Endocrinology to discuss results at an appropriate time.
7. If the patient plans to collect a sample at home, provide container and paper bag and ensure sample will arrive at Andrology Lab within one hour of collection.
8. Document notes in the EMR.
2.8 Testicular Tissue Cryopreservation (TTCP) guidance for biological males

2.8.1 Eligible population for referral to Endocrinology for a TTCP consultation:
1. Pre-pubertal patients at moderate to high risk of infertility (defined as ≥20% risk).
2. Peri-pubertal or post-pubertal patient with inability to ejaculate or poor quality of sperm.
3. At request of patient/family.
4. At discretion of treating team.

Patients may include:
1. Oncology (potential number: 20–30/year).
3. Others may include rheumatology, immunology, neurology.

2.8.2 The role of the oncologist or other treating team:
1. Assessment and discussion with family of risk of infertility attributable to planned chemotherapy, radiation or surgical procedure even if the risk is zero.
2. Consider and discuss possible fertility preservation options as appropriate.
3. Formally document likely impact of treatment on fertility and discussion with the family in EMR using a ‘fertility’ template or just use your own notes.
4. Determine requirements for other co-existent operative procedures (CVL, LP, BMA).
5. Provide written resources to patient/family depending on age and type of FP procedure.
   a. Having Children After Cancer (RCH handout for teens developed by psycho-oncology group).
   d. Male TTCP Information Sheet and Pre-consent Form (for TTCP candidates).

6. Contact the oncofertility coordinator who can invite the patient to participate in the Fertility Preservation Research Audit. Patients can be invited whether they decide to pursue fertility preservation or not. The parent may consent on their behalf, however if the patient is 18 or above, they may complete their own consent form.
   a. 33064 Parent Information and Consent Form if child is <18 years.
   b. 33064 Patient Information and Consent Form if young person is ≥18 years.

7. If a referral is required, contact the oncofertility coordinator and endocrinology team via phone. Use the fertility referral form, which assists with FP decision-making. This should include:
   a. Brief history.
   b. Developmental maturity of patient (tanner stage testicular volume).
   c. Planned treatment, urgency (with clear indication of acceptable timeframe), prognosis, indication of infertility risk (high >80%/medium/low <20%).
   d. Handover of patient/family’s understanding of the diagnosis, prognosis, FP options/response to introductory discussion.
   e. Other procedures planned.
   f. Other factors pertinent to cancer/reproductive potential (e.g. prognosis, likelihood of RT, likelihood of relapse).
   g. Factors that may specifically increase surgical risk such as a mediastinal mass, significant immunosuppression, bleeding disorder.
   h. Social concerns within the family such as custody issues and parental disagreement.
   i. The level of complexity of the young patient’s situation that would necessitate consultant review.
   j. Any specific concerns the treating oncologist has with respect to proceeding with an FP procedure.
   k. Indicate intended date of gonadotoxic treatment and acceptable timeframe for FP.

8. If the referring clinician feels the patient/family should not yet be approached, they will let the endocrinologist know how and when it is best to see the patient. The oncology team should always have introduced the fertility discussion before the endocrine team sees the family for consultation.

2.8.3 Endocrinology consultation for TTCP:
1. Referrals will be made by oncology staff, sometimes other departments (e.g. Renal, Immunology), often at short notice before chemotherapy because they discuss life and death prior to FP. Endocrinology consult needs to fit in with:
   a. Timing of other essential tests (e.g. MRI, CT etc.).
   b. Timing of GA for port insertion.
   c. Availability of staff at the RWH for tissue storage/examination.
2. Patients may be seen a) in the ward or b) brought to endocrine clinic by a CCC staff member or c) sent there by oncology staff or d) at the Day Cancer Centre. Often a preliminary discussion has been made by oncology staff and the oncofertility coordinator and written documents provided. If not the documents are easily available on http://www.rch.org.au/fertility/health-prof/.
3. An endocrinologist needs to discuss issues which differ depending on:
   a. If the boy is pre-pubertal.
   b. Peri-pubertal (e.g. testes 10 ml, pubic hair stage 3).
   c. Post-pubertal.
4. The endocrinology consultation is as follows:
   a. Explanation of role as endocrinologist and purpose of visit.
   b. Assess and discuss young person’s level of infertility risk based upon age, pubertal stage diagnosis and treatment plan.
   c. Explain the level of risk to fertility if no action is taken.
   d. Introduce potential options for preserving fertility.
   e. Clearly describe what is experimental and what is considered standard, and pros and cons of the most appropriate FP plan for the individual.
   f. Details of testicular biopsy and the procedure.

   g. Clear statement that TTCP is currently experimental in humans – has been successful only in animals so far. It is undertaken at RCH as a novel technology.
   h. Explain that we always examine any biopsy at RCH, just in case there is some entirely unexpected abnormality such as no germ cells or spermatogonial cell lines, or malignancy. Please note that for patients with haematological malignancy it is important to have a fertility discussion. However the ability to sift cell lines and remove any potential cancer cells is not perfect at this time. Risks of malignant reseeding can be high and at the current point in time, tissue is not being autografted back into the body at local institutions. Therefore, it is undertaken at RCH as a novel technology.
   i. If the referring clinician feels the patient/family should not yet be approached, they will let the endocrinologist know how and when it is best to see the patient. The oncology team should always have introduced the fertility discussion before the endocrine team sees the family for consultation.

if in future the young man might wish to use the stored tissue to try to propagate sperm, this can only be attempted ex vivo if the technology has advanced enough to allow this to happen. Testicular tissue preservation may be contraindicated in acute lymphoblastic leukemia and lymphoblastic lymphoma and it is best to seek advice from the treating haematologist.

i. Freezing immature testicular tissues is free until 21 years of age at this time if undertaken with RWH. After this time the boy may decide whether to keep the tissue or not — can be a difficult decision at that time due to immaturity — storage may not be required if fertility is proven in the future by semen analysis.

j. For boys who are peri-pubertal at the time of a testis biopsy, tissue may also be dissected to look for mature sperm and any sperm found can be stored for intracytoplasmic sperm injection (seen in those as young as 11-12 years of age). This incurs the same cost as per sperm storage.

k. Please provide information to families as appropriate — available on the intranet under: www.rch.org.au/fertility/health-prof/.

i. Having Children After Cancer (RCH handout for teens developed by psycho-oncology group).

ii. Can I Still Have Children? (Information for Men Having Chemotherapy and Radiotherapy (MIVF Brochure for mature males)).


iv. Male TTCP Information Sheet and Pre-consent Form (for TTCP candidates).

2.8.4 If TTCP is to proceed:

1. Please document in the notes that parent has read and understood the Male Fertility Preservation Information Sheet and Pre-Consent Form (as suggested by RCH Legal). This form goes through the experimental nature of the procedure. Please provide this to families to keep.

2. FP surgical team will obtain surgical consent for FP procedure, which should clearly state its experimental nature.
   b. Consent from parent and assent from patient of sufficient maturity to understand concepts <18.

3. MIVF Testicular Tissue Cryopreservation Form and FPS Database forms to be completed.

4. FP oncology team with assistance of oncofertility coordinator will liaise with family, endocrinology team, lines coordinator, IVF Andrology scientists (RWH (03) 8345 3232 or lab.supervisors@mivf.com.au; Monash (03) 9420 8218) and clinical ethics regarding logistics and booking.

5. Lines coordinator will coordinate with surgeons, theatre and anaesthetists, and endeavor to book cases for the morning as each biopsy takes hours to process by the scientists. There is no reproductive lab service after 3 pm, on weekends or holidays.

6. Oncology and Endocrinology consultant to be notified if there are any delays to expected commencement of cancer treatment due to the FF procedure itself and make decision regarding risk-benefit.

7. The oncofertility coordinator will assist with review of histology (for example to ensure no malignancy on the tissue) and discussion with families, however it is the responsibility of the referring clinician to ensure this has been completed.

2.8.5 Involvement of clinical ethics in decision-making for TTCP:

1. Guidelines are as per the Ethics Checklist for Fertility Preservation Procedures.

2. If a clinical ethics meeting is needed, Clinical Ethics Service Referral Form 3A to be completed for pre-pubertal patients and Form 3B to be completed for post-pubertal patients. This should be regarded as an important medico-legal document for filing in the medical record.

3. If a clinical ethics meeting is held, the expected invitees include a representative from Oncology, Endocrinology and special experts where appropriate (e.g. Haematology/Genetics).

4. If the case issue is that the child is pre-pubertal then an expedited review may occur upon written referral.

2.8.6 Follow-up:

1. The patient/family should be referred after treatment to Paediatric Endocrine Oncology Clinic for further discussion, confirmation of storage arrangements and discussion of the evolving technology. This provides an opportunity to answer questions and manage expectations.

2. Timing of follow-up by Endocrinology is at the discretion of oncologist (around 12 months).

3. The oncofertility coordinator will provide a follow-up summary of fertility care letter to the family and copy to referring oncologist, and local doctor. Transition to an adult facility for discussion with an andrologist is recommended when appropriate.

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Figure 1. Testicular tissue cryopreservation pathway biological males

**Oncology/Clinical team**
- Utilise fertility preservation pathway
- Discuss impact of treatment on fertility (all patients)
- Provide written information to family:
  - Having Children After Cancer
  - Can I Still Have Children? Information for Young Men Having Chemotherapy and Radiotherapy

**Fertility preservation:** NO
- Document fertility note
- Invite onto the Fertility Preservation Research Audit (HREC 33064)

**Fertility preservation:** POSSIBLY
- Contact oncology coordinator
- Complete Fertility Preservation Form section A and Fertility Referral (Male) to Endocrinology
- Invite onto the Fertility Preservation Research Audit (HREC 33064)

**Endocrinology**
- Assess suitability for fertility preservation.
- Complete ‘Fertility Preservation Form’ section B
- Document in EMR that pre-consent form provided

**Fertility preservation:** YES
- Oncology: Coordinate lists with all teams
- Oncofertility coordinator: Review ethics checklist, complete Clinical Ethics referral if required, EPS database forms, IVF forms, coordinate scientists
- Surgeons: Surgical consent, coordinate theatre
- IVF Centre: Collect tissue, and send family report
- Research Team: Enter into Fertility Preservation Database
- Coordinator: Review IVF report, histology with referring team and family, provide summary of care letter

