Management of Germ Cell Tumours of the Testis in Adult Patients. German Clinical Practice Guideline Part I: Epidemiology, Classification, Diagnosis, Prognosis, Fertility Preservation, and Treatment Recommendations for Localized Stages

Kliesch, Sabine; Schmidt, Stefanie; Wilborn, Doris; Aigner, Clemens; Albrecht, Walter; et al; Hermanns, Thomas

Abstract: INTRODUCTION This is the first German evidence- and consensus-based clinical guideline on diagnosis, treatment, and follow-up on germ cell tumours (GCTs) of the testis in adult patients. We present the guideline content in two publications. Part I covers the topic’s background, methods, epidemiology, classification systems, diagnostics, prognosis, and treatment recommendations for the localized stages. METHODS An interdisciplinary panel of 42 experts including 1 patient representative developed the guideline content. Clinical recommendations and statements were based on scientific evidence and expert consensus. For this purpose, evidence tables for several review questions, which were based on systematic literature searches (last search was in March 2018) were provided. Thirty-one experts entitled to vote, rated the final clinical recommendations and statements. RESULTS We provide 161 clinical recommendations and statements. We present information on the quality of cancer care and epidemiology and give recommendations for staging and classification as well as for diagnostic procedures. The diagnostic recommendations encompass measures for assessing the primary tumour as well as procedures for the detection of metastases. One chapter addresses prognostic factors. In part I, we separately present the treatment recommendations for germ cell neoplasia in situ, and the organ-confined stages (clinical stage I) of both seminoma and nonseminoma. CONCLUSION Although GCT is a rare tumour entity with excellent survival rates for the localized stages, its management requires an interdisciplinary approach, including several clinical experts. Quality of care is highly related to institutional expertise and can be reassured by established online-based second-opinion boards. There are very few studies on diagnostics with good level of evidence. Treatment of metastatic GCTs must be tailored to the risk according to the International Germ Cell Cancer Collaboration Group classification after careful diagnostic evaluation. An interdisciplinary approach as well as the referral of selected patients to centres with proven experience can help achieve favourable clinical outcomes.

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Management of Germ Cell Tumours of the Testis in Adult Patients. German Clinical Practice Guideline Part I: Epidemiology, Classification, Diagnosis, Prognosis, Fertility Preservation, and Treatment Recommendations for Localized Stages

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Abstract

Introduction: This is the first German evidence- and consensus-based clinical guideline on diagnosis, treatment, and follow-up on germ cell tumours (GCTs) of the testis in adults. We present the guideline content in two publications. Part I covers the topic’s background, methods, epidemiology, classification systems, diagnostics, prognosis, and treatment recommendations for the localized stages. Methods: An interdisciplinary panel of 42 experts including 1 patient representative developed the guideline content. Clinical recommendations and statements were based on scientific evidence and expert consensus. For this purpose, evidence tables for several review questions, which were based on systematic literature searches (last search was in March 2018) were provided. Thirty-one experts entitled to vote, rated the final clinical recommendations and statements. Results: We provide 161 clinical recommendations and statements. We present information on the quality of cancer care and epidemiology and give recommendations for staging and classification as well as for diagnostic procedures. The diagnostic recommendations encompass measures for assessing the primary tumour as well as procedures for the detection of metastases. One chapter addresses prognostic factors. In part I, we separately present the treatment recommendations for germ cell neoplasia in situ, and the organ-confined stages (clinical stage I) of both seminoma and nonseminoma. Conclusion: Although GCT is a rare tumour entity with excellent survival rates for the localized stages, its management requires an interdisciplinary approach, including several clinical experts. Quality of care is highly related to institutional expertise and can be reassured by established online-based second-opinion boards. There are very few studies on diagnostics with good level of evidence. Treatment of metastatic GCTs must be tailored to the risk according to the International Germ Cell Cancer Collaboration Group classification after careful diagnostic evaluation. An interdisciplinary approach as well as the referral of selected patients to centres with proven experience can help achieve favourable clinical outcomes.

Background

The incidence of germ cell tumours (GCTs) of the testis is low with about 10 out of every 100,000 men in Germany per year [1]. However, the importance arises from the fact that GCTs represent the most common type of cancer in men aged 20–40 years, with 25% of all malignancies diagnosed in this age group [1]. Survival rates are excellent for the localized stages (95–100%) and are still good for the metastasized stages (70–90%) [2]. Clinically, GCTs are classified into seminomas (about 60%) and non-seminomatous GCTs [1, 3].

