

## Core Curriculum

# SCAI Expert Consensus Statement: Evaluation, Management, and Special Considerations of Cardio-Oncology Patients in the Cardiac Catheterization Laboratory (Endorsed by the Cardiological Society of India, and Sociedad Latino Americana de Cardiologia Intervencionista)

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In the United States alone, there are currently approximately 14.5 million cancer survivors, and this number is expected to increase to 20 million by 2020. Cancer therapies can cause significant injury to the vasculature, resulting in angina, acute coronary syndromes (ACS), stroke, critical limb ischemia, arrhythmias, and heart failure, independently from the direct myocardial or pericardial damage from the malignancy itself. Consequently, the need for invasive evaluation and management in the cardiac catheterization laboratory (CCL) for such patients has been increasing. In recognition of the need for a document on special considerations for cancer patients in the CCL, the Society for Cardiovascular Angiography and Interventions (SCAI) commissioned a consensus group to provide recommendations based on the published medical literature and on the expertise of operators with accumulated experience in the cardiac catheterization of cancer patients. © 2015 Wiley Periodicals, Inc.

**Key words:** cardio-oncology; PCI; cancer; malignancy; stent thrombosis

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## INTRODUCTION

Advances in cancer therapy have resulted in a steady decline in cancer-related mortality since the 1990s. In the United States alone, there are currently approximately 14.5 million cancer survivors, and this number is expected to increase to 20 million by 2020 [1]. In view of these trends, as well as the cardiovascular toxicity potential of radiation and chemotherapy, cancer patients are exposed to cardiovascular morbidity and mortality more than ever before, thus generating the call for “onco-cardiology” or “cardio-oncology.” The American College of Cardiology (ACC) recognized cardio-oncology as one of the “top cardiology stories for 2014,” and several healthcare institutions have founded onco-cardiology/cardio-oncology departments and fellowship training programs focusing on these issues.

Anticancer therapies can cause significant injury to the vasculature, resulting in angina, acute coronary syndromes (ACS), stroke, critical limb ischemia, arrhythmias, and heart failure (HF), independently from the direct myocardial or pericardial damage that might occur. Moreover, cancer is generally associated with a hypercoagulable state, which increases the risk of acute thrombotic events. Consequently, the need for invasive evaluation and management in the cardiac catheterization laboratory (CCL) for such patients has been increasing. Unfortunately, there are few data on this patient population, because cancer patients have been excluded from national percutaneous coronary intervention (PCI) registries and most randomized PCI clinical trials.

In recognition of the need for a document on special considerations for cancer patients in the CCL, the Society for Cardiovascular Angiography and Interventions (SCAI) commissioned a writing committee to define the landscape and to provide recommendations (level of evidence C) based on published medical literature and expertise of operators with accumulated experience in the cardiac catheterization of cancer patients. As this document is focused on diagnostic and interventional CCL considerations, chemotherapy- and radiotherapy-induced myocardial dysfunction will not be extensively covered.

## MECHANISMS OF VASCULAR TOXICITIES IN CANCER PATIENTS

### Chemotherapy-Induced Vascular Toxicities

In addition to the known effects on cardiac function, chemotherapeutic agents may injure the vascular system, including coronary and peripheral circulation, causing both acute and long-term consequences (Table I).

The chemotherapeutics notoriously associated with angina and ACS are **5-Fluorouracil (5-FU)** and its oral pro-drug **capecitabine**. The drug 5-FU triggers abnormal vasoreactivity immediately after the initiation of therapy [3], which might be due to endothelial damage and alterations in molecular signaling pathways that control vascular smooth muscle cell tone [6,49–51]. Although myocardial ischemia and arrhythmias are often reversible upon treatment discontinuation, lethal outcomes have been reported.

ACS have also been reported for **paclitaxel** and less for **docetaxel** [9–11]. Vasospasm is the key mechanism, and unrecognized coronary artery disease (CAD) may be a predisposing factor. Unlike 5-FU and capecitabine, cardiac rhythm disturbances including bradycardia are more common than ischemic events [11].

**Cisplatin** has been uniquely associated with acute coronary thrombosis, sometimes in multiple vascular territories [14,16,17]. Endothelial damage, thromboxane production, platelet activation, and platelet aggregation appear to be the main mechanisms [16,52–54]. Patients who receive platinum-based chemotherapies are at a 1.5- to 7-fold greater long-term risk of CAD and myocardial infarction (MI) [55–60].

Often given in combination with cisplatin, **bleomycin** can aggravate endothelial dysfunction and **vinblastine** can induce endothelial apoptosis [61], increasing the vasotoxic potential of these cancer treatment regimens [12,13,15,18,62].

Finally, **cyclophosphamide** causes toxicity to the endothelial cells, with the induction of Prinzmetal’s angina or hemorrhagic peri-myocarditis as the primary presentation [12,63,64].

**Vascular endothelial growth factor (VEGF) signaling pathway inhibitors** are associated with a two- to sixfold increased risk of acute cardiovascular events [19,21,22]. These events might be due to the induction of endothelial dysfunction and the downstream consequences of vasoconstriction, vascular remodeling, inflammation, and platelet activation. Interference with plaque neovessel formation and integrity is another unique aspect of this class of drugs [30,65–71]. Some 70% of patients on sunitinib treatment experience a reduction in coronary flow reserve (on average  $1.8 \pm 0.4$ ), especially with a longer duration of therapy [28]. In experimental models, sunitinib causes microvascular impairment [27] in conjunction with rarefaction of microvascular pericytes and capillaries [72]. Abnormality of the vasofunctional balance due to eNOS uncoupling along with an increase in mitochondrial superoxide production [27,73] and increased endothelin-1 production [74,75] may play an additional role in this alteration.

**TABLE I. Chemotherapeutic Agents Associated with Myocardial Ischemia**

	Incidence	Presentations	FDA-approved cancer therapy
<b>Antimetabolites</b>			
5-Fluorouracil [2–6]	0.1%–19%	Angina, vasospasm, MI, Takotsubo cardiomyopathy	Colorectal, pancreas, gastric, breast, basal cell, and squamous cell cancer of head and neck
Capecitabine [4,7,8]	0.02%–10%	Angina, vasospasm, MI, Takotsubo cardiomyopathy	Colorectal, breast cancer
<b>Anti-microtubule agents</b>			
Paclitaxel [9–11]	0.2%–4%	Angina, vasospasm, MI	Breast, ovarian, non-small lung cancer, Kaposi sarcoma
Vinblastine [12,13]	<5%	Angina, MI	Testicular cancer, Hodgkin’s and non-Hodgkin’s lymphoma, Kaposi’s sarcoma, Mycosis fungoides, breast cancer, and choriocarcinoma
<b>Alkylating agents</b>			
Cisplatin [12–17]	0.2%–12%	Angina, vasospasm, MI, coronary thrombosis, progression of CAD	Bladder, cervical, ovarian, testicular, squamous cell of head and neck, non-small cell lung cancer, and mesothelioma
<b>Antitumor antibiotics</b>			
Bleomycin [12,13,18]	<3%	Angina, vasospasm, MI	Testicular, squamous cell cancer of the vulva, cervix, or head and neck, Hodgkin’s and Non-Hodgkin’s lymphoma
<b>Monoclonal antibodies</b>			
Bevacizumab [19–22]	1%–6%	Angina, MI, Takotsubo cardiomyopathy	Renal cell, colorectal, cervical, non-small cell lung cancer, glioblastoma
Ramucirumab [23]	1.5%–2%	Angina, MI, cardiac arrest	Gastric/gastroesophageal junction adenocarcinoma
Rituximab [24]	Rare	Vasospasm, angina, MI, Takotsubo cardiomyopathy	Non-Hodgkin’s lymphoma, Chronic Lymphocytic Leukemia
Aflibercept	3%	Arterial thromboembolic events	Colorectal cancer
<b>Tyrosine kinase inhibitors</b>			
Sorafenib [25]	1%–2%	Vasospasm, angina, MI	Renal cell, liver, thyroid cancer
Sunitinib [26–31]	1%–13%	Angina, MI, Takotsubo cardiomyopathy, progression of CAD	Renal cell, pancreas cancer, gastrointestinal stromal tumor
Pazopanib [32]	2%–10%	Angina, MI	Renal cell cancer, soft tissue sarcoma
Nilotinib [33–36]	2%–25%	Angina, MI, progression of CAD, peripheral arterial disease	Chronic Myeloid Leukemia (CML)
Ponatinib [37,38]	11%	Angina, myocardial infarction, progression of CAD	CML
<b>Hormone therapy</b>			
Aromatase inhibitors (e.g., anastrozole) [39,40]	1%–2% (12%–17% w/IHD)	Angina, MI	Breast cancer
Anti-androgens (e.g., bicalutamide) [41–44]	2%–5%	Angina, MI, progression of CAD	Prostate cancer
Estrogen/nitrogen mustard [45–47] (Estramustine)	1%–3%	Angina, MI	Prostate cancer
Gonadotropin-releasing hormone agonists [44] (goserelin)	1%–5%	Angina, MI	Prostate cancer
Gonadotropin-releasing hormone antagonists [48] (degarelix)	<1%	MI	Prostate cancer

ACS, acute coronary syndrome; CML, chronic myeloid leukemia; IHD, ischemic heart disease; MI, myocardial infarction.

**Sorafenib** has been also associated with coronary vasospasm and even more profoundly than **sunitinib** and with involvement of multiple vessels [76–78]. Moreover, sorafenib has been associated with progres-

sion of CAD, whereas another report links sunitinib with atherosclerotic plaque rupture due to impaired endothelial healing [79]. In an experimental model, treatment with sorafenib was associated with poorer

**TABLE II. Peripheral Arterial Disease Associated with Radiation Therapy**

Type of radiation	Peripheral arterial disease
Head and neck radiation	CVA/TIA, carotid arterial disease
Supraclavicular and mediastinal radiation	CVA/TIA, carotid, and subclavian arterial disease
Abdominal and pelvic radiation	Renal arterial disease, lower extremity PAD

CVA/TIA, cerebrovascular accident/transient ischemic attack; PAD, peripheral arterial disease.

survival due to the induction of drug-induced myocyte necrosis [25,26]. Moreover, there is an increased bleeding risk in patients treated with VEGF inhibitors [80].

Progression of atherosclerosis and ischemic events has been noted for two tyrosine kinase inhibitors: **nilotinib** and **ponatinib** [36,81,82]. Some patients develop a series of events in various vascular territories, even when they have no cardiovascular risk factors [33–35]. The mechanism of the preferential effect on the peripheral arterial circulation with nilotinib is unclear. In contrast, cardiovascular events seem to be more common with ponatinib than cerebrovascular and peripheral artery disease (PAD) events (6.2%, 4.0%, and 3.6%, respectively). Overall, arterial thrombotic events are three times more frequent than venous occlusive events. Finally, there are several reports of ACS and reversible apical ballooning syndrome (Takotsubo) with **rituximab** therapy [24,83]. The prognosis of Takotsubo cardiomyopathy induced by chemotherapy agents is unclear.

Increased cardiovascular risk is noted in patients with prostate cancer treated with **androgen deprivation therapy (ADT)** in the form of gonadotropin-releasing hormone (GnRH) agonists [41–44]. A 25% increased risk of cardiovascular events was reported for women receiving **aromatase inhibitors** (anastrozole, letrozole, exemestane) [39]. Randomized, placebo-controlled trials have not indicated an increased cardiovascular risk with tamoxifen [40] (it improves metabolic parameters, endothelial function and slows atherosclerosis disease progression [84]) but have shown an increase in thromboembolic event risk [39].

The effect of the chemotherapeutic agents reviewed above on cardiovascular risk, especially those with a key impact on endothelial cells or stent endothelialization and stent thrombosis risk, remains undefined [56]. Drugs similar to vinblastine that stimulate thromboxane production and platelet reactivity have been reported to cause MI, and platelet activation was suppressed only by high-dose clopidogrel [85]. While VEGF-eluting stents have been linked to decreased stent thrombosis rates, it is unknown whether VEGF inhibition is associ-

ated with the opposite effect [86]. Any underlying malignancy by itself may be considered a risk factor for stent thrombosis; some malignancies such as acute promyelocytic leukemia are associated with a high coronary thrombosis risk in general [87–89].

### Radiation-Induced Coronary and Peripheral Arterial Disease

Radiation therapy (RT) is received by over 50% of cancer patients. Ionizing radiation affects non-cancerous cells and among these, endothelial cells are the most vulnerable. Cholesterol plaques and thrombosis can form within a period of days after radiation exposure in experimental models [90,91]. Fibrosis may evolve over time, involving all three layers of the vessel wall and subsequently manifestations can vary from accelerated atherosclerosis to fibro intimal thickening as well as thrombotic occlusion in areas of infarction [92,93].

