Drug-Related Cardiotoxicity for the Treatment of Haematological Malignancies in Elderly

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Abstract: Several publications have focused on the cardiotoxicity of specific classes of haematological therapeutic agents such as anthracyclines and cyclofosfamide. Cardiotoxicity of cancer chemotherapeutics is a problem for patients of all ages, but it increases with age. Toxicity can also be developed months after the last chemotherapy dose, and late reactions can be seen years later when they present new-onset cardiomyopathy. No data are available about the cardiotoxicity of non-chemotherapy agents currently used as preferred therapy for haematological malignancy in elderly. In this review we have provided a summary of the cardiovascular toxic effects produced by different drugs and therapeutic agents. Early identification of patients who are at risk for cardiotoxicity should be a primary goal for haematologists in the development of personalised antineoplastic therapeutic strategies or interventions. Thus, the discovery of new biomarkers to identify patients at a high risk for the development of these complications is a high priority. Although targeted therapies such as imatinib and anti-CD20 antibody, such as rituximab, are considered less toxic and better tolerated by patients compared with classic chemotherapy drugs, certain cardiological complications can be very serious as these agents have been in use for a limited period of time.

Keywords: Haematological malignancies, cardiotoxicity, heart failure, imatinib, rituximab, anthracyclines, cyclofosfamide.

INTRODUCTION

Cardiotoxicity is a frequent side effect of chemotherapeutic agents used in haematological malignancies. Chemotherapy-associated cardiotoxicity varies from transient blood pressure and/or electrocardiographic (EKG) changes to more serious events such as arrhythmias, myocarditis, pericarditis, myocardial infarction and cardiomyopathy, which may end in left ventricular dysfunction (LVD) or congestive heart failure (CHF) [1].

Since cardiotoxicity is often dose-dependent, it has a serious impact on treatment efficacy; cardiac complications of treatment lead to reductions in dose and duration of chemotherapy regimens, potentially compromising clinical efficacy. In fact, aggressive chemotherapy has achieved remission in most types of cancers [2] but its use is often limited or even prohibited in elderly patients because of its cardiotoxicity. In haematological malignancies, anthracycline-based chemotherapy has represented the greatest risk for development of cardiotoxicity in this setting of patients.

The advent of new categories of drugs, like tyrosine-kinase inhibitors, proteosome-inhibitors or monoclonal antibodies has revolutionized the treatment of many blood cancers, such as chronic myeloid leukaemia, lymphoma, myeloma and others. However, although targeted therapies are usually better tolerated than chemotherapy, certain cardiac complications still remain and they can be very serious [3]; moreover, such new drugs are often used in combination with traditional chemotherapy.

Early identification of patients who are at risk for cardiotoxicity should be a primary goal for haematologists in the development of personalized antineoplastic therapeutic strategies or interventions. Purpose of this article is to provide a comprehensive review of the adverse cardiac events associated with the use of the most common (chemo)-therapeutic agents used in haematological elderly patients not eligible for Stem Cell Transplantation (SCT), including new molecular-targeted drugs, and to provide guidance as how to prevent such effects.

A. CHEMOTHERAPY FOR HAEMATOLOGICAL MALIG-NANCES

Haematological cancers represent a heterogeneous setting of diseases that can originate from highly indifferented to mature cells thus giving diseases different characteristics in terms of clinical presentation as well as response to therapy and prognosis. Despite these varieties, analogous chemotherapy has been proven efficacious in clinically different blood cancers that present a common origin. In Table 1 type and duration of the most common therapeutic protocol for treating acute and chronic myeloid and lymphoid neoplasm of adults are reported.

ANTHRACYCLINES

1. Pathogenesis

Anthracyclines have been extensively used for many years to treat a wide spectrum of haematologic malignancies and solid tumours [4]. The anthracyclines, doxorubicin and daunorubicin, are cytostatic antibiotics isolated from fungi belonging to the species Streptomyces. Anthracyclines, such as doxorubicin, and daunorubicin, continue to be some of the most active cytotoxic agents available and have broad activity against a variety of cancers, including acute myeloid and lymphoid leukaemias and solid tumours, such as lymphoma, sarcoma and breast cancer [5-7]. Daunorubicin has demonstrated activity against acute lymphocytic leukaemia and acute myeloid leukaemia.

