





Handbook for Guideline Development

Version 4

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On behalf of the IGHG core group and the PanCare Guidelines Group

This handbook has been developed by a collaborative effort of the International Late Effects of Childhood Cancer Guideline Harmonization Group (IGHG) (www.ighg.org), the PanCare Guidelines Group) (www.pancare.eu) and the Cochrane Childhood Cancer Group (CCG) (ccc.ccchrane.org).

This handbook can be used by the chairs and members of the IGHG guideline panels after they have been assigned to develop a special guideline.

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1 Introduction

1.1 Aim of the handbook

The principle aim of this handbook is to serve as a reference tool and provide guidance to the members of the guideline panels involved in the guideline development work of the International Late Effects of Childhood Cancer Guideline Harmonization Group (IGHG)¹ (www.ighg.org) and the PanCare Guideline Group (www.pancare.eu). The systematic approach outlined in this manual aims to improve the methodological quality of the clinical practice guidelines for the follow-up of survivors of childhood, adolescent and young adult (CAYA) cancer and positively impact on the quality of care CAYA cancer survivors receive.

1.2 Clinical practice guidelines

Clinical practice guidelines (CPGs) are defined by the Institute of Medicine as "statements that include recommendations intended to optimize patient care that are informed by a systematic review of evidence and an assessment of the benefits and harms of alternative care options.²

CPGs aim to provide appropriate recommendations for practice based on a transparent process and informed by evidence. CPGs are essential to ensuring that CAYA cancer survivors receive optimum health care. ^{2,3} However, it is essential to ensure optimum quality of guidelines if they are to improve both the process and outcome of care.

It is critical that CPGs are developed based on the methods of evidence-based medicine (EBM). EBM is "the conscientious, explicit and judicious use of current best evidence in making decisions about the care of individual patients". EBM begins with the formulation of clinically relevant questions based on the Participants, Interventions, Control group & Outcome (PICO) system, followed by a synthesis of the evidence based on an extensive literature search (e.g. systematic review or evidence tables). The data is then used to develop evidence-based clinical policy (recommendations) before applying these policies or CPGs in practice (Figure 1).

EBM is an integration of best research evidence, clinical expertise and patient concerns.

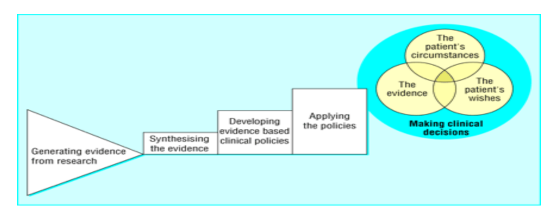


Fig. 1: The path from the generation of evidence to the application of evidence⁵



Members are encouraged to listen to the first web-training conference given by L. Kremer. This provides audio commentary and PowerPoint slides to give useful background information to evidence based clinical practice guidelines. The presentation is available at: https://connect.sunet.se/p5gqc2b67eg/

1.3 Guidelines for the long-term follow-up of CAYA cancer survivors

Several guidelines for the long-term follow-up of CAYA cancer survivors have been developed, published and widely disseminated, including those produced by the US Children's Oncology Group (COG), Dutch Childhood Oncology Group (DCOG), United Kingdom Children's Cancer Study Group (UKCCSG) and Scottish Intercollegiate Guidelines Network (SIGN). A recent survey performed by PCSF found that these guidelines were in widespread use across Europe and that some European nations have also developed additional local guidelines.

However, as existing guideline development groups worked independently, inconsistencies exist in the methodology used and also in the final recommendations of these guidelines. A major consequence of this is uncertainty among clinicians regarding which guidelines to implement.

Therefore, the goal of the IGHG and PanCare Guideline Group is to harmonise efforts and employ a systematic and rigorous methodology to produce clinical consensus in guidelines for long-term follow up of major late adverse effects in CAYA cancer survivors. We aim to promote healthy lifestyles, provide on-going monitoring of health status, facilitate early detection of late effects, and advise about timely intervention strategies to preserve health.

1.4 Structure of this handbook

In this handbook we aim to provide information that may be useful to members of the IGHG/PanCare Guideline Group guideline panels, and the working group leaders in particular as they prepare for and proceed through the guideline development work.

Specifically the handbook will:

- 1. Outline the key steps in the development of clinical practice guidelines.
- 2. Direct members to other important sources of information/documentation integral to the guideline development work.
- 3. Provide practical information regarding the organisation and management of the working groups.

As opposed to fully reproducing information which is documented elsewhere, this handbook provides an overview and directs members to other documents that explain the relevant issues in more detail. Links to these other documents are embedded in the text, and can be accessed by clicking on the document name. These documents include a published methodology paper, protocols from previous guideline topics that have been completed by IGHG/PCSF, and also practical examples from the published guidelines to more clearly illustrate the process.

2 Methodology utilised by IGHG/PCSF to develop evidence based CPG's

Developing a guideline encompasses three phases:

- 1. Preparation phase
- 2. Development phase
- 3. Finalisation phase



Members are encouraged to listen to the second web-training conference given by R Mulder. This provides useful background information to the development of clinical practice guidelines. The presentation is available at:

https://connect.sunet.se/p2a8jwypnwg/

2.1 Preparation phase

The guideline panel

Convening an effective guideline panel is a crucial stage in producing a guideline. Each guideline panel will consist of a working group including:

- Chair(s): leaders in the field
- Coordinator(s): project managers administrating group activities
- Advisors: Leontien Kremer, Melissa Hudson, Renée Mulder, Rod Skinner, Sandy Constine (radiation expert), Hamish Wallace
- Working group leaders: leaders supervising literature reviews of focused clinical questions
- Working group members

Diversity is an essential feature of a guideline panel. Its exact composition should be tailored to the guideline topic and reflect the range of stakeholders involved. At a minimum the panel should comprise at least of content experts, non-expert clinicians, health care providers and methodologists. In addition, patients or their representatives may be eligible members.

Scope of the guideline

It is important to define the scope of the guideline:

- Definition of outcomes / health problem
- Age range of the population of interest:
 - Childhood, adolescent and young adult cancer survivors diagnosed with cancer up to age 30 years; depending on the health problem adaptation of the age range (e.g., 18, 21 or 25 years) may be appropriate.
- Survival time of the population of interest:
 - Childhood, adolescent and young adult cancer survivors 2-years after completion of treatment; depending on the health problem adaptation of the survival time (e.g., immediately following or 5 years post-treatment) may be appropriate.

For every guideline topic, the following key issues can be considered which are important for the final recommendations:

- Does early diagnosis result in better outcomes?
- ➤ Who needs surveillance?
- At what age or time from exposure should surveillance be initiated?
- ➤ At what frequency should surveillance be performed?
- ➤ When should surveillance be stopped?
- What surveillance modality should be used?
- What should be done if abnormalities are identified?

2.2 Development phase

In general, the guideline development process consists of five steps:

- 1. Evaluate concordances and discordances among recommendations in existing guidelines.
- 2. Formulate clinical questions.
- 3. Identify available evidence by systematic literature searches.
- 4. Summarize and grade evidence.
- 5. Formulate and grade recommendations.

Figure 2 outlines the main steps that IGHG & PanCare Guidelines Group will be undertaking in the development of guideline recommendations.

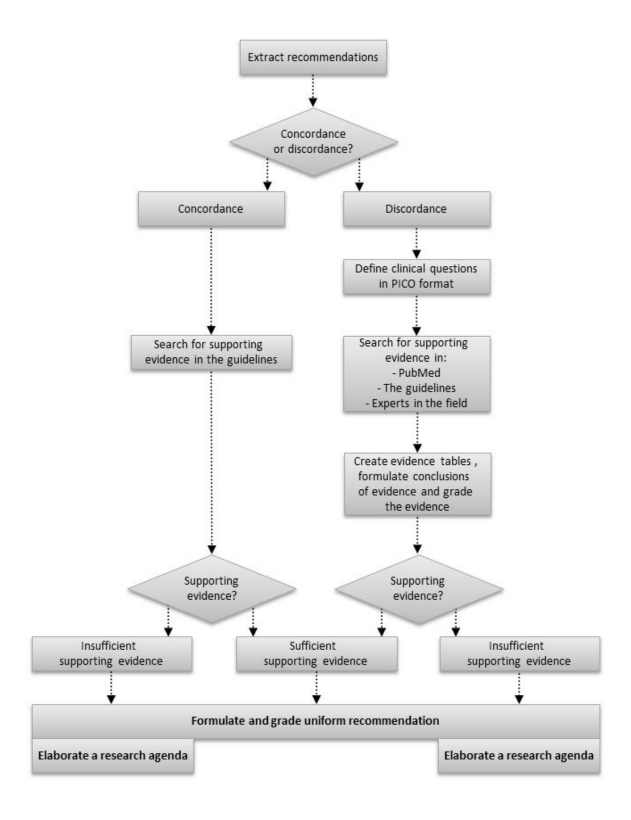


Fig. 2: Key stages in the development of recommendations

Step 1: Evaluate concordances and discordances of current recommendations

The first step is to extract the recommendations for the topic from the existing and more widely disseminated guideline groups (e.g., COG, DCOG, UKCCLG and SIGN guidelines). The level of discordance/concordance between these recommendations is then evaluated.

If recommendations are concordant, the quality of the supporting evidence will be reviewed to determine if it is sufficient or insufficient. Extensive evidence summaries will not be developed for concordant recommendations.

Discordant recommendations will form the basis for the formulation of clinical questions. These clinical questions will clearly state what the evidence aims to answer.

Below is an example of the evaluation of the concordance and discordance regarding the surveillance of breast cancer in survivors (Table 1).

<u>Table 1: Concordance and discordance 'Who needs breast cancer surveillance?'</u>

	cog	DCOG	UKCCLG	Concordant/ discordant
Who needs breast can	cer surveillance?			
At risk Chest radiation	Yes	Yes	Yes	Concordant
± Alkylating agents	Not specified	Not specified	Yes	Discordant
High risk	Not specified	≥7-20 Gy chest radiation (excl. TBI) ≥14-40 Gy abdominal radiation	Not specified	Discordant
Highest risk	≥20 Gy chest radiation	≥20 Gy chest radiation ≥40 Gy abdominal radiation TBI	Not specified	Discordant

Step 2: Formulate clinical questions

Effective and efficient guideline development involves asking and answering key clinical questions. These questions should be clear, focused and closely define the boundaries of the topic. They will serve as a starting point for the systematic literature search that aims to identify all the available evidence. These questions also form the basis of the development of recommendations.

The **PICO** (Participants, Interventions, Control group & Outcome) framework is helpful to identify the main elements of the clinical question. It breaks the question down into four key elements:

- ➤ Who are the **P**articipants you want to study? (e.g., gender, age, disease)
- ➤ What is the Intervention you want to examine? (etiologic/risk factor; e.g., type of treatment)
- What do you want to **C**ompare against your intervention of interest? (e.g., alternative interventions this is not always necessary or relevant)
- What are the Outcomes you want to measure? (e.g., improved quality of life, late effects)

Examples of the formulation of a clinical question is shown in Table 2.

Table 2: Example clinical questions derived from the PICO structure

Does early diagnosis result in better outcomes?

Р	1	С	0	Clinical question
Childhood, adolescent and young adult cancer survivors with a CNS neoplasm	Tumor size, asymptomatic or symptomatic stage	Not applicable	Mortality, recurrence, survival, adverse events, quality of life	Does the detection of a meningioma in a smaller size or asymptomatic stage contribute to a reduced mortality rate in CAYA cancer survivors?

Who needs surveillance?

P	I	С	0	Clinical question
Female	Low dose	Childhood	Breast cancer	What is the risk of breast cancer in
childhood,	chest radiation	cancer	risk	female CAYA cancer survivors treated
adolescent		survivors		with 1-9 Gy chest radiation compared
and young		treated		to survivors treated without chest
adult cancer		without chest		radiation?
survivors		radiation		

At what age or time from exposure should surveillance be initiated?

P	1	С	0	Clinical question
Female childhood, adolescent and young adult cancer survivors	Chest radiation	N/A	Latency time breast cancer	What is the latency time (time of onset) to develop breast cancer in CAYA cancer survivors treated with chest radiation?

At what frequency should surveillance be performed?

P	I	С	0	Clinical question
Female childhood, adolescent and young adult cancer survivors	Chest radiation	N/A	Breast cancer risk over time	Does the breast cancer risk change over time (improve, deteriorate, plateau) in female CAYA cancer survivors treated with chest radiation? What is the timing of such change?

When should surveillance be stopped?

P	I	С	0	Clinical question
Female childhood, adolescent and young adult cancer survivors	Chest radiation	N/A	Breast cancer risk in CAYA cancer survivors aged >50 years	What is the risk of breast cancer in CAYA cancer survivors treated with chest radiation aged >50 years?

What surveillance modality should be used?

Р	1	С	0	Clinical question
Female childhood, adolescent and young adult cancer survivors	MRI	Mammography	Diagnostic value to detect breast cancer	What is the diagnostic value (sensitivity, specificity, predictive value) of a MRI compared to a mammography to detect breast cancer in female CAYA cancer survivors?

What should be done if abnormalities are identified?

P	I	С	0	Clinical question
Childhood, adolescent and young adult cancer survivors	Physical activity training	No physical activity training	Pulmonary outcomes	What are the positive and adverse effects of physical activity on pulmonary outcomes in CAYA cancer survivors?

Step 3: Identify and select the evidence

It is important that the literature search is thorough, objective and rigorous. An inefficient or biased literature search can compromise the validity of the recommendations and the guidelines. The aim is to identify as many relevant studies as possible (within the limits of resources and time). It is also essential that the literature search is transparent, well documented and reproducible.

Where adequate published systematic reviews exist, additional literature searches may be limited to updating, covering the time period since the review was conducted.

Carrying out a literature search to identify and select relevant studies will involve:

- 1. Designing search strategies
- 2. Defining in- and exclusion criteria
- 3. Selecting studies for evidence summaries

1. Design search strategies

Where to search? Searches are carried out in bibliographic databases. There are several that can be searched but Medline and Embase are two of the key international health databases. Although there is significant overlap in these databases, differences do exist. The Cochrane Central Library of Controlled Trials is also a database for systematic reviews that can be searched.

In addition to searching bibliographical databases, papers should also be identified through references in the existing guidelines, as well as important reviews and key papers known to the group members.

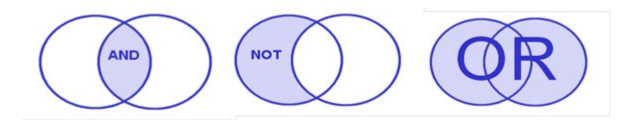
What to search? In order to search for and identify relevant studies, a search strategy must be developed. The search strategy is based on the main concepts in the clinical question identified through the PICO framework e.g. population, intervention, comparison and outcome. The clinical questions should be translated into key words and/or search terms. The Cochrane Childhood Cancer Group (ccg.cochrane.org) will develop the search strategies. However, members of the panel will be asked to suggest appropriate search terms and to check if the final search strategy is comprehensive.

Searches can be conducted in databases using either controlled vocabulary based on Medical Subject Headings (MeSH) or by using free-text/keywords. MeSH headings are useful as they index all articles that use different spellings/words to describe the same concept (e.g. cancer, lymphoma, leukemia, Ewing's sarcoma) under the same subject heading (e.g. Neoplasms). This precludes the need to search for a large list of synonyms. To identify keywords, however, look for the exact word you are searching for within the title and/or abstract of the articles within the database.

It is important to consider and include all of the related terms, variations in spellings and synonyms for each concept included in your search. A combination of subject headings and keywords is usually recommended to ensure that as many relevant records as possible are identified.

In **Appendix 1** standard search strategies as used by the IGHG and PanCare Guideline Group are shown. In addition, an example of a full search strategy taken from the male gonadal dysfunction guidelines protocol can be found in **Appendix 2**.

How to search? In the example in Appendix 2 many of the terms relating to the PICO framework are combined by 'OR'. This is a Boolean operator. Other Boolean operators are 'AND' and 'NOT'. Boolean operators make it possible to combine the results from two or more different searches using controlled vocabulary or keywords.



- AND retrieves only those articles in which all of the terms appear
- **NOT** used to exclude a term from your search
- **OR** retrieves those articles in which either of the terms appear



For an explanation of search strategies and Boolean operators please see Lundh et al (2007). <u>Development of a search strategy</u>.⁶

2. Defining in- and exclusion criteria

It is important to define clear inclusion and exclusion criteria for the selection of studies, based on the **PICO**s. The following criteria should be considered:

• Study population:

- o Childhood, adolescent and young adult cancer survivors
- At least 75% diagnosed with cancer prior to the defined age range (i.e.,18 / 21 / 25 / 30 years)
- At least 50% survived the defined survival time (i.e., immediate end of treatment / ≥
 2 years post-treatment / ≥ 5 years post-treatment)

Outcomes:

- Outcome definition of specific late effect
- Studies investigating one of the following outcomes depending on the clinical question:
 - For 'Who needs surveillance?': risks and risk factors.
 - For 'At what age or time from exposure should surveillance be initiated?: latency time / time of onset after exposure.
 - For 'At what frequency should surveillance be performed?': risks over time.
 - For 'What surveillance modality should be used?': diagnostic value / sensitivity, specificity, positive predictive value, negative predictive value of diagnostic tests.
 - For 'What should be done if abnormalities are identified?': effectiveness of interventions that may result in better outcomes.