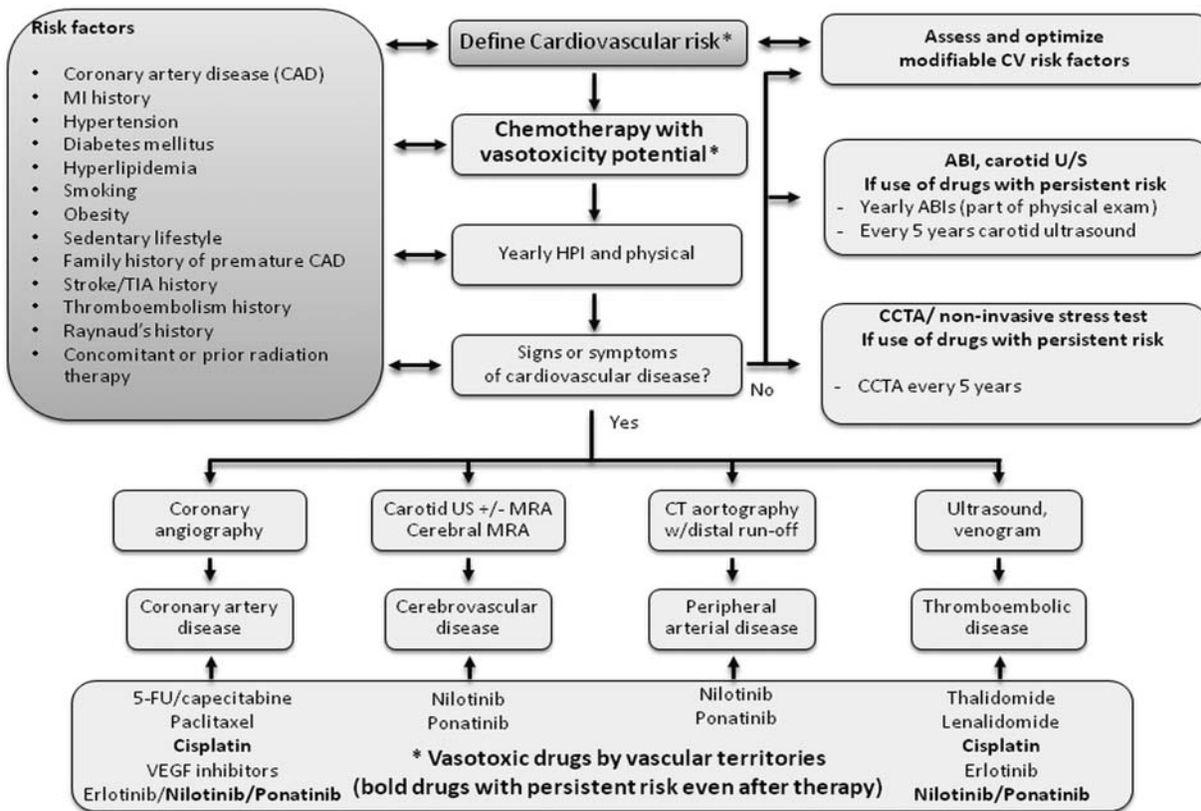
More than two decades after RT for Hodgkin's lymphoma, severe stenosis due to the previously mentioned processes of the ostium of the left main and/or right coronary artery is observed in up to 20% of patients. Some stenosis are even undetected with traditional stress testing [91,94–96]. Even in a younger population with a mean age of just 20 years, coronary artery abnormalities were noted in nearly one in five patients [97]. For breast cancer survivors, RT for left-sided breast cancer has been considered to pose a higher risk of stenosis, which occurs as early as 5 years after therapy [98–101]. In addition to macrovascular disease, RT induces microvascular injury, leading to reduced coronary flow reserve, ischemia, and fibrosis.

As with radiation-induced CAD, PAD remains a concern for patients who receive extracardiac treatments for a variety of malignancies, although their sequelae and complications are less often reported than those of CAD. The central mechanism of post-radiation PAD is similar to that of post-radiation CAD (Table II).

In patients with **head and neck** tumors who have received **RT**, an increased risk of ischemic stroke and carotid arterial disease has been reported [102–105]. These findings across a heterogeneous spectrum of malignancies suggest that a predisposition to more vulnerable and accelerated plaque development in the cerebrovascular system may be aggravated by RT. Case series have shown favorable outcomes for carotid artery stenting (CAS) for radiation-induced carotid artery stenosis [106–109].

In patients with a history of **supraclavicular and mediastinal radiation**, several malignancies have been associated with a higher risk of cerebrovascular events

T2



**Fig. 1. Suggested SCAI algorithm for the cardiovascular screening of patients on chemotherapy.** HPI, history of present illness; TIA, transient ischemic attack; ABI, ankle-brachial index; U/S, Ultrasound; CCTA, cardiac computed tomography angiography. \*Pivotal to the sequence is the determination of baseline cardiac risk, including presence of ischemic heart disease, history of myocardial infarction, cardiovascular risk factor profile, and calculated atherosclerotic cardiovascular disease risk, for example, AHA/ACC ASCVD risk calculator, Framingham risk score, or ESC Score.

and carotid artery disease, particularly head and neck malignancies and lymphomas. A retrospective study on 415 patients with a history of Hodgkin's lymphoma showed a 7.4% prevalence of carotid and/or subclavian artery disease at a median of 17 years after RT [110,111].

Radiation-induced renal artery and lower extremity peripheral vascular disease have been less frequently reported in patients who have received **abdominal radiation** for lymphoma, abdominal sarcomas, and genitourinary malignancies [112–116]. Percutaneous transluminal angioplasty and/or stent placement and surgical interventional strategies have been employed with success, but data are extremely limited. Based on limited case reports and case series, PAD has manifested in patients who have received abdominal radiation for genitourinary malignancies as early as 2 years post-treatment [98].

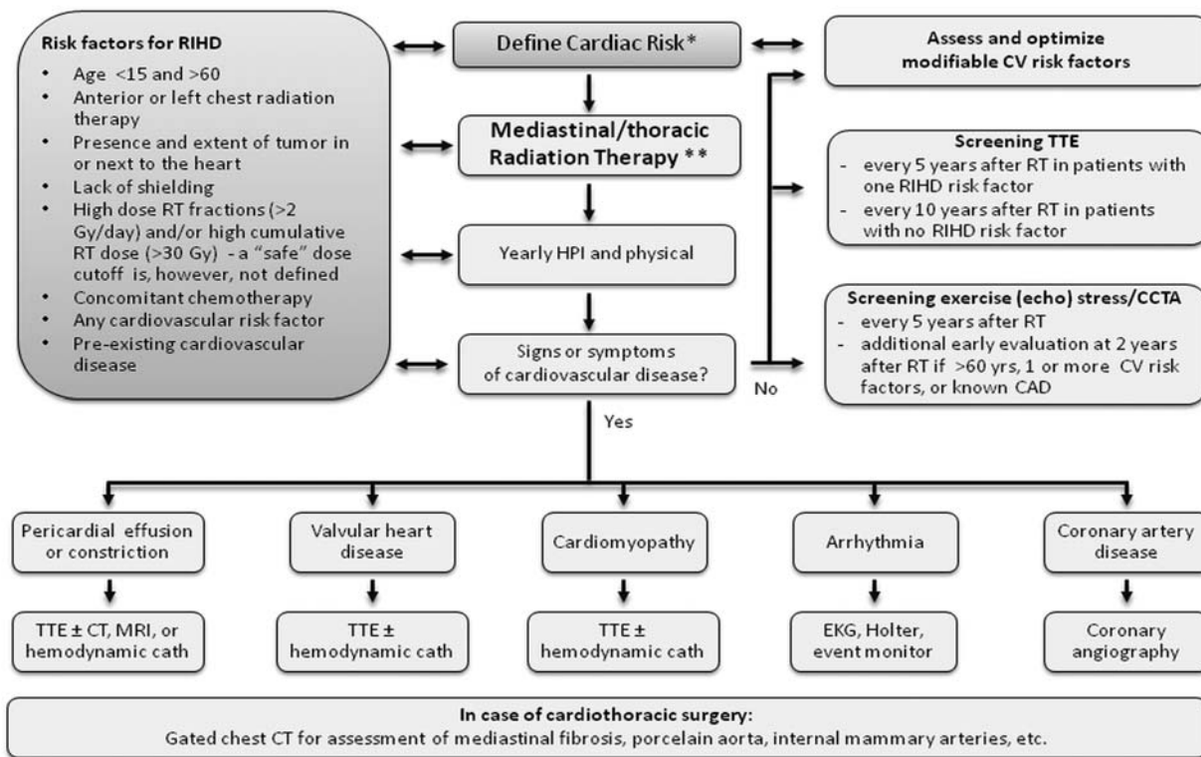
## SCREENING AND PREVENTION OF CARDIOVASCULAR DISEASE IN CANCER PATIENTS

### Screening for Cardiovascular Disease in Patients to Receive Cancer Therapy (Chemotherapy, RT, and Operative Intervention)

Pre-existing cardiovascular risk factors and cardiovascular injury inflicted by chemotherapeutic agents and radiotherapy can have direct effects on coronary and peripheral arteries and the myocardium. This multifactorial insult can lead to an increased risk of developing cardiomyopathy, myocardial ischemia, vascular disease, or conduction abnormalities as well arrhythmias and QT prolongation [117,118].

Risk assessment and treatment for cancer patients with suspected or known cardiovascular disease should generally follow standing ACC/AHA guidelines, with special considerations described in Figs. 1 and 2.

F1  
F2



**Fig. 2. Suggested SCAI algorithm for the cardiovascular screening of patients on radiation therapy.** RIHD, radiation-induced heart disease; HPI, history of present illness; TTE, transthoracic echocardiogram; CCTA, cardiac computed tomography angiography; EKG, electrocardiogram; RT, radiation therapy. \*Pivotal to the sequence is the determination of baseline cardiac risk, including presence of ischemic heart disease, history of myocardial infarction, cardiovascular risk factor profile, and calculated 10 year atherosclerotic cardiovascular dis-

ease (ASCVD) risk (<http://tools.cardiosource.org/ASCVD-Risk-Estimator/>), which remain the cohorts at highest risk for overall and early (<5 years) presentation of acute coronary events during follow-up; if established IHD/CAD or 10-year ASCVD risk  $\geq 5.0\%$  and/or patient >60 years, consider further testing and treatment (moderate-high intensity statin) to define the burden of disease prior to radiation therapy. \*\*Potential sequelae of radiation therapy to the head/neck, abdomen/pelvis should also be assessed as outlined in Table II.

**Pre-chemotherapy cardioprotection.** Although data are limited and such approach remains controversial, the authoring team recommends pre-chemotherapy cardioprotection.

Patients without CAD might benefit from prophylactic treatment with beta-blockers, angiotensin antagonists, statin, or dexrazoxane to reduce cardiotoxicity [119]. The initiation of cardioprotective therapy can be associated with dizziness and fatigue, which might increase due to intravascular volume depletion caused by anorexia, nausea, and vomiting during chemotherapy. Careful volume assessment should be assessed to ensure that the patient remains euvolemic.

For patients with a history of hypertension, blood pressure management is advised as per the eighth Joint National Committee guidelines, with an emphasis on ACE-I and beta blockers (especially carvedilol or nebivolol) as first-line agents. Caution is advised when initiating diuretics and/or angiotensin antagonists due to the propensity to develop electrolyte abnormalities and renal dysfunction.

For patients at intermediate to high cardiovascular risk (based on cardiovascular risk scores) who are potentially undergoing cardiotoxic therapy (i.e., the agents listed in Table I or mediastinal radiation), referral to a cardiologist and/or cardio-oncology program is advised prior to the initiation of treatment.

For patients with established CAD and without contraindications, adding or continuing ACE-I and beta-blockers (preferably carvedilol or nebivolol) might provide additional cardioprotection [120–122].

To identify high-risk patients, intensive and frequent screening for CAD or elevated risk via echocardiographic studies and cardiac biomarkers is encouraged, because the chances of response and recovery are highest with early detection and rapid initiation of therapy [123].

**Pre-radiation therapy cardioprotection.** Aspirin and statin therapy should be encouraged for patients with established CAD or elevated ASCVD risk in keeping with current guidelines. It should be acknowledged that aspirin may not be tolerated in oncologic

patients with high bleeding risk, or statins in patients with impaired liver function or in combination with hepatotoxic chemotherapeutic agents. It is reasonable to further assess cardiovascular risk in intermediate-risk patients at suspected or elevated risk for cardiovascular disease with carotid artery intima-media thickness and ankle-brachial index measurements, coronary artery calcium scoring (CAC), and CCTA as per ACC/AHA guidelines [124]. Exposure of the heart to ionizing radiation during radiotherapy for breast cancer increases the subsequent rate of ischemic heart disease. The increase is proportional to the mean dose to the heart, starting within the first 5 years after radiotherapy and continued into the third decade after radiotherapy. Rates of major coronary events increased linearly with the mean dose to the heart by 7.4% per gray (95% confidence interval, 2.9–14.5;  $P < 0.001$ ), with no apparent threshold. The proportional increase in the rate of major coronary events per gray (Gy) was similar in women with and without cardiac risk factors at the time of radiotherapy.

Older patients with head and neck malignancies or lymphoma who are receiving supraclavicular radiation remain at significantly higher short- and long-term risk for cerebrovascular events [125,126] and thus baseline and more aggressive surveillance via carotid artery screening and subsequent treatment should be advised if the prognosis is favorable from an oncologic standpoint.

Traditional stress testing, although advised for long-term surveillance of RIHD, has its limitations in accuracy and CCTA or coronary angiography may be preferable. In regards to CAC, studies have shown that asymptomatic patients with a coronary artery calcium score of 0 have a “warranty period” of up to 5 years with a very low cardiovascular event rate, although further study regarding CAC progression in this specific patient population is warranted [127–129]. Recent data have suggested that lymphoma survivors with a history of mediastinal radiation and/or anthracycline exposure, have a 40-year cumulative incidence of cardiovascular diseases of 50%, with increased risk of cardiovascular events [130]. Thus, the impetus to optimal screening strategies based on epidemiologic data suggests aggressive surveillance before and after treatment.

### Screening for Cardiovascular Disease in Cancer Survivors

Given the dynamic state of pharmacotherapy for cancer, with generally dramatic improvements in survival as well as new agents in development (with unclear cardiotoxic properties), it is important that patients are made aware of potential short- and long-term consequences as well as the need to follow up

with subspecialists. A retrospective cohort study of the Childhood Cancer Survivor Study demonstrated that adult survivors of childhood malignancies with a history of chemoradiation had a 7-fold higher mortality rate, 15-fold increased rate of HF, 10-fold higher rates of cardiovascular disease, and 9-fold higher rates of stroke compared with their siblings [131]. Cancer survivorship programs and/or cardio-oncology programs provide up-to-date evaluations and appropriate referrals to subspecialists [132]. Periodic annual follow-up in higher risk individuals such as Hodgkin’s lymphoma survivors is also advised, based on the National Comprehensive Cancer Network (NCCN) Clinical Practice Guidelines in Oncology (NCCN). Cancer survivors are at increased risk of secondary and recurrent malignancies as well as cardiotoxic sequelae.

