Incidence and epidemiology

The crude incidence of Hodgkin lymphoma (HL) in the European Union is 2.3, the mortality 0.4 cases/100,000/year. Young adults aged 20–40 years are most often affected. Slightly more men than women are diagnosed with HL. Histologically, classical HL (cHL) accounting for ~95% of all HL cases is distinguished from nodular lymphocyte-predominant HL (NLPHL) representing ~5% of all HL cases.

Diagnosis

Pathological diagnosis should be made according to the World Health Organization (WHO) classification from an excisional lymph node biopsy or a sufficiently large surgical specimen to provide enough material for fresh frozen and formalin-fixed samples. In cHL, the presence of Hodgkin and Reed–Sternberg (HRS) cells is disease-defining, whereas the detection of lymphocyte predominant (LP) cells is required for the diagnosis of NLPHL. The immunophenotype of the malignant cells in cHL and NLPHL differs significantly. In contrast to HRS cells that stain consistently positive for CD30 and CD15, occasionally positive for CD20 and negative for CD45, LP cells are characterised by the expression of CD20 and CD45 but lack CD15 and CD30.

Staging and risk assessment

The diagnostic work-up is shown in Table 1. The medical history including the presence of B symptoms (fever, drenching night sweats, unexplained weight loss >10% over 6 months) and other disease-related symptoms such as fatigue, pruritus and alcohol-induced pain as well as the results of a physical examination should be recorded [1].

Chest X-ray and a contrast-enhanced computed tomography (CT) scan of the neck, the chest and the abdomen are mandatory. In addition, a baseline whole-body positron emission tomography (PET) should be carried out according to the recommendations for staging and response assessment in lymphoma, if this diagnostic tool is available [1, 2].

Given the high sensitivity of PET–CT for bone marrow involvement, a bone marrow biopsy is no longer indicated in patients undergoing PET–CT evaluation [III, B] [1–3]. However, bone marrow biopsy must be carried out if PET–CT is not available.

Full blood cell count, erythrocyte sedimentation rate (ESR) testing and blood chemistry analysis including C-reactive protein (CRP), alkaline phosphatase (AP), lactate dehydrogenase (LDH), liver enzymes and albumin are obligatory. Screening for hepatitis B (HBV), hepatitis C (HCV) and human immunodeficiency virus (HIV) is compulsory [II–III, A].

Staging is carried out according to the Ann Arbor classification in consideration of defined clinical risk factors. After completion of staging, patients are allocated to one of three categories (limited, intermediate and advanced stages) [II–III, A]. Table 2 illustrates the European Organisation for Research and Treatment of Cancer (EORTC)/Lymphoma Study Association (LYSA) and the German Hodgkin Study Group (GHSG) definitions of limited, intermediate and advanced stages.

To identify patients at increased risk for acute and/or long-term complications and to generate baseline values for future...
measurements, cardiac and pulmonary function tests should be carried out before the start of treatment.

As chemotherapy (ChT) and abdominal radiotherapy (RT) can cause permanent infertility, reproductive counselling and consideration of sperm banking, oocyte collection or ovarian tissue cryopreservation should be offered to patients of reproductive age before treatment.

**Treatment of cHL**

**Limited-stage disease (Figure 1)**

Combined-modality treatment consisting of a brief ChT followed by RT was shown to result in superior tumour control compared with RT alone [I, A] [4, 5].

Two or three cycles of doxorubicin/bleomycin/vinblastine/ dacarbazine (ABVD) (Table 3) followed by conventionally fractionated RT represent the standard of care for limited-stage HL. A large multicentre trial in which patients were randomly assigned to either two or four cycles of ABVD followed by either 20 or 30 Gy involved-field RT (IFRT) showed similar freedom from treatment failure (FFTF) and overall survival (OS) rates for all treatment groups. Thus, the least toxic approach consisting of two cycles of ABVD followed by 20 Gy IFRT appears to be sufficient for limited-stage HL [I, A] [6]. Comparable disease control was also observed in a randomised trial comparing IFRT at doses of either 20 or 36 Gy in patients achieving a complete remission after six cycles of ChT with the outdated epirubicin/bleomycin/vinblastine/prednisone (EBVP) protocol [7]. However, the current RT guidelines of the International Lymphoma Radiation Oncology Group (ILROG) recommend involved-site RT (ISRT) after ChT in limited stages. Although ISRT has not been randomly compared with IFRT in a prospective study, there is accumulating evidence of excellent disease control with these smaller RT fields [8].