For all stages, the tumour-specific therapeutic success depends on the correct diagnosis and staging, which is essential to avoid under- but also overtreatment. Interdisciplinary cooperation between urologists, oncologists, radio-oncologists, radiologists, and pathologists is mandatory, especially for multidisciplinary therapy control in the metastatic poor prognosis and relapsed patient groups. In addition, established online-based second-opinion boards enable urologists and oncologists to reassure especially first-line treatment decisions [4–6].
So far, no official evidence-based German clinical practice guideline on this topic has been available. There are interdisciplinary consensus recommendations [7, 8], which were developed by the interdisciplinary German Testicular Cancer Study Group (GTCSG) of the German Cancer Society. However, they are mainly based on expert consensus and do not follow the transparent and high-quality methodology development criteria set up by the Association of Scientific Medical Societies in Germany (AWMF) [9].

Therefore, the German Society of Urology (DGU) together with the interdisciplinary GTCSG initiated the development of an S3 German-language clinical practice guideline on this topic. The guideline category S3 refers to a German guideline classification system in which S3 represents the highest methodological standard. It involves both evidence- and consensus-based concepts to achieve highest quality guideline recommendations from an interdisciplinary panel group [10].

The German S3 guideline is freely available for download in both short and long versions: https://www.awmf.org/leitlinien/detail/ll/043-049OL.html. A separate method report on the guideline development process is additionally available.

Physicians and other medical service providers who are involved in the diagnostics, treatment, and follow-up of GCTs (all stages, outpatient and inpatient care as well as rehabilitation) are the target audience of the present guideline. A patient guideline for patients and lay persons will be published soon. The guideline also comprises quality indicators for measuring the implementation of the guideline recommendations in routine clinical care; these data will be presented in a future publication.

We present the English translation of the main content of this guideline in two separate parts, which includes all clinical recommendations and statements of the original guideline. Part I relates to epidemiology, classification systems, diagnostics, prognosis, and treatment recommendations for the localized stages. Part II covers the treatment recommendations for the metastatic stages as well as relapsed and refractory disease, follow-up schedules, toxicity management, quality of life, and palliative and supportive care.

**Objective**

The overall aim of the present project is to improve the treatment of GCT and thus increase survival of the patients afflicted with the disease. Also, quality of life of surviving patients is intended to be improved by efforts to minimize acute and late toxicity of treatment.

For these goals, we aimed at developing the first German guideline for the management of GCTs in adult patients that is explicitly based on formal scientific evidence and if required on consensus in a working group of dedicated experts.

**Methods**

In the guideline development process, we followed the AWMF methodology for S3 guidelines [10]. This process is illustrated in Figure 1. For details, see the German method report (available at: https://www.awmf.org/leitlinien/detail/ll/043-049OL.html).

**Members and Funding**

The interdisciplinary panel for this guideline consisted of 32 experts (including a patient representative) entitled to vote, as well as several external experts and working group leaders (see Table 1). The project was funded by the German Cancer Aid. Methodological supervision was provided by the German Guideline Program in Oncology [11]. Conflicts of interest were disclosed by all guideline panel members and are available in the guideline report (http://www.awmf.org/leitlinien/detail/ll/043-044.html).
**Table 1. Coordinators, medical societies, and organizations involved**

| Coordinator | Prof. Dr. Sabine Kliesch, DGU (deputy: Prof. Dr. Peter Albers, GTCSG) Janine Weiberg (DGU) |
|-------------|------------------------------------------------------------------------------------------|
| Guideline office | PD Dr. Friedemann Zengerling |
| Scientific Team (DGU) (not entitled to vote) | PD Dr. Jens Bedke Prof. Dr. Jonas Busch PD Dr. Julia Heinzlbecker Prof. Dr. David Pfister PD Dr. Christian Ruf Dr. Stefanie Schmidt Dr. Doris Wilborn Dr. Christian Winter PD Dr. Friedemann Zengerling |