Types of studies:

- Include all study designs except case reports and case series (systematic reviews provide the highest level of evidence followed by randomized controlled trials, observational studies)
 - For 'At what frequency surveillance should be performed?' longitudinal studies with more than one measurement per patient should be included.
 - For 'What surveillance modality should be used?' diagnostic studies should be included.
 - Regarding reviews: During screening of abstracts include all reviews (both systematic and narrative reviews). In cases of systematic reviews, include and use conclusions for generating evidence tables. In cases of narrative reviews, exclude, but screen reference lists in order to check for missing relevant papers.
- Define minimum sample size, for example at least N=20 depending on the clinical problem and availability of evidence.
- o Prioritize, when available, studies that controlled for important confounding factors:
 - Cohort study: multivariable / multiple regression analysis;
 - Case-control study: matching or risk stratification.
- o Limit search to English language publications.
- Define dates of search parameters, e.g., published from a specific date onwards (i.e. 1990).

3. Identify and select studies

Once the literature search of the electronic databases is complete, the following steps should be taken for selecting the studies:

- Two reviewers will assess if publications meet inclusion criteria based on the titles and abstracts of the studies.
- Every abstract will be assessed regarding the appropriateness of study inclusion (i.e., should be included, should be excluded, or inclusion uncertain).
- The results of reviewers' assessments will be compared and discrepancies discussed and resolved.
- The coordinator will obtain all "included" and "uncertain" abstracts in full text and send it to the two reviewers to determine if the inclusion criteria are met.
- Each full text paper will be reviewed and assessed regarding the appropriateness of inclusion of the study (i.e., should be included or excluded). Reason for study exclusion should be noted.
- Identify the clinical question for which the study should possibly be included.
- Discuss discrepancies with companion reviewer to reach consensus.

Besides to the PubMed search additional studies will be identified by:

- References in reviews
- References supporting the existing long-term follow-up guidelines
- Experts in the field

When evidence is lacking for childhood and young adult cancer survivors, we will carefully extrapolate evidence from other populations.

Step 4: Summarize and appraise quality of evidence

1. Evidence tables

The evidence found in the literature should be summarized in evidence tables (see Appendix 3). The evidence tables provide information about study and patient characteristics, primary study outcomes, risk of bias, including selection bias, attrition bias, detection bias and confounding (see Appendix 4), and additional remarks, such as other factors that may bias results.



For an explanation of the different types of bias, please see van Dalen et al (2007) Quality of studies included in a systematic review and associated risk of bias⁷ and the Cochrane Bias Methods Group.

2. <u>Summary of findings tables of the body of evidence</u>

a. <u>Description of studies</u>

For each clinical question a summary of findings table of the body of evidence will be completed. A summary of findings table provides key information of every single study about the main patient characteristics, the magnitude of effects for the defined outcomes and determinants, and the quality of that study (see Appendix 5).

b. Grading the quality of the body of evidence

The Grades of Recommendation, Assessment, Development and Evaluation Working Group (GRADE Working Group) has developed a system for grading the quality of a body of evidence, 8-11 The quality of a body of evidence is defined as the extent to which one can be confident that an identified effect or association is true. Assessing the quality of a body of evidence involves several considerations. There are five reasons for downgrading the quality of a body of evidence:

- Study limitations (risk of bias, **see Appendix 4**): the confidence in the body of evidence decreases when studies have major limitations that may bias the risk estimates. Every study addressing a particular outcome will differ, to some degree, in the risk of bias. The reviewers must make an overall judgement on whether the quality of evidence for an outcome warrants downgrading on the basis of study limitations.
- Inconsistency of the results (heterogeneity): the confidence in the body of evidence decreases
 when there is a degree of inconsistency of effect between or within studies (when studies yield
 widely differing estimates of effect). When heterogeneity exists and affects the interpretation of
 results, but reviewers fail to identify a plausible explanation, the quality of the evidence
 decreases.
- Indirectness of the study population, intervention and outcomes: the confidence in the body of
 evidence decreases when the study population and outcomes from the studies are not
 generalizable to the population and outcome of interest. Reviewers should make judgements
 transparent when they believe downgrading is justified, based on differences in anticipated
 effects in the group of primary interest.
- Imprecision of the effect estimates: the confidence in the body of evidence decreases when the effect estimates are imprecise. This is the case if studies include relatively few patients and few

events and thus have wide confidence intervals. Another criteria to consider is the clinical decision threshold. This is the threshold of the effect size that would change the decision whether or not to adopt a clinical action.

Risk of publications bias: the confidence in the body of evidence decreases when investigators
fail to report studies or outcomes on the basis of results, typically those studies that show no
effect.

There are also circumstances in which the quality of the body of evidence can be upgraded:

- 1. Large magnitude of effect: the confidence in the body of evidence increases when methodologically well-performed observational studies yield large, consistent and precise estimates of the magnitude of effect.
- 2. Dose response gradient: the confidence in the body of evidence increases when there is evidence for a dose response across or within studies, or when inconsistency across studies is explained by a dose response.
- 3. Plausible confounding: the confidence in the body of evidence increases when adjustment for confounding factors would have increased the effect size.

In **Appendix 6** the criteria for grading the quality of the body of evidence is described. The rational for grading the quality of the body of evidence should be described in the summary of findings table (see **Appendix 5**).

The evidence is graded according to four levels:

- $\bigoplus \bigoplus \bigoplus$ High: further research is unlikely to change the confidence in the estimate of effect.
- $\bigoplus \bigoplus \bigoplus \bigoplus$ Moderate: further research is likely to have an important impact on the confidence in the estimate of effect and may change the estimate.
- $\bigoplus \bigoplus \bigoplus \bigoplus$ Low: further research is very likely to have an important impact on the confidence in the estimate of effect and is likely to change the estimate.
- ⊕⊖⊖⊖ Very low: any estimate of effect is very uncertain.

c. Formulation of the conclusions of evidence

Based on the quality of the body of evidence an overall conclusion will be formulated for that specific clinical question (see Appendix 5 and 7).

Grading the evidence gives an impression of the quality of the included studies. It is not related to the importance of the recommendation but to the strength of the supporting evidence.



For an explanation of the different types of factors that may decrease or increase the quality of a body of evidence, please see the in-depth publications on the GRADE website.

Step 5: Formulate recommendations

Once the selection and summary of the evidence is complete, the available evidence must be combined and translated into recommendations. For this purpose we will use the GRADE Evidence to Decision (EtD) framework (Appendix 8). ¹² The EtD framework ensures that all important criteria for making a decision is considered and informs the guideline panel about the relative pros and cons of the interventions or options being considered. It makes the decision making process structured and transparent.

The following criteria should be considered when formulating clinical recommendations:

- 1. Problem: Is the problem a priority?
- 2. Desirable effects: Are the desirable anticipated effects large?
- 3. Undesirable effects: Are the undesirable anticipated effects small?
- 4. Certainty of the evidence: What is the overall certainty of the evidence of effects?
- 5. Values: Is there important uncertainty about or variability in how much people value the main outcomes?
- 6. Balance of effects: Are the desirable effects large relative to the undesirable effects?
- 7. Resources required: Are the resource required small?
- 8. Certainty of evidence of required resources: What is the overall certainty of the evidence of the resources require?
- 9. Cost effectiveness: Is the incremental cost small relative to the net benefits?
- 10. Equity: What would be the impact on health equity?
- 11. Acceptability: Is the intervention/option acceptable to key stakeholders?
- 12. Feasibility: Is the intervention feasible to implement?

In addition, it is important to consider the need to maintain flexibility of application across health care systems.

Based on all the consideration a balance of consequences will be made for all different interventions/options:

- Undesirable consequences clearly outweigh desirable consequences in most settings
- Undesirable consequences probably outweigh desirable consequences in most settings
- The balance between desirable and undesirable consequences is closely balanced or uncertain
- Desirable consequences probably outweigh undesirable consequences in most settings
- Desirable consequences *clearly outweigh* undesirable consequences in most settings



For an explanation of the Evidence to Decision framework, please see the tutorials on <u>GRADEpro website</u>.

How important each of the considerations are for a recommendation can vary. To make a recommendation, the guideline panel must consider the implication and importance of each of the above judgments. In many cases, this will be straightforward and not require detailed consideration. However, when there is uncertainty or disagreement, it can help to explicitly consider this for each criterion. Based on the overall assessment across criteria the guideline panel must reach a conclusion about the direction and the strength of the recommendations.

Recommendations will be classified into three categories: strong recommendation to do (green); moderate recommendation to do (yellow); and recommendation not to do (red) (see Appendix 9).¹³ The guideline panel should provide a justification for the recommendations, based on the criteria used in their assessment.

The recommendations should be a stand-alone text written in a complete sentence. The wording should be unambiguous, clearly defined, easy to translate into clinical practice, and agreed by the complete guideline panel.

The recommendations can include the following items:

- Who needs surveillance?
- At what age or time from exposure should surveillance be initiated?
- At what frequency should surveillance be performed?
- When should surveillance be stopped?
- What surveillance modality should be used?
- What should be done if abnormalities are identified?

A first draft of the recommendations will be prepared by a smaller group (i.e. chairs, advisors, and working group leaders). Next, the recommendations will be discussed and further formulated by the total guideline panel. Additional experts and patients/survivors in the field should be invited to participate in this final discussion.

Below is an example of the recommendations from the breast cancer surveillance guideline (Table 3).

Table 3: Example recommendations

Who needs breast cancer surveillance?

Breast cancer surveillance is recommended for female childhood, adolescent and young adult cancer survivors treated with ≥10 Gy chest radiation (level A evidence, strong recommendation).

Breast cancer surveillance is reasonable for female childhood, adolescent and young adult cancer survivors treated with <u>upper abdominal radiation exposing breast tissue at a young age</u>. The surveillance decision should be an individual one, taking into account additional risk factors² and personal values regarding the harms and benefits of surveillance (see Survivor Information Form) (level B evidence, moderate recommendation).

2.3 Finalisation phase

1. Writing the guideline

All guideline topics will be summarized in a manuscript appropriate for publication in a peer-reviewed journal. The guideline should include the following items:

- Background
- Methods: clinical questions, search strategy, selection of literature
- Results: description of evidence, overall conclusions, quality of the evidence
- Considerations: translation evidence into recommendations, according to the GRADE Evidence to Decision framework
- Recommendations
- Discussion including research agenda
- Reference list

2. External review

After the recommendations have been formulated, there will be a commentary phase where external experts review the guideline for content and implementability. Feedback is sought preferably among the scientific, professional and patient organisations involved. Feedback can also be invited from methodological experts who review the guideline for methodological validity.

3. Updating the guideline

Guidelines should be kept up to date. All IGHG/PCSF guidelines will carry a statement indicating that they will be considered for revision 5 years after publication. Searches for new evidence should be performed and updating of the recommendations might be considered.

3 Roles, publication policy and author contributions

3.1 The IGHG core leadership group

- Melissa Hudson; co-chair, advisor, COG representative
- Leontien Kremer; co-chair, advisor, PanCare and DCOG representative
- Renée Mulder; coordinator, advisor, DCOG representative
- Rod Skinner; advisor, PanCare and UKCCLG representative
- Sandy Constine; advisor and radiation expert, COG representative
- Hamish Wallace, advisor; SIGN representative
- Saro Armenian; COG representative
- Smita Bhatia; COG representative
- Wendy Landier; COG representative
- Gill Levitt; UKCCLG representative
- Kevin Oeffinger; COG representative
- Lars Hjorth; PanCare representative

Roles

- The core leadership group members will set up the methodology.
- The core leadership group members will develop future plans.
- The core leadership group members will organize meetings for the guideline panels.
- The core leadership group members will guide the work of guideline panels.

3.2 The guideline panel

Roles

- The guideline panel consists of: chairs (representing different continents), a coordinator, advisors, working group leaders and working group members.
- The chairs and advisors will appoint the guideline panel.
- The IGHG core leadership group will approve the composition of the guideline panel.
- The guideline group coordinator will facilitate the group's work and telephone meetings.
- The chairs, coordinator and working group leaders will formulate clinical questions with help of the advisors.
- The final clinical questions will be reviewed by the advisors before discussion with the guideline group members.
- The Cochrane Childhood Cancer Group will develop the search strategy together with the chairs, coordinator, working group leaders and advisors.
- The chairs, coordinator and working group leaders will formulate inclusion criteria for evidence selection with help of the advisors.
- The final inclusion criteria will be reviewed/approved by the advisors before discussion with the guideline panel members.
- The chairs, coordinator, working group leaders and members will select the publications meeting the criteria established for evidence selection.

- The chairs, coordinator, working group leaders and members will produce evidence summaries with help of the advisors.
- The final evidence summaries will be reviewed by the advisors.
- The chairs, coordinator and working group leaders will formulate conclusions of evidence with help of the advisors.
- The final conclusions of evidence will be reviewed by the advisors before discussion with the guideline panel members.
- The chairs, coordinator, working group leaders and advisors will prepare a first draft of the recommendations that will be subsequently discussed with the guideline panel members.

3.3 Manuscript writing process

- A primary manuscript of the whole guideline will be drafted that will include a description of the evidence and recommendations.
- The chairs and coordinator of the guideline group will write the first draft of the manuscript.
- The advisors and working group leaders will review/revise the first draft.
- The revised manuscript will be distributed to the guideline panel members.
- The final manuscript summarizing recommendations will be approved by the IGHG core leadership group.
- The development of additional manuscripts describing special aspects of the guideline topic should be discussed with and approved by the advisors.

3.4 Manuscript authorship

- The chairs, coordinator, working group leaders, members and advisors will be authors of the
 manuscript if they are substantially involved in the guideline development process (e.g.,
 participate in the study selection, develop evidence summaries and conclusions, formulate
 recommendations, and write or provide critical input on the manuscript) (see the authorship
 guidelines of the International Committee of Medical Editors: http://www.icmje.org/)
- Authorship criteria should be communicated to members at the beginning of the guideline development process.
- The decision regarding authorship will be made by the chairs of the guideline group in consultation with the advisors.
- The person who drafts the manuscript will be first author of the guideline manuscript; this will be the coordinator or, if the coordinator is not able to write a first draft, one of the chairs. In the event that the coordinator is the first author, the chairs will be 2nd and last author or shared last authors.
- Other working group members and IGHG core leadership group members who have not been substantially involved in the guideline or manuscript development will be acknowledged in the manuscript if their contribution is limited to review and approval of the final manuscript draft.
- If financially supported by PanCare, the collaboration between IGHG and PanCare or other funding sources should be acknowledged in the titles of the published manuscripts.

3.5 Reviewers

- At least two expert reviewers and two patient representatives will be asked to review the final manuscript of the guideline.
- These reviewers will be acknowledged in the manuscript.

3.6 Presentations

- The advisors should be informed if the methods and results of IGHG endeavors are to be submitted for presentation at national and international conferences.
- All presentations in which the results of the guideline harmonization endeavor are highlighted should acknowledge the names of the IGHG core group and the specific guideline group.
- All publications and presentations should acknowledge funding sources.

4 Communication and monitoring of progress

4.1 Expected timeline for guideline development

Please note that the timelines of guideline development work are dependent on many factors, therefore, timelines will differ between topic groups. An example timeline is shown in **Appendix 10.** This provides an illustration of the stages of the work scope that are expected to occur in development work and approximate timelines. However, these timelines will be modified according to the work of each guideline panel and most likely revised as the groups progress through their work.

4.2 Teleconferences

To arrange telephone-conference times, Doodle (www.doodle.com) is useful. Invitations are sent and group members select which days/times they can/cannot attend. Changing on the time-zone support will automatically adapt the time to each participants own time-zone so there is no confusion.

The coordinator can open an account on www.freeconferencecall.com to organize free conference calls with an unlimited number of participants.

4.3 Shared-calendar

It may be useful to set up a shared-calendar system for your working group, or to send calendar invitations for meetings for teleconferences. These can be set-up in Microsoft Outlook. They allow tasks and events to be entered and for reminder alerts to be set. This can be useful to provide a visual representation of the programme of work and for working groups to keep track of tasks and progress.