Table 1. Comparison of biological male fertility compromise guidelines

<table>
<thead>
<tr>
<th>Risk</th>
<th>Stern et al. 201313</th>
<th>COSA guidelines</th>
<th>Green et al. 201414</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>HL—ABVD, OPEA, NOVP, CHOP, COP</td>
<td>HL, NHL—Lower dose alkylating chemotherapy: ABVD (8% risk), OPEA, NOVP, CHOP, COP</td>
<td>Sperrmannogenous less likely when ced &gt;4000mg/m²</td>
</tr>
<tr>
<td></td>
<td>NHL—COP/COPDAM/CYM (± R) RCHOP 2.3</td>
<td>Testicular radiation &lt;0.7Gy RCHOP</td>
<td>No cumulative dose below which azoospermia didn’t occur and above which azoospermia did occur</td>
</tr>
<tr>
<td></td>
<td>ALL—AALL0331 2.0</td>
<td>Temporary azoospermia post treatment</td>
<td></td>
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<tr>
<td>Medium</td>
<td>NHL—COP/COPDAM/CYVE (± R) 4.8</td>
<td>Wilms, NB—Testicular radiation dose 1–6 Gy Ga a result of scatter from abdominal/pelvic radiation</td>
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<tr>
<td></td>
<td>NHL—Abd/pelvic radiation (1–6Gy) with testicular radiation dose= scatter</td>
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<td></td>
<td>GCT—BEP (2/4) 200.0/400.0</td>
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<td></td>
<td>ALL—AALL0331 4.0</td>
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<td>High</td>
<td>Prolonged azoospermia</td>
<td>HSCT—TBI</td>
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<tr>
<td></td>
<td>HL—BEACOPP.7.5,exBEACOPP, CHVP/VPA, EVA, COPP/ABV, COPP/ABV (4/6), COPP/ABV, OPEA/COPP (4)</td>
<td>GC, ALL, NHL sarcoma—≥6Gy radiation to testes</td>
<td></td>
</tr>
<tr>
<td></td>
<td>NHL—HyperCVAD (8)</td>
<td>HL—Protocols containing procarbazine: COPP, MOPP (83% risk), MVPP (97% risk), CHVP, CHVP/EVA, COPP/ABV, COPP/ABV (62% risk) BMT/SCT—Alkylating chemotherapy for transplantation conditioning (cyclophosphamide, busulfan, melphalan) (70% risk)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>NHL testicular radiation &gt;2.5Gy men and &gt;6Gy boys</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>BMT—HSCT containing TBI/ alkylator, cyclo/ busulfan/melphalan</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>BT—SIMB96 (96-03) 16.0, 300</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cranial radiation &gt;40Gy</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ALL—craniospinal radiation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>High</td>
<td>OS—MAP, MAPE 240</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ES—EuroEwings 99 IDE (6)/V19 (8) 102.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>AEWS0331 interval VDC/E/VC 8.4 63.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>RMS/IR 13 V19 23.4, IR 14 V16 26.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>D9803 VAC 30.8, ARIT 03(1) V1 6.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ARST 0331 VAC/V1 8.4, ARST 0431 VDC/E/VC 9.6, 45.0</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Testicular cancer, BMT/SCT, ALL, NHL, sarcoma, NB, HL: Any alkylating agent (eg, procarbazine, nitrogen mustard, cyclophosphamide) + TBI (80-90% risk), pelvic/testicular radiation, HL—BEACOPP (67–80% risk)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sarcoma, NHL, NR</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>ALL—Cyclophosphamide &gt; 7.5 g/m²</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>BT—Cranial/brain radiation ≥ 40 Gy</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Infertility risk and potential recommendations in biological males

<table>
<thead>
<tr>
<th>Age</th>
<th>Risk category</th>
<th>Potential to recommend FP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-pubertal</td>
<td>Low</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>Mod &gt;20% risk</td>
<td>May consider experimental options</td>
</tr>
<tr>
<td></td>
<td>High &gt;80% risk</td>
<td>May consider experimental options</td>
</tr>
<tr>
<td></td>
<td>Uncertain</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>Contraindication ALL LBL</td>
<td>No</td>
</tr>
<tr>
<td>Pubertal</td>
<td>Low</td>
<td>Preserve sperm if able</td>
</tr>
<tr>
<td></td>
<td>Mod</td>
<td>Yes. Consider testicular tissue cryopreservation if unable to produce sample</td>
</tr>
<tr>
<td></td>
<td>High</td>
<td>Yes. Consider testicular tissue cryopreservation if unable to produce sample</td>
</tr>
<tr>
<td></td>
<td>Uncertain</td>
<td>Discuss or consider possible biopsy. Preserve sperm if able</td>
</tr>
<tr>
<td></td>
<td>ALL LBL</td>
<td>Preserve sperm if able</td>
</tr>
</tbody>
</table>

Table 3. Fertility preservation procedure in biological males

**Established methods**

<table>
<thead>
<tr>
<th>Method</th>
<th>Age</th>
<th>Description</th>
<th>Time</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Efficacy</th>
<th>Approximate cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Freezing ejaculated sperm sample</td>
<td>During/after puberty</td>
<td>Sample via masturbation</td>
<td>May need multiple collections</td>
<td>Proven, does not require a partner</td>
<td>Depends on developmental maturity</td>
<td>High pregnancy rate</td>
<td>Needs ICSI with IVF</td>
</tr>
<tr>
<td>Freezing sperm extracted from testis surgically</td>
<td>After puberty</td>
<td>Surgical procedure under anaesthetic if unable to self-collect semen</td>
<td>Time to arrange procedure</td>
<td>Proven, does not require a partner</td>
<td>Anaesthetic only a few sperm</td>
<td>High pregnancy rate</td>
<td>Needs special technologies ICST with IVF</td>
</tr>
<tr>
<td>Donor sperm/Adoption</td>
<td>N/A</td>
<td>Monitor sperm count after treatment</td>
<td>N/A</td>
<td>No intervention to patient</td>
<td>Not biologically fathering a child</td>
<td>Successful methods</td>
<td></td>
</tr>
</tbody>
</table>

**Experimental methods**

| Testicular tissue freezing | Any age | Small sample of testicular tissue surgically removed | Time to arrange procedure | Only option for pre-pubertal boys | Anaesthetic risk of malignant resEDURE | Highly experimental in pre-pubertal boys | Sperm seen in post-pubertal boys? | Free of charge in public hospitals | Future costs uncertain |
3. RCH fertility preservation principles for biological females

3.1 Purpose
To ensure that consistent discussion of infertility risk occurs with all cancer patients and their families, including the discussion of relevant preservation options when suitable.

3.2 Principles
The RCH Children’s Cancer Centre, RCH Paediatric and Adolescent Gynaecology and Reproductive Biology Units will work collaboratively to:
1. Provide education and consultation to all female patients and/or families about potential fertility impairment as a result of treatment in a clear and consistent manner with use of appropriate resource tools.
2. Discuss and consider the following potential fertility preservation (FP) options with newly diagnosed female patients/families where the patient is to receive any chemotherapy, radiation or surgical procedure that could impair fertility (Table 7):
   a. Pubertal females (>12y and ≥ Tanner 3):
      i. Hormone stimulation and oocyte cryopreservation, requires maturity and takes approximately two weeks.
      ii. Ovarian tissue cryopreservation (limited outcome data).
      iii. GnRH agonist (limited outcome data and considered more an adjunct rather than an alternative to other measures).
      iv. Oophoropexy (limited outcome data).
      v. Ovum donation.
   b. Pre-pubertal females:
      i. Ovarian tissue cryopreservation (limited outcome data).
      ii. Oophoropexy (limited outcome data).
      iii. Ovum donation.
3. Discuss FP recommendations based on risk of infertility versus risk of intervention, but all situations are to be considered individually and decisions made by patient/family in consultation with treating team.
4. Involve Clinical Ethics as appropriate.
5. Provide age appropriate discussions with patients when fertility preservation procedure is being considered.
6. Document fertility discussions and use of resources in medical record.
7. Discuss participation in HREC 33064 prospective fertility audit and database (DB 044) for all patients.
8. Provide feedback of results, and appropriate follow up over the treatment journey and during survivorship care.

3.3 Fertility preservation team to include:
1. Oncofertility coordinator.
2. Paediatric Gynaecology (PAG) fellow.
3. PAG consultant.
4. Reproductive Biology Unit liaison.
5. Oncology liaison.
6. Clinical Ethics.
7. Lines coordinator.
8. Surgical team as required.

3.4 Eligibility for fertility discussion
Female patients are to be identified by the Children’s Cancer Centre or treating team staff. The eligible population includes all new female patients with a proven cancer diagnosis with intent to cure.
FP team will also provide consultation at other times in response to:
1. Females with relapsed disease and a new gonadotoxic treatment plan.
2. Referrals from any provider.
3. At request of patients or families.
4. Females who have completed treatment and are having ovarian reserve surveillance.

3.5 Initial discussion by appropriate oncology team clinician
The treating oncologist or other team member has an initial discussion with all eligible patients. This includes:
1. Assessment and discussion of risk of infertility attributable to planned chemotherapy, radiation or surgical procedure even if the risk is zero.
2. If there is intent to cure, Section A of Female Fertility Preservation Form can be completed electronically through EPIC (in notes search ‘fertility’).
3. Written resources may be distributed to the patient and her family, providing an overview of fertility preservation.
   a. Having Children After Cancer Young Women (RCH Handout for teens developed by psych-oncology group).
4. Discussion of possible fertility preservation options as appropriate. The table outlining female fertility preservation procedures may be used to guide discussions.
5. FP consultation can be formally documented in EMR using a template or just use your own notes.
6. Invite the patient to participate in the Fertility Preservation Research Audit. Patients can be invited whether they decide to pursue fertility preservation or not. If the patient is under 18, the parent may consent on their behalf, however if the patient is 18 or above, they may complete their own consent form.
   a. Parent Information and Consent Form if child is <18 years or
   b. Patient Information and Consent Form if young person is ≥18 years.
3.6 Potential indications for FP referrals to Gynaecology:
1. For all pubertal females at any risk of infertility.
2. Pre-pubertal females at moderate to high risk of infertility.
3. At the request of patient or family (investigational procedures are generally not recommended for those at low risk of infertility).
4. At the discretion of the oncology team.

3.7 Procedures for fertility preservation referral biological females
Make a referral to the Paediatric Gynaecology Service at The Royal Children’s Hospital using the Female Oncofertility Referral Form in the EMR. Also please call the oncofertility coordinator and speak directly to the gynaecology fellow or gynaecologist.