The best timing for surveillance initiation for cardiotoxic manifestations is unclear, given the discrepancies in expert opinion and an absence of official guidelines. The Children’s Oncology Group recommends an annual follow up and physical examination for patients with a history of total body irradiation, cardiotoxic chemotherapy, total mediastinal radiation of  $\geq 20$  Gy, or for those who underwent combined chemoradiation. Both serial electrocardiograms and multigated acquisition scan (MUGA) for patients who received treatment  $\geq 5$  years of age and had both chest radiation and a total anthracycline dose  $\geq 300$  mg/m<sup>2</sup>. For lower risk patients, serial echocardiography is recommended every 2–5 years. Stress testing 5–10 years following radiation exposure to the heart should be considered, along with counseling on lifestyle modifications. A cardiology evaluation and monitoring should be provided to women seeking to become pregnant and who have a history of anthracycline or high dose cyclophosphamide therapy. The International Late Effects of Childhood Cancer Guideline Harmonization Group in 2015 unified several international consensus statements and advised cardiomyopathy screening with “strong” recommendations for echocardiographic surveillance of patients with a history of high dose ( $\geq 250$  mg/m<sup>2</sup>) anthracycline therapy, high dose ( $\geq 35$  Gy) chest radiation, or a combination of  $\geq 100$  mg/m<sup>2</sup> cumulative anthracycline and  $\geq 15$  Gy of chest radiation. There were also recommendations to perform screening for CAD on patients with a history of radiation exposure, although concrete recommendations were not made and were planned for future discussions by the group [133].

The American Society of Echocardiography (ASE) has released an expert consensus statement on the use of multimodality imaging with patients with history of radiotherapy. In their document, screening echocardiography is recommended 5 years after exposure for high-risk patients and 10 years after exposure for all

**TABLE III. Cardiovascular Screening Recommendations for Cancer Patients**

Cardiovascular screening recommendations for cancer survivors
Referral to a survivorship center/cardio-oncology program is recommended for cancer survivors who are not being actively followed by hematology/oncologist.
Medical record documentation of the patient's chemotherapy and radiotherapy treatment course with cumulative doses should be retrieved.
Transthoracic echocardiography (TTE) should be performed on patients with a history of significant anthracycline dose exposure (>240 mg/m <sup>2</sup> ) or chest radiation exposure (>30 Gy) starting no later than 2 years after completion of therapy, at 5 years after diagnosis and continued every 5 years thereafter.
In high-risk groups (known coronary artery disease, age >60, one or more CV risk factors) screening after chest radiation therapy should be initiated 2 years after radiation therapy as outlined in Figure 2
Coronary angiography is indicated for symptomatic patients with a history of radiotherapy, risk factors for RIHD, and noninvasive testing (i.e., stress MPI/echo/MRI, CCTA) that suggest a high likelihood of severe ischemic heart disease.
Coronary angiography is reasonable to consider for the evaluation of LV systolic dysfunction after chest radiation and to evaluate for radiation-induced ischemic heart disease.
Right and left heart catheterization is reasonable to evaluate the presence of pericardial constriction and restrictive cardiomyopathy if noninvasive imaging (echocardiography, CT, MR) is insufficient to provide a diagnosis.
Right and/or left heart catheterization and coronary angiography is reasonable to perform as per ACC/AHA guidelines for preoperative planning for patients with severe RIHD.
There is a known association between accelerated coronary artery disease and elevated cardiovascular events and mortality after chest radiation, particularly in high-risk populations such as those with Hodgkin's lymphoma who have undergone mantle field radiation. For these patients, functional imaging and/or CAC/CCTA is reasonable to perform $\geq 5$ years post-radiotherapy, and further workups (e.g., coronary angiography, functional testing) is indicated for risk stratification if there is concern for severe ischemic heart disease.

other patients, with a reassessment every 5 years in asymptomatic patients. Cardiac magnetic resonance imaging (CMR) was recommended if there was a suspicion of pericardial constriction. They also advised stress testing in high-risk patients for CAD detection in asymptomatic individuals 5–10 years after exposure, and a reassessment every 5 years if no new symptoms developed [134].

CAC has attracted interest [135]; however, it is unclear whether these data can be extrapolated to patients with radiation-induced CAD. Another consideration is coronary CT angiography (CCTA), which may be superior to functional stress tests. CCTA offers the advantage of assessing the aorta and internal mammary arteries, which could be affected by RT. Ultimately, the best evaluation technique remains the coronary angiogram with liberal use of intravascular ultrasound (IVUS) and fractional flow reserve (FFR) to reveal diffuse vascular disease and/or lesional physiologic significance respectively. Cardiac catheterization

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should be considered for hemodynamic evaluation of pericardial, myocardial, and valvular heart disease.

Screening for carotid artery disease by carotid ultrasound should be started 5 years after supraclavicular radiation treatment and repeated every 5 years. Earlier screening (i.e., every 2 years) should occur in older patients (>60 years), in symptomatic patients and/or those with a carotid bruit, those with baseline carotid artery disease, or those on drugs with persistent vascular toxicity risk. Patients with a history of supraclavicular radiotherapy who have neurologic symptoms should undergo carotid artery imaging and be managed in accordance with existing guidelines.

Subclavian arterial ultrasound is recommended in symptomatic patients who have received head, neck, supraclavicular, or mediastinal radiation.

Renal ultrasound should be performed on symptomatic patients who have received abdominal and pelvic radiation.

With regards to lower extremity PAD screening in symptomatic patients, ankle-brachial-index (ABI) screening should be performed annually on those who received RT with potential exposure of the lower extremity vasculature (i.e., abdominal or pelvic radiation exposure) (Table III).

T3

### SPECIAL CONSIDERATIONS FOR CANCER PATIENTS WITH THROMBOCYTOPENIA AND ANEMIA

All major clinical trials on antithrombotic therapy have excluded patients with cancer [136]. One reason is the prevalence of thrombocytopenia (TP), which varies from 10% to 25% across the broad range of solid tumor cancer patients treated with intensive chemotherapy (i.e., breast cancer, ovarian, and germ cell) and the majority of acute leukemia, lymphoma, and multiple myeloma patients [137]. Approximately 10% of cancer patients have platelet counts less than 100,000/mm<sup>3</sup>. Baseline TP increases the risk of bleeding and other adverse cardiac events [138]. Acquired TP develops frequently in cancer patients and appears to be different from the TP that occurs after the administration of glycoprotein (GP) IIb/IIIa inhibitors, heparin, thrombolytic therapy, and oral antiplatelet medications, which is strongly associated with ischemic and hemorrhagic complications as well as early mortality [139–148].

TP is not believed to protect cancer patients from ischemic events. In fact, TP is associated with an increased propensity for thrombus formation [149–153]. Clinical experience suggests that platelet function rather than platelet count is the determinant factor [154].

Prophylactic platelet transfusion is not recommended if the platelet count is greater than 10,000/mL (Table III). Transfusion at higher levels may be necessary for

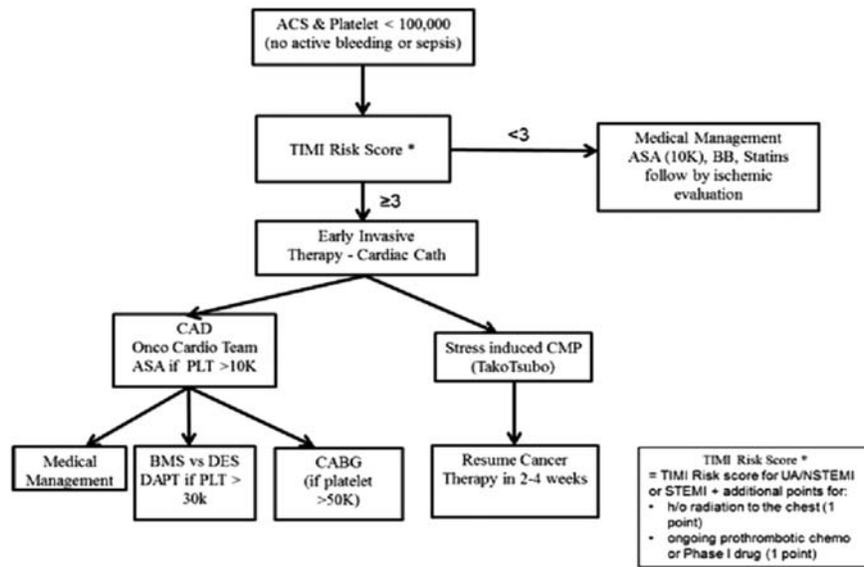


Fig. 3. Recommended revascularization approach used at the MD Anderson Cancer Center.

patients with high fever, hyperleukocytosis, rapid fall in platelet count, or coagulation abnormalities (e.g., acute promyelocytic leukemia) [155]. A prophylactic platelet transfusion should be considered at a threshold of 20,000/mL in patients with solid tumors, who are receiving therapy for bladder, gynecologic, colorectal tumors or melanoma, and for those with demonstrated necrotic tumors, due to the increased risk of bleeding at these sites. When a platelet transfusion is performed, it is critical to repeat the platelet count to ensure that the desired level has been reached. Platelet transfusions should also be available on short notice in case bleeding occurs. For alloimmunized patients, histocompatible platelets must be available.

Because thrombocytopenic patients might have a poor increment after a single transfusion but an excellent response with subsequent transfusions, a diagnosis of refractoriness to platelet transfusion should be made when at least two ABO-compatible transfusions, stored less than 72 hr, result in poor increments. Patients with alloimmune-refractory TP should be managed with platelet transfusions from donors who are HLA-A and HLA-B antigen selected.

On the basis of accumulated clinical experience and a variety of conference consensus documents [156], there is no minimum platelet level that poses an absolute contraindication for a coronary angiogram, and a platelet count of 40,000 to 50,000/mL may be sufficient to perform most interventional procedures with safety, in the absence of associated coagulation abnormalities.

Withholding aspirin (ASA) in thrombocytopenic cancer patients with ACS demonstrated poorer outcomes

[153]. Case series of PCI in thrombocytopenic cancer demonstrated minimal bleeding when meticulous access with micropuncture techniques and careful hemostasis were achieved [155,157]. The initial dose of unfractionated heparin given was lowered to 30–50 U/kg when the platelet count was less than 50K, with additional heparin given if ACT was less than 250 sec. The standard dose of unfractionated heparin 50–70 U/kg or bivalirudin were utilized with platelet counts greater than 50K. In terms of dual antiplatelet therapy (DAPT), the only available experience is with clopidogrel. The recommended revascularization approach currently used at the MD Anderson Cancer Center is shown in Fig. 3.

Various platelet function tests are available; however, there is no data on their value for cancer patients with TP to guide platelet transfusion or DAPT therapy duration and intensity. In patients with significant CAD and platelet counts less than 30,000/mL, and in whom there is concern about intracranial bleed, a normal thromboelastography (TEG) may be considered to determine whether a revascularization strategy would be safe. TEG offers a comprehensive evaluation of both platelet and coagulation function. An abnormal TEG might require initial correction with a platelet transfusion or indicated blood products. This limited experience is available from just a few centers and is extrapolated from the cardiovascular as well as liver transplant surgical literature [158,159].

Anemia is common in cancer patients due to decreased red blood cell (RBC) production or increased RBC loss (bleeding) or destruction (hemolysis) as a direct result of the malignancy or secondary to cancer therapy. The optimal management of those patients

**TABLE IV. Special Considerations for Cancer Patients with Thrombocytopenia**

Special considerations for cancer patients with thrombocytopenia undergoing cardiac catheterization

*Prophylactic platelet transfusion* is not recommended in patients undergoing cardiac catheterization with thrombocytopenia, unless recommended by the oncology/hematology team for one of the following indications:

1. Platelet count <20,000/mL and one of the following: (a) high fever, (b) leukocytosis, (c) rapid fall in platelet count, (d) other coagulation abnormality
2. Platelet count <20,000/mL in solid tumor patients receiving therapy for bladder, gynecologic, or colorectal tumors, melanoma, or necrotic tumors

*Therapeutic platelet transfusions* are recommended in thrombocytopenic patients who develop bleeding during or after cardiac catheterization. Repeat platelet counts are recommended after platelet transfusions.

30–50 U/kg unfractionated heparin is the initial recommended dose for thrombocytopenic patients undergoing PCI who have platelets <50,000/mL. ACT should be monitored.

For platelet counts <30,000/mL, revascularization and DAPT should be decided after a preliminary multidisciplinary evaluation (interventional cardiology/oncology/hematology) and a risk/benefit analysis.

Aspirin administration may be used when platelet counts are >10,000/mL.

DAPT with clopidogrel may be used when platelet counts 30,000–50,000/mL. Prasugrel, ticagrelor and IIB-IIIa inhibitors should not be used in patients with platelet counts <50,000.

If platelet counts are <50,000, the duration of DAPT may be restricted to 2 weeks following PTCA alone, 4 weeks after bare-metal stent (BMS), and 6 months after second or third generation drug-eluting stents (DES) if optimal stent expansion was confirmed by IVUS or OCT.

There is no minimum platelet count to perform a diagnostic coronary angiogram.

with blood product or iron transfusions is unclear and various societies have recommended cut-off levels for transfusion (EORTC, ASH/ASCO) [160–162]. RBC transfusion is generally recommended when hemoglobin is less than 7 g/dL. Consultation with hematology/oncology specialists is recommended for severely anemic cancer patients undergoing cardiac catheterization (Table IV).

T4

**VASCULAR ACCESS CONSIDERATIONS FOR CANCER PATIENTS UNDERGOING CARDIAC CATHETERIZATION**

Complications of vascular access remain the most common cause of morbidity and are also associated with significant mortality [163]. Specific characteristics of the patient may impact access site complications (i) the effect of cancer and cancer treatment on the hematopoietic system [155,157,164–168], (ii) the presence of a hypercoagulable state [20,169–172], and (iii) the potential interactions between cancer and cardiac drugs [173–175]. Even minor vascular complications may lead to prolonged hospitalization and adverse outcomes

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[176,177]. Prior to catheterization, all patients should be evaluated for their bleeding diathesis, the prothrombotic state and the potential for infection due to immunosuppression [174]. Access routes should be carefully assessed and the appropriate steps taken to reduce complications associated with each technique (Table IV) [178–180].