5 Overview of other key sources of information and support

Briefly, others main sources of information available are:

- The <u>methodology paper</u> describing the rationale behind the harmonisation effort and the planned methodology published by Kremer et al (2013)¹
- Recommendations for breast cancer surveillance: a report from the IGHG published by Mulder et al (2013)¹⁴
- Recommendations for cardiomyopathy: a report from the IGHG published by Armenian et al (2015)¹⁵
- Recommendations for premature ovarian insufficiency: a report from the IGHG and PCSF published by van Dorp et al (2016)¹⁶
- Recommendations for male gonadotoxicity: a report from the IGHG and PCSF published by Skinner et al (2017)¹⁷
- Recommendations for thyroid cancer: a report from the IGHG and PCSF published by Clement et al (2018)¹⁸
- Recommendations for ototoxicity: a report from the IGHG and PCSF published by Clemens et al (2019)¹⁹
- Recordings of two one-hour training web-conferences on evidence based guidelines given by Leontien Kremer and Renée Mulder to PCSF WP6 members in June 2012:
 - o https://connect.sunet.se/p5gqc2b67eg/
 - o https://connect.sunet.se/p2a8jwypnwg/
- Protocol from previous guidelines on request

Useful websites are:

- International Guideline Harmonization Group: http://www.ighg.org/
- Cochrane Childhood Cancer Group: http://ccg.cochrane.org/ebch-cochrane-journal/
- Grading of Recommendations Assessment, Development and Evaluation (GRADE) working group: http://www.gradeworkinggroup.org/ and https://gradepro.org/
- Appraisal of Guidelines Research & Evaluation (AGREE): http://www.agreetrust.org/

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Appendix 1

Standard search strategies

Cancer

Cancer OR cancers OR cancer* OR oncology OR oncolog* OR neoplasm OR neoplasms OR neoplasm* OR carcinoma OR carcinom* OR tumor OR tumor OR tumor* OR tumor* OR tumors OR tumors OR tumors OR malignan* OR malignant OR hematooncological OR hemato oncological OR hemato-oncological OR hematologic neoplasms OR hematolo*

Childhood cancer

((leukemia OR leukemi* OR leukaemi* OR (childhood ALL) OR AML OR lymphoma OR lymphom* OR hodgkin OR hodgkin* OR T-cell OR B-cell OR non-hodgkin OR sarcoma OR sarcom* OR sarcoma, Ewing's OR Ewing* OR osteosarcoma OR osteosarcom* OR wilms tumor OR wilms* OR nephroblastom* OR neuroblastoma OR neuroblastom* OR rhabdomyosarcoma OR rhabdomyosarcom* OR teratoma OR teratom* OR hepatoma OR hepatoma OR hepatoma OR neuroblastoma OR hepatoblastoma OR neuroblastoma OR medulloblastoma OR medulloblastom* OR PNET* OR neuroectodermal tumors, primitive OR retinoblastoma OR retinoblastom* OR meningioma OR meningiom* OR gliom*) OR (pediatric oncology OR paediatric oncology) OR (childhood cancer OR childhood tumor OR childhood tumors)) OR (brain tumor* OR brain tumour* OR brain neoplasms OR central nervous system neoplasm OR central nervous system tumor* OR brain cancer* OR brain neoplasm*) OR (leukemia, lymphocytic, acute[mh]) OR (leukemia, lymphocytic, acute*)

Childhood cancer, testis cancer and breast cancer

((leukemia OR leukemi* OR leukaemi* OR (childhood ALL) OR AML OR lymphoma OR lymphom* OR hodgkin OR hodgkin* OR T-cell OR B-cell OR non-hodgkin OR sarcoma OR sarcoma, Ewing's OR Ewing* OR osteosarcoma OR osteosarcom* OR wilms tumor OR wilms* OR nephroblastom* OR neuroblastoma OR neuroblastoma OR rhabdomyosarcoma OR rhabdomyosarcom* OR teratoma OR teratoma OR hepatoma OR hepatoma OR hepatoblastoma OR hepatoblastoma OR neuroblastoma OR medulloblastoma OR medulloblastom* OR PNET* OR neuroectodermal tumors, primitive OR retinoblastoma OR retinoblastom* OR meningioma OR meningiom* OR glioma OR gliom*) OR (pediatric oncology OR paediatric oncology) OR (childhood cancer OR childhood tumor OR childhood tumors)) OR (brain tumor* OR brain tumour* OR brain neoplasms OR central nervous system neoplasm OR central nervous system tumour* OR brain cancer* OR brain neoplasms OR central nervous system tumour* OR brain cancer* OR brain neoplasm* OR intracranial neoplasm*) OR (leukemia, lymphocytic, acute) OR (leukemia, lymphocytic, acute*) OR (breast cancer OR breast cancers OR breast neoplasm OR breast neoplasms OR testicular neoplasms OR testicular cancer OR testis tumor OR cancer of testis OR testis tumour OR testis neoplasm* OR testis tumour OR testis tumour

Childhood brain tumors

PNET OR medulloblastoma OR medulloblastom* OR PNET* OR neuroectodermal tumors, primitive OR brain tumor* OR brain neoplasms OR central nervous system neoplasm OR central nervous system neoplasms OR central nervous system tumor* OR brain cancer* OR brain neoplasm* OR intracranial neoplasm* OR astrocytoma OR astrocytom* OR chondrosarcoma OR chondrosarcoma* OR ependymoma OR ependymom* OR germ cell tumor OR germ cell tumor* OR glioblastoma OR glioblastoma OR glioma OR gliom* OR hemangioma OR hemangioma* OR lipoma* OR meningioma* OR schwannoma OR pineal tumor OR pineal tumor* OR chordoma OR chordom* OR oligodendroglioma OR oligodendrogliom* OR rhabdoid tumor OR rhabdoid tumor* OR

craniopharyngioma OR craniopharyngiom* OR pituitary tumor OR pituitary tumor* OR CNS embryonal tumor OR CNS embryonal tumor* OR pineoblastoma OR pineoblastom* OR medulloepithelioma OR medulloepitheliom* OR ependymoblastoma OR ependymoblastom*

Childhood cancer, excluding brain tumors

leukemia OR leukemi* OR leukaemi* OR (childhood ALL) OR AML OR OR acute lymphocytic leukemia OR acute myeloid leukemia lymphoma OR lymphom* OR hodgkin OR hodgkin* OR T-cell OR B-cell OR non-hodgkin OR sarcoma OR sarcoma OR sarcoma, Ewing's OR Ewing* OR osteosarcoma OR osteosarcom* OR wilms tumor OR wilms* OR nephroblastom* OR neuroblastoma OR neuroblastom* OR teratoma OR teratom* OR hepatoma OR hepatoma OR hepatoblastoma OR retinoblastoma OR retinoblastom* OR rhabdomyosarcoma OR rhabdomyosarcom* OR pediatric oncology OR paediatric oncology OR childhood cancer OR childhood tumor OR childhood tumors

Posterior fossa tumor

(Posterior fossa OR cranial fossa OR "cranial fossa, posterior" [MeSH Terms] OR clivus) AND (Tumor OR cancer OR neoplasm OR neoplasms) OR (infratentorial AND (cancer OR tumor))

Children

Infan* OR toddler* OR minors OR minors* OR boy OR boys OR boyfriend OR boyhood OR girl* OR kid OR kids OR child OR child* OR child* OR schoolchild* OR schoolchild OR school child[tiab] OR school child*[tiab] OR adolescen* OR juvenil* OR youth* OR teen* OR under*age* OR pubescen* OR pediatrics[mh] OR pediatric* OR paediatric* OR school*[tiab]

Children and young adults

Infan* OR toddler* OR minors OR minors* OR boy OR boys OR boyfriend OR boyhood OR girl* OR kid OR kids OR child OR child* OR child* OR schoolchild* OR schoolchild OR school child[tiab] OR school child*[tiab] OR adolescen* OR juvenil* OR youth* OR teen* OR under*age* OR pubescen* OR pediatrics[mh] OR pediatric* OR paediatric* OR school[tiab] OR school*[tiab] OR young adult[mh] OR young adult

Survivors

Survivor OR survivors OR survivor* OR long term survivor OR long term survivor* OR long term survivor* OR survivor* OR survivol [mh]

Late effects

"late effect" OR "late effects" OR "late effect*" OR "late side effect" OR "late side effects" OR "late side effects" OR "late side effects" OR "late adverse effects" OR long term effect[tiab] OR long term effect* OR long term adverse effects[mh] OR aftercare OR follow up studie* OR follow up study

Male

male[tiab] OR males OR boy OR boys OR boyfriend OR boyhood

Female

female[tiab] OR females OR girl OR girls OR girlfriend OR girlhood

Offspring

Offspring OR Descendant OR Generation OR Heir OR Progeny OR Heredity OR Lineage OR Offshoot OR Posterity OR Succession successor OR Progeniture OR Spawn OR Brood OR breed OR baby

Radiotherapy - general

Radiotherapy OR radiation OR radiation therapy OR irradiation OR irradiat* OR radiation injuries OR injuries, radiation OR injury, radiation OR radiation injury OR radiation syndrome OR radiation syndromes OR syndrome radiation OR radiation sickness OR radiation oR radiation OR radiation OR radiations

Radiotherapy - extensive

Radiotherapy OR radiation OR radiation therapy OR irradiation OR irradiat* OR radiation injuries OR injuries, radiation OR injury, radiation OR radiation injury OR radiation syndrome OR radiation syndromes OR syndrome radiation OR radiation sickness OR radiation sicknesses OR sickness radiation OR radiation* OR irradiation OR radiations OR stereotactic RT OR stereotactic radiotherapy[tiab] OR gamma knife OR intensity modulated radiotherapy OR IMRT OR radiotherapy, intensity-modulated[mh] OR (three dimenstional OR 3D OR 3d CRT) OR image guided radiotherapy OR IGRT OR radiotherapy, image-guided[mh] OR photon radiotherapy OR XRT OR "photons/therapeutic use" [Mesh] OR proton radiotherapy OR PRT OR proton therapy OR proton radiation OR proton beam OR carbon ion radiotherapy

Dose-response relationship – combine with radiotherapy search

radiometry OR radiation dosage OR radiation dose OR radiation doses OR radiation dosis OR radiation dosage*
OR radiation dosimetry OR radiation dosimetr* OR dose-response relationship, radiation OR radiometr* OR
radiotherapy dosage OR radiotherapy[sh] OR radiotherapy/adverse effects OR irradiation dose OR radiotherapy
dose OR dose calculation OR near beam dose OR in beam dose OR outside beam dose OR out of beam dose OR
radiation/epidemiology OR Radiation monitoring OR Organs at risk OR radiation effects[sh] OR radiation injury
OR radiation injuries OR radiation OR Radiotherapy/complications[Mesh]

Radiotherapy fields – combine with radiotherapy search

Cranial, head and neck

Cranial OR craniospinal OR OR head[tiab] OR neck[tiab] OR skull

Hypothalamic-pituitary

(Hypothalamus OR Hypothalamus, Middle OR Hypothalamus, Anterior, OR Hypothalamus Posterior OR Pituitary Gland, Posterior OR Skull OR Orbits OR Eye OR Ear OR Nasopharynx)

TBI

TBI OR Total body OR whole body OR total body* OR body whole*

Testes

Testicles OR testicle OR testes OR testis OR testis* OR testicle* OR testes* OR pelvic region OR region, pelvic OR pelvis region OR region pelvis OR pelvic*

Ovaries

Ovary OR Pelvis OR Lesser Pelvis OR Abdomen OR Spine OR Lumbosacral Region OR Urinary Bladder OR Vagina OR Ilium

Thorax

chest OR lung OR axilla OR mediastinal OR mantle OR supraclavicular OR susclavicular OR cranial axis OR total axis OR supra diaphragm[tiab] OR abdominal OR Inverted Y[tiab] OR Left Flank OR Hemiabdomen OR Left upper quadrant OR Paraaortic OR Spleen OR craniospinal

Cranial and craniospinal radiation

cranial irradiation [mh] OR craniospinal irradiation [mh])

Chemotherapy

Antineoplastic Protocols OR Antineoplastic Combined Chemotherapy Protocols OR Chemoradiotherapy OR Chemoradiotherapy, Adjuvant OR Chemotherapy, Adjuvant OR Consolidation Chemotherapy OR Induction chemotherapy OR Maintenance chemotherapy OR Chemotherapy, Cancer, Regional Perfusion OR Antineoplastic agents OR chemotherap*

Alkylating agents

Antineoplastic agents, alkylating* OR antineoplastic alkylating agents OR alkylating agents, antineoplastic OR antineoplastic drugs, alkylating OR antineoplastics, alkylating OR alkylating antineoplastic drugs OR alkylating drugs, antineoplastic OR antineoplastic alkylating drugs OR drugs, antineoplastic alkylating OR alkylating antineoplastic agents OR alkylating Agents OR alkylating agent*

OR busulphan OR busulfan* OR myleran* OR myelosan* OR Carmustine OR BCNU OR Chlorambucil OR ifosfamide OR iphosphamide OR iso endoxan OR isophosphamide OR isofosfamide OR ifosfa* OR iphospha* OR isofosfa* OR cyclophosphamide OR cyclophosphane OR cytophosphan OR endox* OR cyclophospha* OR Lomustine OR CCNU OR lomustine* OR Mechlorethamine OR mechlorethamine*OR Chlormethine OR Mustine OR Chlorethazine OR Procarbazine OR procarbazin* OR Melphalan OR melphalan* OR Thiotepa OR Thio Tepa OR Thiophosphamide OR thiothepa* OR temozolomide OR dacarbazine OR decarbazine OR Fludarabine monophosphate*

Cyclophosphamide

cyclophosphamide OR cyclophosphane OR cytophosphan OR B-518 OR cyclophosphamide monohydrate OR monohydrate, cyclo-phosphamide OR endoxan OR cytoxan OR neosar OR procytox OR sendoxan OR cyclophosphamide, (R)-isomer OR cyclophosphamide, (S)-isomer OR cyclophosphamide, (+)-isomer OR endox* OR cyclophospha*

Ifosfamide

ifosfamide OR iphosphamide OR iso-endoxan OR iso endoxan OR isophosphamide OR isofosfamide OR holoxan OR OR cyclic p-oxides OR ethylamines OR oxazines OR ifosfa* OR iphospha* OR isofosfa* OR isophospha* OR "br cl fosfamide" OR cyfos OR ifex OR "ifo-cell" OR ifolem OR ifomide OR ifosfamidum OR ifosforamide mustard OR ifoxan OR ipambr OR iphosphamid OR isophosphoramide bromide mustard OR mitoxana OR naxamide OR seromida OR tronoxal

Melphalan

Melphalan OR Melfalan OR Alkeran OR L-PAM OR L-Sarcolysin OR Phenylaline mustard OR isopropyl melphalan

Platinum agents

Cisplatin OR Platinum Diamminodichloride OR cis-Platinum OR cis Platinum OR Dichlorodiammineplatinum OR cis-Diamminedichloroplatinum OR cis Diamminedichloroplatinum OR Platinol OR Platinol OR Platinol OR Platinol OR Platinol OR Platinol OR CDDP OR CACP OR cisplatin* OR abiplatin OR neoplatin OR cis-DDP OR Carboplatin OR CBDCA OR Carbosin OR Carbotec OR Ercar OR Neocarbo OR Paraplatin OR Carboplat OR Paraplatine OR Platinwas OR Ribocarbo OR Blastocarb OR Nealorin OR carboplatin* OR Oxaliplatin OR oxaliplatin* OR oxaliplatine OR Eloxatine OR Eloxatin OR eloxatin* OR dacotin OR dacplat OR OR l-ohp OR oxalatoplatinum OR Platinum OR Platinum Compounds OR platinum* OR organoplatinum compounds [mh]

Methotrexate

methotrexate OR methotrex* OR amethopterin OR methotrexate hydrate OR dicesium salt methotrexate OR me xate OR sodium salt methotrexate OR disodium salt methotrexate OR mexat*

Nitrosoureas

Nitrosourea OR Nitrosourea* OR Nitrosourea compounds OR Arabinopyranosyl-N-methyl-N-nitrosourea OR Aranose OR Carmustine OR BCNU OR BiCNU OR Chlorozotocin OR Ethylnitrosourea OR Fotemustine OR Lomustine OR CCNU OR Nimustine OR N-Nitroso-N-methylurea OR NMU OR Ranimustine OR MCNU OR semustine OR streptozocin OR I-nitrosourea

Cytarabine

cytosine* OR citosin* OR cytarabin* OR citarabin* OR arabino* OR arabitin* OR aracytine* OR aracytidin* OR cytin* OR cytidine* OR arac OR arac OR arafcyt OR cytosar* OR cytozar* OR ara-C OR beta-Ara C

Anthracyclines

anthracyclines OR anthracyclin* OR idarubicin OR idarubic* OR epirubicin OR epirubic* OR adriamycin OR doxorubicin OR doxorubic* OR adriamyc* OR daunorubicin OR daunorubic* OR daunoxome OR doxil OR caelyx OR myocet

Mitoxantrone

mitoxantrone OR mitoxantr*

Novel agents

Tyrosine Kinase Inhibitors OR Demethylating agents OR Oxaliplatin OR M-TOR inhibitors OR Akt Inhibitors OR PDK-1 inhibitors OR RAF inhibitors OR MEK inhibitors OR FTS inhibitors OR Drugs targeting HER Receptors OR Drugs Targeting the c-MET Receptor OR Drugs targeting IGF-IR receptor OR SRC -targeting Small Molecule Inhibitors OR Anti VEGF/VEGFR agents OR Vascular disrupting agents OR Heat shock protein inhibitors (HSP-90 inhibitors) OR Inhibitors of Ubiquitin Proteosome System OR PARP Inhibitors OR Classes of Histone Deacetylases Inhibitors OR Anthracyclines OR Monoclonal antibodies OR Tyrosine kinases inhibitors OR Bevacizumab OR avastin OR Etoposide