3.8 Handover from Oncology or referring team to Gynaecology team:
1. The referring clinician needs to relay details relevant to making an FP decision including:
   a. Brief history.
   b. Developmental maturity of patient.
   c. Planned treatment, urgency (with clear indication of acceptable timeframe), prognosis.
   d. Handover of patient/family’s understanding of the diagnosis, prognosis, FP options/response to introductory discussion.
   e. Other procedures planned.
   f. Requirement for menstrual suppression.
   g. Other factors pertinent to cancer/reproductive potential (e.g. prognosis, likelihood of RT, likelihood of relapse).
   h. Factors that may specifically increase surgical risk such as mediastinal mass, battle scarred abdomen, significant immunosuppression, bleeding disorder, presence of a malignant abdominal/retroperitoneal mass.
   i. Social concerns within the family such as custody issues and parental disagreement.
   j. The level of complexity of the young patient’s situation that would necessitate gynaecology consultant review.
   k. Any specific concerns the treating oncologist has with respect to proceeding with an FP procedure.
   l. If the referring clinician feels the patient/family should not yet be approached, they will indicate how and when it is best to see the patient.
2. The oncofertility coordinator will see the family to provide preliminary information and provide an overview of potential fertility care, prior to gynaecology consultation when appropriate.

3.9 Paediatric adolescent gynaecology FP consultation:
1. PAG FP team fellow or consultant will be available for consultation on new patients as soon as possible to further discuss the impact of cancer therapy on the patient’s fertility. Consultations may occur beyond the 24-hour mark if the oncology team indicate that chemotherapy/radiation treatment is less urgent.
2. Those patients who are inpatients, will have this discussion on the ward in an appropriate private space such as a single patient or interview room. Those who are outpatients may have this discussion in the oncology or gynaecology clinic as a drop-in.
3. The gynaecology fellow in association with the oncofertility coordinator and gynaecology consultant will undertake a careful medical/surgical/social risk evaluation. The clinician will research and relay to the family the expected effect of cancer treatment on fertility (via published peer reviewed articles, reputed fertility risk calculators, and discussion with oncology and reproductive teams where appropriate). They will discuss potentially beneficial FP options with the family. Where the Oncology team has not been able to do so, the FP guidance can be utilized. Be aware of considerations for oophorectomy (see appendix).
4. Pre-op bloods may be considered in those who may have already been exposed to gonadotoxic treatment or have a predisposing condition that may impair ovarian function, which could influence the decision to proceed to FP.
5. The consultant and/or fellow can make their own notes in the EMR or use the special ‘fertility’ templates which assist with fertility decision-making.
6. FP gynaecology team can offer potentially beneficial FP options to:
   a. All referred pubertal females (>12y ≥ Tanner 3) and their families when patient is planned to receive any chemotherapy, radiation or surgery that could possibly impair fertility where it is felt that the FP is beneficial to the patient and the risk is acceptable.
   b. Referred pre-pubertal female patients in whom the risk of infertility is moderate or high according to treatment regimen/cumulative drug doses where it is felt that the FP is beneficial to the patient and the risk is acceptable.
   c. Other referred patients where it is felt that the FP is beneficial to the patient and the risk is acceptable.
7. The PAG FP team are to provide age appropriate education and logistical information to the young patient on fertility preservation options. The team is also to address any issues around menstrual management and sexual health.
8. The PAG FP team including oncofertility coordinator will arrange any further consultations if required.
9. PAG FP team will communicate back to the oncology clinician and reproductive team the plan, particularly any impact on the start of treatment.

3.10 Clinical ethics:
1. Clinical ethics must be carefully considered according to the clinical ethics checklist.
2. If a clinical ethics opinion is needed, Clinical Ethics Service, complete Referral Form 3A for pre-pubertal patients and Form 3B for post-pubertal patients. This should be regarded as an important medico-legal document.
3. Expected invitees: a representative from Oncology, Gynaecology, and special experts where appropriate e.g. Haematology/Genetics.
3.11 Fertility preservation procedure to proceed:
1. If OTCP is to proceed, please document in the notes that parent has read and understood RCH information sheet (Ovarian Tissue Pre-Consent Form as suggested by RCH Legal). This form goes through the experimental nature of the procedure. Please provide this to families to keep.
2. Offer storage at any IVF centre of the family’s choice. Both Monash and MIVF documents are on the RCH website.
3. Many families choose RVH as storage which is free until 21 years of age at this time. If so, please complete the MIVF Consent to Ovarian Tissue Storage for Minors.
4. FP gynaecology or surgery team will obtain surgical consent for the fertility preservation procedure which should clearly state its experimental nature.
   b. Consent from parent and assent from patient of sufficient maturity to understand concepts <18.
5. Oncology coordinator will liaise with family, Gynaecology team, lines coordinator, Clinical Ethics and reproductive laboratories.
6. Lines coordinator will liaise with surgeons, theatre and anaesthetists, and endeavour to collect tissue in the morning as each biopsy takes several hours to process by the scientists. There is no reproductive laboratory service after 3 p.m. on weekends or holidays.
7. Gynaecology and oncology consultant to be notified if there are any delays to expected commencement of cancer treatment due to the FP procedure itself and make decision regarding risk-benefit.
8. The oncofertility coordinator will assist with review of histology (for example to ensure no malignancy on the tissue) and discuss reproductive findings with family postoperatively, however it is the responsibility of the referring clinician to ensure this has been completed.

3.12 Follow-up:
1. The oncofertility coordinator will provide a summary of fertility care letter to the family and copy to referring oncologist and local doctor.
2. It is recommended that families have a follow-up discussion after acute treatment with Gynaecology whether they had FP or not. This allows questions to be answered regarding storage, technology, and monitoring of pubertal development, reproductive, sexual and bone health.
3. Timing of follow-up by gynaecology is at the discretion of oncologist (around 12 months) or from around nine years of age.
4. Transition to an adult facility is recommended when appropriate.
### Table 4. Comparison of fertility compromise guidelines biological females

<table>
<thead>
<tr>
<th>Risk</th>
<th>Stern et al. 2013&lt;sup&gt;15&lt;/sup&gt;</th>
<th>Jadoul et al. 2012&lt;sup&gt;16&lt;/sup&gt;</th>
<th>COSA (likelihood of POI)</th>
<th>COSA guidelines (development amenorrhea)</th>
<th>Wallace et al. 2005&lt;sup&gt;16&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>HL — ABVD, DEPA, NOVP, CHOP, COP NH: — COPADCM/CHYM (=&lt;text&gt;3.3&lt;/text&gt;) ALL — AALL0331 &lt;text&gt;2.0&lt;/text&gt;</td>
<td>HL — ABVD, younger patients NH: ALL AML</td>
<td>Sarcoma — (&lt;text&gt;10–40%&lt;/text&gt;) HL — US1% unless intensive (&gt;20-40%) Leukaemias — (&lt;text&gt;30%&lt;/text&gt;)</td>
<td>HL + NH: — ABVD (10%), CHOP(20), COP (in women, 30–35) AML — anthraclycine, cytarabine ALL = multigent therapies V low risk (negligible)</td>
<td>(&lt;text&gt;&lt;20% risk of subfertility&lt;/text&gt;) ALL WT Soft tissue sarcoma stage 1 GCT with no RTx Retinoblastoma BT — cranial radiation &gt;34Gy</td>
</tr>
<tr>
<td>Medium</td>
<td>NHL — COP/ COPADDM/CYVE (&gt;=text&gt;4.8) GCT — BEP (3,4) 200.0/400.0</td>
<td>HL — 40–60% if escalated therapy NH: — 10–40%</td>
<td>HL — BEACOPP Wilms, NB = whole abdo/pelvic RTx 10–15Gy pre-pubertal Spinal tumour, BT, NB, relapsed AML, NHL = whole abdo/ pelvic RTx 5-10Gy post-pubertal girls, spinal ≥25Gy</td>
<td>AML — difficult to quantify Hepatoblastoma ES = nonmet STS — stage II or III NB NHL HD — alternating rx BT — craniospinal radiation therapy, cranial radiation &gt;34Gy</td>
<td>Table 4 continued next page</td>
</tr>
</tbody>
</table>

**Risk**

- **Low**
  - HL — ABVD, DEPA, NOVP, CHOP, COP
  - NHL — COPADCM/CHYM
  - ALL — AALL0331

- **Medium**
  - NHL — COP/ COPADDM/CYVE
  - GCT — BEP (3,4)

**COSA (likelihood of POI)**

- HL — ABVD (10%), CHOP(20)
- COP (in women, 30–35)
- AML — anthraclycine, cytarabine
- ALL — multigent therapies
- V low risk (negligible)

**COSA guidelines (development amenorrhea)**

- HL + NH: — ABVD (10%), CHOP(20), COP (in women, 30–35)
- AML — anthraclycine, cytarabine
- ALL = multigent therapies
- V low risk (negligible)
- Leukaemias — (<text>30%</text>)
- Sarcoma — (<text>10–40%</text>)
- HL — US1% unless intensive (>20-40%)

**Wallace et al. 2005**

- (<text>&lt;20% risk of subfertility</text>)
- ALL
- WT
- Soft tissue sarcoma stage 1
- GCT with no RTx
- Retinoblastoma
- BT — cranial radiation >34Gy

---


### Table 5. Infertility risk and potential recommendations biological females

<table>
<thead>
<tr>
<th>Age</th>
<th>Risk category</th>
<th>Pre-treatment FP recommendation</th>
<th>Contraindication ALL LBL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-pubertal</td>
<td>Low</td>
<td>No</td>
<td>No, unless high risk</td>
</tr>
<tr>
<td></td>
<td>Mod ≥20 risk</td>
<td>Yes, for consideration</td>
<td></td>
</tr>
<tr>
<td></td>
<td>High ≥80% risk</td>
<td>Yes, for consideration</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Uncertain</td>
<td>No</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pubertal</td>
<td>Low</td>
<td>No, but up to patient/family</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Mod</td>
<td>Yes. Consider oocyte harvest if time Consider experimental options</td>
<td></td>
</tr>
<tr>
<td></td>
<td>High</td>
<td>Yes. Consider oocyte harvest if time Consider experimental options</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Uncertain</td>
<td>Up to patient if &gt;16</td>
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</table>