| Medical society or organization | Participants in guideline development |
|---------------------------------|-------------------------------------|
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| German Society of Radiation Oncology (DEGRO) | Prof. Dr. Johannes Claßen (deputy: Prof. Dr. Rainer Souchon) |
| German Society of Andrology (DGA) | PD Dr. Christian Ruf |
| German Society for General and Visceral Surgery (DGAV) | Dr. Stefanie Schmidt |
| German Society of Hematology and Medical Oncology (DGHO) | PD Dr. Anja Lorch (deputy: Dr. Christoph Oing) |
| German Society for Nuclear Medicine (DGN) | Dr. Matthias Gockel |
| German Association for Palliative Medicine (DGP) | PD Dr. Stefan Schweyer (deputy: Prof. Dr. Glen Kristiansen) |
| German Society for Pathology (DGP) | PD Dr. Stefan Schweyer (deputy: Prof. Dr. Glen Kristiansen) |
| German Society of Nursing Science (DGP) | Heinrich Recken, B.A. (deputy: Prof. Dr. Stefanie Seeling) |
| German Society for Thoracic Surgery (DGT) | Prof. Dr. Clements Aigner |
| German Society of Radiology (DRG) | PD Dr. Dirk Beyersdorff (deputy: Prof. Dr. Sascha Kaufmann) |
| Association of German Urologists (BDU) | Dr. med. Bernt Göckel-Beining (deputy: Dipl. Med. Roger Zillmann) |
| Association of German Pathologists (BVP) | Prof. Dr. Glen Kristiansen (deputy: Prof. Dr. Stefan Schweyer) |
| Austrian Society of Urology and Andrology (ÖGU) | PD Dr. Walter Albrecht (deputy: Dr. Renate Pichler) |
| Swiss Society of Urology (SGU) | PD Dr. Thomas Hermanns |
| Pain and Supportive Therapy, Quality of Life, and Palliative Care Working Group (Arbeitskreis Schmerztherapie, Supportivtherapie, Lebensqualität, und Palliativmedizin) | Dr. Matthias Beintker (deputy: Prof. Dr. Oliver Hakenberg) |
| Oncology Working Group (AKO) | Prof. Dr. Oliver Hakenberg |
| Supportive Measures in Oncology, Rehabilitation, and Social Medicine Working Group (ASORS) | Prof. Dr. Oliver Rick (deputy: Prof. Dr. Maike de Wit) |
| Internal Oncology Working Group (AIO) | Prof. Dr. Carsten Bokemeyer (deputy: Dr. Annette Dieng) |
| Andrology Working Group (AKA) | PD Dr. Diemer (deputy: PD Dr. Kathleen Herkommer) |
| Urological and Nephrological Diseases Rehabilitation Working Group (AKR) | Prof. Dr. Ulrich Otto (deputy: Prof. Dr. Dirk-Henrik Zermann) |
| Oncological Pathology Working Group (AOP) | Prof. Dr. Christian Wittekind |
| Oncologic Thoracic Surgery Working Group (AOT) | Prof. Dr. Joachim Schirren |
| Palliative Care Working Group (APM) | Prof. Dr. Karin Oechsle |
| Radiooncology Working Group (ARO) | Prof. Dr. Heinz Schmidberger (deputy: PD Dr. Arndt-Christian Müller) |
| Tumor Classification in Oncology Working Group (ATO) | Prof. Dr. Christian Wittekind |
RCTs [14], SIGN [15], Quadas [16], and Quips [17]. The Oxford performed in March 2018. Cohort studies, clinical trials, systematic reviews, meta-analyses, and international clinical guidelines were considered. We only included publications in English and concluding consensus conference was held in Berlin in May 2018. E-references are listed in the online supplementary eReferences file (see www.karger.com/doi/10.1159/000510407 for all on-line suppl. material).

The following wordings were used for characterizing the strength of the clinical recommendations (see also Table 2):
- A: strong recommendation: “we recommend” or “is recommended/is not recommended”
- B: weak recommendation: “should/should not”
- 0: inconclusive recommendation: “can”

For all studies included, relevant information was extracted in evidence tables, and the risk of bias was assessed according to study design, using AGREE [12], AMSTAR [13], the Cochrane tool for RCTs [14], SIGN [15], Quadas [16], and Quips [17]. The Oxford criteria were used for the level of evidence ratings [18] (Table 2). For the therapeutic key questions, the quality of the evidence was rated according to the Grading of Recommendations, Assessment, Development, and Evaluation (GRADE) approach [19].

The final evidence- and consensus-based recommendations and statements were elaborated, and a formal consensus of the guideline panel was achieved applying the nominal group technique under the guidance of an external moderator. The second and concluding consensus conference was held in Berlin in May 2018. E-references are listed in the online supplementary eReferences file (see www.karger.com/doi/10.1159/000510407 for all on-line suppl. material).