MIBG

131I-Meta-iodobenzylguanidine OR 131I-MIBG OR 131I-metaiodobenzylguanidine OR Iodine-131 Metaiodobenzylguanidine OR Iobenguane (131I) OR (3-Iodo-(131I)benzyl)guanidine OR Iodine Radioisotopes/therapeutic use OR 3-Iodobenzylguanidine/therapeutic use) OR (iodine-131-metaiodobenzylguanidine OR 131I-MIBG therapy OR I-metaiodobenzylguanidine OR I-131-MIBG OR 3-Iodobenzylguanidine[mh] OR (131) I-metaiodobenzylguanidine OR (MIBG AND (treatment OR therapy))

Steroids

dexamethasone OR dexamethasone* OR prednisone OR prednisolone OR prednisolone OR prednisolone OR prednisolone OR glucocorticoids OR glucocorticoid* OR "Steroids" [Mesh:NoExp] OR steroids*

HSCT

Stem cell transplant[mh] OR stem-cell transplant OR stem cell transplant*OR stem cell transplantation OR bone marrow transplantation[mh] OR transplantation, conditioning[mh] OR hematopoetic stem cell transplantation[mh] OR reduced-intensity conditioning regimen OR myeloablative agonists[mh]

Surgery

Surgery OR injury OR injuries OR surgery* OR injury* OR injuries*

Surgery extensive

surgery[tiab] OR surgical procedures, operative[mh] OR general surgery[mh] OR surgery[sh] OR operation OR ((operative OR peroperative OR preoperative OR intraoperative) AND procedure)

Neurosurgery

Neurosurgery OR neurosurger* OR neurosurgical procedure OR neurosurgical procedure* OR (brain[tiab] AND surgery) OR (cranial* AND surger*)

Ovariectomy

Ovariectomy OR ovariectomy[mh] OR ovariectom* OR oophorectomy OR oophorectom* OR unilateral ovariectomy OR partial ovariectomy OR unilateral oophorectomy OR partial ovariectomy

Oophoropexy

Oophoropexy OR Oophoropex* OR ovarium transposition OR ovarian transposition

Orchidectomy

Orchidectomy unilateral OR orchiectomy unilateral OR unilateral orchidectomy OR unilateral orchiectomy* OR unilateral orchidectomy*

Rretroperitoneal lymph node dissection and genitourinary surgery

Lymph Node Excision[mh] OR Lymph Node Excision OR (Lymph Node AND excision) OR Lymph Node Excis* OR Pelvic exenteration[mh] OR ((prostate OR bladder OR bladder neck OR rectum) AND surgery) OR pelvic exenteration

Nephrectomy

Nephrectomy[mh] OR nephrectom* OR postnephrectomy OR partial nephrectomy OR subtotal nephrectomy OR radical nephrectomy OR unilateral nephrectomy OR renal surgery

Pulmonary surgery

pulmonary metastasectomy OR pulmonary lobectomy OR thoracotomy OR sternotomy OR thoracoscopy OR rib resection[tiab] OR spinal surgery OR spinal fusion OR (resection AND (pulmonary wedge OR lung OR claviculae OR scapulae OR muscle tissue on thorax))

Hypdrocephalus VP/CSF shunts

Hydrocephalus OR hydrocephaly OR ventriculomegaly OR ventriculomegal* OR Ventriculoperitoneal Shunt OR Ventriculo peritoneal Shunt OR Cerebrospinal Fluid Shunt OR Cerebro Spinal Fluid Shunt OR ((VP OR CSF) AND shunt*) OR intracranial pressure

Ear or cranial nerve VIII

vestibulocochlear OR cochleovestibular OR statoacoustic OR cranial nerves[mh] OR cranial nerve[tiab]

Brain injury

intracranial pressure[mh] OR intracranial pressur* OR meningitis OR intracranial thrombosis[mh] OR cerebral thrombosis OR cerebral bleeding OR cerebral hemorrhage OR (cerebral[tiab] AND leukemi*[tiab]) OR brain abscess OR brain abscess* OR encephalopathy OR cerebral inflammation OR brain inflammation OR

encephalitis OR (brain AND fungal infection*) OR (vasculitis AND (brain OR central nervous system)) OR (graft versus host disease AND brain) OR hydrocephalus

Hormonal therapy

hormonal therap* OR hormone therap* OR Hormone Replacement Therapy[mh] OR Estrogen Replacement Therapy[mh] OR tamoxifen OR goserelin OR aromatase inhibitors[mh] OR aromatase inhibitor* OR anastrozole OR exemestane OR fadrozole OR formestane OR letrozole OR vorozole OR plomestane OR androgen deprivation therap* OR antiandrogen* OR antiestrogen* OR hormone receptor positive[tiab] OR estrogen receptor positive[tiab] OR fulvestrant OR flutamide OR faslodex

Tobacco smoking

(tobacco OR nicotine OR cigarette OR e-cigarette OR cigar OR pipe OR environmental tobacco smoke OR second hand smoke OR ETS OR waterpipe OR narghile OR arghile OR shisha OR hookah OR marijuana OR joint OR MJ[tiab] OR spice OR thc OR cannabis) AND (smoking OR smoke OR smoke*)

Subsequent neoplasm

Neoplasms, Radiation-Induced [Mesh] OR "Neoplasms, Radiation-Induced" OR "Radiation-Induced Neoplasms" OR "Neoplasm, Radiation-Induced" OR "Radiation Induced Neoplasms" OR "Radiation-Induced Neoplasm" OR "Radiation Induced Cancer" OR "Cancers, Radiation-Induced" OR "Radiation Induced Cancer" OR "Radiation-Induced Cancers" OR "Cancer, Radiation-Induced" OR "Cancer, Radiation Induced" OR Neoplasm, Second Primary [Mesh] OR "Neoplasms, Second Primary" OR "Neoplasm, Second Primary" OR "Second Primary Neoplasms" OR "Metachronous Second Primary Neoplasms" OR "Neoplasms, Metachronous" OR "Second Malignancy" OR "Malignancies, Second" OR "Malignancy, Second" OR "Second Malignancies" OR "Second Neoplasm" OR "Neoplasm, Second" OR "Neoplasms, Second" OR "Second Neoplasms" OR "Second Primary Neoplasms" OR "Metachronous Neoplasms" OR "Metachronous Neoplasm" OR "Therapy Associated Neoplasms' OR "Neoplasms, Treatment-Related" OR "Neoplasms, Treatment Related" OR "Treatment-Related" OR "Neoplasms, Treatment Related" OR "Treatment-Related" OR "Neoplasms, Treatment Related" OR "Treatment Related Neoplasm" OR "Therapy-Related Neoplasms" OR "Therapy Related Neoplasms" OR "Treatment-Associated Neoplasms" OR "Treatment Associated Neoplasms" OR "Treatment-Related Neoplasms" OR "Treatment Related Neoplasms" OR "Neoplasms, Therapy-Related" OR "Neoplasm, Therapy-Related" OR "Neoplasms, Therapy Related" OR "Therapy Related Neoplasm" OR "Therapy Associated Cancer" OR "Cancer, Therapy-Associated" OR "Therapy Associated Cancer" OR "Therapy-Related Cancer" OR "Cancer, Therapy-Related" OR "Cancers, Therapy-Related" OR "Therapy Related Cancer" OR "Therapy-Related Cancers" OR "Treatment-Related Cancer" OR "Cancer, Treatment-Related" OR "Cancers, Treatment-Related" OR "Treatment Related Cancer" OR "Treatment Related Cancers" OR "Treatment-Associated Cancer" OR "Cancer, Treatment-Associated" OR "Treatment Associated Cancer" OR "Treatment-Associated Cancers" OR "Cancer, Second Primary" OR "Cancers, Second Primary" OR "Second Primary Cancer" OR "Second Primary Cancers" OR "Second Primary Cancers" OR "Second Primary Cancers" OR "Second Primary Cancers" OR "Second Primary" OR "Second Primary Cancer" OR "Second Primary Cancer OR " Cancer" OR "Cancer, Second" OR "Cancers, Second" OR "Second Cancers" OR "Neoplasms, Radiation effects" OR "second primary malignancy" OR "second primary malignancies" OR "second malignant neoplasm" OR "second malignant neoplasms" OR "SMN" OR "second neoplasm" OR "second neoplasms" OR "secondary breast cancer" OR "subsequent malignant neoplasm" OR "subsequent malignant neoplasms" OR "subsequent neoplasm" OR "subsequent neoplasms" OR "second malignancy" OR "new malignancy" OR "new malignancies" OR "subsequent primary malignancy" OR "subsequent primary malignancies" OR "subsequent primary neoplasm" OR "subsequent primary neoplasms" OR "subsequent primary tumor" OR "subsequent primary tumors" OR "subsequent malignancy" OR "subsequent malignancies" OR "subsequent tumor" OR "subsequent tumors" OR "secondary cancer" OR "secondary neoplasm" OR "secondary malignancy" OR "secondary tumor" OR "secondary cancers" OR "secondary neoplasms" OR "secondary malignancies" OR "secondary tumors" OR "secondary primary malignancy" OR "second tumor" OR "second tumors" OR "second primary tumor" OR

"second primary tumors" OR "second malignant tumor" OR "second malignant tumors" OR "subsequent malignant tumors"

Breast cancer

breast cancer OR breast cancers OR breast neoplasm OR breast neoplasms OR breast neoplasm* OR mamma carcinoma OR mamma carcinomas OR mammary gland carcinoma or mammary gland carcinomas OR Neoplasm, Breast OR Neoplasms, Breast OR Tumors, Breast OR Breast Tumors OR Breast Tumor OR Tumor, Breast OR Mammary Carcinoma, Human OR Carcinoma, Human Mammary OR Carcinomas, Human Mammary OR Human Mammary Carcinoma OR Mammary Neoplasms, Human OR Human Mammary Neoplasms OR Neoplasms, Human Mammary OR Neoplasms, Human Mammary OR Neoplasms, Human Mammary OR Mammary Neoplasm, Human OR Cancer, Breast OR Cancer of the Breast OR Cancer of Breast

Glioma

(Glioma[mh] OR glioma OR gliomas OR glioma*)

Meningioma

(Meningeal neoplasms[mh] OR meningioma OR meningiomas OR meningioma*)

Other CNS neoplasms

(Pituitary neoplasms[mh] OR pituitary tumor OR pituitary tumors OR neurilemmoma[mh] OR neurilemmoma OR neurilemmoma* OR schwannoma OR schwannomas OR schwannoma* OR craniopharyngioma OR craniopharyngioma* OR pinealoma[mh] OR pineal tumor OR pineal tumors OR choroid plexus neoplasms[mh] OR choroid plexus tumor OR choroid plexus tumors OR supratentorial neoplasms[mh] OR supratentorial tumor OR supratentorial tumors OR opticus glioma OR opticus gliomas OR opticus glioma* OR optic nerve glioma[mh] OR optic nerve glioma OR optic nerve gliomas OR optic nerve glioma* OR medulloblastoma[mh] OR medulloblastoma OR medulloblastomas OR medulloblastoma* OR pilocytic astrocytomas OR oligodendroglioma[mh] OR oligodendroglioma OR oligodendroglioma* OR ganglioglioma* OR ganglioglioma* OR gangliogliomas OR ganglioglioma* OR ependymoma*)

Colorectal cancer

colonic neoplasms[MeSH Terms] OR cecal neoplasms[MeSH Terms] OR colorectal neoplasms[MeSH Terms] OR rectal neoplasms[MeSH Terms] OR ((colonic[tiab] OR colon[tiab] OR large intestine[tiab] OR intestinal[tiab] OR intestinal[tiab] OR cecal[tiab] OR cecal[tiab] OR cecum[tiab] OR caecum[tiab] OR colorectal[tiab] OR rectum[tiab] OR rectal[tiab] OR digestive[tiab] OR bowel[tiab] OR gut[tiab]) AND (cancer[tiab] OR cancers[tiab] OR tumors[tiab] OR tumours[tiab] OR neoplasms[tiab] OR carcinomas[tiab]))

Coronary artery disease

angina OR angina pectoris[mh] OR coronary artery disease OR coronary artery disease[mh] OR myocardial infarction OR myocardial infarction[mh] OR heart attack[tiab] OR heart arrest[mh] OR cardiac arrest OR ischemic heart disease OR ischaemic heart disease OR myocardial ischemia [mh] OR ischemic cardiomyopath* OR ischaemic cardiomyopath* OR coronary ischemia [tiab] OR coronary ischaemia [tiab] OR atherosclerotic heart disease[tiab] OR (coronary AND (vasculopathy OR thrombosis OR occlusion[tiab] OR plaque[tiab] OR occlusive disease* OR atherosclerosis OR artery disease* OR atherosclerotic disease* OR artery calcification* OR angiogram OR angioplasty OR artery bypass OR arteriosclerosis OR aneurysm)) OR coronary angiography[mh] OR angioplasty, balloon, coronary [mh] OR coronary balloon angioplasty OR coronary artery bypass[mh] OR coronary aneurysm[mh] OR (cardiac AND (atherosclerosis OR atherosclerotic*))

Thyroid cancer

(Thyroid carcinoma OR thyroid cancer OR papillary thyroid carcinoma OR follicular thyroid carcinoma OR papillary thyroid cancer OR follicular thyroid cancer OR thyroid neoplasm OR thyroid neoplasms OR non-medullary thyroid carcinoma OR non-medullary thyroid cancer OR thyroid gland OR thyroid adenoma OR thyroid nodule) NOT graves[tiab]

Thyroid ultrasound

Ultrasonography OR ultrasound OR thyroid ultrasound OR thyroid ultrasonography OR thyroid gland/ultrasonography OR thyroid neoplasms/ultrasonography OR thyroid neoplasms/ultrasonography OR neck palpation OR clinical examination OR imaging[tiab] OR diagnosis[sh] OR ultrasonography[sh]

Thyroid cancer diagnostic test

((Mibi OR pertechnate OR pertechnetate OR methoxyisobutylisonitrile) AND (scan OR scintigraphy OR scintigraphy OR scintigraphy OR scintigraphy OR spectron or spec

Thyroid cancer diagnostic test complications

Complications OR complication rate OR bleeding OR haemorrhage OR hemorrhage OR haematomas OR haematoma OR hematomas OR swelling OR infection OR recurrent nerve injury OR tracheal puncture OR vasovagal reaction OR dysphagia

Impaired spermatogenesis

spermatogenesis OR gonadal disorder OR spermiogenesis OR spermatocytogenesis OR spermatogenic failure OR azoospermia OR oligospermia OR asthenozoospermia OR teratozoospermia OR oligoasthenoteratozoospermia OR dysspermia OR normozoospermic OR semen OR semen analysis[text] OR semen quality[text] OR sperm OR sperm count OR sperm motility OR spermatozoa OR progeny OR offspring OR posterity OR fertility OR infertility OR subfertility OR reproduction OR fertilization OR conception OR paternity OR fatherhood OR parenthood OR pregnancy outcome OR fertile OR infertile OR subfertile OR sperm maturation OR aspermia OR spermatozoon abnormality

Testosterone deficiency

Androgen hormone insufficiency OR leydig cell OR cells, leydig failure OR testicular interstitium cell failure OR testicular failure OR gonadal failure OR hypogonadism OR low testosterone OR testosterone deficiency OR androgen deficiency OR low testosterone* OR hypogonadism* OR leydig cell*

Sexual dysfunction

erectile dysfunction OR dysfunction, erectile OR impotence OR male sexual impotence OR impotence, male sexual OR sexual impotence, male OR male impotence OR impotence, male OR impotence male* OR male sexual impotence* OR erectile dysfunction* OR premature ejaculation OR ejaculation, premature OR ejaculations, premature OR premature ejaculations OR ejaculation praecox OR praecox, ejaculation OR ejaculation* OR delayed ejaculation OR retarded ejaculation OR ejaculation dysfunction OR ejaculation dysfunction* OR sexual dysfunctions OR sexual dysfunction OR sexual disorder OR sex disorders OR sexual disorder* OR sexual dysfunction* OR male sexual dysfunction*

Obstructive azoospermia

spermatogenesis OR gonadal disorder OR spermiogenesis OR spermatocytogenesis OR spermatogenic failure OR azoospermia OR oligospermia OR asthenozoospermia OR teratozoospermia OR oligoasthenoteratozoospermia OR dysspermia OR normozoospermic OR semen OR semen analysis[text] OR

semen quality[text] OR sperm OR sperm count OR sperm motility OR spermatozoa OR progeny OR offspring OR posterity OR fertility OR infertility OR subfertility OR reproduction OR fertilization OR conception OR paternity OR fatherhood OR parenthood OR pregnancy outcome OR fertile OR infertile OR subfertile OR sperm maturation OR aspermia OR spermatozoon abnormalit

Quality and yield of sperm

Semen analysis OR semen analysis[mh] OR DNA fragmentation[mh] OR (DNA fragmentation AND sperm) OR sperm count[mh] OR sperm count OR (sperm AND yield)

FSH

Follicle stimulating hormone OR FSH OR follicle-stimulating hormone OR follitropin OR gonadotrope OR luteinizing hormone OR follicle stimulating hormon*

Inhibin B

Inhibin OR inhibins, testicular OR testicular inhibins OR inhibin, testicular OR testicular inhibin OR inhibin* OR inhibin B OR inhib

LH

Luteinizing hormone OR LH OR hormone, luteinizing OR interstitial cell-stimulating hormone OR interstitial cell stimulating hormone OR ICSH OR luteinizing hormon* OR LH*

Testosterone

Testosterone OR testosterone OR 17-beta-Hydroxy-4-Androsten-3-one OR testosterone*