### Table 6. Impact of class of chemotherapeutic agent on ovarian function

<table>
<thead>
<tr>
<th>Agent class</th>
<th>Risk category</th>
<th>Likelihood of livebirth17</th>
<th>Mechanism</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alkylating agents18,19,20,21</td>
<td>High</td>
<td>HR 0.9917</td>
<td>DNA inter-strand cross-linking drugs</td>
</tr>
<tr>
<td>Cyclophosphamide ≥7.4 g/m²</td>
<td>Pre-pubertal patients considered high risk ≥2 g/m² and moderate risk ≥1 g/m²</td>
<td>HR 0.8417</td>
<td></td>
</tr>
<tr>
<td>Ifosfamide ≥2.7 mg/m²</td>
<td>HR 0.8417</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Busulfan &lt;450 mg/m²</td>
<td>HR 0.2017</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Busulfan ≥450 mg/m²</td>
<td>HR 0.1817</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chlorambucil</td>
<td>HR 1.0 [0.85–1.16]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Melphalan</td>
<td>HR 1.0 [0.85–1.16]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nitrogen mustard</td>
<td>HR 1.0 [0.85–1.16]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thiotepa</td>
<td>HR 1.0 [0.85–1.16]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Triazines22</td>
<td>High</td>
<td>HR 0.8717</td>
<td>DNA inter-strand cross-linking</td>
</tr>
<tr>
<td>Procarbazine ≤3.4 mg/m²</td>
<td>HR 0.7817</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Procarbazine ≥5.1 mg/m²</td>
<td>HR 0.7817</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Docetaxel</td>
<td>HR 1.0 [0.85–1.16]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carboplatin</td>
<td>HR 1.0 [0.85–1.16]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Antimetabolites18 (methotrexate, 5 fluorouracil, cytarabine)</td>
<td>Low</td>
<td>HR 0.6017</td>
<td>No DNA damage in human follicles, not gonadotoxic</td>
</tr>
<tr>
<td>Vinca alkaloids18 (vincristine, vinblastine)</td>
<td>Low</td>
<td>HR 0.6017</td>
<td>No DNA damage in human follicles, not gonadotoxic</td>
</tr>
<tr>
<td>Anthracycin antibiotic18 (doxorubicin, Adriamycin)</td>
<td>Low (apart from adriamycin: intermediate)</td>
<td>HR 0.86</td>
<td>Inhibit DNA synthesis, create DNA breaks</td>
</tr>
</tbody>
</table>

---


<table>
<thead>
<tr>
<th>Method</th>
<th>Pubertal status</th>
<th>Description</th>
<th>Time needed</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Efficacy</th>
<th>Approximate cost</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Oocyte (egg) freezing</strong></td>
<td>After puberty</td>
<td>Stimulation of ovaries with daily hormone injections and surgical collection of mature eggs under anaesthetic</td>
<td>a2 weeks</td>
<td>Proven Does not require a partner</td>
<td>Daily hormone injections&lt;br&gt;Delay in start of treatment&lt;br&gt;Requires emotional and physical maturity&lt;br&gt;Poor yield ≤17 years of age</td>
<td>49% clinical pregnancy rate for women ≤34 years of age&lt;br&gt;70% probability of live birth if at least 10 oocytes collected ≥ ≥2000 births-worldwide</td>
<td>Yeasty oocyte storage fee&lt;br&gt;Future assisted reproduction (IVF) costs vary depending on clinic and private health insurance program</td>
</tr>
<tr>
<td><strong>Embryo freezing</strong></td>
<td>After puberty</td>
<td>Stimulation of ovaries with daily hormone injections and surgical collection of mature eggs under anaesthetic, mixed with sperm in the lab to create embryos</td>
<td>≥2 weeks</td>
<td>Proven</td>
<td>Daily hormone injections&lt;br&gt;Delay in start of treatment&lt;br&gt;Requires emotional maturity&lt;br&gt;Requires sperm</td>
<td>Pregnancy rates 30–37% per transfer&lt;br&gt;Most established method</td>
<td>Yeasty embryo storage fee&lt;br&gt;Future assisted reproduction (IVF) costs vary depending on clinic and private health insurance plan</td>
</tr>
<tr>
<td><strong>Donor oocytes/embryo surrogacy adoption</strong></td>
<td>N/A</td>
<td>Surrogacy if unable to carry pregnancy&lt;br&gt;Consider adoption</td>
<td>N/A</td>
<td>No pre-treatment intervention to patient&lt;br&gt;Not biologically related to child</td>
<td>Successful methods&lt;br&gt;Vary depending on choice</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Ovarian suppression (GnRH agonist)</strong></td>
<td>After puberty</td>
<td>Hormone injections to switch off ovaries. This may decrease ovarian damage from chemotherapy</td>
<td>Immediate</td>
<td>May stop or decrease menstruation during treatment</td>
<td>Monthly to 3 monthly injection&lt;br&gt;No benefit with radiation&lt;br&gt;Menopausal symptoms</td>
<td>May decrease oocyte loss by 40%&lt;br&gt;Conflicting results: probably overall benefit in adults but the benefit is likely small&lt;br&gt;Not considered a replacement for other options</td>
<td>£340/ month for injection&lt;br&gt;Partially covered through some drug plans&lt;br&gt;Expenses covered via drug committee at RCH</td>
</tr>
</tbody>
</table>
4. Clinical ethics checklist for all fertility preservation procedures

This document refers to surgical procedures to retrieve reproductive tissue from a child who has cancer, for the purpose of attempting to preserve fertility by freezing the tissue for future use. This refers to ovarian tissue from a girl, and testicular tissue from a boy.

4.1 Basic ethical requirements for a child of any age:

1. Informed consent of parents: In all cases where fertility preservation procedures are contemplated, parents should be provided with comprehensive written information, including clear and accurate information about the storage of reproductive tissue, including place costs of storage, what will be done with the tissue if the child does not survive, who has the right to access the tissue and for what purposes. It is crucial that parents understand that this procedure offers only a theoretical possibility of fertility preservation, for which there is not yet any evidence of success, and is no means a guarantee that the child will have fertility in adulthood. Parents must also be aware that the offer of a fertility preservation procedure does not imply that child’s survival into adulthood is certain.

2. Assent of child (where child is old enough). The child should be given a developmentally-appropriate explanation of the procedure that will be done and its purpose. Ideally, the child should be in agreement with the procedure.

4.2 Formal clinical ethics review — when required

4.2.1 Clinical ethics review is NOT required in the following circumstances:

1. This procedure will not delay or interfere with the cancer treatment.
2. This procedure is itself of minimal risk, and will be performed under a GA which is required in any case for treatment for the cancer.
3. The treatment for cancer is being undertaken with the intent of cure or long-term survival.
4. Survival into adulthood is sufficiently probable that it is appropriate to undertake procedures aimed at promoting quality of life long-term.
5. There is risk of loss of fertility due to chemotherapy.
6. The procedure will leave one gonad intact, so that if the gonad from which tissue is taken ends up being damaged or completely removed, and the chemotherapy does not in fact cause loss of fertility, there is still one functioning gonad with good chance for natural fertility.
7. There are none of the following complicating factors involved:
   a. The treatment for cancer is not being undertaken with the intent of cure or long-term survival.
   b. The child has an intellectual disability.
   c. The child or young person objects to having a fertility preservation procedure, but parents still want to go ahead.
   d. The parents are unwilling to inform the child about the procedure, when the child is at developmental stage where they would be able to understand at least the basic idea of the procedure.

4.2.2 Clinical ethics review for a fertility preservation procedure IS REQUIRED if one or more of the following apply:

1. The risks of the FP procedure:
   a. The procedure will delay or interfere with the cancer treatment.
   b. The procedure is itself of greater than minimal risk (e.g. because of a co-morbidity which makes the procedure more risky than usual).
   c. The procedure has a significant risk of not leaving one gonad intact (e.g. if the child has only one gonad to begin with).
2. The potential benefits:
   a. The risk of loss of fertility due to chemotherapy is low.
   b. The potential for retrieving tissue that might be usable in the future is lower than usual, for any reason.
3. Other factors:
   a. The treatment for cancer is not being undertaken with the intent of cure or long-term survival.
   b. The child has an intellectual disability.
   c. The child is pre-pubertal.
   d. The child or young person objects to having a fertility preservation procedure, but parents still want to go ahead.
   e. The parents are unwilling to inform the child about the procedure, when the child is at developmental stage where they would be able to understand at least the basic idea of the procedure.
   f. Any treating clinician has an ethical question or concern about the procedure.

4.3 Clinical ethics checklist:

1. Pre-pubertal child — use Referral Form 3A (FP Pre-pubertal).
2. Post-pubertal — use checklist below. If one or more items ticked below, clinical ethics meeting will be held — use Referral Form 3B (FP Post-pubertal).
3. If no items ticked, no clinical ethics referral needed, no meeting required:
   a. The procedure will delay or interfere with the cancer treatment.
   b. The procedure is itself of greater than usual risk (e.g. because of a co-morbidity).
   c. The procedure has a significant risk of not leaving one gonad intact (e.g. if the child has only one gonad).
   d. The risk of loss of fertility due to chemotherapy is low.
   e. The potential for retrieving tissue that might be usable in the future is lower than usual, for any reason.
   f. The treatment for cancer is not being undertaken with the intent of cure or long-term survival.
   g. The child or adolescent is unlikely to be able to use any stored tissue for fertility purposes in the future, but parents still want the procedure done.
   h. The child or adolescent objects to having the fertility preservation procedure, but parents still want to go ahead.
   i. The parents are unwilling to inform the child or adolescent about the procedure (where developmentally appropriate to inform), but want the procedure done.
   j. Any treating clinician has an ethical question or concern about the procedure.

Clinical Ethics Referral Forms 3A and 3B can be found on the clinical ethics website: http://www.rch.org.au/bioethics/clinical_ethics_service/
5. RCH fertility preservation principles for transgender patients

5.1 Purpose
To ensure that consistent discussion of infertility risk occurs with all gender diverse patients and their families, including the discussion of relevant preservation options when suitable.

5.2 Principles
The Gender Service Clinic at the Royal Children’s Hospital provides assessment, support and medical treatments to around 250–300 new trans and gender diverse young people per year in Victoria. Medical gender affirmation treatment may impact the future fertility of trans and gender diverse young people by affecting maturity of the gonads. This has necessitated fertility preservation counselling prior to the initiation of gender affirming hormone treatment (an international standard of care). There is a paucity of data around the impact of treatment on actual fertility. Recent data suggests that some gonadal function can be preserved but the evidence is limited. Over 50% of adult trans patients desire fertility.34,35,36,37,38,39,40

The RCH principles are:
1. Consistent discussion of infertility risk with all trans and gender diverse patients and their families, including the discussion of relevant fertility preservation options when suitable.
2. Discussion of potential fertility preservation (FP): collection of collection and storage of ovarian or testicular material, or eggs or sperm for future use).
3. Involving Clinical Ethics as appropriate.
4. Document discussions in the medical record.
5. Discuss participation in monitoring of safety and efficacy.
6. Provide feedback of results and appropriate follow-up care.