The question of whether RT can be omitted in selected patients with complete metabolic response at interim PET is a matter of debate. Several randomised trials addressing this issue have been conducted within the last years. The available data consistently demonstrate a progression-free survival (PFS) advantage for patients treated with combined-modality approaches despite a negative interim PET (defined as a Deauville score $\leq 2$ within the RAPID and H10 studies). Thus, a patient group that can be safely treated with ChT alone could not yet be defined [I, A] [9, 10]. However, as patients treated with ChT alone still have a good overall prognosis, this approach may be offered to individual patients when the late risk of delivering RT is thought to outweigh the...
short-term benefit of improved disease control. Early treatment intensification appears to improve the prognosis of patients with a positive interim PET (defined as a Deauville score/C21 within the H10 study). A large randomised study including patients with limited- and intermediate-stage HL revealed a significantly reduced relapse rate in those patients with a positive interim PET after two cycles of ABVD who completed ChT with two cycles of bleomycin/etoposide/doxorubicin/cyclophosphamide/vincristine/procabazine/prednisone in escalated dose (BEACOPPescalated) (Table 4) instead of one (limited stages) or two (intermediate stages) additional cycles of ABVD before involved-node RT (INRT) [10]. However, the study was not powered to analyse patients with limited- and intermediate-stage disease separately. Patients with a

**Table 3. The ABVD regimen**

<table>
<thead>
<tr>
<th>Dose (mg/m²)</th>
<th>Administration</th>
<th>Days</th>
</tr>
</thead>
<tbody>
<tr>
<td>Doxorubicin 25</td>
<td>i.v.</td>
<td>1 + 15</td>
</tr>
<tr>
<td>Bleomycin 10</td>
<td>i.v.</td>
<td>1 + 15</td>
</tr>
<tr>
<td>Vinblastine 6</td>
<td>i.v.</td>
<td>1 + 15</td>
</tr>
<tr>
<td>Dacarbazine 375</td>
<td>i.v.</td>
<td>1 + 15</td>
</tr>
</tbody>
</table>

Recycle: day 29.

ABVD, doxorubicin/bleomycin/vinblastine/dacarbazine; i.v., intravenous.
two (intermediate stages) additional cycles of ABVD before cycles of BEACOPPescalated instead of one (limited stages) or PET after two cycles of ABVD who completed ChT with two patients with limited- and intermediate-stage HL revealed a score of patients with a positive interim PET (defined as a Deauville score). Early treatment intensification appears to improve the prognosis towards a better OS with BEACOPPescalated when compared with ABVD. However, several non-randomised studies have suggested that patients with advanced HL who have a positive interim PET (defined as a Deauville score ≥ 4 within the HD18 and SWOG S0816 studies and ≥ 3 within the HD0801 study) have a better prognosis after switching from ABVD to intensified protocols than after continued treatment with ABVD [12]. A recent randomised study demonstrated an improved modified 2-year PFS after six cycles of brentuximab vedotin in combination with AVD (A-AVD) as compared with standard ABVD. However, A-AVD was associated with an increased rate of neuropathy and haematological toxicity. Thus, longer follow-up is required to draw final conclusions in terms of the A-AVD regimen [13, 18].

Intermediate-stage disease (Figure 2)

Intermediate-stage HL is usually treated with combined-modality approaches.