Evidence and Recommendation Development

In January 2017, the scope of work and the work packages were defined during the first consensus conference held in Berlin, Germany. Seventy-two key questions were defined, and afterwards several systematic literature searches were conducted using the databases Medline (via Ovid) and the Cochrane Library. The search period was from January 2010 to May 2017. A search update was performed in March 2018. Cohort studies, clinical trials, systematic reviews, meta-analyses, and international clinical guidelines were considered. We only included publications in English and German. Case reports, case series, case-control studies, editorials, commentaries, conference abstracts, and studies available in other languages were excluded.

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Results

Chapter 3: Quality of Care
See also Table 3.

1 We recommend that patients with a metastatic GCT belonging to the poor prognosis group according to the International Germ Cell Cancer Collaboration Group (IGCCCG) are treated at centres with proven experience (GRADE of recommendation A, level of evidence 2b) [4, 6].

2 We recommend that GCT patients with post-chemotherapeutic residual tumours receive residual tumour resection only after multidisciplinary consultation at centres with proven experience, which fulfil the prerequisites for multidisciplinary surgical interventions (expert consensus).

Chapter 4: Epidemiology, Risk Factors, Screening, and Prevention

1 In men aged 20–44 years, GCT is the most common malignant disease, accounting for about 25% of all cancers (statement, 2b) [1].

2 Within the last decades, the incidence of GCT has increased to currently 10 out of every 100,000 men per year in all industrialized countries including Germany (raw disease rate) (statement, 2b) [1].

3 GCTs are among the cancers with highest survival rates. The patient’s prognosis essentially depends on tumour histology, clinical stage, age, and quality of care (statement, 2b) [1, 6].
Accepted pathogenetic risk factors for the development of GCTs are previous contralateral GCT, history of maldescensus testis, positive family history, and male infertility (statement, 2b) [e20–e25].

General screening for the presence of a GCT is not recommended (A, 5) [e26–e28].

Regular self-examination of the testicles should be recommended especially to young men, as it can lead to early diagnosis (expert consensus).

In the presence of risk factors, the possibility of a GCT should be investigated (expert consensus).

Chapter 5: Pathological Classification

We recommend the 2016 WHO classification for the histopathological evaluation of GCTs [e29] (expert consensus).

Chapter 6: Diagnostics, Stages, and Classification Systems

See also Tables 3 and 4.

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Table 2. Levels of evidence according to the Oxford Centre for Evidence-Based Medicine 2009 classification system (from [18])

| Level of evidence | Diagnostics                                                                 | Therapy                                                                 |
|-------------------|-----------------------------------------------------------------------------|------------------------------------------------------------------------|
| Ia                | SR (with homogeneity*) of level 1 diagnostic studies; clinical decision rule with 1b studies from different clinical centres | SR (with homogeneity*) of RCTs                                           |
| Ib                | Validating cohort study with good reference standards or clinical decision rule tested within 1 clinical centre | Individual RCT (with narrow confidence interval)                       |
| Ic                | Absolute SpPins and SnNouts                                                  | All or none principle                                                  |
| IIa               | SR (with homogeneity*) of level >2 diagnostic studies                        | SR (with homogeneity) of cohort studies                                |
| IIb               | Exploratory cohort study with good reference standards, clinical decision rule after derivation, or validated only on split-sample or databases | Individual cohort study (including low quality RCT)                    |
| IIc               |                                                                             | Outcomes research, ecological studies                                  |
| IIIa              | SR (with homogeneity*) of 3b and better studies                             | SR (with homogeneity) of case-control studies                          |
| IIIb              | Non-consecutive study, or without consistently applied reference standards   | Individual case-control study                                           |
| IV                | Case-control study, poor or non-independent reference standard              | Case series (and poor-quality cohort and case-control studies)         |
| V                 | Expert opinion without explicit critical appraisal, or based on physiology, bench research or first principles | Expert opinion without explicit critical appraisal, or based on physiology, bench research or first principles |

RCT, randomized control trial; SR, systematic review. * By homogeneity, we mean a systematic review that is free of worrisome variations (heterogeneity) in the directions and degrees of results between individual studies. Not all systematic reviews with statistically significant heterogeneity need be worrisome, and not all worrisome heterogeneity need be statistically significant.