5.3 International guidelines provide these additional principles of care:
1. Programs treat all requests for assisted reproduction without regard to gender identity status (American Society for Reproductive Medicine 2015).41
2. Trans and gender diverse young people have equitable access to reproductive counselling prior to the onset of hormone therapy (ASRM 2015; WPATH 2012; Endocrine Society Taskforce 2017; Australian Standards of Care 2018).41,42,43,44 As many will not cease pubertal suppression once it has been commenced, these discussions are advocated prior to onset of puberty blockers.
3. Programs without sufficient resources to offer care have an ethical duty to assist in referral to providers equipped to manage such patients (ASRM 2015).41

5.3.1 International guidelines:
1. Standards of Care for the Health of Transsexual, Transgender, and Gender Nonconforming People, The World Professional Association for Transgender Health (WPATH) 2012.42
2. ESHRE Task Force on Ethics and Law 23: medically assisted reproduction in singles, lesbian and gay couples and transsexual people.45
3. Australian Standards of Care and Treatment Guidelines: for Trans and Gender Diverse Children and Adolescents (The Royal Children’s Hospital) 2018.46
4. ASRM Guideline: Access to fertility services by transgender persons: an Ethics Committee opinion.41
5. Endocrine Society appointed task force (international): Endocrine Treatment of Gender-Dysphoric/Gender-Incongruent Persons: An Endocrine Society Clinical Practice Guideline.47

5.4 The fertility team can include:
1. Gender Service physician.
2. Gender Service endocrinologist.
3. Paediatric Gynaecology (PAG) fellow.
4. PAG consultant.
5. Oncofertility coordinator.
6. Reproductive Biology Unit liaison.
7. Clinical Ethics.
8. Surgical team as required.
5.5 Eligibility for fertility discussion:

1. Not all gender diverse youth seek gender affirmation treatment or identify as the opposite sex. Thus goals of medical care are highly individualised and decisions made by the multidisciplinary Gender team.

2. Transgender youth opting for Stage I (puberty suppression) should have a fertility discussion. Gonadotropin releasing hormone analogues (GnRHαs), prescribed by a paediatric or paediatric endocrinologist, suppress the development of secondary sex characteristics whilst providing time for cognitive and emotional development. As they are reversible in their effects, should an adolescent wish to stop taking them at any time, puberty will resume in the biological sex with resultant return of fertility.

3. Transgender youth opting for Stage II (gender affirming hormone treatment) should have a fertility discussion with discussion of potential fertility preservation measures that might be available. A trans female would be commenced on oestrogen and a trans male would start testosterone in order to induce secondary sex characteristics congruent with their gender identity. These are only partially reversible in their effects. The degree to which fertility is permanently impacted by testosterone or oestrogen treatments is currently unknown. No studies address whether fertility can be gained, either naturally or with exogenous gonadotropins, in transgender individuals who undergo pubertal suppression in adolescence, followed directly by gender-affirming hormone therapy.46 Age and pubertal development will determine efficacy of the FP procedures and whether they are regarded as investigational or not.

5.6 Initial discussion by gender physician and endocrinologist:

1. Consistent discussion of impact of treatment on fertility with all trans and gender diverse patients and their families starting GnRH analogues by the multidisciplinary Gender Service.

2. Further fertility counselling if young person is opting for stage II treatment. The complexities in fertility counselling and decision making in trans and gender diverse youth arise due to uncertainties regarding the impact of hormone treatment on gonadal function, the limited efficacy of some fertility preservation technologies, (with tissue preservation procedures being investigational), the factors influencing decision-making (such as dysphoria) and the developmental stage of the young person. While young people may not see parenting as important, it is described as an important life goal in adulthood.

3. Discussions can include the impact of hormone treatment on fertility, the uncertainties and gaps in knowledge around some FP technologies (including in vitro maturation versus grafting), the potential use of donor gametes, and surrogacy, the physical and psychological impact of discontinuing hormone therapy,46 how to use reproductive tissue in the future and the impact of fertility treatments (such as transvaginal ultrasound, pelvic examinations, hormonal side effects such as breast tenderness); future disclosure to offspring, psychological assistance, ART laws, alternate forms of parenting.46

4. Documentation in the EMR.

5. Provision of written resources which can be found at http://www.rch.org.au/fertility/health-prof/

6. Contact onc fertility coordinator for those wishing referral.

5.7 Indications for referral of birth assigned males

5.7.1 Sperm collection

Post-pubertal patients who are comfortable producing a sample can be referred directly to Andrology for sperm collection, follow sperm collection pathway).

5.7.2 Referrals for discussion of testicular tissue cryopreservation (to Gynaecology/Surgery):

1. Referrals may be appropriate for patients who do not wish to produce a sample or if patient/family request.

2. For those who have had puberty suppressed mature sperm may be absent in the tissue, and the procedure is considered investigational and approved as a special procedure under novel technologies, clinical ethics and research governance.

3. Consider factors that assist with fertility decision making:
   a. Developmental maturity of patient (Tanner stage testicular volume).
   b. Patients understanding of discussion and experimental nature of procedures.
   c. Factors that may increase surgical risk.
   d. Social concerns or discordance within the family regarding medical care.

4. Discuss with family your role, purpose of visit, explain level of risk if no action taken (oestrogen can impair spermatogenesis and affect development of gonad, but studies are small).

5. Clearly describe what FP procedures are experimental and what is standard, and pros and cons.

6. Discuss details of testicular biopsy and the procedure.

7. Clear statement that peri-pubertal TTCP is currently experimental in pre-pubertal humans — has been successful only in animals so far. It is possible to not undertake fertility preservation and await adulthood. The impact of gender-affirming hormones on fertility is unclear. The difficulty lies in the fact that many young people don’t wish to cease treatment in the future. In older adolescents (Tanner 4/5) who have not started gender-affirming hormone treatment sperm are more likely to be found and TTCP is an appropriate standard of care.

8. Freezing testicular tissue (germ cells) is free until 21 years of age.

9. The tissue may be dissected to look for mature sperm, and if found may offer real hope of future fertility. This will incur same cost as sperm storage.

10. Please provide information to families (The Royal Children’s Hospital Gender Service [RCHGS] information sheet: Testicular Biopsy and Tissue Harvest for Possible Fertility Preservation).

5.7.3 If TTCP is to proceed:

1. Please document in the notes that parent has read and understood RCHGS Information sheet. This form goes through the experimental nature of the procedure. Please provide this to families to keep.

2. FP surgical team to obtain surgical consent for FP procedure, which should clearly state its experimental nature in pre-pubertal patients, or in those who have commenced hormone treatment.

3. Consent from patient if patient ≥18, or in younger patient if Gillick competent.

4. Consent from parent and assent from patient of sufficient maturity to understand concepts and <18.

5. MIVF Testicular Tissue Cryopreservation Form and FPS Database forms to be completed.

6. TG team with assistance of onc fertility coordinator will liaise with family, IVF Andrology scientists (RWH 8345 3232 or lab.supervisors@mivf.com.au; Monash (03) 9420 8218) and clinical ethics and surgeons regarding logistics and booking.

7. There is no reproductive lab service after 3pm, on weekends or holidays.

8. The onc fertility coordinator will assist with review of histology and discussion with families, however it is the responsibility of the referring clinician to ensure this has been completed.

5.4 Involvement in clinical ethics in decision-making for TTCP:

1. Guidelines are as per the Ethics Checklist for Fertility Preservation Procedures for TG patients.
2. If a clinical ethics meeting is needed, Clinical Ethics Service, Referral Form 3A to be completed for pre-pubertal patients and Form 3B to be completed for post-pubertal patients. This should be regarded as an important medico-legal document for filing in the medical record.
3. If a clinical ethics meeting is held, the expected invitees include a representative from transgender team, endocrinology and special experts where appropriate.

5.5 Follow-up:

1. The oncofertility coordinator will provide a summary letter of fertility care to the family and copy to referring team.
2. Oncofertility team will monitor safety and efficacy in the oncofertility registry (DB044).

5.8 Indications for referral for birth assigned females to Gynaecology:

1. Patients and families about to start gender affirming hormone treatment who desire further sexual and reproductive health information. According to Medicare data, >60 men give birth per year in Australia.41,42
2. Gender affirming hormone treatment treatment with testosterone may induce a phenotype similar to polycystic ovary syndrome, however does not affect egg numbers in the ovary, and the effect is reversible on cessation. Thus there is no urgency for oocyte collection or ovarian tissue preservation.
3. Most will resume menses after 3–6 months of cessation of testosterone. Natural pregnancies have been seen while still on testosterone.43 Testosterone is teratogenic, and men should be advised to use contraception if they have a male partner.
4. Case series of oocyte collection in adolescents demonstrate low oocyte yield and in the very young and adverse psychological effects. Dysphoria can be triggered by using the wrong pronoun, using standard ‘female’ medical illustrations during consultations, pelvic examinations and transvaginal ultrasounds. Oocyte collection in adult males has been tolerated.44,45

5. Indications for referral for birth assigned males to Gynaecology:

1. Patients and families about to start gender affirming hormone treatment who desire further sexual and reproductive health information. According to Medicare data, >60 men give birth per year in Australia.41,42
2. Gender affirming hormone treatment treatment with testosterone may induce a phenotype similar to polycystic ovary syndrome, however does not affect egg numbers in the ovary, and the effect is reversible on cessation. Thus there is no urgency for oocyte collection or ovarian tissue preservation.
3. Most will resume menses after 3–6 months of cessation of testosterone. Natural pregnancies have been seen while still on testosterone.43 Testosterone is teratogenic, and men should be advised to use contraception if they have a male partner.
4. Case series of oocyte collection in adolescents demonstrate low oocyte yield and in the very young and adverse psychological effects. Dysphoria can be triggered by using the wrong pronoun, using standard ‘female’ medical illustrations during consultations, pelvic examinations and transvaginal ultrasounds. Oocyte collection in adult males has been tolerated.44,45


5. Ovarian tissue preservation: Normal ovarian histology and physiology has been seen 58 weeks after androgen treatment.44 Normal in vitro maturation of androgen exposed oocytes has been seen, which might one day also be an option for fertility in the future through ovarian tissue preservation although this is currently experimental. There are no studies on the health of offspring conceived from testosterone exposed oocytes.

6. Decisions for gonadectomy or gender affirming surgery should be deferred until adulthood.

7. Provide written information (RCHGS information sheet on impact of testosterone treatment and fertility).

8. The oncofertility coordinator will provide a summary letter of fertility care to the family and copy to referring team.