Four cycles of ABVD followed by conventionally fractionated RT at 30 Gy are widely considered standard of care for intermediate-stage HL [I, A] [5]. In patients ≤ 60 years who are eligible for a more intensive treatment, this standard is challenged by a protocol consisting of two cycles of BEACOPPescalated followed by two cycles of ABVD and RT at 30 Gy. After a median follow-up of 43 months, FFTF with this protocol was superior in comparison with four cycles of ABVD followed by RT. An advantage in OS could not be shown [I, B–C] [11].

Although no results of a randomised study comparing both RT fields are available to date, the ILROG guidelines recommend ISRT instead of IFRT after ChT in intermediate stages [8].

The question of whether RT is dispensable in intermediate-stage patients with complete metabolic response at interim PET is unanswered. A large randomised study failed to demonstrate non-inferiority of ChT alone as compared with combined-modality treatment in patients with a negative interim PET (defined as a Deauville score ≤ 2 within the H10 study) [I, A] [10]. However, as patients treated with ChT alone still have a good overall prognosis, this approach may be offered to individual patients when the late risk of delivering RT is thought to outweigh the short-term benefit of improved disease control. Early treatment intensification appears to improve the prognosis of patients with a positive interim PET (defined as a Deauville score ≥ 3 within the H10 study). A randomised study including patients with limited- and intermediate-stage HL revealed a significantly reduced relapse rate in patients with a positive interim PET after two cycles of ABVD who completed ChT with two cycles of BEACOPPescalated instead of one (limited stages) or two (intermediate stages) additional cycles of ABVD before INRT [10]. Patients with a positive interim PET after two cycles of ABVD should be treated with two cycles of BEACOPPescalated before ISRT [I, A].

Due to the relevant bleomycin-induced toxicity observed in older individuals receiving more than two cycles of ABVD, bleomycin should be discontinued after the second ChT cycle in patients > 60 years [III, B–C] [12].

Advanced-stage disease (Figure 3)

Advanced-stage HL is usually treated with ChT alone. Additional RT is confined to patients with residual disease after ChT.

Patients ≤ 60 years are treated with either ABVD (six cycles) or BEACOPPescalated (four to six cycles), optionally followed by localised RT [I, A] [13, 14]. When ABVD is applied, the omission of bleomycin, i.e. the use of doxorubicin/vinblastine/dacarbazine (AVD) in cycles 3–6 in the case of a negative interim PET (defined as a Deauville score ≤ 3 within the RATHL study) after two cycles of ChT should be considered, especially in elderly patients and those at an increased risk for lung toxicity, although a randomised multicentre study was not able to exclude a PFS difference of > 5% at 3 years [I, A] [15]. The question of whether consolidating RT can be safely omitted in patients who have a negative PET after two cycles of ABVD or at the end of ChT has also not yet been definitively answered. There is no randomised study evaluating the role of early treatment intensification in advanced-stage patients who have a positive interim PET after two cycles of ABVD. However, several non-randomised studies have suggested that patients with advanced HL who have a positive interim PET (defined as a Deauville score ≥ 4 within the RATHL and SWOG S0816 studies and ≥ 3 within the HD0801 study) have a better prognosis after switching from ABVD to intensified protocols than after continued treatment with ABVD [II, B] [16–17].

A recent randomised study demonstrated an improved modified 2-year PFS after six cycles of brentuximab vedotin in combination with AVD (A-AVD) as compared with standard ABVD. However, A-AVD was associated with an increased rate of neuropathy and haematological toxicity. Thus, longer follow-up is required to draw final conclusions in terms of the A-AVD regimen [I–II, C] [18].