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Imaging procedures in the context of local diagnostics and/or staging diagnostics:

1. On clinical suspicion of a GCT, we recommend to perform a physical examination as well as a bilateral testicular ultrasonography conducted with a high-resolution transducer involving at least 7.5 MHz (expert consensus).
2. We recommend that men with newly diagnosed GCT receive a contrast-enhanced computed tomography (CT) scan of the abdomen/pelvis and chest for staging (A, 5) [e30, e31].
3. In patients with newly diagnosed GCT and contraindications to iodine-containing contrast media, such as allergy, impaired renal function, or thyroid dysfunction, MRI of the abdomen/pelvis is recommended to replace a CT (A, 5) [e30]. In case of the above-mentioned contraindications, a non-contrast-enhanced CT scan of the chest should be performed additionally to investigate pulmonary involvement (B, 5).
Patients of the IGCCCG poor prognosis group as well as patients with excessive beta human chorionic gonadotropin (β-hCG) values, multiple pulmonary metastases, or neurological symptoms should receive an additional cranial MRI (B, 5) [e28].

An FDG-PET/CT is not recommended for routine use in primary staging diagnostics (A, 1a) [e32].

FDG-PET/CT can be considered in seminoma patients with residual tumours larger than 3 cm in diameter and with normal or normalized serum tumour markers after therapy completion (0, 1a) [e32]. The CT as part of the FDG-PET/CT examination should preferably be performed as contrast-enhanced CT (expert consensus).

Timing of FDG-PET/CT for the assessment of serum tumour marker-negative residual tumours in patients with seminoma is at the earliest recommended at 6 weeks after the end of the last cycle of chemotherapy (expert consensus).

FDG-PET/CT is not recommended in patients with a non-seminomatous GCT (expert consensus).

Serum tumour markers (primary diagnostics and staging diagnostics):

We recommend determination of the serum tumour markers α-fetoprotein (AFP), β-hCG, and lactate dehydrogenase [33].

Table 3. Prognostic classification of metastatic GCT [2]a

| Good prognosis group | All of the following: |
|---------------------|-----------------------|
| Non-seminomatous GCT (56% of cases) | Testis/retroperitoneal primary No non-pulmonary visceral metastases AFP <1,000 ng/mL hCG <5,000 IU/L (1,000 ng/mL) LDH <1.5× ULN |
| Seminoma (90% of cases) | Any primary site No non-pulmonary visceral metastases Normal AFP Any hCG Any LDH |

| Intermediate prognosis group | All of the following: |
|-----------------------------|-----------------------|
| Non-seminomatous GCT (28% of cases) | Testis/retroperitoneal primary No non-pulmonary visceral metastases Any of the following: AFP ≥1,000 and ≤10,000 ng/mL or hCG ≥5,000 and ≤50,000 IU/L or LDH ≥1.5 and ≤10× ULN |
| Seminoma (10% of cases) | Any primary site Any non-pulmonary visceral metastases Normal AFP Any hCG Any LDH |

| Poor prognosis group | (At least) one of the following: |
|---------------------|---------------------------------|
| Non-seminomatous GCT (16% of cases) | Mediastinal primary Any non-pulmonary visceral metastases AFP >10,000 ng/mL or hCG >50,000 IU/L (10,000 ng/mL) or LDH >10× ULN |
| Seminoma | No patients classified as poor prognosis |

IGCCCG, International Germ Cell Cancer Collaboration Group; GCT, germ cell tumour; AFP, α-fetoprotein; hCG, human chorionic gonadotropin; LDH, lactate dehydrogenase; CT, computed tomography; IGCCCG, International Germ Cell Cancer Collaboration Group; LH, luteinizing hormone; FSH, follicle-stimulating hormone. We recommend advising all patients on the cryopreservation of sperm preferably before the primary surgical intervention and, if necessary, arrange consultation with an andrologist or a fertility centre.

Table 4. Recommended examinations for primary staging

| Test | Patients |
|------|----------|
| Physical examination | All patients |
| Ultrasonography of the testes | All patients |
| Serum tumour markers | All patients |
| AFP | |
| β-hCG | |
| LDH | |
| Chest/abdominal/pelvic CT | Symptomatic patients and patients in the IGCCCG poor prognosis group, or patients with multiple pulmonary metastases and/or high β-hCG values (>5,000 IU/L) |
| Cranial MRI | Patients with unfinished family planning, active or postponed desire for offspring as well as for ruling out/detecting hypogonadism |
| Fertility tests | |
| Sperm analysis/cryopreservation | |
| Testosterone | |
| LH | |
| FSH | |

AFP, α-fetoprotein; hCG, human chorionic gonadotropin; LDH, lactate dehydrogenase; CT, computed tomography; IGCCCG, International Germ Cell Cancer Collaboration Group; LH, luteinizing hormone; FSH, follicle-stimulating hormone.
hydrogenase in patients with suspected GCT prior to orchiectomy (expert consensus).