Anthracycline-induced cardiotoxicity is categorised by the time of presentation as acute, early-onset or late-onset [8, 9]. In acute anthracycline cardiotoxicity, symptoms manifest within hours or days of administration, often presenting as disturbances in intracardiac conducion and arrhythmias [10, 11]. One of the best predictors of cardiotoxicity is the total cumulative dose of anthracyclines [12, 13]; generally, doses of > 400 mg/m² are associated with the greatest risk of cardiac injury. The incidence of doxorubicin-induced early HF is almost 3% at a cumulative dose of 400 mg/m², 7% at a dose of 550 mg/m² and 18% at 700 mg/m² [14, 15]. In addition to the total cumulative dose, the incidence of acute and chronic cardiotoxicity may depend on the rate of anthracycline administration during each session [16], the schedule of delivery [17], the type of anthracycline used [18] and additional underlying risk factors, such as younger age in children and an age of > 65
years in adults, African-American ancestry, female sex and trisomy 21 [19].

The cause of anthracycline-induced cardiotoxicity is probably multifactorial. Free radical-mediated myocyte damage is one of the most thoroughly studied mechanisms by which anthracyclines have been proposed to cause cardiotoxicity [20, 21]. Pathologic findings in chronic anthracycline cardiotoxicity include loss of myofibrils within myocytes, vascular and mitochondrial degeneration and interstitial fibrosis [22, 23]. As myocytes undergo apoptosis and their cardiac growth responses to increasing somatic growth in children become inadequate, the LV walls become thinner, and the remaining cells compensate by becoming hypertrophic to create an LV mass that is as close as possible to normal for body surface area. This compensation means that many anthracycline-treated survivors with abnormal LV structure and function by echocardiography may show only histologic evidence of individual cardiomyocyte hypertrophy [24]. Anthracycline damage to all cardiac structures may begin with the first anthracycline dose [25]. Cardiotoxicity that occurs soon after anthracycline administration may be a harbinger of later toxicity; relatively little data exist about late-onset anthracycline cardiotoxicity in long-term survivors of adult malignancy [24].

Various methods have been recommended for the monitoring of cardiotoxicity in oncology. Recently, the applicability of cardiac troponins and creatine kinase MB CK-MB mass has been investigated in this setting and the results of clinical studies are inconsistent [26]. Horacek et al. [27] suggest that GPBB could be a new promising marker for detection of anthracycline-related cardiotoxicity and probably superior to cardiac troponins.

Table 1. Cardiac Toxicity and Proposed Prevention for the Most Commonly Used Chemo and Biological Therapy in Blood Cancers