Ultrasound guidance, micropuncture needles, and fluoroscopic guidance all contribute to the best possible outcome (Figs. 4 and 5) [180–183].

F4  
F5

**Femoral Access Site**

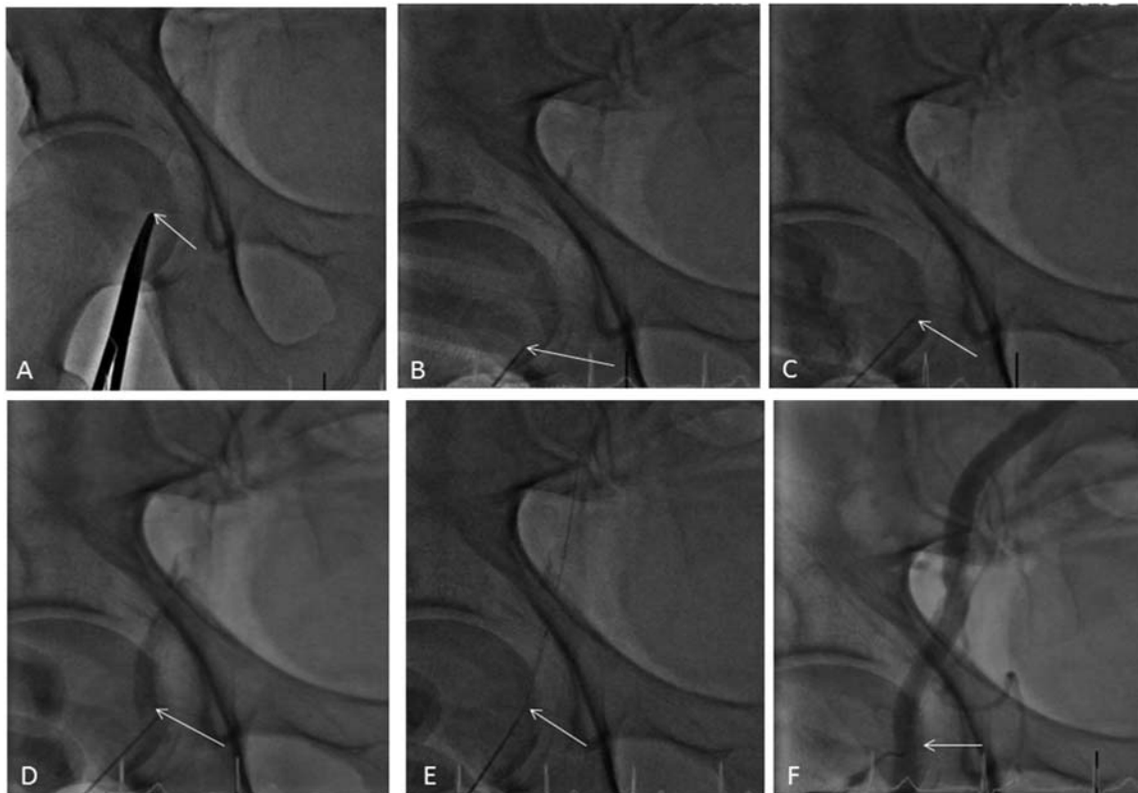
Femoral artery access has been associated with a higher risk of bleeding, even with the use of vascular closure devices as compared with radial artery access [184]. A controversy exists concerning the need for performing the Allen test before transradial procedure [182]. Oncologic patients who have failed the Allen’s test in both arms, who are on hemodialysis, with multiple previous radial procedures or arterial lines, and patients who have undergone bilateral mastectomy are probably better candidates for the femoral approach. Femoral access gives the operator greater flexibility for complex coronary interventions, rotational atherectomy, and the use of mechanical assist devices.

Puncture of the common femoral artery (CFA) at its mid-section should be the goal for optimal vascular access. Access outside that zone may lead to retroperitoneal hemorrhage (RPH), pseudoaneurysm, arteriovenous fistula, thrombosis, or excessive bleeding [185–187], which may be fatal in cancer patients. Therefore, meticulous identification of the inguinal ligament and “lower” access should be preferred to prevent bleeding. Vascular closure devices do not seem to decrease bleeding compared with manual compression, and should be avoided in immunocompromised patients due to the higher risk of local infection or delayed endothelialization [188].

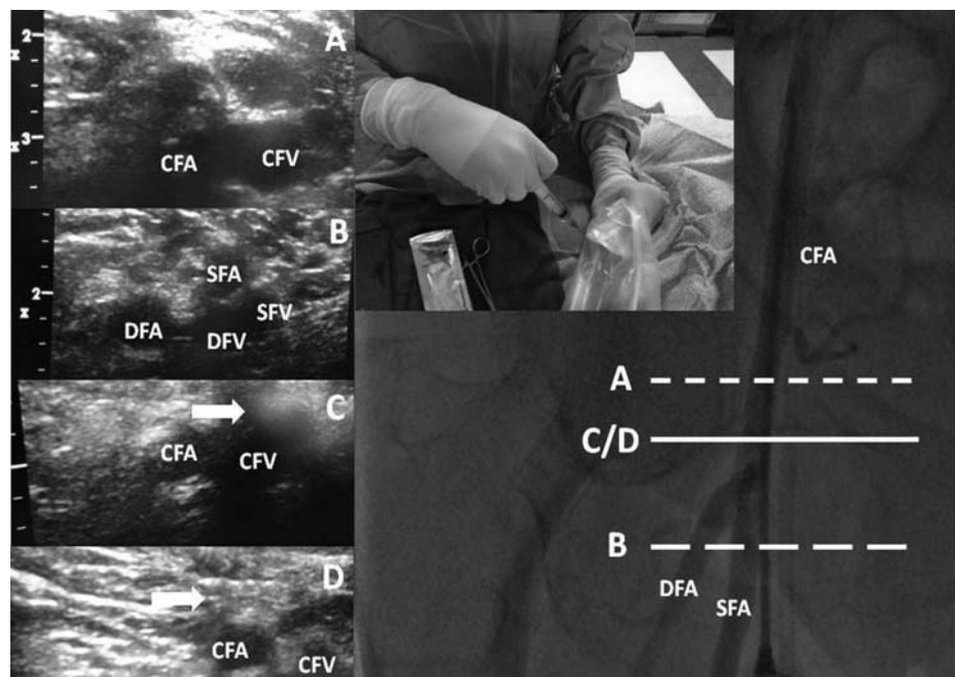
**Radial Access Site**

The radial artery is favored because of the lower bleeding risk and increased patient satisfaction [189]. Reductions in bleeding complications can be attained even in patients with TP who are receiving anticoagulant and antiplatelet therapy [190–192]. Early ambulation after radial access site catheterization favors fewer thrombosis complications. The need for anticoagulant administration remains a limitation for oncologic patients undergoing transradial diagnostic catheterizations.

Depending on the operator’s skills, the advantages of radial access may be balanced in some cases by technical difficulty, increased fluoroscopy time, and



**Fig. 4.** Use of micropuncture to gain femoral access. (A) Anatomic localization of the femoral head (*white arrow*). (B, C) Micropuncture needle access. (D) Contrast injection through the micropuncture needle. (E) Micropuncture wire placement in the common iliac artery. (F) Common femoral angiogram. [Color figure can be viewed in the online issue, which is available at [wileyonlinelibrary.com](http://wileyonlinelibrary.com).]



**Fig. 5.** Ultrasound-guided transfemoral access technique and correlation with femoral angiography. Panel A: Visualization of both the common femoral artery and vein. Panel B: Visualization of vascular structures below the bifurcation of the common femoral artery and vein. Panel C: Ultrasound guided access of the common femoral vein using a micropuncture needle (*arrow*). Panel D: Ultrasound guided access of the common femoral artery using a micropuncture needle

(*arrow*). CFA, common femoral artery; CFV, common femoral vein; DFA, deep femoral artery (profunda femoris artery); SFA, superficial femoral artery; DFV, deep femoral vein (profunda femoris vein); SFV, superficial femoral vein (greater saphenous vein). Special thanks to Dr. Timothy Canan for assistance in providing demonstration. [Color figure can be viewed in the online issue, which is available at [wileyonlinelibrary.com](http://wileyonlinelibrary.com).]

**TABLE V. Access Considerations for Cancer Patients Undergoing Cardiac Catheterization**

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Access considerations for cancer patients undergoing cardiac catheterization

For cancer patients who are excellent candidates for both access types, the radial artery is preferred. Femoral access is the preferred approach for cancer patients on hemodialysis, those with abnormal Allen's tests in both arms, multiple radial procedures or a-lines, bilateral mastectomy or when a complex intervention is anticipated.

The use of smaller sheath sizes, prompt removal of sheaths and early ambulation is recommended.

A lower dose of intra-arterial or intravenous unfractionated heparin at a dose of 50 U/kg or 3,000 units is recommended for cancer patients with thrombocytopenia and platelet count <50k undergoing cardiac catheterization via radial access.

A femoral angiogram is recommended after transfemoral access to promptly identify and address potential access complications.

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increased radiation exposure [193,194]. Smaller hydrophilic sheaths and catheters should be used for further reductions in the risk of bleeding [183]. Patent hemostasis must be achieved to reduce the risk of radial artery occlusion, incidence of which is approximately 1%–3% in the general population. The patency of the radial artery should be preserved in case the artery is needed as a conduit for intra-arterial pressure monitoring, coronary artery bypass surgery, or hemodialysis access [195].

Taking into consideration the above special circumstances, for cancer patients who are excellent candidates for both types of access, transradial access should be preferred. Meticulous hemostasis as well as frequent catheter and sheath flushing are required, as cancer patients face the challenge of concomitant increased risk of both bleeding and thrombosis (Table V). Ultimately, operator clinical judgment is paramount in the final decision for the optimal access site.

**INVASIVE EVALUATION AND MANAGEMENT OF CAD IN CANCER PATIENTS**

Effective clinical risk stratification of patients undergoing cancer surgery must be undertaken prior to the consideration of cardiac catheterization. Breast, endocrine, reconstructive, gynecologic and minor urologic operations are considered low risk; abdominal and urologic operations are considered intermediate risk. The American College of Surgeons National Surgical Quality Improvement Program (ACS-NSQIP) risk calculator has been emphasized in the 2014 ACC/AHA guidelines on perioperative cardiovascular evaluation as an excellent risk assessment tool, and it can be used to stratify cancer patients with underlying cardiovascular disease. Overall, the urgency of cancer surgery and the need for PCI or coronary artery bypass surgery (CABG)

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**TABLE VI. Special Considerations for Cancer Patients Undergoing Cardiac Catheterization**

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Special considerations for cancer patients undergoing cardiac catheterization for CAD

Decision making regarding revascularization in patients with active cancer must take into consideration the overall prognosis of the patient.

**For cancer patients with an acceptable prognosis**, the general revascularization criteria for appropriate use must be carefully evaluated and only the most appropriate indications (scores 7 and above) should be considered [225].

**For cancer patients with an expected survival <1 year**, percutaneous revascularization may be considered for patients with acute STEMI and high-risk NSTEMI. For patients with stable angina, every effort must be made to maximally optimize medical therapy before resorting to an invasive strategy. This approach must include addressing other cancer-related comorbidities that potentially exacerbate ischemia, such as anemia, infection, hypoxia, etc. Should the patient continue to experience persistently severe angina (CCS Class III or IV), consideration may be given to percutaneous revascularization as a palliative option.

FFR is recommended before non-urgent PCI to justify the need for revascularization.

**When invasive approach is indicated:**

- Balloon angioplasty should be considered for cancer patients who are not candidates for DAPT (Platelets <30,000/mL) or when a non-cardiac procedure or surgery is necessary as soon as possible.
- BMS should be considered for patients with platelet counts >30,000/mL who need a non-cardiac procedure, surgery or chemotherapy which can be postponed for >4 weeks.
- Newer generation DES should be considered for patients with platelet counts >30,000/mL who are not in immediate need for a non-cardiac procedure, surgery or chemotherapy.
- Bivalirudin and/or radial approach should be considered to minimize the risk of bleeding.

**Post intervention:**

Intravascular imaging such as IVUS or optical coherence tomography (OCT) is recommended after stent placement to ensure optimal expansion and an absence of complications given the potential for early DAPT interruption.

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should be guided by a collaborative clinical evaluation by oncologists and cardiologists [177]. Furthermore, it is reasonable to consider a staged approach, with an initial coronary anatomy (invasive coronary angiography or coronary CT angiography) and physiology assessment (stress test, cardiac PET, FFR), followed by an interdisciplinary meeting (medical or surgical oncology, RT, and cardiology) to outline the optimal plan (Table VI).

Patients with stable angina can be treated medically, as PCI does not offer a survival advantage in most cases [196]. For patients with angina despite medical therapy, the severity of the cardiac disease, stage of the malignancy, and the condition of the patient should determine the strategy for PCI versus CABG. PCI is preferred when the malignancy is aggressive or widespread. CABG can be considered when the malignancy is potentially curable or when the estimated prognosis is acceptable [197].

T5

T6

Cancer patients with STEMI have higher mortality and morbidity compared with those without cancer. Patients with recently diagnosed cancer (<6 months) undergoing primary PCI had threefold higher cardiac mortality compared with those with a prior diagnosis and a control group (adjusted HR 3.3; CI 1.5–7) [165]. Similar findings were reported from the NHLBI registry of ACS patients undergoing PCI where cancer was one of the strongest independent predictors of in-hospital death (OR 3.2; CI 1.12–9.4) and 1 year mortality (OR 2.15; 1.3–3.4) [198].