Premature ovarian insufficiency

premature menopause* OR early menopause* OR menopausal status* OR ovarian failure* OR premature ovarian failure* OR acute ovarian failure* OR imminent ovarian failure* OR ovarian insufficiency* OR ovarian function* OR ovarian damage* OR Gonadotropin-Resistant Ovary Syndrome* OR Female Genital Diseases* OR Female infertility* OR primary ovarian insufficiency* OR gonadotoxicity OR gonado toxicity* OR gonadal damage* OR hypergonadotropic amenorrhoea* OR gonad dysfunction* OR gonadal function* OR gonadal effects* OR ovarian reserve* OR gonadal hormone deficiency*

FSH, AMH, AFC

Follicle Stimulating Hormone, Human* OR Follicle Stimulating Hormone* OR Gonadotrophs OR Estradiol OR Estrogen OR Anti-Mullerian Hormone* OR Ovarian Follicle* OR Antral Follicle Count* OR AFC

Live birth

Live birth [mh] OR live birth OR live birth* OR live-birth* OR progeny OR offspring OR posterity OR ovarian reserve OR fertile OR fertilization OR fertility OR birth OR parturition OR childbirth OR live birth[mh] OR reproduction[tiab] OR reproduction* OR reproduction[mh:noexp] OR ovulation[mh] OR gravidity OR term birth[mh] OR term birth [tiab] OR ((full term[tiab] OR term) AND (birth[tiab] OR births[tiab]))

Ototoxicity

deafness OR hearing loss OR Loss, Hearing OR hearing disorder OR hearing disorders OR auditory OR hearing impairment OR hearing impairment* OR audiologic OR audiometry OR audiometr* OR audiogram OR audiography OR ototoxicology OR ototoxic* OR hypoacusis OR hypoacuses OR

hypoacus* OR ototoxicity OR deaf* OR cochleotoxicity OR tinnitus[mh] OR tinnitus[tiab] OR tinnitus* OR unable to hear

Ototoxicity diagnostic test

pure tone audiometry OR audiometry, pure-tone[mh] OR extended high frequency audiometry[tiab] OR speech audiometry OR audiometry, speech[mh] OR otoscopy OR otoscopic* OR otologic* OR diagnostic techniques, otological[mh] OR ear microscopy OR speech discrimination test OR speech discrimination tests[mh] OR speech reception threshold test[mh] OR speech reception threshold test OR evoked potentials, auditory, brain stem[mh] OR acoustic evoked brain stem potential* OR auditory brain stem evoked response OR auditory brain stem respons* OR brain stem auditory evoked potential OR brainstem auditory evoked potential OR brainstem auditory evoked potential* OR auditory brainstem response OR auditory brainstem respons* OR otoacoustic emissions, spontaneous[mh] OR otoacoustic emission OR otoacoustic emissions OR tinnitus evaluation OR acoustic impedance test OR Audiometry[mh] OR tympanometry

Ototoxicity interventions

hearing aid OR hearing aids OR hearing aids[mh] OR ear mold OR earmold OR ear mould OR (cochlear AND (implant OR implantation OR implants OR prosthes*)) OR auditory prosthes* OR (tinnitus AND (mask OR masking OR mask*)) OR acoustic stimulation[mh] OR acoustic stimulation OR (acoustic AND (stimulation* OR implant OR implants)) OR (mainstreaming OR "mainstreaming (education)" [Mesh] OR persons with hearing impairments OR (acoustics AND classroom) OR speech therapy[mh] OR language therapy[mh] OR ((speech OR remedial) AND therapy[tiab]) OR sign language OR aural rehabilitation OR ((aural OR auditory) AND rehabilitation[tiab]) OR vocational guidance OR special education[tiab] OR education, special[mh] OR acoustic stimulation OR tinnitus retraining therapy OR (tinnitus AND (training [tiab] OR management[tiab])) OR ((counseling[tiab] OR counselling[tiab]) AND (structured[tiab] OR education[tiab])) OR speech perception OR speech acoustics OR auditory threshold OR linguistics) OR ((vocational AND (rehabilitation OR accommodations OR modifications)) OR noise management OR noise protection OR ear protective device OR ear protective devices[mh] OR (hearing AND (protection OR protector OR conservation OR impairment OR preservation)) OR hearing loss prevention OR amplification system OR hearing assistive technology)

Hepatic toxicity

liver fibrosis OR liver cirrhosis OR liver disease OR liver diseases OR liver diseases OR liver dysfunction OR liver dysfunctions OR liver damage OR liver failure OR liver enzyme[all fields] OR liver enzymes[all fields] OR liver enzym* OR liver toxicity OR liver disfunction OR radiation-induced liver disease OR radiation induced liver disease OR RILD OR liver function test OR liver function tests OR liver insufficiency OR Hepatic Cirrhosis OR Cirrhoses, Hepatic OR Cirrhosis, Hepatic OR Hepatic Cirrhoses OR Cirrhosis, Liver OR Cirrhoses, Liver OR Liver Cirrhoses OR Fibrosis, Liver OR Fibroses, Liver OR Liver Fibroses OR Disease, Liver OR Diseases, Liver OR Dysfunction, Liver OR Dysfunctions, Liver OR Liver Dysfunctions OR Function Test, Liver OR Function Tests, Liver OR Liver Function Test OR Test, Liver Function OR Tests, Liver Function OR Insufficiency, Hepatic OR Liver Insufficiency OR Insufficiency, Liver OR hepatic dysfunction OR hepatic dysfunctions OR hepatic cirrhosis OR hepatic failure OR hepatic function[all fields] OR liver function[all fields] OR radiation hepatitis OR hepatitis irradiation OR impaired liver function OR hepatic fibrosis OR hepatic fibroses OR drug induced hepatitis OR toxic hepatitis OR hepatitides OR ASAT OR ALAT OR SGPT OR SGOT OR GGT OR alanine transaminase OR Transaminase, Alanine OR Glutamic-Alanine Transaminase OR Glutamic Alanine Transaminase OR Transaminase, Glutamic-Alanine OR Alanine-2-Oxoglutarate OR Aminotransferase OR Alanine 2 Oxoglutarate Aminotransferase OR Aminotransferase, Alanine-2-Oxoglutarate OR Alanine Aminotransferase OR Aminotransferase, Alanine OR Glutamic-Pyruvic Transaminase OR Glutamic Pyruvic Transaminase OR Transaminase, Glutamic-Pyruvic OR gamma Glutamyltransferase OR Glutamyl Transpeptidase OR

Transpeptidase, Glutamyl OR GGTP OR gamma-Glutamyl Transpeptidase OR Transpeptidase, gamma-Glutamyl OR gamma Glutamyl Transpeptidase OR gammaglutamyltransferase OR Aspartate Aminotransferases OR Aminotransferases, Aspartate OR Aspartate Apoaminotransferase OR Apoaminotransferase, Aspartate OR Aspartate Transaminase OR Transaminase, Aspartate OR Glutamic-Oxaloacetic Transaminase OR Glutamic Oxaloacetic Transaminase OR Transaminase, Glutamic-Oxaloacetic OR L-Aspartate-2-Oxoglutarate Aminotransferase OR Aminotransferase, L-Aspartate-2-Oxoglutarate OR L Aspartate 2 Oxoglutarate Aminotransferase OR Aspartate Aminotransferase OR Aminotransferase, Aspartate OR Glutamate-Aspartate Transaminase OR Glutamate Aspartate Transaminase OR Transaminase, Glutamate-Aspartate OR Serum Glutamic-Oxaloacetic Transaminase OR Glutamic-Oxaloacetic Transaminase, Serum OR Serum Glutamic Oxaloacetic Transaminase OR Transaminase, Serum Glutamic-Oxaloacetic OR hepatotoxicity OR hepatotoxic OR hepatotoxic* OR Veno-occlusive disease OR VOD OR Veno occlusive disease OR hepatic veno-occlusive disease OR Disease, Hepatic Veno-Occlusive OR Hepatic Veno-Occlusive Diseases OR Sinusoidal Obstruction Syndrome OR Syndrome, Sinusoidal Obstruction OR Hepatic Veno Occlusive Disease OR Veno-Occlusive Disease, Hepatic OR Veno Occlusive Disease, Hepatic OR bilirubin OR bilirubins OR bilirubin* OR Bilirubin IX alpha OR Bilirubin, (4E)-Isomer OR Bilirubin, (4E,15E)-Isomer OR Hematoidin OR Bilirubin, Disodium Salt OR Disodium Salt Bilirubin OR Bilirubin, Monosodium Salt OR Monosodium Salt Bilirubin OR delta-Bilirubin OR delta Bilirubin OR Bilirubin, (15E)-Isomer OR Bilirubin, Calcium Salt OR Calcium Salt Bilirubin OR Salt Bilirubin, Calcium OR Calcium Bilirubinate OR Bilirubinate, Calcium OR albumin OR albumins OR albumin* OR prothrombin OR prothrombins OR prothrombin* OR Factor II OR Blood Coagulation Factor II OR Differentiation Reversal Factor OR Factor, Differentiation Reversal OR Coagulation Factor II OR Factor II, Coagulation OR II, Coagulation Factor OR Alkaline phosphatase

Iron overload

(iron overload OR hemosiderosis OR siderosis OR heamosiderosis OR haemosiderosis) OR (Hemosideroses OR Overload, Iron)

Hypothalamic-pituitary dysfunction general

hypothalamic diseases OR pituitary diseases OR hypopituitarism OR pituitary hormones, anterior[mh] OR anterior pituitary hormone deficiency OR endocrine system diseases[mh] OR endocrine disorder OR hypophyseal deficiency OR (pituitary AND (dysfunction* OR hypofunction)) OR panhypopituitarism OR combined pituitary hormone deficiency OR multiple pituitary hormone deficiency OR (("endocrine system"[MeSH Terms] OR (endocrine AND system) OR endocrine system OR endocrine) AND ("complications"[Subheading] OR complications OR sequelae))

Growth hormone deficiency

(growth hormone OR GHD OR GH) AND (deficien* OR disorder OR impairment OR deviation)

Growth hormone deficiency diagnostic tests

insulin-like growth factor I OR insulin-like growth factor 1 OR IGF-I OR IGF-I OR insulin like growth factor I OR insulin like growth factor 1 OR somatomedin C OR insulin-like somatomedin peptide I OR insulin like somatomedin peptide I OR insulin like somatomedin peptide 1 OR insulin-like growth factor binding protein 3 OR insulin like growth factor binding protein 3 OR IGF-binding protein 3 OR IGF binding protein 3 OR IGFBP-3 OR ((growth hormone stimulation OR GH stimulation) AND (test[tiab] OR testing)) OR glucagon stimulation test[tiab] OR arginine test[tiab] OR clonidine test [tiab] OR ((GHRH OR growth hormone releasing hormone) AND (test[tiab] OR assay)) OR insulin tolerance test[tiab] OR ITT OR (propranolol AND (I dopa OR levodopa)) OR human growth hormone/blood[mh] OR growth hormone overnight

Treatment

("therapy"[Subheading] OR "therapy"[All Fields] OR "treatment"[All Fields] OR "therapeutics"[MeSH Terms] OR "therapeutics"[All Fields] OR treatment OR intervention*)

Central hypothyroidism

((Thyroid stimulation hormone OR TSH OR Thyrotropin releasing hormone OR TRH OR thyroxine OR tri iodothyronine OR triiodothyronine) AND (deficiency OR disease)) OR (thyroid hormones[mh] AND (deficiency OR disease)) OR thyroid diseas* OR hypothyroidism[mh] OR hypothyroidism* OR thyroxine OR levo-thyroxine OR tri-iodothyronine

Central hypothyroidism diagnostic tests

FT4 OR TSH OR FT3 OR FT3 FT4 ratio* OR (TRH AND (test[tiab] OR surge OR value* OR peak OR nadir OR measurement OR assay OR concentration*)) OR Thyroxine/blood[mh] OR thyrotropin/blood[mh] OR thyroid hormones/blood[mh]

Central hypocortisolism

((adrenocorticotropic hormone[mh] OR adrenocorticotropic hormone* OR ACTH OR hydrocortisone) AND (insufficiency OR deficiency)) OR ((adrenal OR hydrocortisone) AND (insufficiency OR deficiency OR crisis)) OR hypocortisolism OR hypocortisolism* OR Addison disease[mh]

Central hypocortisolism diagnostic tests

Cortisol OR hydrocortisone OR ACTH OR adrenocorticotropic hormone OR morning glucose OR ITT OR insulin tolerance test[tiab] OR synacthen test[tiab] OR synacthen tests OR metyrapone

Hypogonadotropic hypogonadism

hypogonadotropic hypogonadism OR hypogonadism[mh] OR (infertility AND pituitary) OR ((follicle stimulating hormone OR FSH OR gonadotropin* OR luteinizing hormone OR LH OR estrogen OR testosterone) AND deficiency) OR ((puberty OR menarche OR sexual development) AND delay*) OR ((gonadotropin OR estrogen OR testosterone) AND (deficiency OR insufficiency))

Central Precocious puberty

(precocious AND (sexual OR puberty pubarche OR menarche)) OR CPP OR puberty, precocious[mh] OR premature puberty OR breast development puberty OR skeletal maturation OR epiphyseal plate closure OR bone age* OR skeletal age*) OR thelarche

Hypogonadotropic hypogonadism and central Precocious puberty diagnostic tests

(tanner[tiab] AND (scale* OR stage* OR staging)) OR bone age* OR skeletal age* OR ((follicle stimulating hormone OR FSH OR gonadotropin* OR luteinizing hormone OR LH OR estrogen OR testosterone OR LHRH OR gonadotropin-releasing hormone OR GnRH OR triptorelin OR GnRHa OR Gonadotropin-Releasing Hormone Agonist) AND (assay OR level OR test[tiab])) OR pelvic ultrasound OR gonadotropin-releasing hormone/blood[mh] OR (growth velocity AND puberty)

Hypertension

Hypertension[mh] OR hypertension[tiab] OR blood pressure[mh] OR blood pressure[tiab] OR arterial pressure[mh]

Hyperlipidemia

hyperlipidemias[mh] OR hyperlipidemia* OR "triglycerides" [MeSH Terms] OR tryglyceride* OR cholesterol OR Cholesterol, HDL[mh] OR Cholesterol, LDL[mh] OR hypertriglyceridemia[mh] OR hypertriglyceridemia* OR hypercholesterolemia OR hypercholesterolemia* OR hypercholesterolemia* OR hypercholesterolemia* OR hypercholesterolemia* OR hypercholesterolemia* OR hydroxymethylglutaryl-coa reductase inhibitors[mh] OR lipoproteins[mh] OR lipoprotein*

Impaired glucose metabolism

diabetes mellitus[tiab] OR diabetes mellitus[mh] OR prediabetic state[mh] OR prediabetes[tiab] OR fasting glucose OR glucose tolerance[tiab] OR glucose intolerance[mh] OR glucose intolerance[tiab] OR insulin resistance[mh] OR insulin resistance[tiab] OR hyperinsulinaemia OR insulin sensitivit* OR homeostasis model assessment[tiab] OR HOMA OR hemoglobin a, glycosylated[mh] OR HbA1c OR glycosylated hemoglobin a OR glycated haemoglobin[tiab] OR hypoinsulinaemia

Obesity

Obesity[mh] OR obesity[tiab] OR obese[tiab] OR overweight[mh] OR overweight[tiab] OR body mass index[mh] OR body mass index OR BMI OR waist circumference OR waist hip ratio OR waist height ratio OR fat body[mh] OR fat body OR body fat[tiab] OR adipose tissue[mh] OR adipose

Metabolic syndrome

metabolic syndrome OR metabolic syndrome X[mh] OR insulin resistance[mh] OR insulin resistance[tiab]