<table>
<thead>
<tr>
<th>Type of Blood Cancer</th>
<th>Protocols’ Acronymous</th>
<th>Type of Cardiotoxicity</th>
<th>Prevention of Cardiotoxicity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute Myeloid Leukemia</td>
<td>3+7</td>
<td>congestive heart failure</td>
<td>cardioprotective agents in conjunction with anthracyclines</td>
</tr>
<tr>
<td>Acute Lymphoid Leukemia</td>
<td></td>
<td>congestive heart failure</td>
<td>cardioprotective agents in conjunction with anthracyclines</td>
</tr>
<tr>
<td>Chronic Myeloid Leukemia</td>
<td>Imatinib-dasatinib-nilotinib</td>
<td>Oedema, dyspnea, left ventricular contractile dysfunction</td>
<td>Careful cardiac monitoring</td>
</tr>
<tr>
<td>Non Hodgkin Lymphomas</td>
<td>R-CHOP</td>
<td>supraventricular arrhythmia, tachycardia, complete atrioventricular block</td>
<td>Careful cardiac monitoring</td>
</tr>
<tr>
<td></td>
<td>R-CVP</td>
<td>supraventricular arrhythmia, tachycardia, complete atrioventricular block</td>
<td>Careful cardiac monitoring</td>
</tr>
<tr>
<td></td>
<td>R-ICE</td>
<td>supraventricular arrhythmia, tachycardia</td>
<td>Careful cardiac monitoring</td>
</tr>
<tr>
<td>Chronic Lymphatic Leukemia</td>
<td>FCR</td>
<td>Acute onset heart failure and/or myocarditis</td>
<td>Diuretics, angiotensin-converting enzyme inhibitor</td>
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<tr>
<td>Hodgkin lymphoma</td>
<td>ABVD</td>
<td>Acute onset heart failure</td>
<td>Use of liposomal encapsulation</td>
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<tr>
<td>Myeloma</td>
<td>MP</td>
<td>bradycardia</td>
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<td></td>
<td>MP-THAL</td>
<td>bradycardia</td>
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<td></td>
<td>THAL-DEX</td>
<td>bradycardia</td>
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<td></td>
<td>LEN</td>
<td>atrial fibrillation</td>
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3+7: Cytarabine 100 mg/m² i. v. , Daunorubicin 45 mg/m² i. v.  
R-CHOP: Rituximab 375 mg/m² i. v. , Cyclophosphamide 750 mg/m² i. v. ; Adriamycin 50 mg/m² i. v.; Prednisone 100 mg/m² oral  
R-CVP: Rituximab 375 mg/m² EV; Cyclophosphamide 750 mg/m² EV; Vincristine 1. 4 mg/m² i. v. , Prednisone 100 mg/m² oral  
R-ICE: Rituximab 375 mg/m² i. v. ; Ifosfamide 1 gr/m² (dmax 1, 75g) i. v. ; Cisplatin 25 mg/m² i. v. ; Etoposide 100 mg/m² i. v.  
FCR: Rituximab 375 mg/m²(day 1); Fludarabine 25 mg/m²; Cyclophosphamide 750 mg/m²  
ABVD: Adriamycin 25mg/m² i. v. ; Bleomycin 10U/m² EV; Vinblastine 6 mg/m² i. v. ; Dacarbazine 75 mg/m² i. v.  
LEN: lenalidomide  
MP: Melphalan 10 mg/m² oral; Prednisone 60 mg/m² oral  
MP/Thal: Melphalan 10 mg/m² oral; Prednisone 60 mg/m² oral; Thalidomide 50-100 mg oral  
MPV: Melphalan 9 mg/m² oral; Prednisone 60 mg/m² oral; Bortezomib 1, 3 mg/m² i. v.  
Thal/Dex: Thalidomide 200 mg; Dexamethasone 40 mg
mours, e.g. osteogenic sarcoma, [32] Ewing's sarcoma and Wilm's tumour, [33] and breast carcinoma [34].

The risk of doxorubicin induced congestive heart failure (CHF) increases with the cumulative dose. Cardiotoxicity is reported in 14% to 49% of patients treated for lymphoma, and among patients with NHL, the risk of CHF increases with the patient's age and history of coronary heart disease, valvular heart disease, hypertension, diabetes, cigarette smoking, or obesity [35, 36].

Hershman et Colleagues [37] found that hypertension, a known risk factor for CHF, increases the risk of doxorubicin-related cardiotoxicity. Authors also demonstrated that selection factors may appropriately contribute to the receipt of doxorubicin-based chemotherapy in the elderly. Although late cardiac toxicity is increased with both longer duration of doxorubicin therapy and known predictors of HD, such as advanced age, comorbid conditions, prior heart disease, and diabetes, these other risks did not seem to potentiate the cardiotoxic effects of doxorubicin in Elderly Patients With Diffuse B-Cell Non-Hodgkin’s Lymphoma.

2. How to Prevent Anthracyclines Cardiotoxicity

To maximize the beneficial effects of anthracyclines, much effort has been spent devising new strategies to decrease or prevent their cardiotoxic effects. In addition to various dosing schedules aimed at lowering peak plasma concentrations, an alternative strategy has been liposomal encapsulation. The most commonly used formulation is pegylated liposomal doxorubicin (Doxil). This has a reduced tendency to accumulate in myocardial cells, with a resultant decrease in cardiotoxicity. Unlike traditional doxorubicin, pegylated doxorubicin has lower concentrations of free drug in the bloodstream and limited concentration in myocardial cells [38].

The non-pegylated liposomal anthracycline formulation (Myocet) had at least three randomized controlled trials with promising cardioprotective results. Myocet improves the therapeutic index of doxorubicin by significantly reducing cardiotoxicity and grade 4 neutropenia and provides comparable antitumour efficacy, when used in combination with cyclophosphamide as first-line therapy for MBC [39].