9. Oncofertility team will monitor safety and efficacy in the oncofertility registry (DB044).

Figure 3. FP pathway transgender patients

Transgender paediatric or endocrinologist

- Utilise fertility preservation pathway
- Discuss impact of fertility with all patients, both prior to puberty blockers and again prior to gender affirming hormone therapy (as applicable)
- Provide appropriate written information to family
  - RCH Gender Service information sheet Testicular Biopsy and Tissue Harvest for possible fertility preservation
  - RCH Gender Service information sheet Ovarian Tissue Harvest and Egg Collection for possible fertility preservation
  - MIVF forms
  - Reproductive Services Information Sheet for Transgender Patients
  - Young people with cancer

Gynaecology

- See transmales and transfemales

Fertility preservation: YES

Gender Service physician/endocrinologist:

- Completes documentation, reviews ethics checklist, refers to surgeon/lines coordinator, notifies oncofertility coordinator
- Coordinates ethics, FPS Database forms, MIVF storage consent forms, theatre booking, liaises with surgical team to get date for theatre, coordinates scientists, checks histology with referring team, completes summary of fertility care letters for FP patients, updates oncofertility database for safety and efficacy data.
- Theatre bookings/surgical team:
  - Notifies oncofertility coordinator of theatre date, surgical consent

Fertility preservation: NO

• Document fertility note
• Invite onto the Fertility Preservation Audit (HREC 33064)

Information Sheet for Transgender Patients

• Utilise fertility preservation pathway
• Discuss impact of fertility with all patients, both prior to puberty blockers and again prior to gender affirming hormone therapy (as applicable)
• Provide appropriate written information to family
  - RCH Gender Service information sheet Testicular Biopsy and Tissue Harvest for possible fertility preservation
  - RCH Gender Service information sheet Ovarian Tissue Harvest and Egg Collection for possible fertility preservation
  - MIVF forms
  - Reproductive Services Information Sheet for Transgender Patients
  - Young people with cancer
6. Clinical ethics checklist for fertility preservation procedures in transgender youth

This document refers to surgical procedures to retrieve reproductive tissue from an adolescent transgender youth for the purpose of attempting to preserve fertility by freezing the tissue for future use. This refers to ovarian tissue from a trans male, and testicular tissue from a trans female.

6.1 Basic ethical requirements

Informed consent of young person: In all cases where fertility preservation procedures are contemplated, the young person and parents should be provided with comprehensive written information, including clear and accurate information about the storage of reproductive tissue, including place and costs of storage, what will be done with the tissue if the child passes, who has the right to access the tissue and for what purposes. It is crucial that they understand that this procedure offers only a theoretical possibility of fertility preservation, for which there is not yet any evidence of success, and is by no means a guarantee that the child will have fertility in adulthood.

6.2 Formal clinical ethics review — when required

Clinical ethics review is required in the following circumstances:
1. This procedure will interfere with medical treatments.
2. This procedure is itself of greater than minimal risk (e.g. because of a co-morbidity which makes the procedure more risky than usual).
3. The procedure has a significant risk of not leaving one gonad intact (e.g. if the child has only one gonad to begin with). So that if the gonad from which tissue is taken ends up being damaged or completely removed, there is no functioning gonad with good chance for natural fertility.
4. The risk of loss of fertility due to medical treatment is low.
5. The potential for retrieving tissue that might be useable in the future is lower than usual, for any reason.
6. Consensus between young person and family about the fertility preservation procedure cannot be reached.
7. Any treating clinician has an ethical question or concern about the procedure.

6.3 Clinical ethics checklist for transgender youth

1. Post-pubertal — use checklist below If one or more items ticked below, clinical ethics meeting will be held — use Referral Form 3B (FP Post-pubertal).
2. If no items ticked, no clinical ethics referral needed, no meeting required:
   - The procedure is itself of greater than usual risk (e.g. because of a co-morbidity).
   - The procedure has a significant risk of not leaving one gonad intact (e.g. if the child has only one gonad).
   - The risk of loss of fertility due to treatment is low.
   - The potential for retrieving tissue that might be useable in the future is lower than usual, for any reason.
   - The adolescent is unlikely to be able to use any stored tissue for fertility purposes in the future, but parents still want the procedure done.
   - The adolescent objects to having the fertility preservation procedure, but parents still want to go ahead.
   - The parents are unwilling to inform the adolescent about the procedure (where developmentally appropriate to inform), but want the procedure done.
   - Any treating clinician has an ethical question or concern about the procedure.

Clinical Ethics Referral Forms 3A and 3B can be found on the clinical ethics website: http://www.rch.org.au/bioethics/clinical_ethics_service/.
7. Deceased patients

7.1 Purpose
To ensure that the RCH can support families around decision making with respect to the stored tissue, and assist compliance with state legislation around tissue storage.

7.2 Principles
Tissue from minors is required to be discarded by law for deceased patients and cannot be used by others for fertility purposes nor for research.\textsuperscript{55,56} We have developed a pathway for notification to the Women’s laboratory and a stepped care protocol for discussion with families who may opt to cremate or bury the tissue if they desire. Ultimately this pathway is the responsibility of the oncology consultant.

Principles for data sharing with the Reproductive Services laboratory:
1. When patients store tissue, they become a patient of the RCH and the adult institution.
2. Families have to let the IVF centres know about the passing of their child but this does not always occur.
3. Eventually we will have a shared EMR so clinicians from all sites can access information as required for seamless integration of care.
4. Until then we will provide this information to the adult storage centre for those who have signed consent for data linkage.
5. We have also modified the IVF forms to let families know that we may notify the laboratories in the event of their child’s passing. For those who sign this new form we will share this data.
6. Families have to sign a consent to disposal form before the tissue is discarded. This does not occur if the family is lost to follow up and the storage period of 20 years is exceeded in which case the decision to discard the tissue is made at an executive level at The Royal Women’s Hospital.
7. The Women’s laboratory will notify the RCH prior to disposal of any tissue of RCH patients who are lost to follow up.

8. Monash Children’s

8.1 Purpose
To provide support to Victorian children and families having treatment under the statewide Pediatric Integrated Cancer Service attending Monash Children’s Hospital.

8.2 Principles:
1. The Pediatric Integrated Cancer Service is a statewide service endorsing fertility care for Victorian children in statewide guidance.57,58
2. Monash Children’s offers FP onsite to children aged 13 years and above.
3. Until such time as Monash Children’s establishes laboratory and clinical ethics governance for pre-pubertal children, The RCH team have provided support, by sharing of fertility protocols and guidance, advice to clinicians, and patient care.
4. In the past three years clinicians have referred five families for onsite fertility consultation and care at the RCH. In 2019, five referrals were received of whom one proceeded to FP. These numbers are expected to increase with a projection of 10 referrals per year to the RCH for consultation.
5. In the event that a Monash clinician requests FP consultation and care:
   i. They will complete an RCH Fertility Referral Form, provide timelines for start date of cancer treatment and speak with Dr Leanne Super, Monash Liaison.
   ii. The oncofertility coordinator and the relevant RCH team (Gynaecology or Endocrinology or Surgery) will be notified who will seek advice from executive.
   iii. RCH Executive will endeavor to provide an answer within 24 hours so as not to delay the onset of cancer treatment, or to allow time for review at another centre if feasible e.g. with IVF specialists at the Women’s.


Appendix

Attachments: revised or new documents
A. Information sheet on TTCP for RCH patients
B. Information sheet on OTCP for RCH patients
C. RCH Gender Service information sheet for patients having TTCP
D. RCH Gender Service information sheet for trans masculine young people, trans and gender diverse or non-binary people with ovaries
E. Summary of care of letters
About the procedure

During the procedure a small part of one testicle is removed and frozen. It is important to understand that there is no guarantee that the banking of testicular tissue will lead to successful sperm production.

The surgery is performed under a general anaesthetic, preferably at the same time that another procedure is required. After removing part of a testicle, the scrotum may be painful for a few days. The tissue is collected by IVF scientists and stored at IVF laboratories. Currently the scientists from the Reproductive Services Department at The Royal Women’s Hospital (RWH) are able to collect the tissue at surgery and process it at their centre. You do not need to have tissue stored at this centre if you don’t want to, in which case we will do everything possible to arrange storage at another IVF centre of your choice and to inform you of any costs involved. If your child’s tissue is stored at RWH, it does not mean that he will be required to have future fertility care at that centre (Melbourne IVF, RWH). He may go wherever he chooses. Currently there is no storage fee for the tissue up until your child turns 21. However, a storage charge may be introduced by Melbourne IVF at any point. If the tissue also contains mature sperm an annual storage fee may be requested. Doctors at The Royal Children’s Hospital have nothing to do with the tissue storage after it is collected by the scientists, and all future dealings regarding the stored tissue would be between yourself and the IVF centre.

Who is eligible

Theoretically there is no lower age limit for testicular tissue biopsy. We may advise against the procedure in certain situations such as cancers where there may be a risk of reintroducing the cancer back into the body from transplantation in the future. For example, in leukemia if tissue is collected it is currently deemed too dangerous to put back into the body and cannot be currently used for fertility purposes.

We also need to determine that your child is well enough for surgery and to ensure that the procedure does not delay cancer treatment. Bleeding disorders or serious immune deficiency may prevent your child from having the procedure done. We also take into account differing views within a family about such procedures. Sometimes we consult with the Clinical Ethics Committee to assist the decision making for the procedure in the event that the decision is not so clear cut. This is undertaken urgently so as not to delay treatment.

Risks and benefits

It remains experimental to collect testicular tissue from boys to be stored frozen in the hope of future fertility. The surgery for the biopsy itself is not experimental as this procedure is performed routinely by surgeons for other indications.

Fertility Preservation is not necessarily offered only if your child has an excellent chance of survival. It can still be offered when the oncology doctors are hoping for cure, regardless of the chance of survival.

Expected risks of the surgical procedure:

1. Risk of a general anaesthetic. There may be situations where your child’s medical situation may present specific increases in the risk of anaesthesia. In these situations you the anaesthetist and the other treating teams will need to discuss the risk versus benefit issues. The safety of your child in the short term is a very important factor. Mostly the risk relating to the anaesthesia for the extra fertility procedure for your child will be so small it would be difficult to estimate. The anaesthetist can clarify if your child has any special aspects of their condition that could influence the risk of surgery.

2. Risk of surgery: infection, bleeding and damage to internal structures. In the rare instance that there is a complication such a haematoma (collection of blood) a second surgery to manage a complication might be required. These risks are likely to be higher during cancer therapy.