In patients receiving BEACOPPescalated, treatment can be safely reduced to a total of only four cycles in the case of a negative interim PET (defined as a Deauville score ≤ 2 within the HD18 study) compared with a total of six cycles for PET-positive patients [I, A] [19]. In addition, RT can be restricted to the patients with PET-positive (defined as a Deauville score ≥ 3 within the HD15 study and most of the HD18 study and a Deauville score ≥ 4 within a part of the HD18 study) residual lymphoma ≥ 2.5 cm after four and six cycles of BEACOPPescalated, respectively [I, A] [14, 19]. Several trials randomly comparing ABVD and BEACOPPescalated have shown a superior tumour control and a non-significant trend towards a better OS with BEACOPPescalated [20, 21]. A network meta-analysis including 9993 patients also revealed a significantly better OS with BEACOPPescalated when compared with ABVD. The survival benefit was 10% at 5 years [I, A] [22]. However, given the relevant acute toxicity of BEACOPPescalated, appropriate surveillance and supportive care must be available when this protocol is used. In patients > 60 years, the BEACOPP regimen should not be given, as an increased rate of treatment-related mortality has
been observed in this age group [II, A] [23]. Thus, ABVD-based ChT represents the standard of care for older HL patients who are fit enough for multi-agent ChT. However, due to the relevant bleomycin-induced toxicity observed in older individuals receiving more than two cycles of ABVD, bleomycin should be discontinued after the second ChT cycle in this patient group [III, B–C] [12].

**Relapsed disease**

For most patients with refractory or relapsed HL, the treatment of choice consists of high-dose ChT (HDCT) followed by autologous stem cell transplantation (ASCT) [I, A] [24]. High-risk patients may benefit from tandem ASCT [III, B] [25]. Consolidating treatment with the antibody-drug conjugate brentuximab vedotin following HDCT and ASCT was shown to improve the tumour control in patients presenting with at least one of the following risk factors: primary disease progression, early disease recurrence < 12 months after the end of first-line treatment and extranodal disease at the time of relapse [II, B] [26].

Salvage regimens such as dexamethasone/high-dose cytarabine/cisplatin (DHAP), ifosfamide/gemcitabine/vinorelbine (IGEV) or ifosfamide/carboplatin/etoposide (ICE) are given to reduce the tumour burden and mobilise stem cells before HDCT and ASCT [II–III, A] [27–29]. In some patients, single-agent brentuximab vedotin results in a negative PET and may therefore be sufficient as salvage therapy before HDCT and ASCT [III, B] [30]. Achieving a negative PET should be the goal of salvage therapy, irrespective of the applied protocol, because a complete metabolic response before HDCT and ASCT was shown to be associated with an improved clinical outcome [III, B] [31]. The role of RT before HDCT and ASCT is not defined. However, its use may be discussed in patients with single PET-positive lymph nodes after salvage therapy [IV, C] [32].

**Figure 2.** Therapeutic algorithm for newly diagnosed, intermediate-stage HL in patients ≤ 60 years. The figure includes one approach not guided by interim PET, based on the GHSG HD14 study (left) and one PET-guided approach based on the EORTC/LYSA/FIL H10 study (right).

ABVD, doxorubicin/bleomycin/vinblastine/dacarbazine; BEACOPPesc, bleomycin/etoposide/doxorubicin/cyclophosphamide/vincristine/procarbazine/prednisone in escalated dose; CT, computed tomography; EORTC, European Organisation for Research and Treatment of Cancer; FIL, Fondazione Italiana Linfomi; GHSG, German Hodgkin Study Group; HL, Hodgkin lymphoma; ISRT, involved-site radiotherapy; LYSA, Lymphoma Study Association; PET, positron emission tomography.
Figure 3. Therapeutic algorithm for newly diagnosed, advanced-stage HL in patients ≤ 60 years.
The figure includes one approach not guided by interim PET (left) and two PET-guided approaches based on the GHSG HD18 study (middle) and the RATHL study (right).
ABVD, doxorubicin/bleomycin/vinblastine/dacarbazine; AVD, doxorubicin/vinblastine/dacarbazine; BEACOPPesc, bleomycin/etoposide/doxorubicin/cyclophosphamide/vincristine/procarbazine/prednisone in escalated dose; CT, computed tomography; GHSG, German Hodgkin Study Group; HL, Hodgkin lymphoma; PET, positron emission tomography; RT, radiotherapy.
Annals of Oncology

The use of brentuximab vedotin represents an option in patients failing ASCT. After a pivotal phase II study including 102 cHL patients with relapse after HDCT and ASCT had demonstrated an overall response rate (ORR) of 75% with single-agent brentuximab vedotin, the drug was approved for the treatment of such patients [III, B]. A recent follow-up analysis of the study revealed a 5-year OS estimate of 41% for the patients included in the study. However, most patients received additional treatment following brentuximab vedotin. The proportion of patients who achieved long-term remission exceeding 5 years without further treatment was 9% [III, B] [33, 34].