One of the largest series to date, with 456 cancer patients (88% advanced cancer, 61% solid tumors) with ACS (15% STEMI, 85% NSTEMI) from a single institution without onsite catheterization laboratory demonstrated that the presenting symptoms of ACS in cancer patients are dyspnea (44%), followed by chest pain (30.3%), and hypotension (23%) [199]. The majority of those patients were treated medically (ASA 46%; beta blockers 48%; statins 21%) and only 15 (3.3%) underwent percutaneous revascularization. One-year survival was only 26%. The use of aspirin (HR 0.77; 0.6–0.98) and beta blockers (HR 0.64; 0.51–0.81) were independently predictive of improved survival, whereas there was a trend toward improved survival with catheter-based revascularization (HR 0.57; 0.29–1.10;  $P=0.09$ ). However, selection bias should be taken into consideration as patients who received antiplatelet therapy and PCI were probably at lower risks than those that did not (bleeding risk, comorbidities, etc.). In another small report from Japan, 18 cancer patients (15 with advanced cancer) with acute MI undergoing PCI (15 BMS; 2 PTCA only; 1 DES) were compared with 59 controls. Both groups had similar procedural success (>97%). There were four in-hospital deaths in the cancer group (1 cardiac; 3 cancer related) and three patients had major bleeding [200]. These studies as well as a small cohort study from Russia [201] suggest the relative safety and efficacy of primary PCI in acute MI patients with advanced cancer.

When performing PCI, it is important to balance lesion characteristics and cancer stage and therapy, recognizing that cancer as a prothrombotic and proinflammatory state is associated with a higher risk of stent thrombosis and possibly in-stent restenosis. We recommend the use of BMS or newer-generation DES, which may have lower rates of stent thrombosis (ST) than BMS. Attempts should be made to avoid bifurcation and overlapping stents, both of which increase the risk of ST. High pressure ( $\geq 16$  atm), non-compliant balloon inflations and the use of IVUS or optical coherence tomography (OCT) is recommended to assure adequate stent expansion, apposition and lack of edge

dissection. Although the use of drug-eluting balloons, bio-absorbable polymers, or scaffolds may reduce the need for DAPT, these devices are not currently available in the United States. The initial experience at a major cancer center using OCT in cancer patients with DES placed for less than 12 months and requiring premature discontinuation of DAPT suggested that many patients had incomplete stent coverage or apposition, underexpansion, or already developed in-stent restenosis. Patients with optimal stent apposition and coverage without in-stent restenosis had no adverse events after premature DAPT discontinuation. However, the recently reported DAPT study suggests that patients treated with either BMS or DES may benefit from 30 weeks of therapy [202], although it is unclear whether a patient population with cancer or ultrasound-guided stent placement would derive a similar benefit. In fact, unrecognized cancer was a significant problem in this study, highlighting the merit of cardio-oncology awareness.

All patients undergoing PCI should receive anticoagulant agents to maintain an activated clotting time of greater than 250 sec during the procedure [182,203]. In oncologic patients with severe TP (<50k platelets), lower doses of unfractionated (50 U/kg) may achieve a therapeutic activated clotting time [182]. Patients with heparin-induced TP should receive intravenous bivalirudin.

If PCI is necessary in patients awaiting cancer surgery, balloon angioplasty without stenting or implantation of BMS is recommended (Table VI), although newer generation DES may also be acceptable. Any interruption of DAPT may lead to in-stent thrombosis, especially in types of cancer with increased propensity for thrombosis.

With chemotherapy, DAPT may need to be extended due to the delayed re-endothelialization of the stent [177]. Some agents are thrombogenic, such as cisplatin and thalidomide, and require an antithrombotic regimen. Others might induce TP, which hampers the use of DAPT. When urgent surgery is needed shortly after PCI, at least one antiplatelet agent should be continued if at all possible. If oral antiplatelet agents must be discontinued, a short acting intravenous IIb/IIIa receptor blocker could be considered until shortly before non-cardiac surgery; however, data are non-existent and this approach remains controversial. Clopidogrel should be restarted after surgery with a loading dose of 300 mg [177].

Digestive tract tumors can pose different problems. In patients undergoing colonoscopy, the presence of CAD is an independent predictor for advanced colon carcinoma [204,205]. In addition, digestive tract tumors may bleed. When antiplatelet therapy must be stopped

due to gastrointestinal bleeding, the cardiac complication rate after PCI increased from 2.4% to 5.8% [206]. Initial treatment with balloon angioplasty followed by delayed stenting after recovery from cancer surgery may be an alternative, but this option has less predictable results [177].

### USE OF FFR, IVUS, AND OCT TO AVOID UNNECESSARY CORONARY INTERVENTIONS IN CANCER PATIENTS

The limitations of the coronary angiogram in defining the hemodynamic significance of a lesion are well known [207–210]. For example, coronary angiography may overestimate the significance of ostial or side-branch lesions, leading to unnecessary complex interventions [209,210]. FFR has emerged as a powerful tool to determine the functional importance of the lesion [204–206,209,211].

Experience from an unpublished small case series at MD Anderson suggests that deferring cancer patients with FFR greater than 0.75 in order to expedite cancer care (chemotherapy, radiation, or surgery) may not be associated with increased cardiovascular mortality within 1 year. A similar approach in cancer patients with significant LM disease who were evaluated with FFR or IVUS resulted in improved quality of life, early cancer therapy initiation, and reduced hospital stay and costs.

IVUS and OCT should be liberally used to assure adequate stent expansion, apposition and lack of edge dissection [212,213].

### SPECIAL CONSIDERATIONS FOR CANCER PATIENTS WHO HAVE RECEIVED RT

After RT, the distribution of CAD has been associated with the location of radiotherapy: for example left breast/chest wall radiation has been associated with disease of the mid and distal left anterior descending artery and distal diagonal branch [214]. Ostial lesions are also more common. The mean time interval for the development of radiation induced CAD in relation to radiotherapy is approximately 82 months (range 59–104) [93] and it generally presents at younger age than the general population, especially in survivors of childhood and adolescence malignancies treated with mediastinal radiation [215]. There are no specific guidelines in the management of patients with radiation induced CAD. The decision on medical therapy or revascularization depends on patient's symptoms, cancer stage, expected survival, and comorbidities. Regarding revascularization both percutaneous intervention and coronary artery bypass graft (CABG) have been used.

During PCI, the use of orbital or rotational atherectomy should be considered for heavily calcified lesions. Surgical revascularization may pose difficulties in these patients because of mediastinal fibrosis, with high incidence of complications. In addition, the use of internal mammary artery as a graft may not always be possible due to radiation disease with this vessel itself [94].

### SPECIAL CONSIDERATIONS FOR CABG WITH CANCER PATIENTS

When considering CABG, it is essential to consider the tumor stage and general condition of the patient. CABG is intended to reduce cardiac complications during or after noncardiac surgery [177]. CABG and cancer surgery can be performed simultaneously as a one- or two-stage procedure. If a two-stage procedure is preferred, a recovery of 4–6 weeks should be anticipated [216–218]. Pulmonary tumors can be treated simultaneously with CABG through the same incision. This is not necessarily the case for tumors of the digestive tract, due to the risk of mediastinitis [218–220]. Simultaneous CABG and tumor resection has advantages including reducing hospitalization and costs, repeat thoracotomy, complications, and delay in treating the malignancy [218–220]. When available, minimally-invasive and off-pump CABG is preferred, to shorten the recovery period.

Hematological malignancies, such as chronic lymphatic leukemia (CLL), are associated with a dysfunctional immunological state, bleeding, the need for transfusion, a risk of infection, and mortality after cardiac surgery [221–223]. However, cardiac surgery did not result in a long-term negative impact on the course of this malignancy and CLL is not a contraindication for heart surgery [224].

### NON-CORONARY INTERVENTIONAL PROCEDURES IN CANCER PATIENTS

#### Right Heart Catheterization

Right heart catheterization can accurately assess the presence of left ventricular systolic or diastolic heart failure (HF), restriction or constriction physiology [226] and valvular dysfunction [227] (Table VII). It should be used to diagnose and differentiate complications of cancer therapy (i.e., hypoalbuminemia, intralveolar hemorrhage, renal failure) and to monitor the left and right ventricular filling pressures. Prior to catheterization, patients with evidence of tumor invasion of the inferior or superior vena cava should undergo thorough imaging studies. In patients with evidence of pulmonary embolism, pulmonary capillary wedge pressure measurements should be avoided on the side of the

T7

**TABLE VII. Indications for Non-Coronary Interventional Procedures in Cancer Patients**

Procedure	Indications
Right heart catheterization	Evaluation of heart failure, constrictive or restrictive cardiomyopathy, valvular heart disease, pulmonary hypertension, and pericardial disease.
Endomyocardial biopsy	Evaluation of intracardiac tumors, unexplained heart failure associated with suspected anthracycline cardiomyopathy, infiltrative cardiomyopathies, and myocarditis.
Pericardiocentesis	Evaluation of pericardial effusion and symptomatic relief.
Balloon pericardiotomy	Prevention of large malignant pericardial effusion, especially in poor surgical candidates
Balloon aortic valvuloplasty and TAVR	Palliative measure for symptomatic AS (or as a bridge for SAVR/TAVR)

pulmonary embolus (if the side is known). If the procedure is done in the catheterization laboratory, the use of the micropuncture technique, ultrasound-guided access, and the use of a 5F (French) Swan catheter from the right forearm should be considered to reduce the risk of bleeding complications [228].

**Endomyocardial Biopsy (EMB)**

EMB is widely used for the surveillance of cardiac allograft rejection and for the diagnosis of unexplained ventricular dysfunction or fulminant myocarditis. Current AHA/ACCF/ESC guidelines recommend EMB for the diagnosis of cardiac tumors (with the exception of cardiac myxomas) if four specific criteria are met: (i) a diagnosis cannot be made in any other way, (ii) the diagnosis with EMB will alter therapy, (iii) the success of a biopsy is believed to be reasonably likely, and (iv) the biopsy will be performed by an experienced operator. Tissue samples should be sent in both formalin and gluteraldehyde for hematoxylin and eosin staining and electron microscopy (EM), as unique features of anthracycline injury are seen only on EM. On Congo Red staining for cardiac amyloidosis, EM identifies specific features that confirm the diagnosis of amyloidosis. Frozen sections should be examined to ensure adequate tissue samples for analysis. The use of transthoracic echocardiographic, transesophageal echocardiography (TEE), or intracardial echocardiography (ICE) guidance in combination with fluoroscopy might result in better quality samples, the use of less radiation and early detection of complications [229]. Data on ICE for this purpose is currently limited.

EMB is reasonable in the setting of unexplained HF with suspected anthracycline cardiomyopathy (Class IIa

indication, Level of evidence C) as it may result in changes in the chemotherapeutic dose and/or regimen [229]. EMB should not be used for routine monitoring after anthracycline or other chemotherapy treatment.

**Pericardiocentesis**

The drainage of malignant pericardial effusion is frequently performed on cancer patients for symptom relief or diagnostic purposes. Pericardiocentesis alone is generally inadequate for malignant effusions, given that only one-third of the effusions are controlled and re-accumulation frequently occurs within 24–48 hr [230]. We recommend that pericardiocentesis should be performed in the catheterization laboratory under fluoroscopic and echocardiographic guidance. The access site should be determined by echocardiography based on the route with the shortest and easiest access to the pericardial space. Echocardiographic assistance also allows for the detection of early complications. If a subxyphoid approach is used, care should be taken to avoid trauma to the left lobe of the liver. For intercostal approaches, placement of the needle above the specific rib margin is necessary to avoid damage to the intercostal areas [231]. The use of micro-puncture needle and small sheath size is recommended to minimize procedural risks. The pericardial drain should be maintained for a minimum of 3 days (optimally 5 days) [232].

**Balloon Pericardiotomy**

Malignant pericardial effusion has a high recurrence rate after pericardiocentesis. Percutaneous balloon pericardiotomy is a simple, safe technique that can be effective in the prevention of recurrence in patients with large malignant pericardial effusion, especially in poor surgical candidates. Several centers have actually adopted that strategy as the initial preferred treatment with low complication rates similar to simple aspiration. The subxyphoid is the standard approach. A dilating balloon containing 30% radiographic contrast medium is advanced over the guide wire to the pericardial border and it is manually inflated to create the window [233,234]. Post-procedure echocardiography and chest radiography are recommended to monitor for possible re-accumulation of pericardial fluid or development of an iatrogenic left pleural effusion.

**Balloon Aortic Valvuloplasty and TAVR**

Balloon aortic valvuloplasty (BAV) was once the only percutaneous option for severe, symptomatic aortic stenosis (AS). Today BAV is mainly used in combination or as a bridge to surgical aortic valve replacement (SAVR), or transcatheter aortic valve replacement (TAVR). Data

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**APPENDIX A. Author Relationships with Industry and Other Entities (Relevant)**

Committee Member	Consultant	Speakers Bureau	Ownership/ Partnership/ Principal	Personal Research	Institutional, Organizational or Other Financial Benefit	Expert Witness
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Cindy Grines, MD	None	None	None	None	None	None
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Eric Yang, MD	None	None	None	None	None	None
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Konstantinos Charitakis, MD	None	None	None	None	None	None
Massoud Leesar, MD	None	None	None	None	None	None

This table presents the relevant healthcare relationships of committee members with industry and other entities that were reported by authors at the time this document was under development. The table does not necessarily reflect relationships with industry at the time of publication. A person is deemed to have a significant interest in a business if the interest represents ownership of  $\geq 5\%$  of the voting stock or share of the business entity, or ownership of  $\geq \$10\,000$  of the fair market value of the business entity; or if funds received by the person from the business entity exceed 5% of the person's gross income for the previous year. Relationships no financial benefit is also included for the purpose of transparency. Relationships in this table are modest unless otherwise noted.

on TAVR in cancer patients are currently unavailable and cancer patients are excluded in most TAVR programs [235] A recent case series of six cancer patients demonstrated the feasibility of BAV when urgent, non-cardiac surgery was necessary [236]. TAVR may be a viable option in cancer patients with acceptable prognoses and severely symptomatic AS. Furthermore, cancer patients may be at higher perioperative risk of mortality with traditional surgical aortic valve replacement due to prohibitive anatomy (i.e., mediastinal fibrosis, severe lung disease, porcelain aorta, and prior thoracic surgeries).