10 In patients with preoperatively elevated serum tumour marker levels, we recommend to monitor AFP, β-hCG, and lactate dehydrogenase postoperatively every 5–7 days until normal values or nadir is reached, or until serum tumour markers increase (expert consensus).

Surgically determined diagnosis/inguinal exploration of the testis (including organ preservation):

11 If a GCT is suspected, we recommend inguinal testicular exploration and, in case of evidence for a malignancy, radical orchiectomy of the respective testis is recommended (A, 5) [e28, e33].

12 In the presence of a healthy contralateral testis, organ-preserving excision is not recommended in case of a unilateral malignant GCT (with the exception of a teratoma without accompanying germ cell neoplasia in situ [GCNIS]) (expert consensus).

13 Organ-preserving tumour excision is recommended in patients with a bilateral GCT, with monorchid tumour, in case of stromal or other benign tumours (epidermoid cyst and monodermal teratoma), respectively (expert consensus).

Organ-preserving tumour resection in GCT:

14 For eradication of GCNIS after organ-preserving surgery in monorchid tumour patients, we recommend adjuvant radiation of the affected testis with 18–20 Gy. As the development of a solid GCT from GCNIS may take several years, we recommend the discussion of regular sonographic controls with patients who declined radiotherapy for the purpose of paternity and to those who are under surveillance (expert consensus).

15 We recommend that patients with increased risk for contralateral GCNIS undergo contralateral biopsy at the time of orchiectomy after appropriate counselling (A, 4) [e34].

Pathological examination of the testicular tissue (including GCNIS):

16 For patients with ultrasonographic testicular microlithiasis and no other risk factors, testicular biopsy is not recommended (A, 2a). A testicular biopsy can be considered in patients with microlithiasis presenting one of the following additional parameters: infertility, previous tumour disease of the testicle, first-degree relative with GCT, history of maldescended testis, or testicular atrophy with a sonographic testis volume <12 mL (0) [e35].

17 We recommend that the pathohistological report on the testicular specimen includes the following information: labelling of side, testicular size, maximum tumour size (in 3 dimensions), macroscopic features of the epididymis, spermatic cord and tunica vaginalis, tumour in the surgical margins (yes/no), histological subtype with specification of individual components and percentage according to WHO 2016, tumoral venous and/or lymphatic invasion (yes/no), invasion of the tunica albuginea (yes/no), rete testis (yes/no), soft tissue of the hilum (yes/no) and the epididymis or spermatic cord (yes/no), GCNIS in non-tumorous parenchyma (yes/no), and pT category according to the 2017 TNM classification (expert consensus).

Fertility and cryopreservation of spermatozoa:

18 GCT therapy impairs the patients’ fertility to a variable extent, depending on the treatment regimen applied and the initial situation (statement, expert consensus).

19 We recommend that if a GCT is suspected, cryopreservation of spermatozoa is offered to the patient before treatment initiation (before orchiectomy, at the latest before chemotherapy or radiotherapy) (A, 5) [e28, e36].

20 We recommend that patients with azoospermia at the time of scheduled cryopreservation prior to therapy are offered bilateral testicular sperm extraction (if possible simultaneously with orchiectomy) (A, 5) [e36].

Staging diagnostics and classification:

21 To characterize the anatomical spread of the GCT, we recommend the use of the current TNM classification (expert consensus).

Chapter 7: Prognosis

1 The stage-independent 5-year survival rate in patients with GCT (in Germany) is 97.9% for seminoma and 94.9% for non-seminomatous GCT (statement, 2b) [1].

2 For stage I GCT patients, the cancer-specific 10-year survival rate is 99.7% and the 10-year overall survival rate is 95–99% (statement, 2b) [e37].

3 The 5-year survival rates for metastatic GCT are 86–95% for patients in the good prognosis group, 72–85% for the intermediate prognosis group, and 48–64% for the poor prognosis group (statement, 2b) [2, e38].

4 In non-seminomatous GCTs, lymphovascular invasion represents the only prospective multivariate analysis-validated risk factor for recurrence in clinical stage I (cSI) disease (statement, 2b). In seminoma, tumour size correlates positively with the occurrence of recurrences in patients with cSI disease. Data on the impact of stromal rete testis infiltration are inconsistent. With a tumour size of less than 4 cm and a lack of stromal rete testis infiltration, the risk of recurrence is minimal (<5%) (statement, 1b) [e39–e46].

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We recommend the classification of metastatic GCT to be based on the prognostic criteria of the IGCCCG (expert consensus).