Antibodies targeting the programmed cell death protein 1 (PD-1) represent another novel treatment option for patients with multiple relapses. Early-phase studies evaluating anti-PD-1 antibodies have shown high response rates and durable remissions in a relevant proportion of patients with disease recurrence after HDCT followed by ASCT and brentuximab vedotin therapy [III, B] [35, 36]. On the basis of these results, the anti-PD-1 antibodies nivolumab and pembrolizumab were approved by the Food and Drug Administration (FDA) and the European Medicines Agency (EMA) for the treatment of such patients.

Allogeneic stem cell transplantation represents a potentially curative treatment option for patients failing HDCT and ASCT. This approach should be considered and discussed in young, chemosensitive patients in good general condition after careful evaluation of the risk–benefit ratio [III, C] [37, 38].

In patients with multiple relapses who have no other treatment options, acceptable remission rates, satisfying quality of life and prolonged survival can be achieved with gemcitabine-based palliative ChT and/or regional RT.

In general, patients with multiple relapses should be enrolled in clinical trials evaluating novel agents whenever possible.

**Treatment of NLPHL**

**Stage IA without risk factors**

ISRT at 30 Gy alone is the standard treatment for stage IA NLPHL patients presenting without clinical risk factors [III, A] [39]. Although data from prospective studies are only available for IFRT, the current ILROG guidelines recommend the use of ISRT [8]. Of note, the ISRT fields irradiated in this RT alone approach are larger than the ISRT fields in combined-modality approaches to include potential microscopic regional disease.

**Other stages**

Usually, NLPHL is treated identically to cHL in all patients except for those with stage IA disease presenting without clinical risk factors [III, B] [40]. However, as the malignant LP cells of NLPHL consistently express CD20, the addition of an anti-CD20 antibody may improve treatment efficacy, but prospective data addressing this issue are pending. The largest retrospective study evaluating the combination of an anti-CD20 antibody and conventional ChT revealed promising results with the rituximab/cyclophosphamide/doxorubicin/vincristine/prednisone (R-CHOP) protocol [V, B] [41].

**Relapsed NLPHL**

Even more importantly than in cHL, a renewed biopsy should be obtained in patients with suspected NLPHL relapse before salvage therapy is initiated, because transformation into aggressive non-Hodgkin lymphoma (NHL) must be excluded. According to newer analyses, transformation rates appear to be higher than previously reported [IV, A] [42].

Localised NLPHL relapses can be effectively treated with anti-CD20 antibodies such as rituximab or ofatumumab given as single agent [III, B] [43, 44].

Patients with more disseminated disease at relapse and additional poor-risk features may require more aggressive salvage ChT, possibly combined with an anti-CD20 antibody [III, B] [45]. However, salvage therapy should be chosen individually and be based on factors such as time to relapse, extent of disease at relapse and prior treatment [III, B] [46].

Given the lack of CD30 on the malignant LP cells in NLPHL, brentuximab vedotin does not represent a treatment option in this entity.

**Response evaluation**

If no PET-guided treatment is intended, interim response evaluation by contrast-enhanced CT scan should be carried out before RT in limited and intermediate stages and after four cycles of ChT as well as before RT in advanced stages. If treatment is guided by interim PET, patients who receive ABVD should undergo an interim PET–CT scan after two cycles of ChT, irrespective of the stage at diagnosis. Patients with advanced HL receiving ABVD should also have a PET–CT after the end of ChT. In patients with advanced HL who are treated with BEACOPPescalated, interim PET–CT scans should be carried out after two cycles of ChT and after the end of ChT.