In recent years, novel anthracycline analogues have been developed with the hope that they will be as effective as conventional anthracyclines, but without the risk of cardiotoxicity. Idarubicin, a synthetic derivative of daunorubicin, is more lipophilic and has a longer half-life than daunorubicin [40]. It is more lipophilic and can be administered orally as well as intravenously. Its main metabolite, idarubicinol, is as active as the parent drug. Idarubicin is effective at treating lymphoid and myeloid leukaemias, non-Hodgkin’s lymphoma and advanced breast cancer. Similar to epirubicin, idarubicin causes less cardiotoxicity than doxorubicin [41].

Another approach to preventing anthracycline-induced cardiotoxicity is to use cardioprotective agents in conjunction with anthracyclines to mitigate their cardiotoxic effects. Dexrazoxane (ICRF-187) is a cardioprotective agent approved by the FDA. It is a bisphosphonate, a nonpolar derivative of ethylenediaminetetraacetic acid and a water-soluble positive enantiomer of the racemic drug razoxane. The suggestion that Fe plays an important role in anthracycline cardiotoxicity has been strengthened by observation that the chelator, dexrazoxane (ICRF-187), has a potent cardioprotective effect [42]. Despite the beneficial effects of dexrazoxane, concerns exist about the lack of information on its effect on late onset progressive cardiomyopathy, the lack of conclusive evidence that it reduces overall morbidity and mortality in paediatric patients, and its possible interference with the antitumour efficacy of anthracyclines [39].

ALKYLATING AGENTS

1. Cyclophosphamide

Cyclophosphamide (CY) is a non-cell-cycle-specific alkylating agent. It is a broadly active antineoplastic and immunosuppressant agent used in combination chemotherapy for non-Hodgkin’s lymphoma, leukaemia, Hodgkin’s disease, Burkitt’s lymphoma, multiple myeloma, endometrial cancer, lung cancer and breast cancer [2]. Cardiotoxicity due to CY was reported in 1972 as a complication of bone marrow transplantation [43]. CY causes a direct damage to the vascular endothelium, followed by transudation of the toxic metabolite and the resulting damage to myocytes, interstitial haemorrhage, oedema, and intercapillary microthrombi responsible for ischaemic myocardial damage [44]. Cardiac toxicity caused by CY seems to be due to pharmacokinetics and metabolism of the drug showing tenfold interindividual variation, most probably because of the impact of genetic polymorphism on the activity of liver microsomal enzymes [39]. In contrast to doxorubicin-induced cardiotoxicity, high-dose CY causes acute cardiac dysfunction that is independent of cumulative dose. CY doses greater than 1.5 g/m2 may be associated with cardiotoxic risk [40]. Cardiac toxicity is manifested by acute on set heart failure and/or myocarditis occurring within 1-10 days after the administration of the first dose of CY. Some reports indicated that prior radiation therapy to the mediastinum or chest wall is an independent predictor of cardiotoxicity in patients with lymphoma undergoing high-dose chemotherapy [45]. A total dose ranging from 180 to 200 mg/kg over 2 to 4 days has been reported to cause symptomatic cardiomyopathy [46]. However, a dose as low as 120 mg/kg may also result in cardiotoxicity.

Goldberg et al., found that doses based on body surface area rather than body weight correlate well with incidence of cyclophosphamide-induced cardiotoxicity [47]. The fact that young children have a relatively higher body surface area probably explains the lower incidence and severity of cyclophosphamide-induced cardiotoxicity in them compared to adolescents and adults [47]. Several widely employed combinations of High-Dose (HD) cyclophosphamide are not associated with an increased risk of cardiotoxicity over single-agent HD cyclophosphamide. However, the risk for HD cyclophosphamide-associated cardiac toxicity may be increased by the concomitant administration of cytarabine or mitoxantrone [48].

Very little is known about cardiac toxicity of the treatment regimen with CY as part of stem cell collection followed by two autologous HSCT treatments with myeloablative dosages of MEL given in a very short time. Same et al. [49] showed that MM patients treated with tandem autologous HSCT develop transient, mostly reversible and clinically non- overt neurohumoral activation of heart failure in each phase of treatment.