Chapter 8: Primary and First-Line Therapy

Here, in part I, we outline the primary surgical treatment of the testis and the treatment of the localized stages. Part II of the publication will summarize all advanced disease stages.

Inguinal Exposure

In case of a testicular suspicious mass, the surgical inguinal exposure of the testis constitutes both diagnostics and therapy. Depending on the initial findings, it is performed with or without frozen section-based intraoperative assessment of the dignity of the tumour for either an organ-preserving approach (enucleation) or complete removal of the testis (orchietomy), including the spermatic cord. In life-threatening disease due to metastatic spread, orchiectomy can be postponed and performed later in the course of the disease.

Rare giant tumours with a diameter of more than 10 cm are an exception. In these cases, an inguino-scrotal approach including resection of the respective scrotal skin area should be applied [e47].

There are no data available that would justify scrotal orchiectomy as an exclusion criterion for a subsequent surveillance strategy. Cryopreservation of sperm should be performed before taking the first therapeutic step, which usually is orchiectomy (see above).

The insertion of a silicone implant during orchiectomy procedure should be discussed with and offered to the patient. This is not associated with an increased risk of infection. The overall surgical complication rate is about 7% [e48]. The satisfaction rates (assessed by structured patient interviews) exceed 80% among these patients [e49, e50].

Therapy of GCNIS

1 In monorchid GCNIS, local radiotherapy with 18–20 Gy attains eradication of GCNIS cells in more than 95% of cases (statement, 2b) [e53, e54].

2 In case of active surveillance of GCNIS, an invasive GCT develops within 5 years in 50% of cases (statement, expert consensus).

3 In case of unilateral GCNIS in the presence of a healthy contralateral testis, either orchiectomy or sonographic follow-up (active surveillance) are recommended (expert consensus).

4 In case of bilateral GCNIS, both testes should be irradiated (expert consensus).

5 After adjuvant chemotherapy, more than 50% of GCNIS cases develop into a manifest tumour. Additional radiotherapy should be performed (B, 2b) [e54, e55].

6 Radiotherapy of GCNIS-affected testes causes sterility. Prior cryopreservation of sperm (from the ejaculate or in the presence of azoospermia by means of a testicular sperm extraction) is recommended to be discussed with the patient (expert consensus).

7 After local radiotherapy with 18–20 Gy, long-term hypogonadism can arise in 30% of cases. Regular controls

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Table 5. Recurrence rates in studies with non-seminomatous GCT patients in cSI [26–31]

| Reference | N  | Therapy    | Recurrence rate, % | Mean follow-up, months |
|-----------|----|------------|---------------------|------------------------|
| [26]      | 43 | 2× PEB     | 2.5                 | 42                     |
| [27]      | 114| 2× PEB     | 1.7                 | 48                     |
| [28]      | 42 | 2× PEB     | 4.8                 | 79                     |
| [29]      | 58 | 2× PEB     | 1.7                 | 93                     |
| [30]      | 40 | 2× PEB     | –                   | 113.2                  |
| [31]      | 142| 2× PEB     | 0.7                 | 79                     |

GCT, germ cell tumour; cSI, clinical stage I.

Table 6. Recurrence rates of chemotherapy versus surveillance in studies with non-seminomatous GCT patients in cSI [22–24, 32, 33]

| Reference | N  | Therapy     | Recurrence rate, % |
|-----------|----|-------------|---------------------|
| [22]      | 234| Surveillance| 30.0                |
|           | 148| 1–2× PEB    | 4.0 (6.5+3.6)       |
| [23]      | 231| 2× PEB (hr) | 0.8                 |
|           | 358| surveillance (lr) | 19.0            |
| [32]      | 60 | 2× CVB (hr) | 1.6                 |
|           | 124| surveillance | 12.9               |
|           | 40 | 1× CVB (lr) | 10.0                |
| [33]      | 258| 1× PEB (hr) | 3.1                 |
|           | 255| 1× PEB/surveillance (lr) | 1.6          |
| [24]      | 167| 2× PEB (hr) | 1.2                 |
|           | 287| surveillance (lr) | 16.7            |

GCT, germ cell tumour; cSI, clinical stage I; hr, high risk; lr, low risk.
of the serum testosterone level are recommended (A, 2b) [e53, e54].

8 Three or more cycles of cisplatin-based chemotherapy lead to eradication of GCNIS in 60–70% of cases. A control biopsy should be performed before radiotherapy or, if the subsequent procedure depends on the result, 2 years after chemotherapy at the earliest (B, 2b) [e54, e55].