3. Delay to treatment: the procedure will usually be timed with other operative procedures necessary for treatment of the disease. As far as possible the surgery is performed within a few days of diagnosis, and if so should not impact prognosis. We will try and find the earliest date to undertake the procedure, however we will defer to your treating doctor (e.g. oncologist) if he or she deems that it is too unsafe to wait for this date. In this event the fertility procedure will need to be cancelled, as safety is our priority.
4. The biopsied testis may not develop fully as a result of the biopsy so may remain smaller than the non-biopsied
testis. This is additional to the damage that chemotherapy or radiation treatment produce to reduce the size of
the testes by damaging sperm production.

Other options:
1. Delay intervention until fertility is required. Sperm may still develop in the boy as an adult. But if not, it is
possible to have a biopsy or similar procedure as an adult to collect or mature sperm. In some boys, although the
chemo- or radiotherapy may be damaging to the testis, sperm production may still develop in adulthood. Even if
this is at lower levels than normal, there may be sufficient sperm produced to allow for fatherhood either naturally
or with IVF assistance. The degree of testicular damage depends on the extent of cancer treatment, which is
usually not known for any boy before treatment starts.
2. Sperm donation in future.
3. Adoption.

Other issues to consider:
1. Cost of procedure: this is currently free but it may change in future.
2. Storage of tissue: occurs for at least 20 years after which time you have to renew the request. This is an
arrangement between you and the IVF centre and does not involve The Royal Children’s Hospital.
3. Costs of future IVF treatment and tissue storage.
4. The tissue can only be used by your child and in the unfortunate event of death the tissue must be disposed of.
   In this instance a member of the team would contact you to discuss arrangements.
5. Due to current legal restrictions, the tissue cannot be donated for research. The tissue may never be utilized by
   anyone other than your child.
Outcomes so far

Only around 140 pregnancies have been reported worldwide using ovarian harvest technology so it is considered experimental. Two live births have been reported in women who have had their tissue stored in childhood. The ovaries of young people, especially children contain immature eggs. It is difficult to mature the eggs after they are thawed. Furthermore there are high rates of loss of eggs during the freezing and thawing process. The procedure is offered in the hope that by the time your child has achieved adulthood and wishes to have a baby, the procedures may be more successful.

We need to also determine that your child is well enough for surgery. Multiple abdominal scars, bleeding disorders or serious immune deficiency may preclude your child from having the procedure done. We also take into account the differing views within a family about such procedures. Sometimes we consult with the Clinical Ethics Committee to assist the decision making in the event that the decision is not so clear cut. This is undertaken urgently so as not to delay treatment.

Risks and benefits

The surgery (laparoscopy and ovarian tissue harvest or removal of one ovary) is not experimental as this procedure is performed routinely by gynaecologists and surgeons for other indications. However, the use of immature ovarian tissue to attempt pregnancy in the future is considered experimental.

Fertility Preservation is offered where there is a hope of cure, irrespective of the chance of survival. Theoretically there is no lower age limit for ovarian tissue harvest. We may advise against the procedure in certain situations such as certain cancers where there may be a risk of reintroducing the cancer back into the body in the future. For example, in leukemia if tissue is collected it is deemed too dangerous to put back into the body at this stage. Technology may advance to allow the tissue to be matured outside the body so that mature eggs can be collected from that tissue for IVF, however the technology is in very early stages.

Technology has any special aspects of their condition that could influence the risk of surgery. The tissue cannot be donated to research or be utilized by anyone other than your child.

Who is eligible

If your child has a diagnosis of cancer, there is a risk that the tissue might contain cancer cells which could be reintroduced back into the body when the tissue is implanted.

Who is eligible

The surgery (laparoscopy and ovarian tissue harvest or removal of one ovary) is not experimental as this procedure is performed routinely by gynaecologists and surgeons for other indications. However, the use of immature ovarian tissue to attempt pregnancy in the future is considered experimental.

Fertility Preservation is offered where there is a hope of cure, irrespective of the chance of survival.

Expected risks of the surgical procedure:

1. Risk of a general anaesthetic. There may be situations where your child’s medical situation may present specific increases in the risk of anaesthesia, for example age less than one year. In these situations yourself, the anaesthetist and the other treating teams will need to discuss the risk versus benefit issues. The safety of your child in the short term is the very important factor. Mostly the risk relating to the anaesthesia for the extra fertility procedure for your child will be so small it would be difficult to estimate. The anaesthetist can clarify if your child has any special aspects of their condition that could influence the risk of surgery.

2. Risk of laparoscopy in general are: infection (around 7%), bleeding, damage to internal structures (bladder, bowel, blood vessels, 1-3/1000) which may occasionally require performing an open operation. These risks are likely to be higher during cancer therapy.

3. Risk of changing from keyhole surgery to a larger incision (laparotomy) 0.5-2%.

4. Death 4/100 000.

5. In the event of young age: The ovaries will usually be very small, it is highly possible that one entire ovary may need to be removed. We are not sure if the removed ovarian tissue or the remaining ovarian tissue will be functional in the future.

6. Delay to treatment: the procedure will usually be timed with other operative procedures necessary for treatment of the disease. As far as possible the surgery is performed within a few days of diagnosis, and if so should not impact prognosis. We will try and find the earliest date to undertake the procedure, however we will defer to your treating doctor (e.g. oncologist) if he or she deems that it is too unsafe to wait for this date. In this event the fertility procedure will need to be cancelled, as safety is our priority.

Other options:

1. Monitoring of ovarian function when your child is older occurs regardless of whether they have fertility preservation now or not, with a view to having egg freezing. However, if there is a high chance of ovarian failure we may not have the opportunity to undertake this.

2. Oocyte retrieval now if age appropriate and time permitting.

3. Egg donation from mother, sibling or other donor in the future.

4. Adoption.

5. Use of a hormone which suppresses ovarian function (Zoladex) and may protect the ovary, however studies on this are very conflicting in terms of success. This is only offered to girls who have already gone through puberty.

Other issues to consider:

1. Cost of procedure: this is currently free but it may change.

2. Storage of tissue: occurs for 20 years after which time you have to renew the request. This is an arrangement between you and the IVF centre and does not involve The Royal Children’s Hospital.

3. Costs of future IVF treatment and tissue storage costs.

4. The tissue can only be used by your child and in the unfortunate event of death the tissue must be disposed of. In this instance a member of the team will contact you to discuss arrangements.

5. The tissue cannot be donated to research or be utilized by anyone other than your child.
When might testicular biopsy for future fertility preservation be appropriate?

Testicular biopsy to store immature testicular tissue for future fertility preservation is experimental, and is only undertaken at the RCH under special approvals. The only situations at The RCH where we may find it acceptable to store germ cells from immature testes in a child, is if the surgical procedure is going to be very low surgical risk, and if the medical therapy proposed has a significant risk of infertility. Most young TGD people who are otherwise well and who are about to commence oestrogen as gender affirming hormone therapy will meet these criteria, which is why we discuss this option with you. In addition, a biopsy may also be appropriate for TGD adolescents who are in the mid-later stages of puberty when they are about to start puberty blockers, particularly if they can’t provide a semen sample and oestrogen therapy is envisaged in the future. Your doctor will discuss the best timing for you to consider a biopsy if this is something you would like to do.

About the procedure

During the surgical procedure a part of one testicle is removed and frozen. This is done under a general anaesthetic (you are put to sleep for the procedure). An incision of 2-3 cm is made in the scrotum and a small sample of one testis is taken. The testicle and skin are repaired with dissolving sutures. A small dressing or special skin glue is used to protect the wound. You normally require some Paracetamol or Neurofen for a few days after it aches. Normal activities can be resumed a few days after the biopsy and no routine surgical appointments are needed after surgery. There is usually no significant difference in the size of the testes afterwards and it is unusual to see the scar when it heals.

What might be found at biopsy and what happens to samples taken?

The tissue taken at biopsy is collected by in vitro fertilization (IVF) scientists and stored at IVF laboratories. Currently the scientists from the Reproductive Services Department at The Royal Women’s Hospital (RWH) are able to collect the tissue at surgery and process it at their centre. Depending on your stage of puberty, it may be possible that mature sperm (that could be frozen/’banked’) directly may be found in a biopsy sample. This is more likely if you are in mid-later stages of puberty as that is when sperm production becomes established. If this is the case, these sperm will be frozen and stored. This would potentially be more beneficial to you, as there is more chance that mature sperm will be useful for a pregnancy in future. In addition, if there is any remaining tissue that has immature germ cells, this will be frozen. If the biopsy sample only contains immature germ cells, this tissue will be stored by freezing it. It is important that you understand that at present, any immature testicular tissue that is removed and stored cannot currently be used for fertility purposes. There is some hope that the technology may progress by the time you are ready to try to have a baby; however this will require future advancements in scientific processes. There is no guarantee of future fertility with stored testicular tissue. This is why the procedure (testicular tissue cryopreservation) is not considered standard practice. Until these scientific techniques are successfully developed for humans, the best way of storing testicular tissue from biopsies is not yet known. If it turns out in future that freezing is not the best storage method, sperm production from the immature tissue may not be possible either.

If your tissue is stored at RWH, it does not mean that you will be required to have future fertility care at that centre (Melbourne IVF, RWH). You may go wherever you choose when the time comes.

Who is eligible?

At RCH Gender Service (RCHGS), testicular biopsy for tissue cryopreservation is offered to young people who are about to undertake gender affirming hormone therapy with oestrogen. It is also offered to adolescents who are already in mid-late stages of puberty when they are starting puberty blockers, as mature sperm that could be ‘banked’ may be found in this instance. For those who start puberty blockers in the early stages of puberty where mature sperm are unlikely to be present, we typically defer testicular biopsy until such a time as oestrogen therapy may be considered in future, as there is no benefit in doing it early.
We also need to determine that you are well enough for surgery and to ensure that the procedure does not interfere with any other treatment you are having. We also take into account differing views within a family about such procedures. Sometimes we consult with the Clinical Ethics Committee to assist the decision making for the procedure in the event that the decision is not so clear cut.

When might a testicular biopsy not be low risk/in your best interests?

In some people, the procedure may have higher risks than typical and this should be discussed in more detail. Please let your doctor know if you have/have ever had:

- Any significant problem with a previous general anaesthetic.
- Any known problem with clotting (eg haemophilia/previous significant bleeding after surgery) or serious immune system disorders.
- Any significant problem with a previous general anesthetic.

Risks and benefits

It remains experimental to collect and store frozen testicular tissue in the hope of future fertility. The surgery for the biopsy itself is not experimental as this procedure is performed routinely by surgeons for other indications. All surgical procedures have some associated risks. It is likely that you would be having a general anaesthetic for this procedure that you would not otherwise need. General anaesthetics are generally considered safe; however some people may have side effects. This needs to be weighed against the potential benefits of keeping fertility options open in the future, although we do not know whether this will indeed be the case for you.