Nephrotoxicity

glomerular filtration rate OR GFR OR Filtration Rate, Glomerular OR Filtration Rates, Glomerular OR Glomerular Filtration Rates OR Rate, Glomerular Filtration OR Rates, Glomerular Filtration OR glomerular OR glomerul* OR tubular OR tubula* OR renal tubular acidosis OR RTA OR Acidosis, Renal Tubular OR Renal Tubular Acidosis, Type I OR Type I Renal Tubular Acidosis OR Distal Renal Tubular Acidosis OR Acidosis, Renal Tubular, Type I OR Classic Distal Renal Tubular Acidosis OR Renal Tubular Acidosis, Distal, Autosomal Dominant OR Renal Tubular Acidosis, Type II Acidosis, Renal Tubular, Type II OR Renal Tubular Acidosis, Proximal, with Ocular Abnormalities OR Proximal Renal Tubular Acidosis OR Type II Renal Tubular Acidosis OR renal acidosis OR renal insufficiency OR Renal Insufficiencies OR Kidney Insufficiency OR Insufficiency, Kidney OR Kidney Insufficiencies OR microalbuminuria OR microalbumin* OR hypophosphatemia OR hypophosp hypomagnes* OR hypomagnesemia OR hypomagnesaemia OR magnesium OR phosphorus OR Hyponatremia OR Hyponatremias OR hyponatraemia OR hyponatraemias OR hyponatraem* OR hyponatraem* OR Potassium OR Potassium Ion Level OR Ion Level, Potassium OR Level, Potassium Ion OR Hypokalemia OR Hypokalemias OR Hypopotassemia OR Hypopotassemias OR hypokalemic OR hypokalem* OR hypokalaemic OR hypokalaem* OR Hypocalcemia OR hypocalcemias OR hypocalciuria OR hypocalciuri* OR hypocalcem* OR hypocalce* OR "hypocarbia" OR Proteinuria OR proteinurias OR proteinuri* OR albuminuria OR albuminurias OR albuminuri* OR Aminoaciduria OR Renal Aminoaciduria OR Renal Aminoacidurias OR Aminoacidurias, Renal OR Aminoaciduria, Renal OR aminoacidur* OR Glucosuria OR glucosurias OR glucosur* OR glycosuria OR glycosurias OR glycosuria, renal OR low molecular weight OR LMW OR alpha 1 microglobulin OR a1 microglobulin OR beta 2 microglobulin OR b2 microglobulin OR 2-Microglobulin, beta OR creatinine OR Creatinine Sulfate Salt OR Salt, Creatinine Sulfate OR Sulfate Salt, Creatinine OR 60-27-5[rn] OR inulin OR 9005-80-5[rn] OR "(51) Cr EDTA" OR 51chromium edetic acid OR "(99) Tc DTPA" OR Tc DTPA OR 65454-61-7[rn] OR Technetium Tc 99m Pentetate OR (99m)Tc-DMSA OR 99mTc(V)DMSA OR DMSA OR dimercaptosuccinic acid OR Technetium Tc 99m Dimercaptosuccinic Acid OR 65438-08-6[rn] OR 99Tc-Succimer OR 99Tc Succimer OR 99mTc-Dimercaptosuccinate OR 99mTc Dimercaptosuccinate OR renal scan OR "kidney size" OR cystatin c OR gamma-Trace OR gamma Trace OR Cystatin 3 OR cystatins OR cystatin* OR renal failure OR kidney failure OR Failure, Kidney OR Failures, Kidney OR Kidney Failures OR Failure, Renal OR Failures, Renal OR Renal Failures OR renal plasma flow OR Plasma Flow, Renal OR Flow, Renal Plasma OR RPF OR ERPF OR Renal clearance OR reabsorption OR re-absorption OR nephrotoxicity OR nephrotox* OR rickets*

Osteoporosis

"Osteoporosis" [Mesh] OR "Fractures, Bone" [Mesh] OR "Bone Density" [Mesh] OR "Bone Remodeling" [Mesh] OR "Bone Demineralization, Pathologic" [Mesh] OR osteoporos* [tiab] OR osteopenia* [tiab] OR bone mineral densit* [tiab] OR bone densit* [tiab] OR bone loss* [tiab] OR bone health [tiab] OR bone turnover [tiab] OR bone morbidit* [tiab] OR bone fragil* [tiab] OR bone content [tiab] OR bone mass* [tiab] OR bone deformit* [tiab] OR fracture* [tiab] OR bone* [tiab]

Osteonecrosis

"Osteonecrosis" [Mesh] OR osteonecros*[tiab] OR avascular necros*[tiab] OR bone necros*[tiab] OR bone aseptic necros*[tiab]

DXA

DXA OR DEXA OR absorptiometry, photon[mh] OR photon absorptiometry OR dual energy x ray absorptiometry OR bone density[mh] OR bone densitometry

Obstetric problems

hypertension, pregnancy-induced[mh] OR pregnancy induced hypertension OR PIH OR pre-eclampsia[mh] OR pre eclampsia OR hellp syndrome[mh] OR HELLP OR placentation[mh] OR placental development OR ((placenta AND (accreta OR increta OR previa)) OR placenta accreta[mh] OR placenta previa[mh] OR vasa previa OR fetal malpresentation OR fetal malposition OR pregnancy outcome[mh] OR pregnancy complications[mh] OR breech presentation[mh] OR breech OR obstetrics labor complication[mh] OR obstetric complication* OR transverse lie[tiab] OR shoulder presentation[tiab] OR cord presentation[tiab] OR birth weight[mh] premature birth[mh] OR fetal growth retardation[mh] OR obstetric labor, premature[mh] OR fetal development[mh] OR ((fetal OR intrauterine) AND (growth OR development OR restriction)) OR small for gestational age[tiab] OR SGA OR birth weight[mh] OR low birth weight OR premature birth OR infant, premature[mh] OR premature delivery[tiab] OR immature delivery OR dysmature OR miscarriage OR spontaneous abortion OR abortion OR abortion, spontaneous[mh] OR postpartum hemorrhage[mh] OR post partum hemorrhage OR hemorrhage post-partum OR (manual removal[tiab] AND placenta) OR placenta, retained[mh] OR lactation OR lactat* OR breast feeding[mh] OR breast feeding[tiab] OR breastfeeding* OR ((vaginal ultrasonography OR pulsatility index[tiab]) AND (gestation OR pregnancy)) OR stillbirth

Pulmonary disease

Pulmonary Fibrosis OR lung fibrosis OR (scarring AND (lung OR lungs*)) OR interstitial lung disease OR acute respiratory distress syndrome [tiab] OR ARDS OR respiratory distress syndrome OR shock lung[tiab] OR pneumonia OR COP[tiab] OR pneumonitis[tiab] OR pulmonitis[tiab] OR (lung AND (cancer OR carcinoma OR tumor)) OR lung neoplasms[mh] OR (lung AND (infection OR disease)) OR lung diseases[mh] OR (chest wall AND (abnormalit* OR disease)) OR kyphoscoliosis OR fibrothorax OR bronchitis OR bronchiectasis OR emphysema OR fibroelastosis OR Bronchiolitis OR BOS[tiab] OR BOOP OR cryptogenic organizing pneumonia[mh] OR cryptogenic organizing pneumonia[tiab] OR pulmonary disease OR pulmonary disease, chronic obstructive[mh] OR COPD OR pulmonary complications OR OSA OR respiratory tract diseases[mh] OR respiratory disease* OR low infectious respiratory disease OR respiratory defect OR apnea OR asthma

Pulmonary symptoms

dyspnea OR cough OR mucus OR sputum OR hypoxia OR oygen requirement[tiab] OR exercise intolerance[tiab] OR respiratory sounds[mh] OR wheeze OR wheeze* OR breathlessness[tiab] OR shortness of breath OR chest pain OR chest discomfort[tiab] OR snore OR snoring OR hemoptysis OR oxygen requirement

Pulmonary diagnostic tests

respiratory function tests[mh] OR (function test AND (lung OR pulmonary OR respiratory)) OR spirometry OR bronchospasmolysis OR plethysmography OR DLCO OR diffusion capacity OR breath washout OR pulsoxymetry OR therapeutic irrigation[mh] OR broncho alveolar lavage[tiab] OR bronchoscopy OR blood gas analysis OR FEV1 OR forced expiratory volume OR LCI OR lung clearance index OR TLC OR total lung capacity OR FVC OR forced vital capacity OR PEF OR peak expiratory flow OR forced expiratory flow OR FEF OR maximum expiratory flow OR MEF OR KCO OR diffusion capacity OR maximal inspiratory pressure OR maximal expiratory pressure OR respiratory muscle pressure OR ((HR-CT OR MRI OR X-ray OR Biopsy OR lavage) AND (lung OR pulmonary OR chest OR thorax)) OR (transfer factor AND lung)

Fatigue

fatigue[mh] OR fatigue OR fatigu* OR tired[tiab] OR tiredness[tiab] OR tired* OR asthenia[mh] OR asthenia OR asthenia OR exhausted OR exhaust* OR loss of energy[tiab] OR energy loss[tiab] OR loss of vitality OR (vital* AND loss) OR weary[tiab] OR weariness[tiab] OR weakness OR apathy[mh] OR apath* OR lassitude[tiab] OR lethargy[mh] OR letharg* OR sleep OR sleep deprivation OR sleepiness[tiab] OR drowsy[tiab] OR drowsiness[tiab] OR chronic fatigue syndrome OR CFS OR (CF AND syndrome[tiab])

Screening

(screening[tiab] OR "diagnosis"[Subheading] OR "diagnosis"[All Fields] OR (tool OR tools) AND (diagnostic OR screening OR assessment) OR questionnaire OR test[tiab] OR measure[tiab] OR scale[tiab]

Validation

validation OR reliability OR validity OR sensitivity OR specificity OR psychometric* OR psychometrics[mh]

Employment

employment [MeSH terms] OR absenteeism [MeSH Terms] OR sick leave [MeSH Terms] OR rehabilitation, vocational [MeSH Terms] OR vocational guidance [MeSH Terms] OR occupation[MeSH Terms] OR occupation* OR return to work[mh] OR work performance[mh] OR employment OR unemploy* OR job*[tiab] OR vocational OR vocation* OR career OR income OR benefits use[tiab] OR socioeconomic status[tiab] OR insurance, disability[mh] OR disability insurance[tiab] OR poverty OR salaries and fringe benefits[mh] OR salaries and fringe benefits OR social security

Education

educational status[mh] OR education[mh] OR education* OR level of education

Mental health problems

depressive disorder[mh] OR depression OR depression* OR (mental AND (distress OR health OR disorder*)) OR mental disorders[mh] OR antidepressive agents[mh] OR antidepressant OR antidepressant* OR (psychological AND (outcome OR distress OR functioning OR stress*)) OR (psychosocial AND (outcomes OR impact OR challenge* OR issue* OR difficulties OR care OR need* OR profile OR functioning OR adjustment OR problem* OR health)) OR anxiety[mh] OR anxiety OR suicide OR suicide* OR (conduct AND (disorder* OR problem*)) OR behavioral symptoms[mh] OR post traumatic stress[tiab] OR PTSS

Mental health problems diagnostic tests

Brief Symptom Inventory[tiab] OR BSI OR SCL-90 OR Pediatric Symptom Checklist OR PSC OR "Checklist/methods"[MeSH Terms] OR "Checklist/standards"[MeSH Terms] OR

"Psychometrics/methods" [MeSH Terms] OR "Psychometrics/standards" [MeSH Terms] OR "Surveys and Questionnaires/methods"[MeSH Terms] OR "Surveys and Questionnaires/standards"[MeSH Terms] OR Child Behavior Checklist OR CBCL OR Emotion Thermometer[tiab] OR distress thermometer OR Depression anxiety stress scale[tiab] OR DASS OR hospital anxiety depression scale[tiab] OR HADS OR behavior assessment system[tiab] OR BASC OR ((Structured Clinical Interview[tiab] OR SCID) AND DSM) OR NIH PROMIS OR Pediatric Quality of Life Inventory[tiab] OR PedsQL OR Strengths Difficulties Questionnaire OR SDQ OR kidscreen OR SF-36 Health Survey[tiab] OR SF-12 Health Survey[tiab] OR Patient Health Questionnaire 9[tiab] OR PHQ 9[tiab] OR Child Depression Inventor* OR CDI OR (Beck Depression Inventor* AND Youth) OR BDI Youth OR beck anxiety inventor* OR BAI[tiab] OR ((Manifest Anxiety Scale[mh] OR Manifest Anxiety Scale) AND (children* AND revised)) OR RCMAS 2 OR (Multidimensional Anxiety Scale[tiab] AND children) OR MASC[tiab] OR State-Trait Anxiety Inventor* OR STAI OR phobia inventor* OR SPAI OR (Beck[tiab] AND Suicide Ideation) OR (BSS AND suicide) OR Suicidal Ideation Questionnaire OR (SIQ AND suicide) OR SNAP IV OR (Behavior Rating Inventor* AND Executive Function) OR (BRIEF AND executive function) OR Connors[tiab] OR Conduct Disorder Scale OR Disruptive behavior disorder rating scale OR ("attention deficit and disruptive behavior disorders" [MeSH Terms] AND rating scale) OR (DBD AND disruptive) OR Impact of Event Scale-Revised[tiab] OR IES-R OR Posttraumatic Stress Disorder Reaction Index[tiab] OR PTSD-RI OR (PTSD AND Reaction Index) OR Posttraumatic Diagnostic Scale[tiab] OR (PDS AND posttraumatic) OR Somatoform Dissociation Questionnaire OR SDQ-20 OR K-SADS-PL OR Psychiatric Status Rating Scales[mh]

Fertility preservation general

(preservation AND (sperm OR ovarian OR semen OR fertility OR oocyte)) OR (cryopreservation AND (sperm OR ovarian OR semen OR oocyte)) OR (vitrification AND (sperm OR ovarian OR semen OR oocyte)) OR ovarium transposition OR ovarian transposition OR Gonadotropin-releasing hormone OR GNRH OR (suppression AND (ovarian OR hormonal OR hormone OR gonadotropin OR gonadotrophin)) OR fertility preservation[mh]

Fertility preservation male

(preservation AND (sperm OR semen)) OR (cryopreservation AND (sperm OR semen)) OR (vitrification AND (sperm OR semen)) OR (suppression AND (hormonal OR hormone OR gonado)) OR fertility preservation[mh] OR (cryopreserv* AND (masturbation OR testicular tissue)) OR electroejaculation OR electro ejaculation OR EEJ OR testicular sperm extraction OR TESE[tiab] OR sperm banks[mh] OR sperm banking OR sperm bank* OR penile vibration OR penile vibrat* OR (spermatogonial AND (stem cell* OR stem cells[mh])) OR SSC* OR testicular shield* OR Testis/radiation effects[mh] OR Radiation Protection/methods[mh]

Fertility preservation female: ovarian, embryo and oocyte cryopreservation

ovarian cryopreservation OR embryo cryopreservation OR oocyte cryopreservation OR ((cryopreservation[mh] OR cryopreservation) AND ovarian[tiab]) OR ((cryopreservation[mh] OR cryopreservation) AND (female[mh] OR female*[tiab])) OR ((vitrification[mh] OR vitrification) AND (ovarian OR embryo OR oocyte)) OR ((freezing[mh] OR freezing) AND (ovarian OR embryo OR oocyte))

Fertility preservation female: in vitro maturation

in vitro techniques[mh] OR (in vitro AND maturation) OR in vitro maturation OR in vitro maturation* OR (in vitro AND (maturation OR techniques)) OR Reproductive Techniques, Assisted[mh]

Gonadotropin-releasing hormone (GnRH) analogues

gonadotropin-releasing hormone [MeSH Terms] OR (gonadotropin-releasing AND hormone) OR gonadotropin-releasing hormone OR (gonadotropin AND releasing AND hormone) OR gonadotropin releasing hormone OR gonadotropin-releasing hormone OR gonadotropin-releasing hormone OR gonadotropin-releasing hormone analogue* OR GnRH analogue* OR GnRHa OR GnRHa OR Luteinizing Hormone-Releasing Hormone OR LH-FSH Releasing Hormone OR LH-Releasing Hormone OR Gonadoliberin OR Gonadorelin

Immunomodulators

immunologic factors[MeSH Terms] OR immunomodulator* OR adjuvants, immunologic[MeSH Terms] OR AS101 OR sphingosine 1-phosphate OR sphingosine kinase* OR S1P OR biological response modifiers

Oral contraceptive pill

contraceptive OR contraceptive* OR contraceptive agents,female[mh] OR contraceptives,oral[mh] OR birth control pill OR birth control pill* OR hormonal contraception OR ((mini OR hormone) AND (pill OR pills))

Donor eggs

Donor, ovum[mh] OR Donors, ovum[mh] OR donor eggs OR oocyte donation

Surrogacy

Surrogacy OR surrogate pregnancy OR surrogate mother[mh]

Complications

Complication* OR side effect* OR side-effect* OR complications[sh] OR sequelae* OR adverse effect* OR adverse effect[sh]

Complications to pregnancy

((Preterm OR premature OR still) AND birth) OR stillbirth OR fetal death OR pregnancy outcome[mh] OR perinatal outcome OR gestational age[mh] OR premature birth[mh] OR stillbirth[mh] OR infant, low birth weight[mh] OR low birth weight OR gestational age[mh] OR gestational age* OR SGA OR fetal macrosomia[mh] OR fetal macrosomia* OR ((intrauterine growth OR fetal growth) AND retardation OR restriction) OR preeclampsia OR eclampsia

Cost-benefit

Cost benefit analysis[mh] OR cost benefit OR cost benefit OR costs* benefit OR cost effectiveness OR health care costs OR cost and cost analysis OR cost saving OR cost savings

Risk factors

(Risk factors OR predictors OR clinical characteristics OR patient characteristics OR clinical symptoms OR ultrasound features OR ultrasound characteristics OR age[tiab] OR gender OR sex OR family history OR stridor OR cough OR lymphadenectomy OR firm texture OR fixed mass OR hoarse voice OR pain OR shortness of breath OR dysphagia)

Attitudes, information needs, satisfaction

Attitudes[mh] OR belief* OR perception* OR attitude of health personnel[mh] OR choice behavior[mh] OR cooperative behavior[mh] OR decision making[mh] OR clinical decision making[mh] OR informed consent[mh] OR professional-family relations[mh] OR Physician-Patient Relations[mh] OR psychology[sh] OR choice behavio* OR clinical support technique* OR cognitive aspect* OR collaboration* OR compliant behavio* OR consensus OR informed consent* OR cooperative behavio* OR co-operative behavio* OR disput* OR disput*