Seminoma in non-metastatic cSI:

9 In seminoma, the tumour size correlates positively with the presence of occult metastases in cSI. Data on stromal rete testis infiltration are inconsistent. With a tumour size of <4 cm and absence of rete testis infiltration, the risk of occult metastases is minimal (<5%) (statement, 2a) [e46].

10 All treatment options (active surveillance, adjuvant chemotherapy with carboplatin, and adjuvant radiotherapy) achieve the same survival rates with stage-adapted treatment of the disease in case of recurrence (statement, GRADE: very low quality of evidence) [e56–e63].

11 Age, comorbidities, patient preferences, and patient compliance are recommended to be considered for the choice of therapy (expert consensus).

12 Patients with seminoma and cSI are recommended to be followed up by active surveillance and in case of recurrence, they are treated according to stage-specific guidelines (A, GRADE: very low quality of evidence) [e57, e59, e60, e64–e67].

13 In patients with a tumour diameter >4 cm, adjuvant therapy can be considered in individual cases, especially if compliance concerns or psychological distress of the patient are suspected. One to 2 cycles of carboplatin (dosed according to AUC 7) or, alternatively, radiotherapy of the para-aortic region with 20 Gy are options for adjuvant therapy. Both therapeutic modalities offer specific advantages and disadvantages, which are recommended to be discussed in detail with the patient (0, 2b) [e63, e68].

Non-seminomatous GCT in non-metastatic cSI:

14 The lymphovascular invasion of tumour cells in the primary tumour is the most important risk factor for occult retroperitoneal metastases. In the low-risk situation (without lymphovascular invasion), the risk of occult metastases is about 15%, and in the high-risk situation (with lymphovascular invasion), it is up to 50% (statement, 2b) [e37, e39–e44, e45, e69].

15 In the low-risk situation, active surveillance is recommended (A, 2b) [e70–e72].

16 In the high-risk situation, the treatment modalities including 1 cycle of PEB versus active surveillance should be discussed with the patient. One cycle of PEB reduces the risk of recurrence from 50 to 3%. The overall survival of both groups is the same. In case of individual patient’s reluctance to surveillance and adjuvant chemotherapy, or in case of contraindications against chemotherapy, primary nerve-sparing retroperitoneal lymph node dissection with modified field boundaries should be considered alternatively. It is recommended to perform this intervention at centres with proven experience (B, GRADE: moderate quality of the evidence) [e72–e74].

See Tables 5–8 for the corresponding recurrences rates.

Stage I disease with serum tumour marker increase:

17 In patients with persistently elevated serum tumour markers after orchiectomy (cSI), it is recommended to
rule out non-GCT-related causes for serum tumour marker elevation (expert consensus). 18 Patients with persistent tumour marker elevations after orchiectomy and negative imaging, in whom other causes for serum tumour marker elevation have been ruled out, should be closely monitored prior to treatment decision-making until either an unequivocal continuous marker increase of AFP or β-hCG becomes evident or a detectable lesion upon imaging becomes evident (expert consensus).

Key Messages of Part I

- GCTs have an incidence of 10:100,000 men/year but represent 25% of cancer cases diagnosed in the age group of 20–40 years.
- The patient’s prognosis essentially depends on tumour histology, clinical stage, age, and quality of care.
- Stage I GCTs have excellent 5-year survival rates with 97.9% for seminoma and 94.9% for non-seminomatous GCT.
- Five-year survival rates for metastatic GCTs are 86–95% in the good prognosis group, 72–85% for the intermediate prognosis group, and 48–64% for the poor prognosis group.
- We recommend the 2016 WHO classification for histopathological evaluation of GCTs.
- We recommend the classification of metastatic GCTs based on the prognosis criteria of the IGCCCG classification.
- The patients’ fertility is impaired by GCT therapy to variable extent, depending on the treatment modality and the initial situation. Therefore, we recommend that cryopreservation of spermatozoa is offered to the patient before any treatment.
- FDG PET/CT is not recommended for routine use in primary staging.
- In monorchid GCNIS, local radiotherapy with 18–20 Gy eradicates GCNIS cells in more than 95% of cases.
- CNIS develop invasive cancer. In these cases, additional radiotherapy should be performed.
- In seminoma cSI, all treatment options (active surveillance, adjuvant chemotherapy with carboplatin, and adjuvant radiotherapy) achieve the same survival rates.
- Patients with seminoma cSI are recommended to be followed up by active surveillance and treated stage-adapted in the case of recurrence.

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