**CONCLUSION**

The cardio-oncology patient population has been increasing in recent years, with better quality of life and improved survival rates. Invasive cardiac assessment is important for the evaluation and management of concomitant heart disease. This SCAI consensus document aims to indicate special considerations to be addressed by interventional cardiologists when managing this frail patient subgroup. Collaboration between cardiologists and hematologists/oncologists is of prime importance. Further research involving cancer patients is also needed to optimize the care of oncology patients in the CCL.

**REFERENCES**

1. Cancer Data and Statistics. 2011. <http://www.cdc.gov/cancer/dpcp/data/>. Accessed on 10 June 2015.
2. Gianni M, Dentali F, Lonn E. 5 fluorouracil-induced apical ballooning syndrome: A case report. *Blood Coagul Fibrinolysis* 2009;20:306–308.
3. Sudhoff T, Enderle MD, Pahlke M, et al. 5-Fluorouracil induces arterial vasoconstrictions. *Ann Oncol* 2004;15:661–664.

4. Polk A, Vaage-Nilsen M, Vistisen K, Nielsen DL. Cardiotoxicity in cancer patients treated with 5-fluorouracil or capecitabine: A systematic review of incidence, manifestations and predisposing factors. *Cancer Treat Rev* 2013;39:974–984.
5. Shahrokni A, Rajebi MR, Harold L, Saif MW. Cardiotoxicity of 5-fluorouracil and capecitabine in a pancreatic cancer patient with a novel mutation in the dihydropyrimidine dehydrogenase gene. *JOP* 2009;10:215–220.
6. Stewart T, Pavlakis N, Ward M. Cardiotoxicity with 5-fluorouracil and capecitabine: More than just vasospastic angina. *Intern Med J* 2010;40:303–307.
7. Lestuzzi C, Tartuferi L, Corona G. Capecitabine (and 5 fluorouracil) cardiotoxicity. Metabolic considerations. *Breast J* 2011; 17:564–565; author reply 566–567.
8. Ambrosy AP, Kunz PL, Fisher GA, Witteles RM. Capecitabine-induced chest pain relieved by diltiazem. *Am J Cardiol* 2012; 110:1623–1626.
9. Schrader C, Keussen C, Bewig B, von Freier A, Lins M. Symptoms and signs of an acute myocardial ischemia caused by chemotherapy with Paclitaxel (Taxol) in a patient with metastatic ovarian carcinoma. *Eur J Med Res* 2005;10:498–501.
10. Shah K, Gupta S, Ghosh J, Bajpai J, Maheshwari A. Acute non-ST elevation myocardial infarction following paclitaxel administration for ovarian carcinoma: A case report and review of literature. *J Cancer Res Ther* 2012;8:442–444.
11. Rowinsky EK, McGuire WP, Guarnieri T, et al. Cardiac disturbances during the administration of taxol. *J Clin Oncol* 1991;9: 1704–1712.
12. Stefenelli T, Kuzmits R, Ulrich W, Glogar D. Acute vascular toxicity after combination chemotherapy with cisplatin, vinblastine, and bleomycin for testicular cancer. *Eur Heart J* 1988;9:552–556.
13. Samuels BL, Vogelzang NJ, Kennedy BJ. Severe vascular toxicity associated with vinblastine, bleomycin, and cisplatin chemotherapy. *Cancer Chemother Pharmacol* 1987;19:253–256.
14. Berliner S, Rahima M, Sidi Y, et al. Acute coronary events following cisplatin-based chemotherapy. *Cancer Invest* 1990;8: 583–586.
15. Doll DC, List AF, Greco FA, et al. Acute vascular ischemic events after cisplatin-based combination chemotherapy for germ-cell tumors of the testis. *Ann Intern Med* 1986;105:48–51.
16. Jafri M, Protheroe A. Cisplatin-associated thrombosis. *Anti-cancer Drugs* 2008;19:927–929.

17. Karabay KO, Yildiz O, Aytekin V. Multiple coronary thrombi with cisplatin. *J Invasive Cardiol* 2014;26:E18–E20.
18. Schwarzer S, Eber B, Greinix H, Lind P. Non-Q-wave myocardial infarction associated with bleomycin and etoposide chemotherapy. *Eur Heart J* 1991;12:748–750.
19. Chen XL, Lei YH, Liu CF, et al. Angiogenesis inhibitor bevacizumab increases the risk of ischemic heart disease associated with chemotherapy: A meta-analysis. *PLoS One* 2013;8:e66721.
20. Scappaticci FA, Skillings JR, Holden SN, et al. Arterial thromboembolic events in patients with metastatic carcinoma treated with chemotherapy and bevacizumab. *J Natl Cancer Inst* 2007;99:1232–1239.
21. Schutz FA, Je Y, Azzi GR, Nguyen PL, Choueiri TK. Bevacizumab increases the risk of arterial ischemia: A large study in cancer patients with a focus on different subgroup outcomes. *Ann Oncol* 2011;22:1404–1412.
22. Ranpura V, Hapani S, Chuang J, Wu S. Risk of cardiac ischemia and arterial thromboembolic events with the angiogenesis inhibitor bevacizumab in cancer patients: A meta-analysis of randomized controlled trials. *Acta Oncol* 2010;49:287–297.
23. Fuchs CS, Tomasek J, Yong CJ, et al. Ramucirumab monotherapy for previously treated advanced gastric or gastro-oesophageal junction adenocarcinoma (REGARD): An international, randomised, multicentre, placebo-controlled, phase 3 trial. *Lancet* 2014;383:31–39.
24. Arunprasath P, Gobu P, Dubashi B, Satheesh S, Balachander J. Rituximab induced myocardial infarction: A fatal drug reaction. *J Cancer Res Ther* 2011;7:346–348.
25. Duran JM, Makarewich CA, Trappanese D, et al. Sorafenib cardiotoxicity increases mortality after myocardial infarction. *Circ Res* 2014;114:1700–1712.
26. Di Lorenzo G, Autorino R, Bruni G, et al. Cardiovascular toxicity following sunitinib therapy in metastatic renal cell carcinoma: A multicenter analysis. *Ann Oncol* 2009;20:1535–1542.
27. Kappers MH, van Esch JH, Sluiter W, et al. Hypertension induced by the tyrosine kinase inhibitor sunitinib is associated with increased circulating endothelin-1 levels. *Hypertension* 2010;56:675–681.
28. Sen F, Yildiz I, Basaran M, et al. Impaired coronary flow reserve in metastatic cancer patients treated with sunitinib. *J Buon* 2013;18:775–781.
29. Choueiri TK, Schutz FA, Je Y, Rosenberg JE, Bellmunt J. Risk of arterial thromboembolic events with sunitinib and sorafenib: A systematic review and meta-analysis of clinical trials. *J Clin Oncol* 2010;28:2280–2285.
30. Numico G, Sicuro M, Silvestris N, et al. Takotsubo syndrome in a patient treated with sunitinib for renal cancer. *J Clin Oncol* 2012;30:e218–e220.
31. Ropert S, Vignaux O, Mir O, Goldwasser F. VEGF pathway inhibition by anticancer agent sunitinib and susceptibility to atherosclerosis plaque disruption. *Invest New Drugs* 2011;29:1497–1499.
32. Lim WT, Ng QS, Ivy P, et al. A Phase II study of pazopanib in Asian patients with recurrent/metastatic nasopharyngeal carcinoma. *Clin Cancer Res* 2011;17:5481–5489.
33. Tefferi A. Nilotinib treatment-associated accelerated atherosclerosis: When is the risk justified? *Leukemia* 2013;27:1939–1940.
34. Coon EA, Zalewski NL, Hoffman EM, Tefferi A, Flemming KD. Nilotinib treatment-associated cerebrovascular disease and stroke. *Am J Hematol* 2013;88:534–535.
35. Tefferi A, Letendre L. Nilotinib treatment-associated peripheral artery disease and sudden death: Yet another reason to stick to imatinib as front-line therapy for chronic myelogenous leukemia. *Am J Hematol* 2011;86:610–611.
36. Levato L, Cantaffa R, Kropp MG, Magro D, Piro E, Molica S. Progressive peripheral arterial occlusive disease and other vascular events during nilotinib therapy in chronic myeloid leukemia: A single institution study. *Eur J Haematol* 2013;90:531–532.
37. Cortes JE, Kantarjian H, Shah NP, et al. Ponatinib in refractory Philadelphia chromosome-positive leukemias. *N Engl J Med* 2012;367:2075–2088.
38. Cortes JE, Kim DW, Pinilla-Ibarz J, et al. A phase 2 trial of ponatinib in Philadelphia chromosome-positive leukemias. *N Engl J Med* 2013;369:1783–1796.
39. Amir E, Seruga B, Niraula S, Carlsson L, Ocana A. Toxicity of adjuvant endocrine therapy in postmenopausal breast cancer patients: A systematic review and meta-analysis. *J Natl Cancer Inst* 2011;103:1299–1309.
40. Goss PE, Ingle JN, Martino S, et al. A randomized trial of letrozole in postmenopausal women after five years of tamoxifen therapy for early-stage breast cancer. *N Engl J Med*. Nov 6 2003;349:1793–1802.
41. Nanda A, Chen MH, Braccioforte MH, Moran BJ, D'Amico AV. Hormonal therapy use for prostate cancer and mortality in men with coronary artery disease-induced congestive heart failure or myocardial infarction. *Jama* 2009;302:866–873.
42. Nguyen PL, Chen MH, Goldhaber SZ, et al. Coronary revascularization and mortality in men with congestive heart failure or prior myocardial infarction who receive androgen deprivation. *Cancer* 2011;117:406–413.
43. Parekh A, Chen MH, Graham P, et al. Role of androgen deprivation therapy in early salvage radiation among patients with prostate-specific antigen level of 0.5 or less. *Clin Genitourin Cancer* 2014;13:e1–e6.
44. Ziehr DR, Chen MH, Zhang D, et al. Association of androgen-deprivation therapy with excess cardiac-specific mortality in men with prostate cancer. *BJU Int* 2014;116:358–365.
45. Carles Galceran J, Bastus Piulats R, Martin-Broto J, et al. A phase II study of vinorelbine and estramustine in patients with hormone-resistant prostate cancer. *Clin Transl Oncol* 2005;7:66–73.
46. Nakagami Y, Ohori M, Sakamoto N, et al. Safety and efficacy of docetaxel, estramustine phosphate and hydrocortisone in hormone-refractory prostate cancer patients. *Int J Urol* 2010;17:629–634.
47. Lubiniecki GM, Berlin JA, Weinstein RB, Vaughn DJ. Thromboembolic events with estramustine phosphate-based chemotherapy in patients with hormone-refractory prostate carcinoma: Results of a meta-analysis. *Cancer* 2004;101:2755–2759.
48. Albertsen PC, Klotz L, Tombal B, et al. Cardiovascular morbidity associated with gonadotropin releasing hormone agonists and an antagonist. *Eur Urol* 2014;65:565–573.
49. Polk A, Vistisen K, Vaage-Nilsen M, Nielsen DL. A systematic review of the pathophysiology of 5-fluorouracil-induced cardiotoxicity. *BMC Pharmacol Toxicol* 2014;15:47.
50. Grunwald MR, Howie L, Diaz LA, J. Takotsubo cardiomyopathy and Fluorouracil: Case report and review of the literature. *J Clin Oncol* 2012;30:e11–e14.
51. Smith SA, Auseon AJ. Chemotherapy-induced takotsubo cardiomyopathy. *Heart Fail Clin* 2013;9:233–242.
52. Togna GI, Togna AR, Franconi M, Caprino L. Cisplatin triggers platelet activation. *Thromb Res*. 2000;99:503–509.
53. Dieckmann KP, Gerl A, Witt J, Hartmann JT. Myocardial infarction and other major vascular events during chemotherapy for testicular cancer. *Ann Oncol* 2010;21:1607–1611.
54. Ito D, Shiraiishi J, Nakamura T, et al. Primary percutaneous coronary intervention and intravascular ultrasound imaging for coronary thrombosis after cisplatin-based chemotherapy. *Heart Vessels* 2012;27:634–638.

Catheterization and Cardiovascular Interventions DOI 10.1002/ccd.

Published on behalf of The Society for Cardiovascular Angiography and Interventions (SCAI).