Final staging should be carried out after the completion of treatment. Physical examination, laboratory analyses and contrast-enhanced CT are mandatory. If available, PET–CT should replace CT at final staging according to the guidelines for staging and response assessment in lymphoma [1, 2].

**Prognosis**

With modern treatment strategies, 80%–90% of HL patients achieve permanent remission and can be considered cured.

**Follow-up, long-term implications and survivorship**

History, physical examination and laboratory analysis including full blood cell count, ESR testing and blood chemistry should be carried out every 3 months for the first half year, every 6 months until the fourth year and once a year thereafter [V, B].

CT scans and previously pathological radiographic tests must be carried out once to confirm the remission status. Thereafter, the patients should be followed clinically. Surveillance scans are not indicated unless clinical symptoms occur [1, 2].

The thyroid function (thyroid-stimulating hormone) should be evaluated once a year if the neck had been irradiated. Furthermore,
**Table 5. Summary of recommendations**

**Diagnosis**
- The presence of HRS cells is disease-defining in cHL, the detection of LP cells is required for the diagnosis of NLPHL

**Staging and risk assessment**
- Chest X-ray and a contrast-enhanced CT scan of the neck, chest, and abdomen are mandatory
- A baseline PET should be conducted if this diagnostic tool is available
- A bone marrow biopsy is not indicated in patients undergoing PET–CT evaluation (III, B) but must be carried out if PET–CT is not available
- Full blood cell count, ESR testing and blood chemistry analysis are obligatory. Screening for HBV, HCV and HIV is compulsory (II–III, A)
- Staging is carried out according to the Ann Arbor classification and patients are allocated to one of three categories (limited, intermediate and advanced stages) (II–III, A)
- After staging examinations are completed, HL patients are allocated to distinct risk groups depending on their clinical stage and the presence of clinical risk factors
- Cardiac and pulmonary function tests should be carried out before the start of treatment
- Reproductive counselling and consideration of sperm banking, oocyte collection or ovarian tissue cryopreservation should be offered to patients of reproductive age before treatment

**Treatment of HL**
- HL patients should be treated within clinical trial protocols whenever possible

**Treatment of cHL**
- First-line treatment of cHL patients usually consists of combined-modality approaches (limited-stage and intermediate-stage disease) or ChT alone (advanced-stage disease). Intensity of treatment depends on the patient’s risk profile at diagnosis and the result of an interim PET–CT evaluation (if PET is available)

**Limited-stage disease**
- Combined-modality treatment consisting of a brief ChT followed by RT was shown to result in superior tumour control compared with RT alone (I, A)
- Two or three cycles of ABVD followed by conventionally fractionated RT represent the standard of care for limited-stage HL
- ISRT is recommended instead of IFRT after ChT in limited stages

**Intermediate-stage disease**
- Four cycles of ABVD followed by conventionally fractionated RT at 30 Gy are widely considered standard of care for intermediate-stage HL (I, A). Two cycles of BEACOPPesc followed by two cycles of ABVD and RT at 30 Gy can be proposed to the patients ≤ 60 years who are eligible for a more intensive treatment
- ISRT is recommended instead of IFRT after ChT in intermediate stages

**Limited- and intermediate-stage disease**
- ChT alone may be offered to the individual patients when the late risk of delivering RT is thought to outweigh the short-term benefit of improved disease control
- Patients with a positive interim PET after two cycles of ABVD should be treated with two cycles of BEACOPPesc before ISRT (I, A)
- Bleomycin should not be given for more than two cycles in patients > 60 years (III, B–C)