2. How to Prevent Cyclofosfamide Cardiotoxicity

Although there are no specific predictive parameters of cardiotoxicity, cardiac monitoring in high-risk patients undergoing high dose CY therapy may help to prevent serious cardiotoxicity [50]. It has been reported that cyclosporin-A, administered with CY, prevents cardiotoxicity [30], suggesting that CY induced cardiotoxicity is associated with mitochondrial membrane permeability to Ca2+ and that cyclosporine-A may restore mitochondrial permeability and prevent apoptosis of cardiomyocytes [51, 52]. Therapy with diuretics should be started in the first instance. The addition of an angiotensin-converting enzyme inhibitor in case of ECG and/or two-dimensional echocardiogram (2D echo) evidence of impaired left ventricular contraction, the patient not being hypotensive, should be considered according to established guidelines [53].
B. NON CHEMOTHERAPIC AGENTS

Tyrosin-Kinase Inhibitors (Imatinib, Dasatinib and Nilotinib)

Imatinib, an example of a molecular-targeted drug, has revolutionised treatment of chronic myeloid leukaemia (CML), and is now recommended as standard treatment for the accelerated and chronic-phase disease [54]. The cardiac effect of imatinib is controversial [54, 55]. Cardiac toxicity may range from asymptomatic subclinical abnormalities such as electrophysiological changes and left ventricular ejection fraction decline to life threatening events like congestive heart failure and acute coronary syndrome.

Clinical trials of the agent have reported a relatively high incidence of peripheral oedema (63-66%), some of which has been classified as severe (4-5%). In addition, dyspnea has been reported in 12-16% of treated individuals and has been classified as severe in 4-5% [56]. Cardiomyocytes from subjects who are being treated with imatinib appear to have endoplasmic reticula that have been activated in response to cellular stress, collapsed mitochondrial membrane potential, reduced ATP content, and to be more prone to cell death [56]. However, whether the observed cardiomyocyte cell damage represents clinical cardiotoxicity is still controversial [57], as is the role of the protooncogene abl (a target of imatinib and similar compounds) in the physiology of the cardiomyocyte [56, 57].

Kerkela et al. [58] report ten individuals who developed severe congestive heart failure while on imatinib and they show that imatinib-treated mice develop left ventricular contractile dysfunction. With imatinib treatment, cardiomyocytes in culture show activation of the endoplasmic reticulum (ER) stress response, collapse of the mitochondrial membrane potential, release of cytochrome c into the cytosol, reduction in cellular ATP content and cell death.

It has been reported that individuals treated with imatinib developed severe CHF due to myocyte contractile dysfunction [58]. All patients had their left ventricular ejection fraction (LVEF) calculated by radionuclide imaging before the onset of treatment and after they developed symptoms of heart failure. In a series of 103 patients with CML treated with imatinib [59] and 57 patients with CML not treated with imatinib, no statistical difference was observed between the two groups regarding cardiac symptoms and signs, BNP levels, and echocardiographic measurements. However, peripheral oedema was more frequent in the group that received imatinib.

The early detection and application of the treatment for left ventricular dysfunction is of key importance for the prevention of irreversible myocardial injury. Careful cardiac monitoring and assessment by a cardiologist throughout the course of treatment with those TKIs that exert cardiac toxic effect is of primary importance [3].

Dasatinib is a multitargeted kinase inhibitor against Bcr-Abl, cKit, PDGFR-a and b and the SRC family of kinases. Although clinical trials report high rates of peripheral oedema [60], only a 2% incidence of congestive heart failure as well as arrhythmias (including tachycardia) has been associated with Dasatinib treatment [61]. Nilotinib is an inhibitor of Bcr-Abl, c-KIT and PDGFRa and b receptors; Except for QT prolongation on ECG, no other cardiac event has been reported [62].

RITUXIMAB

Rituximab is a recombinant, chimeric anti-CD20 antibody that has emerged in recent years as an effective therapy for non-Hodgkin’s lymphoma (NHL) and other B-cell malignancies [63]. Rituximab binds to the CD20 antigen expressed on almost all malignant B-cells and leads to their destruction; it is used usually in combination of various protocols of chemotherapy. Treatment with rituximab is generally well tolerated and can be combined with chemotherapy, improving response rate, response duration, and in some cases overall survival of patients with B-cell lymphomas.