55. Meinardi MT, Gietema JA, van der Graaf WT, et al. Cardiovascular morbidity in long-term survivors of metastatic testicular cancer. *J Clin Oncol* 2000;18:1725–1732.
56. Gietema JA, Meinardi MT, Messerschmidt J, et al. Circulating plasma platinum more than 10 years after cisplatin treatment for testicular cancer. *Lancet* 2000;355:1075–1076.
57. Huddart RA, Norman A, Shahidi M, et al. Cardiovascular disease as a long-term complication of treatment for testicular cancer. *J Clin Oncol* 2003;21:1513–1523.
58. van den Belt-Dusebout AW, Nuver J, de Wit R, et al. Long-term risk of cardiovascular disease in 5-year survivors of testicular cancer. *J Clin Oncol* 2006;24:467–475.
59. Haignes HS, Wethal T, Aass N, et al. Cardiovascular risk factors and morbidity in long-term survivors of testicular cancer: A 20-year follow-up study. *J Clin Oncol* 2010;28:4649–4657.
60. Feldman DR, Schaffer WL, Steingart RM. Late cardiovascular toxicity following chemotherapy for germ cell tumors. *J Natl Compr Canc Netw* 2012;10:537–544.
61. Gallagher H, Carroll WM, Dowd M, Rochev Y. The effects of vinblastine on endothelial cells. *Endothelium* 2008;15:9–15.
62. Panella M, Ross JE, Garvin K, Martin A. Cardiac sudden death as a result of acute coronary artery thrombosis during chemotherapy for testicular carcinoma. *J Forensic Sci* 2010;55:1384–1388.
63. Gottdiener JS, Appelbaum FR, Ferrans VJ, Deisseroth A, Ziegler J. Cardiotoxicity associated with high-dose cyclophosphamide therapy. *Arch Intern Med* 1981;141:758–763.
64. Katayama M, Imai Y, Hashimoto H, et al. Fulminant fatal cardiotoxicity following cyclophosphamide therapy. *J Cardiol* 2009;54:330–334.
65. Folkman J. Angiogenesis inhibitors: A new class of drugs. *Cancer Biol Ther* 2003;2:S127–133.
66. Moulton KS, Heller E, Konerding MA, et al. Angiogenesis inhibitors endostatin or TNP-470 reduce intimal neovascularization and plaque growth in apolipoprotein E-deficient mice. *Circulation* 1999;99:1726–1732.
67. Moulton KS, Vakili K, Zurakowski D, et al. Inhibition of plaque neovascularization reduces macrophage accumulation and progression of advanced atherosclerosis. *Proc Natl Acad Sci U S A* 2003;100:4736–4741.
68. Herrmann J, Lerman LO, Mukhopadhyay D, Napoli C, Lerman A. Angiogenesis in atherogenesis. *Arterioscler Thromb Vasc Biol* 2006;26:1948–1957.
69. Holm PW, Slart RH, Zeebregts CJ, Hillebrands JL, Tio RA. Atherosclerotic plaque development and instability: A dual role for VEGF. *Ann Med* 2009;41:257–264.
70. Isenberg JS, Martin-Manso G, Maxhimer JB, Roberts DD. Regulation of nitric oxide signalling by thrombospondin 1: Implications for anti-angiogenic therapies. *Nat Rev Cancer* 2009;9:182–194.
71. Franco TH, Khan A, Joshi V, Thomas B. Takotsubo cardiomyopathy in two men receiving bevacizumab for metastatic cancer. *Ther Clin Risk Manag* 2008;4:1367–1370.
72. Chintalgattu V, Rees ML, Culver JC, et al. Coronary microvascular pericytes are the cellular target of sunitinib malate-induced cardiotoxicity. *Sci Transl Med* 2013;5:187ra169.
73. Winnik S, Lohmann C, Siciliani G, et al. Systemic VEGF inhibition accelerates experimental atherosclerosis and disrupts endothelial homeostasis—implications for cardiovascular safety. *Int J Cardiol* 2013;168:2453–2461.
74. Kappers MH, Smedts FM, Horn T, et al. The vascular endothelial growth factor receptor inhibitor sunitinib causes a preeclampsia-like syndrome with activation of the endothelin system. *Hypertension* 2011;58:295–302.
75. Kappers MH, de Beer VJ, Zhou Z, et al. Sunitinib-induced systemic vasoconstriction in swine is endothelin mediated and does not involve nitric oxide or oxidative stress. *Hypertension* 2012;59:151–157.
76. Arima Y, Oshima S, Noda K, et al. Sorafenib-induced acute myocardial infarction due to coronary artery spasm. *J Cardiol* 2009;54:512–515.
77. Porto I, Leo A, Miele L, et al. A case of variant angina in a patient under chronic treatment with sorafenib. *Nat Rev Clin Oncol* 2010;7:476–480.
78. Naib T, Steingart RM, Chen CL. Sorafenib-associated multivesel coronary artery vasospasm. *Herz* 2011;36:348–351.
79. Pantaleo MA, Mandrioli A, Saponara M, et al. Development of coronary artery stenosis in a patient with metastatic renal cell carcinoma treated with sorafenib. *BMC Cancer* 2012;12:231
80. Je Y, Schutz FA, Choueiri TK. Risk of bleeding with vascular endothelial growth factor receptor tyrosine-kinase inhibitors sunitinib and sorafenib: A systematic review and meta-analysis of clinical trials. *Lancet Oncol* 2009;10:967–974.
81. Aichberger KJ, Herndlhofer S, Scherthaner GH, et al. Progressive peripheral arterial occlusive disease and other vascular events during nilotinib therapy in CML. *Am J Hematol* 2011;86:533–539.
82. Quintas-Cardama A, Kantarjian H, Cortes J. Nilotinib-associated vascular events. *Clin Lymphoma Myeloma Leuk* 2012;12:337–340.
83. Armitage JD, Montero C, Benner A, Armitage JO, Bociek G. Acute coronary syndromes complicating the first infusion of rituximab. *Clin Lymphoma Myeloma* 2008;8:253–255.
84. Stamatelopoulos KS, Lekakis JP, Poulakaki NA, et al. Tamoxifen improves endothelial function and reduces carotid intima-media thickness in postmenopausal women. *Am Heart J* 2004;147:1093–1099.
85. Lynch DR, Kickler TS, Rade JJ. Recurrent myocardial infarction associated with gefitinib therapy. *J Thromb Thrombolysis* 2011;32:120–124.
86. Swanson N, Hogrefe K, Javed Q, Malik N, Gershlick AH. Vascular endothelial growth factor (VEGF)-eluting stents: In vivo effects on thrombosis, endothelialization and intimal hyperplasia. *J Invasive Cardiol* 2003;15:688–692.
87. van Werkum JW, Heestermans AA, Zomer AC, et al. Predictors of coronary stent thrombosis: The Dutch Stent Thrombosis Registry. *J Am Coll Cardiol* 2009;53:1399–1409.
88. Cahill TJ, Chowdhury O, Myerson SG, et al. Myocardial infarction with intracardiac thrombosis as the presentation of acute promyelocytic leukemia: Diagnosis and follow-up by cardiac magnetic resonance imaging. *Circulation* 2011;123:e370–e372.
89. Sargsyan Z, Higgins C, Alexandrescu S, Ott DA, Jain SK. Acute promyelocytic leukemia as a cause of intracoronary drug-eluting-stent thrombosis. *Tex Heart Inst J* 2012;39:416–419.
90. Stewart FA, Heeneman S, Te Poele J, et al. Ionizing radiation accelerates the development of atherosclerotic lesions in ApoE<sup>-/-</sup> mice and predisposes to an inflammatory plaque phenotype prone to hemorrhage. *Am J Pathol* 2006;168:649–658.
91. Lee MS, Finch W, Mahmud E. Cardiovascular complications of radiotherapy. *Am J Cardiol* 2013;112:1688–1696.
92. Brosius FC, 3rd, Waller BF, Roberts WC. Radiation heart disease. Analysis of 16 young (aged 15 to 33 years) necropsy patients who received over 3,500 rads to the heart. *Am J Med* 1981;70:519–530.
93. Veinot JP, Edwards WD. Pathology of radiation-induced heart disease: A surgical and autopsy study of 27 cases. *Hum Pathol* 1996;27:766–773.
94. Orzan F, Brusca A, Conte MR, Presbitero P, Figliomeni MC. Severe coronary artery disease after radiation therapy of the chest and mediastinum: Clinical presentation and treatment. *Br Heart J* 1993;69:496–500.
95. Mulrooney DA, Nunnery SE, Armstrong GT, et al. Coronary artery disease detected by coronary computed tomography

- angiography in adult survivors of childhood Hodgkin lymphoma. *Cancer* 2014;120:3536–3544.
96. Rademaker J, Schoder H, Ariaratnam NS, et al. Coronary artery disease after radiation therapy for Hodgkin's lymphoma: Coronary CT angiography findings and calcium scores in nine asymptomatic patients. *AJR Am J Roentgenol* 2008;191:32–37.
  97. Kupeli S, Hazirolan T, Varan A, et al. Evaluation of coronary artery disease by computed tomography angiography in patients treated for childhood Hodgkin's lymphoma. *J Clin Oncol* 2010;28:1025–1030.
  98. Taylor CW, Nisbet A, McGale P, Darby SC. Cardiac exposures in breast cancer radiotherapy: 1950s-1990s. *Int J Radiat Oncol Biol Phys* 2007;69:1484–1495.
  99. Correa CR, Litt HI, Hwang WT, et al. Coronary artery findings after left-sided compared with right-sided radiation treatment for early-stage breast cancer. *J Clin Oncol* 2007;25:3031–3037.
  100. Darby S, McGale P, Correa C, et al. Effect of radiotherapy after breast-conserving surgery on 10-year recurrence and 15-year breast cancer death: Meta-analysis of individual patient data for 10,801 women in 17 randomised trials. *Lancet* 2011;378:1707–1716.
  101. Mulrooney DA, Yeazel MW, Kawashima T, et al. Cardiac outcomes in a cohort of adult survivors of childhood and adolescent cancer: Retrospective analysis of the Childhood Cancer Survivor Study cohort. *BMJ* 2009;339:b4606.
  102. Lee CC, Su YC, Ho HC, et al. Increased risk of ischemic stroke in young nasopharyngeal carcinoma patients. *Int J Radiat Oncol Biol Phys* 2011;81:e833–e838.
  103. Lam WW, Leung SF, So NM, et al. Incidence of carotid stenosis in nasopharyngeal carcinoma patients after radiotherapy. *Cancer* 2001;92:2357–2363.
  104. Dorresteijn LD, Kappelle AC, Boogerd W, et al. Increased risk of ischemic stroke after radiotherapy on the neck in patients younger than 60 years. *J Clin Oncol* 2002;20:282–288.
  105. Cheng SW, Ting AC, Lam LK, Wei WI. Carotid stenosis after radiotherapy for nasopharyngeal carcinoma. *Arch Otolaryngol Head Neck Surg* 2000;126:517–521.
  106. Al-Mubarak N, Roubin GS, Iyer SS, et al. Carotid stenting for severe radiation-induced extracranial carotid artery occlusive disease. *J Endovasc Ther* 2000;7:36–40.
  107. Ting AC, Cheng SW, Yeung KM, et al. Carotid stenting for radiation-induced extracranial carotid artery occlusive disease: Efficacy and midterm outcomes. *J Endovasc Ther* 2004;11:53–59.
  108. Houdart E, Mounayer C, Chapot R, Saint-Maurice JP, Merland JJ. Carotid stenting for radiation-induced stenoses: A report of 7 cases. *Stroke* 2001;32:118–121.
  109. Benitez RP, Armonda RA, Harrop J, Thomas JE, Rosenwasser RH. Carotid angioplasty and stenting for recurrent and radiation-induced stenosis: Preliminary experience. *Neurosurg Focus* 1998;5:e14.
  110. Hull MC, Morris CG, Pepine CJ, Mendenhall NP. Valvular dysfunction and carotid, subclavian, and coronary artery disease in survivors of Hodgkin lymphoma treated with radiation therapy. *Jama* 2003;290:2831–2837.
  111. De Bruin ML, Dorresteijn LD, van't Veer MB, et al. Increased risk of stroke and transient ischemic attack in 5-year survivors of Hodgkin lymphoma. *J Natl Cancer Inst* 2009;101:928–937.
  112. Milutinovic J, Darcy M, Thompson KA. Radiation-induced renovascular hypertension successfully treated with transluminal angioplasty: Case report. *Cardiovasc Intervent Radiol* 1990;13:29–31.
  113. Saka B, Bilge AK, Umman B, et al. Bilateral renal artery stenosis after abdominal radiotherapy for Hodgkin's disease. *Int J Clin Pract* 2003;57:247–248.
  114. Jurado JA, Bashir R, Burket MW. Radiation-induced peripheral artery disease. *Catheter Cardiovasc Interv* 2008;72:563–568.
  115. Saliou C, Julia P, Feito B, Renaudin JM, Fabiani JN. Radiation-induced arterial disease of the lower limb. *Ann Vasc Surg* 1997;11:173–177.
  116. Moutardier V, Christophe M, Lelong B, Houvenaeghel G, Delpero JR. Iliac atherosclerotic occlusive disease complicating radiation therapy for cervix cancer: A case series. *Gynecol Oncol* 2002;84:456–459.
  117. Jones LW, Haykowsky MJ, Swartz JJ, Douglas PS, Mackey JR. Early breast cancer therapy and cardiovascular injury. *J Am Coll Cardiol* 2007;50:1435–1441.
  118. Herrmann J, Lerman A. An update on cardio-oncology. *Trends Cardiovasc Med* 2014;24:285–295.
  119. Vejpongsa P, Yeh ET. Prevention of anthracycline-induced cardiotoxicity: Challenges and opportunities. *J Am Coll Cardiol* 2014;64:938–945.
  120. Kaya MG, Ozkan M, Gunebakmaz O, et al. Protective effects of nebivolol against anthracycline-induced cardiomyopathy: A randomized control study. *Int J Cardiol* 2013;167:2306–2310.
  121. Elitok A, Oz F, Cizgici AY, et al. Effect of carvedilol on silent anthracycline-induced cardiotoxicity assessed by strain imaging: A prospective randomized controlled study with six-month follow-up. *Cardiol J* 2014;21:509–515.
  122. Bosch X, Rovira M, Sitges M, et al. Enalapril and carvedilol for preventing chemotherapy-induced left ventricular systolic dysfunction in patients with malignant hemopathies: The OVERCOME trial (preventiOn of left Ventricular dysfunction with Enalapril and carvedilol in patients submitted to intensive Chemotherapy for the treatment of Malignant hEmopathies). *J Am Coll Cardiol* 2013;61:2355–2362.
  123. Cardinale D, Colombo A, Sandri MT, et al. Prevention of high-dose chemotherapy-induced cardiotoxicity in high-risk patients by angiotensin-converting enzyme inhibition. *Circulation* 2006;114:2474–2481.
  124. Greenland P, Alpert JS, Beller GA, et al. 2010 ACCF/AHA guideline for assessment of cardiovascular risk in asymptomatic adults: A report of the American College of Cardiology Foundation/American Heart Association Task Force on Practice Guidelines. *J Am Coll Cardiol* 2010;56:e50–e103.
  125. Hong JC, Kruser TJ, Gondi V, et al. Risk of cerebrovascular events in elderly patients after radiation therapy versus surgery for early-stage glottic cancer. *Int J Radiat Oncol Biol Phys* 2013;87:290–296.
  126. Darby SC, Ewertz M, McGale P, et al. Risk of ischemic heart disease in women after radiotherapy for breast cancer. *N Engl J Med* 2013;368:987–998.
  127. Al Rifai M, Cainzos-Achirica M, Blaha MJ. Establishing the warranty of a coronary artery calcium score of zero. *Atherosclerosis* 2015;238:1–3.
  128. Heidenreich PA, Schnittger I, Strauss HW, et al. Screening for coronary artery disease after mediastinal irradiation for Hodgkin's disease. *J Clin Oncol* 2007;25:43–49.
  129. Min JK, Lin FY, Gidseg DS, et al. Determinants of coronary calcium conversion among patients with a normal coronary calcium scan: What is the "warranty period" for remaining normal? *J Am Coll Cardiol* 2010;55:1110–1117.
  130. van Nimwegen FA, Schaapveld M, Janus CP, et al. Cardiovascular disease after Hodgkin lymphoma treatment: 40-year disease risk. *JAMA Intern Med* 2015;175:1007–1017.
  131. Oeffinger KC, Mertens AC, Sklar CA, et al. Chronic health conditions in adult survivors of childhood cancer. *N Engl J Med* 2006;355:1572–1582.
  132. Steingart RM, Yadav N, Manrique C, Carver JR, Liu J. Cancer survivorship: Cardiotoxic therapy in the adult cancer patient; cardiac outcomes with recommendations for patient management. *Semin Oncol* 2013;40:690–708.
  133. Armenian SH, Hudson MM, Mulder RL, et al. Recommendations for cardiomyopathy surveillance for survivors of

Catheterization and Cardiovascular Interventions DOI 10.1002/ccd.