**Advanced-stage disease**
- Advanced-stage HL is usually treated with ChT alone. Additional RT is confined to the patients with residual disease after ChT
- Patients ≤ 60 years are treated with either ABVD (six cycles) or BEACOPPesc (four to six cycles), optionally followed by localised RT (I, A)
- After two cycles of ABVD, the omission of bleomycin in cycles 3–6 in the case of a negative interim PET should be considered, especially in elderly patients and those at an increased risk for lung toxicity (I, A)
- Patients with advanced HL who have a positive interim PET after two cycles of ABVD could switch from ABVD to BEACOPPesc (II, B)
- After two cycles of BEACOPPesc, PET-negative patients can safely receive only two more cycles compared with PET-positive patients who need four more cycles (I, A)
- RT can be restricted to the patients with PET-positive residual lymphoma ≥ 2.5 cm after four or six cycles of BEACOPPesc (I, A)
- The BEACOPP regimen should not be given to the patients > 60 years (II, A)
- ABVD-based ChT represents the standard of care for older HL patients who are fit enough for multi-agent ChT. Bleomycin should be discontinued after the second ChT cycle in this patient group (III, B–C)

**Relapsed disease**
- For most patients with refractory or relapsed HL, the treatment of choice consists of HDCT followed by ASCT (I, A)
- High-risk patients may benefit from tandem ASCT (III, B)
- Consolidating treatment with brentuximab vedotin following HDCT and ASCT is recommended in patients presenting with defined poor-risk factors (II, B)
- DHAP, IGEV or ICE can be given before HDCT and ASCT (II–III, A)
- In some patients, single-agent brentuximab vedotin may be sufficient as salvage therapy before HDCT and ASCT (III, B)
- Achieving a negative PET should be the goal of salvage therapy irrespective of the applied protocol (III, B)
- RT before HDCT and ASCT may be discussed in patients with single PET-positive lymph nodes after salvage therapy (IV, C)

Continued
testosterone and oestrogen levels should be monitored, particularly in younger patients who had intensive ChT [V, B] [47].

Patients should be asked about symptoms indicating the existence of long-term toxicity, especially affecting the heart and lungs. Cancer screening should be conducted regularly due to the persistently increased risk for the development of haematological and solid second malignancies after HL treatment [48, 49]. Particular attention should be paid to breast cancer screening in female patients who had received chest or axillary irradiation before the age of 40 years. These patients should have a mammography once a year starting 8–10 years after RT. Individuals who were ≤30 years at the time of chest irradiation should have a breast magnetic resonance imaging (MRI) in addition to mammography [V, A] [47].

Methodology

These Clinical Practice Guidelines were developed in accordance with the ESMO standard operating procedures for Clinical Practice Guidelines development http://www.esmo.org/Guidelines/ESMO-Guidelines-Methodology. The relevant literature has been selected by the expert authors. A summary of recommendations is provided in Table 5. Levels of evidence and grades of recommendation have been applied using the system shown in Table 6. Statements without grading were considered justified standard clinical practice by the experts and the ESMO Faculty. This manuscript has been subjected to an anonymous peer review process.

Disclosure

MA has reported being an advisory board member of and received travel grants from Takeda and Bristol-Myers Squibb and research grant from Takeda; MH has reported being an advisor for and received research support from Takeda; TI received honoraria from Takeda; AE received honoraria from Takeda, Bristol-Myers Squibb and Amgen and research funding from Takeda,
Clinical Practice Guidelines

Table 6. Levels of evidence and grades of recommendation (adapted from the Infectious Diseases Society of America–United States Public Health Service Grading System)

<table>
<thead>
<tr>
<th>Levels of evidence</th>
<th>Grades of recommendation</th>
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<tbody>
<tr>
<td>I</td>
<td>A: Strong evidence for efficacy with a substantial clinical benefit, strongly recommended</td>
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<tr>
<td></td>
<td>B: Moderate evidence for efficacy with a limited clinical benefit, generally recommended</td>
</tr>
<tr>
<td></td>
<td>C: Insufficient evidence for efficacy or benefit does not outweigh the risk or the disadvantages (adverse events, costs, ...), optional</td>
</tr>
<tr>
<td></td>
<td>D: Strong evidence against efficacy or for adverse outcome, generally not recommended</td>
</tr>
<tr>
<td></td>
<td>E: Moderate evidence against efficacy or for adverse outcome, never recommended</td>
</tr>
</tbody>
</table>

By permission of the Infectious Diseases Society of America [50].

References


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