The cardiotoxicity of Rituximab has been reported when administered with chemotherapy or as monotherapy. Cardiac complications involving the conduction system, such as supraventricular arrhythmia and tachycardia, frequently occur in rituximab/CHOP therapy [64]. In these cases, it may be difficult to correctly determine which medication, rituximab or anthracycline, cause them. Some authors reported a frequency of cardiac toxicity, manifested as grade 3 and 4 arrhythmias and supraventricular arrhythmias (such as tachycardia and auricular fibrillation or flutter) as high as 5.9% when rituximab is used with CHOP (cyclophosphamide, doxorubicin, oncovin, and prednisone) versus a level of 1.0% in patients who received chemotherapy alone [64]. During events of heart failure, higher levels of tumour necrosis factor-α (TNF-α) and interleukin-6 (IL-6) were observed, though this association might be related to a greater extent to their vasodilating compensating effect, unable to repair the endothelial damage seen in these situations [65].

Other clinical experiences reported unusual cardiac toxicity during rituximab treatment. Cervera et al. [66] described a case of elderly woman treated with Rituximab in monotherapy who experienced a complete atioventricular block (CAVB) after the fifth administration. Authors explained such event likely due to cytokine release by anti-CD 20. Arai Y et al. [67] report herein a patient with mantle cell lymphoma (MCL) presenting ventricular tachycardia during infusion of rituximab. He received rituximab intravenously at a dose of 375 mg/m2 weekly for eight consecutive courses. During the rituximab infusion, electrocardiogram monitoring was performed. The second to seventh courses were eventless, although the first course was associated with pyrexia. However, transient ventricular tachycardia occurred during the infusion of the eighth course. Foran reported that 7% of 131 patients with MCL, immunocytoma, or small B-cell lymphocytic lymphoma who received rituximab as a single agent developed arrhythmia, including brady- and tachyarrhythmias [68]. Cardiac toxicity of Rituximab has been reported when administered with chemotherapy or as monotherapy. The cardiotoxicity of Rituximab has been reported when administered with chemotherapy or as monotherapy. Cardiac complications involving the conduction system, such as supraventricular arrhythmia and tachycardia, frequently occur in rituximab/CHOP therapy [64]. In these cases, it may be difficult to correctly determine which medication, rituximab or anthracycline, cause them. Some authors reported a frequency of cardiac toxicity, manifested as grade 3 and 4 arrhythmias and supraventricular arrhythmias (such as tachycardia and auricular fibrillation or flutter) as high as 5.9% when rituximab is used with CHOP (cyclophosphamide, doxorubicin, oncovin, and prednisone) versus a level of 1.0% in patients who received chemotherapy alone [64]. During events of heart failure, higher levels of tumour necrosis factor-α (TNF-α) and interleukin-6 (IL-6) were observed, though this association might be related to a greater extent to their vasodilating compensating effect, unable to repair the endothelial damage seen in these situations [65].

However, such findings were not reported by others. Kilickap et al. [70] evaluated whether the addition of rituximab to an anthracycline-based regimen (CHOP) resulted in an increase in subclinical cardiac toxicity. Patients who were to receive CHOP or rituximab plus CHOP (R-CHOP) combination chemotherapy with a diagnosis of NHL were included in the study. Diastolic and systolic cardiac parameters, by using two-dimensional echocardiography, conventional pulsed-wave Doppler echocardiography and tissue Doppler echocardiography, were monitored before and after chemotherapy. There were 28 (M/F; 14/14) patients on CHOP and 33 (M/F; 16/17) patients on R-CHOP. Parameters of systolic function such as LVEF and FS did not significantly change in any patient. Doppler parameters of diastolic function such as lateral E and septal E velocity of mitral annulus decreased significantly after therapy (P < 0.001). However, the decrease in diastolic function was similar in both arms (P > 0.05). The addition of rituximab to CHOP chemotherapy does not significantly increase the risk of doxorubicin-induced cardiotoxicity during administration and in the follow-up. Siano M et colleagues have shown that the administration of rituximab with a maximal dose of 700 mg over 60 minutes without steroid premedication is feasible and safe, provided one well-tolerated rituximab dose had been given in the previous 3 months. Patients with any type of B-cell lymphoma were eligible; Cohorts of at least three patients were assigned to rituximab with or without concomitant chemotherapy. An extensive cardiologic evaluation was done concomitantly to evaluate whether the increased infusion rate induced cardiotoxicity. Only one patient showed totally asymptomatic ECG alterations (repolarization disorder) of still uncertain origin. This alteration was correlated to rituximab infusion—although a coronary heart disease was not formally ruled out—and after a reduction
of the infusion rate, no further ECG alterations were reported. In all cases, normal troponin values along with unchanged echocardiograms and Holter ECGs confirmed the lack of cardiotoxic effect of a fast infusion rate. Zaja et al. [71] investigate rituximab efficacy in previously untreated adult ITP patients with a platelet count of 20 x 10^9/L or less. One hundred three patients were randomly assigned to receive 40 mg/d dexamethasone for 4 days with or without 375 mg/m² rituximab weekly for 4 weeks. One patient experienced supraventricular tachycardia during the first administration of rituximab, and 1 patient experienced seizure during the salvage treatment: both were probably related to study drug, and both patients were discontinued from the study. One patient with a previous history of vascular disease had a transitory ischaemic attack not related to study drug during salvage therapy.