Expected risks of the surgical procedure:

1. Risk of a general anaesthetic. There may be situations where your medical situation may present specific increases in the risk of anaesthesia. In these situations the anaesthetist and the surgical/other treating teams will need to discuss the risk versus benefit issues. Your safety is a very important factor. Mostly the risk relating to the anaesthesia will be very small. The anaesthetist can clarify if there are any special aspects of your condition that could influence the risk of surgery.

2. Risk of surgery: infection, bleeding and damage to internal structures. In the rare instance that there is a complication such as a haematoma (collection of blood) a second surgery to manage a complication might be required.

3. The biopsied testis may not develop fully as a result of the biopsy so may remain smaller than the non-biopsied testis.

What are the costs involved?:

- The procedure itself is currently offered without charge at RCH but this may change in future. If you wait to have procedure done as an adult, there would likely be costs involved.

Other issues to consider:

1. Storage of testicular tissue/immature germ cells/sperm: occurs for 20 years if it collected prior to 18 years of age, after which time you have to renew the request. This is an arrangement between you and the IVF centre and does not involve The Royal Children’s Hospital. You will be provided with a summary of what tissue has been stored and contact details for the relevant facility for your future use. If your details (e.g. contact details/home address) change over time it will be important to let the IVF centre know.

2. Costs of future IVF treatment and tissue storage.

3. The tissue can only be used by you and, in the unfortunate event of your death in future, the tissue must be disposed of. In this instance a member of the team would contact your family to discuss arrangements.

4. Due to current legal restrictions, the tissue cannot be donated for research. The tissue may never be utilized by anyone other than you.
The Royal Children’s Hospital Gender Service

Information sheet — Trans masculine young people and other trans, gender diverse or non-binary people with ovaries

Fertility is the ability to produce biologically related children. Many trans and gender diverse (TGD) youth will want to be parents one day and may wish to be biologically related to their children.

There are a few different ways to become a parent. Some people will have a child they are biologically related to. Other people will be foster-parents, step-parents or adoptive-parents. There is no right or wrong way to become a parent. Some adults will choose not to be parents and will prefer to pass on their wisdom to the next generation by being an uncle, aunt or other significant person in a young person’s life. All of these choices are valid and many young people do not know which they will pursue in the future. Many adults make these decisions together with a life partner which adds another layer of unknowns to the decision making. The Gender Service at the RCH keeps options open for our TGD youth so they can make their own fertility decisions in the future.

To create a baby, an egg from an ovary and a sperm from a testicle must meet together. This can occur either in a human body or in a lab with in vitro fertilisation (IVF). Together, the egg and sperm become an embryo. An embryo can develop into a baby if it grows inside a person’s uterus. Once the baby is ready to be born, it can come out through the birth canal (vagina) or via an operation on the abdomen (caesarean section). The eggs and sperm carry genetic information from the person whose ovary or testis made them. This genetic information is what makes people biologically related. It is the reason that parents and children often look similar.

For people with ovaries, the process of fertility begins when the ovaries first develop. All the eggs a person will ever have are produced before birth. The eggs are stored in the ovaries and after puberty the eggs can become active and be used to create a baby. The eggs are activated a few at a time by hormones from the brain and this occurs as part of a menstrual cycle. Activation and release of an egg is called ovulation. Ovaries usually run out of eggs at around age 50 but after age 40 the eggs are lower quality.

For TGD youth, gender affirming hormone treatment may be used: i) to stop the further progression of female puberty (with puberty blockers/GnRHa therapy) or ii) to masculinise physical features (with testosterone). Some TGD youth will also use hormonal medication such as norethisterone to stop bleeding (periods). All of these hormonal medications affect the ovaries while they are being taken. Puberty blockers and testosterone stop the ovaries from producing hormones and reduce the number of ovulations that occur. If someone stops taking puberty blockers or testosterone, the ovaries will start producing hormones and ovulating again. Taking puberty blockers or testosterone does not reduce the quality of the eggs over time, but age does, so a person in their 40s who stops testosterone will have less chance of making a baby compared to someone in their 20s or early 30s.

Regardless of being TGD or not, around 10-15% of the Australian population will have reduced fertility. Some people are never able to have biologically related children, even if they strongly wish to. Medical treatments can be used to increase fertility and these are called artificial reproductive technologies (ART). ART includes ovulation induction, IVF, egg freezing (oocyte cryopreservation), ovarian cryopreservation and donor gametes. Donor gametes is the name given to eggs and sperm which are given (donated) from one person to another to allow the recipient to have a child.

Ovulation induction is when hormone medication is used to make the ovaries activate some eggs. These activated eggs can be collected via a surgical procedure (through the vagina) and be frozen (oocyte cryopreservation) or used to create an embryo via IVF. The hormone medication used for ovulation induction makes a large surge of oestrogen in the body which can have mental or physical health impacts. Egg collection is usually performed after age 17.

Ovarian cryopreservation is when a small segment of ovary (tissue) is surgically removed and frozen. It can be re-implanted into the same person’s body and then used to harvest eggs. Scientists are trying to develop this technology so that eggs can be collected from this tissue in a lab, but it isn’t possible to do this yet. Ovarian cryopreservation is often considered when someone will be having chemotherapy for cancer which can damage the ovaries. There have been 150 births using this technique worldwide and it is considered experimental. The surgical procedure used to collect the piece of ovary has rare but significant risks associated with it, such as a four in 100,000 risk of death.

If tissue or eggs are collected, they will be stored at an IVF centre. There may be a cost involved for processing and storage that is determined by the IVF centre. The tissue or eggs stored from teenagers can’t be donated to anyone else. They can be stored for 20 years, and after that point need to be discarded unless an extension is sought. The RCHGS is not an IVF centre. Tissue collection procedures can be undertaken at the RCH, but we refer our young people on to an IVF centre of their choice if egg freezing is requested.

It is important to note that whilst testosterone reduces the number of ovulations that occur, it does not stop them completely in some people. This means that people taking testosterone need to use contraception (such as condoms and/or an intrauterine device) if they are having sex that could lead to pregnancy (eg. penis in vagina).

Some TGD people may choose to have gender affirming surgery as adults which can significantly affect fertility, such as removal of the uterus or ovaries. This surgery is not performed at the RCH.

Information updated by the RCH Gender Service, RCH July 2019.
Dear Parent/Guardian of: 
Date of birth: 

Fertility preservation procedure: OVARIAN TISSUE FREEZE 
Date of procedure: 
Number of slices frozen: 
Location where ovarian tissue is being stored: The Royal Women’s Hospital/Melbourne IVF (Reproductive Services Unit)
Contact telephone number: (03) 8345 3242
Storage facility email address: lab.supervisors@mivf.com.au
RCH fertility preservation contact: Rafael Serrano
RCH fertility preservation contact email address: fertility@rch.org.au

Please remember that at the time of this letter, the fees for storage of the ovarian tissue from your child have been waived until the age of 21, however the IVF centre may change this in the future. According to Victorian Law, your child will take ownership of the frozen ovarian tissue once she reaches the age of 18.

It is important that you contact either the facility where your child’s ovarian tissue is stored or the RCH contact person if the following circumstances happen:
• Change of address or telephone number.
• Your child turns 21 years of age.
• If the tissue is not going to be used or no longer required.

After treatment is over, members of the Oncofertility team can answer questions about growth development and fertility. Your oncologist can organise a referral to the Oncofertility team when you and your child are ready.

Kind regards,
Rafael Serrano Real
Oncofertility Coordinator
Dear Parent/Guardian of:

Fertility preservation procedure: SPERM BANKING

Date of procedure:

Number of straws frozen:

Location where sperm is stored: Andrology Unit/The Royal Women's Hospital

Andrology Contact number: (03) 8345 3992

RCH fertility preservation contact: Rafael Serrano

RCH fertility preservation contact email address: fertility@rch.org.au

Please be advised that the fee for the storage of the sperm samples in the Andrology department will be covered by the charity My Room for the first two years. You will be contacted by the Andrology department after this period.

It is important that you contact either the facility where your child’s sperm sample is stored or the RCH contact person if the following circumstances happen:

• Change of address or telephone number.
• Your child turns 21 years of age.
• If the sperm sample is not going to be used or no longer required.

After treatment is over, members of the Oncofertility team can answer questions about growth development and fertility. Your oncologist can organise a referral to the Oncofertility team when you and your child are ready.

Kind regards,

Rafael Serrano Real
Oncofertility Coordinator

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Dear Parent/Guardian of:

Fertility preservation procedure: TESTICULAR TISSUE FREEZE

Date of procedure:

Number of vials frozen:

Location where testicular tissue is being stored: The Royal Women's Hospital/Melbourne IVF (Reproductive Services Unit)

Contact telephone number: (03) 8345 3232

Storage facility email address: lab.supervisors@mivf.com.au

RCH fertility preservation contact: Rafael Serrano

RCH fertility preservation contact email address: fertility@rch.org.au

Please remember that at the time of this letter, the fees for storage of the testicular tissue from your child have been waived until the age of 21, however the IVF centre may change this in the future. According to Victorian Law, your child will take ownership of the frozen testicular tissue once he reaches the age of 18.

It is important that you contact either the facility where your child’s testicular tissue is stored or the RCH contact person if the following circumstances happen:

• Change of address or telephone number.
• Your child turns 21 years of age.
• If the tissue is not going to be used or no longer required.

After treatment is over, members of the Oncofertility team can answer questions about growth development and fertility. Your oncologist can organise a referral to the Oncofertility team when you and your child are ready.

Kind regards,

Rafael Serrano Real
Oncofertility Coordinator
To the Parent/Guardian of:

Fertility preservation procedure:

Date of injection:

The medication your child has received is a long-acting injection usually given every three months. It works by shutting down the activity of the ovaries. Some research on adults suggests these medications might have a slight protective effect on fertility, but the results are not consistent. It can also be prescribed to suppress menstruation, which can be advantageous during chemotherapy treatment, in order to reduce heavy periods.

Some of the side effects of the Zoladex® injection include hot flushes and mood changes if they used for over six months. Your doctor will let you know if you have to use it for longer.

Date for the next Zoladex® injections (if needed):

After treatment is over, members of the Oncofertility or Gynaecology team can answer questions about growth development and fertility if required. Your oncologist can organise a referral to the Oncofertility team when you and your child are ready.

For any questions related to this medication or to the fertility preservation service, please do not hesitate to contact the Oncofertility Coordinator.

RCH fertility preservation contact: Rafael Serrano Real
RCH fertility preservation contact email address: fertility@rch.org.au

Kind regards,

Rafael Serrano Real
Oncofertility Coordinator