OR doctor patient relation* OR doctor-patient relation* OR educational technology OR emotional aspect* OR health attitude* OR health education OR health information OR health literacy OR illness behavio* OR informed choice* OR informed decision* OR negotiati* OR nursing role* OR (nurse* AND role*) OR professional* patient* communic* OR patient professional communic* OR patient acceptance OR patient adherence OR patient attitude* OR patient compliance OR patient cooperation OR patient education OR patient education[mh] OR patient involvement OR patient non adherence OR patient non compliance OR patient nonadherence OR patient non-adherence OR patient noncompliance OR patient participation OR patient* preference* OR patient preference[mh] OR patient satisfaction OR patient satisfaction[mh:noexp] OR physician attitude OR physician patient relation* OR physician patient relation* OR professional patient disagreement* OR professional-family disagreement* OR professional-family relation* OR professional-patient disagreement* OR shared decision* OR sharing decision* OR staff attitude* OR treatment refusal* OR uncertainty OR health priorities[mh]

Barriers (social, legal, ethical, financial, religious, access) in general

((Barrier OR barrier*) AND (treatment* OR therapeutic[mh])) OR perceived barrier* OR ((barrier* OR obstacle* OR issue* OR impediment*) AND (social OR legal OR ethic* OR financ* OR cost* OR insurance OR resource* OR religious OR access OR patient* OR famil* OR institution* OR organization* OR hospital* OR cancer center* OR cancer centre* OR cultural OR education* OR psycholog* OR practical OR clinician* OR oncolog* OR nurs* OR fertility specialist* OR referral* OR referral and consultation[mh] OR health care professional*)) OR communication barriers[mh] OR communic* barrier* OR written communication[mh] OR verbal communication[mh] OR refusal participat* OR patient nonadherence OR patient non-adherence OR patient noncompliance

Decision tools, educational strategies, organizational strategies

decision making,computer assistance[mh] OR decision support OR decision support system OR (Decision AND (making OR tool* OR technique* OR management OR aid* OR toolkit* OR instrument* OR framework OR strateg*)) OR decision-aid* OR pamphlet* OR brochure* OR booklet* OR patient education[mh] OR teaching material[mh] OR educational strateg* OR organizational strateg* OR (online OR web-based OR educational OR organizational AND (decision-aid* OR decision aid* OR tool OR instrument OR framework OR strateg*))

Ethical considerations

Autonom* OR personal autonomy [mh] OR principle-based ethics[mh] OR beneficienc* OR maleficence OR informed consent OR declaration helsinki OR ethic* OR ethics[mh] OR ethic*[tiab] OR ethics[sh] OR ethics, medical[mh] OR medical oncology,ethics[mh]

Legal considerations

legal OR legal guardians[mh] OR social justice[mh] OR future death OR terminally ill OR postmortem paternity OR postmortem maternity OR posthumous sperm procurement OR legislation[pt] OR legal cases[pt] OR social control, formal[mh]

For **neurocognitive outcomes** the following search terms were used in MEDLINE (Ovid):

- exp Reaction Time/
- 2. *Mental Processes/
- 3. exp *Cognition Disorders/
- 4. exp Intelligence/
- 5. exp Psychomotor Performance/

- 6. ((neurocognitive or neuropsychological or neurodevelopmental or developmental or intellectual or mental) adj2 (function* or development* or follow?up or outcome*)).tw.
- 7. ((neurocognitive or neuropsychological or neurodevelopmental or developmental or intellectual or mental) adj2 (function* or development* or follow?up or outcome*)).kf.
- 8. ((neurocognitive or neuropsychological or neurodevelopmental or developmental or intellectual or mental or academic) adj2 (deficit* or disabilit* or disorder* or delay? or impairment* or difficult* or disturbanc*)).tw.
- ((neurocognitive or neuropsychological or neurodevelopmental or developmental or intellectual or mental or academic) adj2 (deficit* or disabilit* or disorder* or delay? or impairment* or difficult* or disturbanc*)).kf.
- 10. (intelligence or attention or (executive adj function*) or (working adj memory) or cognit* or learning or memory or reading or (reaction adj time*) or (response adj time*) or (processing adj speed) or vigilance or (executive adj control) or (problem? adj solving) or dyslexia or dyscalculia or impulsiveness or impulsivity).tw.
- 11. (intelligence or attention or (executive adj function*) or (working adj memory) or cognit* or learning or memory or reading or (reaction adj time*) or (response adj time*) or (processing adj speed) or vigilance or (executive adj control) or (problem? adj solving) or dyslexia or dyscalculia or impulsiveness or impulsivity).kf.
- 12. exp Arousal/
- 13. exp Memory/
- 14. communication/ or language/ or exp verbal behavior/
- 15. exp Education/
- 16. exp Academic Performance/
- 17. or/26-42
- 18. (neurocogniti* adj2 outcome?).ti.
- 19. 25 or 43
- 20. 42 or 44

Transition of care

transition of care OR (transition* AND care) OR transition to adult care[mh]

Randomized controlled trials and clinical controlled trials

Randomized controlled trial[pt] OR controlled clinical trial[pt] OR randomized[tiab] OR placebo[tiab] OR drug therapy[sh] OR randomly[tiab] OR trial[tiab] OR groups[tiab]) NOT (animals [mh] NOT humans [mh]

Systematic review

systematic review[tiab] OR review literature[mh] OR review[tiab] OR reviews[tiab] OR review[pt] OR systematic literature review[tiab] OR meta analysis[pt] OR meta analysis[mh] OR meta analysis[tiab] OR meta analysis[tiab] OR meta analysis[tiab]

Clinical practice guidelines

guideline[pt] OR guideline OR guideline* OR Guidelines as Topic[Mesh] OR Health Planning Guidelines[Mesh]

Appendix 2 Example search strategy for male gonadal dysfunction

Search 1: Patient	(((leukemia OR leukemi* OR leukaemi* OR (childhood ALL) OR AML OR hodgkin OR hodgkin* OR T-cell OR B-cell OR non-hodgkin OR sarcoma OR OR Ewing* OR osteosarcoma OR osteosarcom* OR wilms tumor OR wil neuroblastoma OR neuroblastom* OR rhabdomyosarcoma OR rhabdom teratom* OR hepatoma OR hepatoma* OR hepatoblastoma OR hemedulloblastoma OR medulloblastom* OR PNET* OR neuroectode retinoblastoma OR retinoblastom* OR meningioma OR meningiom* OR gli oncology OR paediatric oncology) OR (childhood cancer OR childhood tum (brain tumor* OR brain tumour* OR brain neoplasms OR central nervous	sarcom* OR sarcoma, Ewing's Ims* OR nephroblastom* OR nyosarcom* OR teratoma OR patoblastom* OR PNET OR rmal tumors, primitive OR oma OR gliom*) OR (pediatric nor OR childhood tumors)) OR system neoplasm OR central	
	nervous system neoplasms OR central nervous system tumor* OR central brain cancer* OR brain neoplasm* OR intracranial neoplasm*) OR testinal of the strain o	stis neoplasm OR neoplasm,	
	testicular OR testicular neoplasm OR testicular neoplasms OR testis cancer tumor OR testicular cancer OR cancer of testis OR testis tumour OR testis OR testis tumor* OR (leukemia, lymphocytic, acute[mh]))		
Search 2: Patient	male[tiab] OR males OR boy OR boys OR boyfriend OR boyhood		
Search 3: Intervention	Radiotherapy OR radiation OR radiation therapy OR irradiation OR irradiation OR injury, radiation OR radiation injury OR radiationsyndromes OR syndrome radiation OR radiation sickness OR radiation OR radiation* OR irradiation OR radiations	tion syndrome OR radiation	
Search 4: Intervention	Testicles OR testicle OR testes OR testis OR testis* OR testicle* OR testes pelvic OR pelvis region OR region pelvis OR pelvis* OR pelvic*	* OR pelvic region OR region,	
Search 5: Intervention	Brains OR brain OR encephalon OR encephalons OR brain* OR encephalon*	k	
Search 6: Intervention	total body OR whole body OR total body* OR body whole*		
Search 7: Outcome			
Search 8: Outcome	androgen hormone insufficiency OR leydig cell OR cells, leydig failure Of failure OR testicular failure OR gonadal failure OR hypogonadism OR low		
0 1449:50	deficiency OR androgen deficiency OR low testosterone* OR hypogonadism	n* OR leydig cell*	
	AND 3 AND (4 OR 5 OR 6) AND (7 OR 8) d in the last 20 years; Humans	= 488 hits	

Appendix 3

Example of an empty evidence table for 'Who needs surveillance?'

Clinical question Author et al. Title. Journal year;volume:pages Study design **Participants** Treatment era **Treatment** Main outcomes **Additional remarks** Years of follow-up 1. Study design 1. Type and number of 1. Chemotherapy 1. Outcome definitions 1. Strengths participants 2. Radiotherapy 2. Limitations 2. Treatment era 2. Diagnoses 2. Results 3. Surgery 3. Risk of bias A. Selection bias 3. Age at diagnosis 4. Other treatments Low risk/High risk/Unclear 3. Follow-up 4. Age at follow-up Reason: 5. Controls (if applicable) B. Attrition bias Low risk/High risk/Unclear Reason: C. Detection bias Low risk/High risk/Unclear Reason: D. Confounding Low risk/High risk/Unclear Reason:

Who needs premature ovarian insufficiency surveillance?

Byrne et al. Early menopause in long-term survivors of cancer during adolescence. Am J Obstet Gynecol 1992;166:788-793

Study design	-	-	_	
Study design Treatment era	Participants	Treatment	Main outcomes	Additional remarks
Years of follow-up	Turcie partes	Treatment	Wall decomes	Additional Temarks
Study design:	Type and number of	Chemotherapy only:	Outcome definitions:	<u>Strengths</u>
Multi-center cohort	participants	68 (6.5%)	- Amenorrhea: woman's report of whether she was still	- study sample
study	1048 female CCS ≥21 years of		having menstrual periods	
	age at study entry; 954 were	Alkylating agents and		<u>Limitations</u>
Treatment era:	menstruating before study	radiotherapy above	Amenorrhea:	- Self-reported outcome
1945-1974	entry and 94 became	diaphragm:	- 123/954 (12.9%) menopausal after study entry	- Control group not representative
	menopausal before they were	38 (3.6%)	- 831/954 (87.1%) still menstruating	for general population
Follow-up:	eligible for the cohort			
>19 yr after cancer		Alkylating agents and	Age-specific relative risks for amenorrhea survivors vs.	Risk of bias
diagnosis	<u>Diagnoses:</u>	radiotherapy below	controls:	A. Selection bias: Unclear
	Female genital cancer (n=90),	<u>diaphragm:</u>	- All survivors aged 21-25: RR 4.32, 95% CI 2.28-8.17	Reason: unclear how many patients
	Hodgkin's disease (n=206), non-	79 (7.5%)	- All survivors aged 26-30: RR 1.61, p>0.05	were included in the original cohort
	Hodgkin's lymphoma (n=31),		- All survivors aged 31-40: RR 0.78, p>0.05	of survivors
	soft tissue sarcoma (n=115),	Radiotherapy only:	- All survivors aged 41+: RR 0.98, p>0.05	
	leukaemia (n=15), brain or CNS	261 (24.9%)	- Alkylating agents alone aged 21-25: RR 9.17, 95% CI	B. Attrition bias: Low risk
	tumour (n=133), bone cancer		2.67-31.49	Reason: 90% of eligible survivors
	(n=65), other (n=393)	Surgery only:	- Radiotherapy below diaphragm and alkylating agents	completed follow-up assessment.
		493 (47.0%)	aged 21-25: RR 27.39, 95% CI 12.42-60.35	At follow-up, 10% of the survivors
	Age at diagnosis:		- Radiotherapy below diaphragm and alkylating agents	and 1% of the controls had died
	Mean 13.6 yr	Sterilizing surgery and	aged 26-30: RR 4.64, p<0.01	
		chemotherapy and	- Radiotherapy alone aged 21-25: RR 3.66, 95% CI 1.34-	C. Detection bias: Unclear
	Age at follow-up:	<u>radiotherapy:</u>	9.99	Reason: unclear if the outcome
	Mean 32.3 yr	25 (2.4%)	- Radiotherapy alone aged 26-30: RR 2.41, p<0.05	assessors were blinded for
			- Radiotherapy alone aged 31-40: RR 0.90, p>0.05	important determinants related to
	Controls:	Other treatments:	- Radiotherapy alone aged 41+: RR 1.22, p>0.05	the outcome
	1596 menstruating siblings at	84 (8.0%)	- Aged 0-12 at diagnosis aged 21-30 at follow-up: RR 0.62,	
	age 21 yr; Mean age at follow-		p>0.05	D. Confounding: High risk
	up 33.0 yr		- Aged 13-19 at diagnosis aged 21-30 at follow-up: RR	Reason: Controls not matched to
			2.32, 95% CI 1.63-3.291	cases

Example of an empty evidence table for 'What surveillance modality should be used?'

Clinical question Author et al. Title. Journal year; volume: pages Study design Treatment era **Participants** Diagnostic tests Main outcomes Additional remarks Years of follow-up 1. Study design 1. Type and number of 1. Diagnostic test(s) 1. Diagnostic outcomes (sensitivity, specificity, participants PPV, NPV, ROC) 2. Outcome definitions 2. Age at diagnosis 2. Treatment era 3. Age at follow-up 3. Follow-up 4. Cancer treatment 5. Prevalence/risk of late effect

What surveillance modality should be used to detect impaired spermatogenesis?

Green et al. Lack of specificity of plasma concentrations of inhibin B and follicle-stimulating hormone for identification of azoospermic survivors of childhood cancer: a report from the St Jude Lifetime Cohort Study. J Clin Oncol 2013;31:1324-1328

Jade Elletime Conort 3	ludy. J Cliff Officor 2013;31:1324-1328			
Study design Treatment era Years of follow-up	Participants	Diagnostic tests	Main outcomes	Additional remarks
1. Study design	1. Type and number of	1. Diagnostic test	Azoospermia	298 out of 485 (61.4%) eligible
Single-centre cohort	<u>participants</u>	Inhibin B, FSH and inhibin	<u>Sensitivity</u>	survivors participated in this study;
study	275 male CCS who received	B:FSH ratio	- Inhibin B: 100%	23 treated with ≥40 Gy
	gonadotoxic treatment, were not		- FSH: 78.1%	hypothalamic/pituitary irradiation
2. Treatment era	receiving exogenous androgens,	2. Cut-off levels for	- Inhibin B:FSH ratio: 75.3%	and/or tumour in
Not reported	had received <40 Gy	<u>azoospermia</u>		hypothalamic/pituitary region
	hypothalamic/pituitary irradiation	- Inhibin B ≤31 ng/L	Specificity	excluded.
3. Follow-up	and were ≥18 years of age at	- FSH >11.5 IU/L	- Inhibin B: 45.0%	
≥10 years from	follow-up	- Inhibin B:FSH ratio ≤2.52	- FSH: 74.1%	Inhibin B was measured in 238
diagnosis		pg/mIU	- Inhibin B:FSH ratio: 74.5%	patients and FSH was measured in
	2. Age at diagnosis			275 patients.
	Range 0-21 years	ROC analysis determined the	Negative predictive value	
		optimal cut-off levels	- Inhibin B: 100%	Patient sample divided into a
	3. Age at follow-up		- FSH: 84.6%	learning set (n=140) and a validation
	Median 30.5 (19.7-59.1) years		- Inhibin B:FSH ratio: 83.5%	set (n=135) by random assignment: diagnostic values were similar.
	4. Gonadotoxic treatment		Positive predictive value	Results are shown for the combined
	275 (100%);		- Inhibin B: 52.1%	data sets.
	Alkylating agents, direct testicular		- FSH: 65.1%	
	irradiation or <40 Gy		- Inhibin B:FSH ratio: 63.8%	
	hypothalamic/pituitary irradiation			
	(≥40 Gy hypothalamic/pituitary		Area under the ROC curve	
	irradiation specifically excluded		- Inhibin B: 0.72	
	from analysis)		- FSH: 0.83	
			- Inhibin B:FSH ratio: 0.83	
	5. Prevalence azoospermia:			
	105 (38.2%)			

Appendix 4

<u>Risk of bias assessment of observational studies</u>

	Internal validity
Study group	 Selection bias - Low risk/high risk/unclear Is the study group representative? Low risk if: the study group consisted of more than 75% of the original cohort of childhood cancer survivors or it was a random sample with respect to the cancer treatment
Follow-up	Attrition bias - Low risk/high risk/unclear Is the follow-up adequate? Low risk if: • the outcome was assessed for more than 75% of the study group
Outcome	 Detection bias - Low risk/high risk/unclear Are the outcome assessors blinded for important determinants related to the outcome? Low risk if: the outcome assessors were blinded for important determinants related to the outcome
Risk estimation	 Confounding - Low risk/high risk/unclear Are the analyses adjusted for important confounding factors? Low risk if: important prognostic factors (i.e. age, gender, co-treatment, follow-up) were taken adequately into account