**PROTEASOME INHIBITORS**

Bortezomib (Velcade, formerly PS-341) is a selective proteasome inhibitor that shows strong activity in vitro and in vivo against solid and haematologic malignancies [72]. Preclinical study results show that bortezomib suppresses tumour cell growth, induces apoptosis, overcomes resistance to standard chemotherapy agents and radiation therapy, and inhibits angiogenesis. Phase I study results established the antitumour activity of bortezomib, administered alone or in combination with standard chemotherapy agents, in patients with advanced haematologic malignancies or solid tumours, usually without additive toxicities [73].

Enrico O et al. [74] reported cardiotoxicity in haematological patients treated with bortezomib. Eight (11. 6 %) of 69 patients developed serious cardiac side effects requiring medication, hospitalisation or an implant of a pacemaker. All eight patients who experienced cardiotoxicity underwent at least four cycles of bortezomib, strengthening the hypothesis of a direct connection with the drug. Authors explained that bortezomib may simultaneously cause atherosclerotic plaque progression and tendency to rupture, and facilitate ischaemic heart complications by reducing/abrogating myocardial preconditioning.

Takamatsu et al. [75] reported a case of a 79-year-old female patient with multiple myeloma who had no prior cardiac disease history developed an acute myocardial infarction on day 5 after receiving bortezomib and dexamethasone (BD). After treatment of coronary stenoses by stents, she received another course of BD therapy and developed angina pectoris on day 5 after the therapy.

Additionally, there is much experimental evidence to support the crucial role of a deficient proteasome activity in impairing cardiac function by several mechanisms, the most important of which is putatively the accumulation of unfolded, damaged and undergraded proteins inside myocytes (as extensively reviewed by Willis & Patterson, 2006).

Zheng Hu et al. [76] tested the therapeutic efficacies as well as adverse effects of low dose imatinib mesylate (IM) in combination with proteasome inhibitor, Bortezomib (BOR) or proteasome inhibitor I (PSI), in two CML murine models, and investigated possible mechanisms of action on chronic myeloid leukaemia (CML) cells. Their results demonstrated that low dose IM in combination with BOR exerted satisfactory efficacy in prolongation of life span and inhibition of tumour growth in mice, and did not cause cardiotoxicity or body weight loss.

**ANTIANGIOGENETIC DRUGS**

**Thalidomide, Lenalidomide, Pomalidomide**

Angiogenesis is a vital process resulting in the formation of new blood vessels [77]. While several studies have found moderate improvements when treated with angiogenesis inhibitors, greater success is being seen when the inhibition of angiogenesis is combined with other traditional forms of available therapy [78]. Thalidomide, an immunomodulatory and antiangiogenic agent, is useful in the treatment of some haematologic (mainly myeloma and other plasma-cell related disorders) and oncologic diseases. Thalidomide-mechanisms of action are multifactorial based on immunomodulation, antiangiogenesis and cytokines regulation, particularly tumour necrosis factor-alpha. It has been reported that up to 6. 8% of thalidomide-treated patients present bradycardia [79].