Risk of bias assessment criteria for diagnostic studies

	Internal validity
Study group	Selection bias - Low risk/high risk/unclear
	Is the study group representative? yes/no/unclear
	Low risk if:
	 the study group consisted of more than 75% of the original
	cohort of childhood cancer survivors
	• or it was a random sample
Index test	Index test bias - Low risk/high risk/unclear
	Are the index test results interpreted without knowledge of the
	results of the reference standard? yes/no/unclear
	Low risk if:
	• the index test results were interpreted without knowledge of the
	results of the reference standard in all patients
Reference test	Reference test bias - Low risk/high risk/unclear
	Are the reference test results interpreted without knowledge of
	the results of the index test? yes/no/unclear
	Low risk if:
	 the reference test results were interpreted without knowledge
	of the results of the index test in all patients
Flow and	Verification bias - Low risk/high risk/unclear
timing	Was there an appropriate interval (to be determined by the group)
	between index test(s) and reference standard? yes/no/unclear
	Low risk if:
	 there was an appropriate interval between the index test(s) and
	reference standard in all patients
	Attrition bias - Low risk/high risk/unclear
	Did all patients receive the same tests (i.e. same reference
	standard and same index test)? yes/no/unclear
	Low risk if:
	 More than 75% of the study group received the same tests

Risk of bias assessment criteria for randomized clinical trials

	Internal validity
Study group	Selection bias - Low risk/high risk/unclear Is the study group representative? yes/no/unclear Low risk if:
	there was random sequence allocation and allocation concealment
Follow-up	Attrition bias - Low risk/high risk/unclear Is the follow-up adequate? yes/no/unclear Low risk if: • the outcome was assessed for more than 90% in each treatment arm
Outcome	 Performance bias - Low risk/high risk/unclear Are the participants and personnel blinded from knowledge of which intervention was received? yes/no/unclear Low risk if: the participants and personnel were blinded from knowledge of which intervention was received
	<u>Detection bias</u> - Low risk/high risk/unclear Are the outcome assessors blinded from knowledge of which intervention was received? yes/no/unclear Low risk if:
	 the outcome assessors were blinded from knowledge of which intervention was received

Appendix 5

Example of a Summary of findings table of the body of evidence for the risk of premature ovarian insufficiency after procarbazine

Outcome	Study	No. of participants	Follow up (median/mean, range) yr	Alkylating agents Radiotherapy to ovaries	Events	Effect size	Risk of bias
1.5 Risk POI after procarbazine (n=4 studies)	Chemaitilly 2006*	3,390 CCS	>5 yr after cancer diagnosis	Alkylating agents: 49.7%; RT to ovaries: 24.5%	215/3390 (6.3%) amenorrhea within 5 yr after their cancer diagnosis	Odds ratio (95% CI) for amenorrhea age at diagn 0-12 yr Procarbazine yes vs. no: OR 3.2 (1.3-7.3)	SB: high risk AB: low risk DB: unclear CF: low risk
	Thomas-Teinturier 2013*	706 CCS	>5 yr after cancer diagnosis	Any alkylating agent: 47.7%; Procarbazine: 7.2%; RT to ovaries: 56.7%	62/706 (8.9%) nonsurgical menopause; 15/706 (2.1%) nonsurgical premature menopause <age 40="" td="" yr<=""><td>Relative risk (95% CI) for nonsurgical menopause Procarbazine dose per g/m²: RR 2.5 (1.4-5.8)</td><td>SB: high risk AB: low risk DB: unclear CF: low risk</td></age>	Relative risk (95% CI) for nonsurgical menopause Procarbazine dose per g/m ² : RR 2.5 (1.4-5.8)	SB: high risk AB: low risk DB: unclear CF: low risk
	Thomas-Teinturier 2015*	108 CCS	>3 yr after cancer treatment	Any alkylating agent: 100%; Procarbazine: 21.9%; RT to ovaries: 17.6%	8/108 (7.6%) altered ovarian function (↑ FSH, ↓ AMH and amenorrhoea)	Mean FSH Procarbazine dose: β 0.012, p<0.001; (Each unit increase in procarbazine dose, mean FSH values increased by 0.012 IU/L)	SB: high risk AB: low risk DB: unclear CF: low risk
	Levine 2018*	2,930 CCS	>5 yr after cancer diagnosis	Alkylating agents: 46.5%; Procarbazine: 201 (7.2%); RT to ovaries: 55.4%	110/2930 (3.8%) nonsurgical premature menopause <age 40="" td="" yr<=""><td>Odds ratio (95% CI) for nonsurgical premature menopause Procarbazine dose <4000 mg/m² vs. 0: OR 3.07 (0.76-12.43); Procarbazine dose ≥4000 mg/m² vs. 0: OR 8.96 (5.02-16.00)</td><td>SB: high risk AB: low risk DB: unclear CF: low risk</td></age>	Odds ratio (95% CI) for nonsurgical premature menopause Procarbazine dose <4000 mg/m² vs. 0: OR 3.07 (0.76-12.43); Procarbazine dose ≥4000 mg/m² vs. 0: OR 8.96 (5.02-16.00)	SB: high risk AB: low risk DB: unclear CF: low risk
GRADE assessment:		astiva sabart studios				0.0	
Study design: Study limitations: Consistency: Directness:	Consistency: 0 No important inconsistency, all show effect of procarbazine						
Precision: Publication bias:	0 Unlikely	,		total number of events	and narrow confidence inter	vals	
Effect size: 0 No large magnitude of effect Dose-response: +1 Dose response relationship as higher doses are associated with an increased risk as compared to lower doses Plausible confounding: 0 No plausible confounding Quality of evidence: ⊕⊕⊕ HIGH Conclusion: Increased risk of POI after procarbazine vs. no procarbazine in female cancer survivors diagnosed before age 25 years. (4 studies significant effect; 7,134 participants; 395 events; 4 multivariable analyses)							

Abbreviations: AB, attrition bias; AMH, anti-Müllerian hormone; CCS, childhood cancer survivors; CF, confounding; DB, detection bias; FSH, follicle stimulating hormone; SB, selection bias; yr, year.

^{*} Overlap in included patients in studies of Chemaitily 200 and Levine 2018; and Thomas-Teinturier 2013 and 2015.

Appendix 6

GRADE quality assessment

Initial score based on type of evidence

- +4: RCTs/ SR of RCTs
- +2: CCTs or observational evidence (e.g., cohort, case-control) for intervention questions
- +4: Observational evidence for etiologic, prognostic and diagnostic questions

Factors that might decrease the quality of the body of evidence

- 1. Study limitations: risk of bias based on selection bias, attrition bias, detection bias and confounding as defined in the risk of bias table.
 - 0: No problems
 - -1: Problem with 1 element
 - -2: Problem with 2 elements
 - -3: Problem with 3 or more elements
- 2. Consistency: degree of consistency of effect between or within studies
 - 0: All/most studies show similar results
 - -1: Lack of agreement between studies (statistical heterogeneity / conflicting result, e.g. effect sizes in different directions)
- 3. <u>Directness:</u> the generalizability of population and outcomes from each study to the population of interest
 - 0: Population and outcomes broadly generalizable
 - -1: Problem with 1 element (population different from the defined inclusion criteria OR outcomes different from the defined inclusion)
 - -2: Problem with 2 elements (population and outcomes)
- 4. Precision: the precision of the results
 - 0: No important imprecision when studies include many patients and many events and thus have narrow confidence intervals; Determine with the chairs and advisors what is seen as many patients, many events and narrow confidence intervals
 - -1: Important imprecision when studies include relatively few patients and few events and thus have wide confidence intervals (especially when the confidence interval cross the 0).
 Another criteria to consider is the clinical decision threshold. This is the threshold of the effect size that would change the decision whether or not to adopt a clinical action.
 Downgrade if the effect estimate and confidence intervals cross the clinical decision threshold. Determine with the chairs and advisors the clinical decision threshold.
 OR if only one study has been identified

- -2: If there is important imprecision (see -1) AND if only one study has been identified
- <u>5.</u> <u>Publication bias:</u> if investigators fail to report studies and outcomes (typically those that show no effect)
 - 0: Publication bias unlikely
 - -1: Risk of publication bias when for example published evidence is limited to industry funded trials

Factors that might increase the quality of the body of evidence

1. Magnitude of effect:

- +1: Large magnitude of effect; all studies show significant effect sizes (point estimate) >2
 or <0.5
- +2: Very large magnitude of effect; all studies show significant effect sizes (point estimate) >5 or <0.2

2. Dose response gradient:

• +1: Evidence of clear relation with increases in the outcome with higher exposure levels across or within studies

3. Plausible confounding:

+1: If adjustment for confounders would have increased the effect size; for example the
estimate of effect is not controlled for the following possible confounders: smoking,
degree of education, but the distribution of these factors in the studies is likely to lead to
an underestimate of the true effect

Total score

- ⊕⊕⊕ High quality evidence
- ⊕⊕⊕ Moderate quality evidence
- $\bigoplus \bigoplus \bigoplus \bigoplus$ Low quality evidence
- ⊕⊖⊖ Very low quality evidence

Appendix 7

Overall conclusions of evidence table for male gonadotoxicity survveillance

Who needs surveillance?	
Risk of impaired spermatogenesis in male cancer survivors diagnosed before age 25 years	Quality of evidence
Increased risk after alkylating agents vs. no alkylating agents	⊕⊕⊕ HIGH ¹⁻⁵
Increased risk after increasing doses of alkylating agents	⊕⊕⊕⊕ HIGH ¹⁻⁵
Increased risk after cyclophosphamide vs. no cyclophosphamide	⊕⊕⊕⊕ HIGH ^{1,3-5}
Increased risk after increasing doses of cyclophosphamide	⊕⊕⊕⊕ HIGH ^{1,3-5}
Increased risk after <i>procarbazine</i> and <i>mechlorethamine</i> (given as part of multi-agent treatment) vs. no procarbazine and mechlorethamine	⊕⊖⊖⊖ VERY LOW ²
Increased risk after <i>increasing doses of procarbazine</i> and <i>mechlorethamine</i> (given as part of multi-agent treatment)	⊕⊖⊖⊖ VERY LOW ²
Unknown risk after dacarbazine	No studies
No significant effect of dacarbazine dose	⊕⊖⊖ VERY LOW ²
Unknown risk after busulfan, chlorambucil, ifosfamide, melphalan, thiotepa, carmustine (BCNU), lomustine (CCNU)	No studies
Unknown risk after antimetabolites	No studies
Unknown risk after <i>platinum compounds</i>	No studies
Increased risk after <i>radiotherapy exposing the testes</i> vs. no radiotherapy exposing the testes	⊕⊖⊖ VERY LOW ^{3,6}
Unknown risk after higher vs. lower doses of radiotherapy exposing the testes	No studies
Unknown risk after <i>gonadotoxic chemotherapy combined with radiotherapy exposing</i> the testes	No studies
Unknown risk after <i>unilateral orchiectomy combined with radiotherapy exposing the testes</i>	No studies
Unknown risk after novel agents (tyrosine kinase inhibitors, demethylating agents, oxaliplatin)	No studies

Appendix 8 Evidence to decision (EtD) framework - template

	Criteria	Judgements	Research evidence	Additional
				considerations
PROBLEM	Is the problem a priority?	☐ No ☐ Probably no ☐ Uncertain ☐ Probably yes ☐ Yes		
	What is the overall certainty of this evidence?	☐ No included studies ☐ Very low ☐ Low ☐ Moderate ☐ High		
BENEFITS AND HARMS	Is there important uncertainty about how much people value the main outcomes?	☐ Important uncertainty or variability ☐ Possibly important uncertainty or variability ☐ Probably no important uncertainty or variability ☐ No important uncertainty or variability ☐ No important uncertainty or variability ☐ No known undesirable outcomes		
38	desirable anticipated effects large?	☐ No ☐ Probably no ☐ Uncertain ☐ Probably yes ☐ Yes ☐ Varies		
	Are the undesirable anticipated effects small?	☐ No ☐ Probably no ☐ Uncertain ☐ Probably yes ☐ Yes ☐ Varies		
	Are the desirable effects large relative to undesirable effects?	☐ No ☐ Probably no ☐ Uncertain ☐ Probably yes ☐ Yes ☐ Varies		

	Are the	□ No	
	resources	☐ Probably no	
ш	required small?	☐ Uncertain	
		☐ Probably yes	
US		☐ Yes	
RESOURCE USE		☐ Varies	
JUC	Is the	□ No	
(ES	incremental cost	\square Probably no	
Æ	small relative to	☐ Uncertain	
	the net	\square Probably yes	
	benefits?	☐ Yes	
		□Varies	
	What would be	☐ Increased	
	the impact on	☐ Probably	
>	health	increased	
EQUITY	inequities?	☐ Uncertain	
EQ		☐ Probably	
		reduced	
		☐ Reduced	
		□ Varies	
T	Is the option	□No	
3ILI	acceptable to key	☐ Probably no	
TAI	stakeholders?	☐ Uncertain	
ACCEPTABILITY	stakeriolaers:	☐ Probably yes	
AC		☐ Yes	
	la tha antion	□ Varies	
_	Is the option feasible to	□ No	
FEASIBILITY	implement?	☐ Probably no	
	implement:	☐ Uncertain	
ΕΑ		☐ Probably yes☐ Yes	
4			
		☐ Varies	

Overall conclusions

	Balance of consequences				
Undesirable	Undesirable	The balance between	Desirable	Desirable	
consequences	consequences	desirable and	consequences	consequences	
clearly outweigh	probably outweigh	undesirable	probably outweigh	clearly outweigh	
desirable	desirable	consequences	undesirable	undesirable	
consequences	consequences	is closely balanced or	consequences	consequences	
in most settings	in most settings	uncertain	in most settings	in most settings	

Appendix 9

Criteria for grading the recommendations

Grade of Recommendation Conclusions of evidence according	Strong recommendation to do Benefits >>> risk & harms	Moderate recommendation to do Benefits > or = risk & harms	Recommendation not to do No benefit / Potentially harm
to GRADE High quality of evidence Consistent evidence from well performed and high quality studies or systematic reviews (low risk of bias, direct, consistent, precise).	Strong recommendation based on high quality evidence	Moderate recommendation based on high quality evidence	Recommendation not to do based on high quality evidence
Moderate quality of evidence Evidence from studies or systematic reviews with few important limitations.	Strong recommendation based on moderate quality evidence	Moderate recommendation based on moderate quality evidence	Recommendation not to do based on moderate quality evidence
Low to very low quality of evidence Evidence from studies with serious flaws, only expert opinion, or standards of care.	Strong recommendation based on expert opinion	Moderate recommendation based on (very) low quality evidence Diverging expert opinions	Recommendation not to do based on expert opinion
	Wording in recommend is recommended	dations: Is reasonable	Is not recommended

Appendix 10

Example timeline for guideline development work

Tasks to be finished	By whom	Estimated time to complete task
Preparation phase		
Compose working group	Chairs and advisors	Allow 2 months for this
Identify coordinator and WG leaders		before:
Conference call: introduction and composition of	Total group	
working groups		
Development phase		
Step 1		
Develop protocol	Chairs, coordinator, WG	Allow 2 months for this
Define scope of the guideline, i.e. outcomes and	leaders and advisors	before:
population of interest		
Evaluate concordances/discordances		
Formulate clinical questions		
Step 2		
Send scope of the guideline and clinical questions to	Coordinator	Allow 1 month for this
WG members		before:
Conference call: discuss scope of the guideline and	Total group	
clinical questions		
Step 3		
Finalize clinical questions	Chairs, coordinator, WG	Allow 2 months for this
Develop search strategy	leaders and advisors	before:
Define in- and exclusion criteria		
Conference call: discuss search strategy and in- and	Total group	
exclusion criteria		
Perform literature search	Cochrane Childhood Cancer	Allow 2 months for this
	Group	before:
Conference call: discuss steps for evidence selection	Total group	
Send results literature search and instructions to WG	Coordinator	
leaders and members		
Select evidence based on search	Coordinator, WG leaders and	Allow 2-3 months for this
Send final inclusion of eligible studies to coordinator	members	depending on number of articles
Step 4		ar treres
Conference call: discuss steps for summarizing the	Total group	
evidence	. 3tai 61 aap	
Make evidence tables	Coordinator, WG leaders and	Allow 1-2 months for this
iviance evidence tables	members	before:
Conference call: if necessary to discuss difficulties	Total group or separate WGs	Allow 1 month to make
conference can. If thecessary to discuss difficulties	Total group of separate 11 cs	modifications before:
Circulate evidence tables to the whole group	Coordinator	
Each WG checks evidence tables (missing studies,	WG leaders to coordinate	
completeness, etc)	within their WG	Allow 1 month for this
Return comments evidence tables	Within their We	before:
Agree final evidence tables	Total group	before.
Conference call: discuss and agree final evidence	Total group	
tables and outline next steps for formulating overall		
conclusions of the evidence	Chaire as and in 1990	
Develop conclusion of evidence tables	Chairs, coordinator, WG	
	leaders and advisors	Allow 2-3 months for this
Circulate conclusions of evidence tables to the	Coordinator	before:
working group members		

Conference call: discuss and agree final conclusions of evidence tables	Total group	Allow 2 months to make modifications before:
Step 5		
Formulate draft recommendations	Chair, coordinator, advisors and WG leaders	Allow 2 months before:
Conference call: discuss draft recommendations	Total group	Allow 2 months to make modifications before:
Discuss and develop final recommendations,	IGHG, PCSF and external	
preferably in a face-to-face meeting	experts	