Lopez-de la cruz et al. [78] reported their experience with Thalidomide in patients with haematologic diseases. In a 34-month period, 33 patients with different haematologic diseases (multiple myeloma [MM], 20; myelodysplastic syndrome, eight; Waldenström macroglobulinemia, two; non-Hodgkin's lymphoma, two; malignant histiocytosis, one) were treated with thalidomide. Of them, five (15. 1%) had bradycardia, all with MM. Bradycardia was detected with a daily thalidomide dose ranging from 100 to 300 mg and the time patients received thalidomide before cardiac event went from one to 18 months. Palladini et al. [79] evaluated the combination of thalidomide and intermediate-dose dexamethasone (T-Dex) in 31 patients with AL who did not respond to, or whose disease relapsed after, first-line therapy. In this study, symptomatic bradycardia emerged as a common (26%) adverse reaction to thalidomide in patients with AL. Bradycardia was reported in 1 of 16 patients by the Boston group [80] and in 1 of 12 patients by the Mayo Clinic Group [81]. A recent retrospective study performed on 96 myeloma patients treated with thalidomide showed a decrease in heart rate below 60 beats/min in 53% of patients and symptomatic bradycardia in 19% of patients, 23 which initially went unrecognized because studying the heart rate was not within the objectives of the original Trial [82].

Interestingly, despite these findings, few studies suggested the role for thalidomide in the management of chronic heart failure (CHF) in addition to traditional cardiovascular medications. In fact, Thalidomide is a drug with potential immunomodulating and matrix-stabilizing properties, that can improve left ventricular (LV) function in patients with CHF secondary to idiopathic dilated cardiomyopathy (ICDM) or coronary artery disease (CAD). Gullestad et al. [83] reported in their double-blind, placebo controlled study, that thalidomide increases LVEF and that these changes are accompanied by a decrease in LVEDV, suggesting that thalidomide can alleviate adverse myocardial remodelling and improve LV function in CHF. Importantly, this improvement in LV function occurred in the patients who were clinically stable and who, in the majority of cases, were receiving angiotensin-converting enzyme inhibitors or angiotensin II and _-blockers in addition to diuretics.

The use of thalidomide is limited by adverse effects of sedation, constipation, neuropathy and thromboembolism. In order to overpass such side-effects thus producing more potent and less toxic immunomodulators, thalidomide chemical structure has been modified and new compounds such as lenalidomide and pomalidomide were produced. Lenalidomide is approved by the US FDA for the treatment of patients with low-risk myelodysplastic syndrome (MDS) with deletion 5q cytogenetic abnormality [84]. Lenalidomide, unlike thalidomide, does not appear to cause peripheral neuropathy, but it does provoke cardiac arrhythmia such as atrial fibrillation (18% versus 11% with placebo) and had severe haematological effects [85]. In two randomised, doubleblind, multicenter studies, the combination of lenalidomide and dexamethasone (LD) was compared with placebo and dexamethasone (PD) in patients with MM who had received at least one prior therapy [86]. The most clinically relevant grade 3 and 4 adverse events that occurred more frequently in the LD arm were neutropenia, thrombocytopenia, deep vein thrombosis, pulmonary embolism, and atrial fibrillation.

Immunomodulatory drugs (IMiDs) including thalidomide and lenalidomide have a proven role in the treatment of multiple myeloma (MM) and have significant activity in myelofibrosis (MF). Pomalidomide is the newest IMiD and appears to be promising for use in both MM and MF. IMiDs are hypothesised to act through multiple mechanisms including effects on angiogenesis and cyto-
CONCLUSIONS

Introduction of high-dose (HD) chemotherapy has improved the clinical outcome of patients with chemosensitive tumours, such as non-Hodgkin’s lymphoma, Hodgkin’s disease, multiple myeloma and leukaemias [88, 89]. However, cardiac toxicity has been frequently reported, particularly in HD cyclophosphamide-containing regimens, thus limiting their use especially in older patients.

Management of cardiac toxicity of chemotherapeutic agents is crucial since the increasing pool of long-term haematological cancer survivors. Preventative strategies that have met with some success have included the use of less cardiotoxic analogues such as epirubicin and liposomal anthracycline preparations. The role of investigational markers of myocardial injury, such as troponin T or brain natriuretic peptide, remains of great interest but still without robust evidences. In table 1 cardiotoxicity and its prevention in the most common antineoplastic regimens are reported.

The development of the so-called “targeted therapies”, particularly drugs that inhibit the activity of tyrosine kinases, has become a remarkable progress in the treatment of neoplastic diseases. Recent data has shown that some of these therapies are associated with certain cardiotoxicity ranging from asymptomatic mild left ventricular dysfunction to congestive heart failure through different mechanisms.

In conclusion, old and new therapies for managing haematological malignancies are associated with cardiotoxicity; knowledge of any potential effect on cardiac function is crucial for the choice of the best regimen specially in elderly patients.

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