Published on behalf of The Society for Cardiovascular Angiography and Interventions (SCAI).

- childhood cancer: A report from the International Late Effects of Childhood Cancer Guideline Harmonization Group. *Lancet Oncol* 2015;16:e123–e136.
134. Lancellotti P, Nkomo VT, Badano LP, et al. Expert consensus for multi-modality imaging evaluation of cardiovascular complications of radiotherapy in adults: A report from the European Association of Cardiovascular Imaging and the American Society of Echocardiography. *J Am Soc Echocardiogr* 2013;26:1013–1032.
  135. Detrano R, Guerci AD, Carr JJ, et al. Coronary calcium as a predictor of coronary events in four racial or ethnic groups. *N Engl J Med* 2008;358:1336–1345.
  136. Amsterdam EA, Wenger NK, Brindis RG, et al. 2014 AHA/ACC guideline for the management of patients with non-ST-elevation acute coronary syndromes: A report of the American college of cardiology/American heart association task force on practice guidelines. *J Am Coll Cardiol* 2014;64:2645–2687.
  137. Elting LS, Rubenstein EB, Martin CG, et al. Incidence, cost, and outcomes of bleeding and chemotherapy dose modification among solid tumor patients with chemotherapy-induced thrombocytopenia. *J Clin Oncol* 2001;19:1137–1146.
  138. Hakim DA, Dargas GD, Caixeta A, et al. Impact of baseline thrombocytopenia on the early and late outcomes after ST-elevation myocardial infarction treated with primary angioplasty: Analysis from the Harmonizing Outcomes with Revascularization and Stents in Acute Myocardial Infarction (HORIZONS-AMI) trial. *Am Heart J* 2011;161:391–396.
  139. Kereiakes DJ, Berkowitz SD, Lincoff AM, et al. Clinical correlates and course of thrombocytopenia during percutaneous coronary intervention in the era of abciximab platelet glycoprotein IIb/IIIa blockade. *Am Heart J* 2000;140:74–80.
  140. Makoni SN. Acute profound thrombocytopenia following angioplasty: The dilemma in the management and a review of the literature. *Heart* 2001;86:E18.
  141. McClure MW, Berkowitz SD, Sparapani R, et al. Clinical significance of thrombocytopenia during a non-ST-elevation acute coronary syndrome. The platelet glycoprotein IIb/IIIa in unstable angina: Receptor suppression using integrilin therapy (PURSUIT) trial experience. *Circulation* 1999;99:2892–2900.
  142. Harrington RA, Sane DC, Califf RM, et al. Clinical importance of thrombocytopenia occurring in the hospital phase after administration of thrombolytic therapy for acute myocardial infarction. The Thrombolysis and Angioplasty in Myocardial Infarction Study Group. *J Am Coll Cardiol* 1994;23:891–898.
  143. Manor SM, Guillory GS, Jain SP. Clopidogrel-induced thrombotic thrombocytopenic purpura-hemolytic uremic syndrome after coronary artery stenting. *Pharmacotherapy* 2004;24:664–667.
  144. Best PJ, Mathew V, Markovic SN. Clopidogrel-associated autoimmune thrombocytopenic purpura. *Catheter Cardiovasc Interv* 2004;62:339–340.
  145. Nikolsky E, Sadeghi HM, Effron MB, et al. Impact of in-hospital acquired thrombocytopenia in patients undergoing primary angioplasty for acute myocardial infarction. *Am J Cardiol* 2005;96:474–481.
  146. Sideris SK, Bonios MJ, Eftihiadis EE, et al. Severe thrombocytopenia after heparin therapy in a patient with unstable angina and recent stent implantation. *Hellenic J Cardiol* 2005;46:242–246.
  147. Bovill EG, Terrin ML, Stump DC, et al. Hemorrhagic events during therapy with recombinant tissue-type plasminogen activator, heparin, and aspirin for acute myocardial infarction. Results of the Thrombolysis in Myocardial Infarction (TIMI), Phase II Trial. *Ann Intern Med* 1991;115:256–265.
  148. Merlini PA, Rossi M, Menozzi A, et al. Thrombocytopenia caused by abciximab or tirofiban and its association with clinical outcome in patients undergoing coronary stenting. *Circulation* 2004;109:2203–2206.
  149. Luzzatto G, Schafer AI. The prethrombotic state in cancer. *Semin Oncol* 1990;17:147–159.
  150. Bobbio-Pallavicini E, Gugliotta L, Centurioni R, et al. Antiplatelet agents in thrombotic thrombocytopenic purpura (TTP). Results of a randomized multicenter trial by the Italian Cooperative Group for TTP. *Haematologica* 1997;82:429–435.
  151. Alarcon-Segovia D, Sanchez-Guerrero J. Correction of thrombocytopenia with small dose aspirin in the primary antiphospholipid syndrome. *J Rheumatol* 1989;16:1359–1361.
  152. Fayyad AM, Brummitt DR, Barker HF, Spooner SF. May-heggin anomaly: The role of aspirin in the treatment of this rare platelet disorder in pregnancy. *Bjog* 2002;109:223–224.
  153. Sarkiss MG, Yusuf SW, Warneke CL, et al. Impact of aspirin therapy in cancer patients with thrombocytopenia and acute coronary syndromes. *Cancer* 2007;109:621–627.
  154. Kroll MH, Feng S. Targeting shear stress-induced platelet activation: Is lesion-specific antiplatelet therapy a realistic clinical goal? *Expert Rev Cardiovasc Ther* 2005;3:941–951.
  155. Yusuf SW, Iliescu C, Bathina JD, Daher IN, Durand JB. Antiplatelet therapy and percutaneous coronary intervention in patients with acute coronary syndrome and thrombocytopenia. *Tex Heart Inst J* 2010;37:336–340.
  156. Schiffer CA, Anderson KC, Bennett CL, et al. Platelet transfusion for patients with cancer: Clinical practice guidelines of the American Society of Clinical Oncology. *J Clin Oncol* 2001;19:1519–1538.
  157. Iliescu C, Durand JB, Kroll M. Cardiovascular interventions in thrombocytopenic cancer patients. *Tex Heart Inst J* 2011;38:259–260.
  158. Mahla E, Suarez TA, Bliden KP, et al. Platelet function measurement-based strategy to reduce bleeding and waiting time in clopidogrel-treated patients undergoing coronary artery bypass graft surgery: The timing based on platelet function strategy to reduce clopidogrel-associated bleeding related to CABG (TARGET-CABG) study. *Circ Cardiovasc Interv* 2012;5:261–269.
  159. Kwak YL, Kim JC, Choi YS, et al. Clopidogrel responsiveness regardless of the discontinuation date predicts increased blood loss and transfusion requirement after off-pump coronary artery bypass graft surgery. *J Am Coll Cardiol* 2010;56:1994–2002.
  160. Leibovitz A, Baumohl Y, Walach N, et al. Medical staff attitudes: Views and positions regarding blood transfusion to terminally ill cancer patients. *Am J Clin Oncol* 2004;27:542–546.
  161. Goksu SS, Gunduz S, Unal D, et al. Use of blood transfusion at the end of life: Does it have any effects on survival of cancer patients? *Asian Pac J Cancer Prev* 2014;15:4251–4254.
  162. Bokemeyer C, Aapro MS, Courdi A, et al. EORTC guidelines for the use of erythropoietic proteins in anaemic patients with cancer: 2006 update. *Eur J Cancer* 2007;43:258–270.
  163. Ellis SG, Bhatt D, Kapadia S, et al. Correlates and outcomes of retroperitoneal hemorrhage complicating percutaneous coronary intervention. *Catheter Cardiovasc Interv* 2006;67:541–545.
  164. Matthai WH, J. Thrombocytopenia in cardiovascular patients: Diagnosis and management. *Chest* 2005;127:46S–52S.
  165. Velders MA, Boden H, Hofma SH, et al. Outcome after ST elevation myocardial infarction in patients with cancer treated with primary percutaneous coronary intervention. *Am J Cardiol* 2013;112:1867–1872.
  166. Gustavsson A, Osterman B, Cavallin-Stahl E. A systematic overview of radiation therapy effects in Hodgkin's lymphoma. *Acta Oncol* 2003;42:589–604.
  167. Beneficial and harmful effects of anthracyclines in the treatment of childhood acute lymphoblastic leukaemia: A systematic review and meta-analysis. *Br J Haematol* 2009;145:376–388.

168. Kupeli S. Risks and diagnosis of coronary artery disease in Hodgkin lymphoma survivors. *World J Cardiol* 2014;6:555–561.
169. Falanga A, Russo L, Verzeroli C. Mechanisms of thrombosis in cancer. *Thromb Res* 2013;131:S59–S62.
170. Franchini M, Minno D, Coppola MNA. Disseminated intravascular coagulation in hematologic malignancies. *Semin Thromb Hemost* 2010;36:388–403.
171. De Cicco M. The prothrombotic state in cancer: Pathogenic mechanisms. *Crit Rev Oncol Hematol* 2004;50:187–196.
172. Lee JM, Yoon CH. Acute coronary stent thrombosis in cancer patients: A case series report. *Korean Circ J* 2012;42:487–491.
173. Lenihan DJ, Cardinale D, Cipolla CM. The compelling need for a cardiology and oncology partnership and the birth of the International CardiOncology Society. *Prog Cardiovasc Dis* 2010;53:88–93.
174. Mann DL, Krone RJ. Cardiac disease in cancer patients: An overview. *Prog Cardiovasc Dis* 2010;53:80–87.
175. Hong RA, Iimura T, Sumida KN, Eager RM. Cardio-oncology/onco-cardiology. *Clin Cardiol* 2010;33:733–737.
176. Patel IJ, Davidson JC, Nikolic B, et al. Consensus guidelines for periprocedural management of coagulation status and hemostasis risk in percutaneous image-guided interventions. *J Vasc Interv Radiol* 2012;23:727–736.
177. Krone RJ. Managing coronary artery disease in the cancer patient. *Prog Cardiovasc Dis* 2010;53:149–156.
178. Bertrand OF, Belisle P, Joyal D, et al. Comparison of transradial and femoral approaches for percutaneous coronary interventions: A systematic review and hierarchical Bayesian meta-analysis. *Am Heart J* 2012;163:632–648.
179. Kedev S, Kalpak O, Dharma S, et al. Complete transitioning to the radial approach for primary percutaneous coronary intervention: A real-world single-center registry of 1808 consecutive patients with acute ST-elevation myocardial infarction. *J Invasive Cardiol* 2014;26:475–482.
180. Bangalore S, Bhatt DL. Femoral arterial access and closure. *Circulation* 2011;124:e147–e156.
181. Ben-Dor I, Maluenda G, Mahmoudi M, et al. A novel, minimally invasive access technique versus standard 18-gauge needle set for femoral access. *Catheter Cardiovasc Interv* 2012;79:1180–1185.
182. Rao SV, Tremmel JA, Gilchrist IC, et al. Best practices for transradial angiography and intervention: A consensus statement from the society for cardiovascular angiography and intervention's transradial working group. *Catheter Cardiovasc Interv* 2014;83:228–236.
183. Abdelaal E, Rimac G, Plourde G, et al. 4Fr in 5Fr sheathless technique with standard catheters for transradial coronary interventions: Technical challenges and persisting issues. *Catheter Cardiovasc Interv* 2015;85:816–817.
184. Sciahbasi A, Fischetti D, Picciolo A, et al. Transradial access compared with femoral puncture closure devices in percutaneous coronary procedures. *Int J Cardiol* 2009;137:199–205.
185. Belli AM, Cumberland DC, Knox AM, Procter AE, Welsh CL. The complication rate of percutaneous peripheral balloon angioplasty. *Clin Radiol* 1990;41:380–383.
186. Kim D, Orron DE, Skillman JJ, et al. Role of superficial femoral artery puncture in the development of pseudoaneurysm and arteriovenous fistula complicating percutaneous transfemoral cardiac catheterization. *Cathet Cardiovasc Diagn* 1992;25:91–97.
187. Sherev DA, Shaw RE, Brent BN. Angiographic predictors of femoral access site complications: Implication for planned percutaneous coronary intervention. *Catheter Cardiovasc Interv* 2005;65:196–202.
188. Krishnasamy VP, Hagar MJ, Scher DJ, et al. Vascular closure devices: Technical tips, complications, and management. *Tech Vasc Interv Radiol* 2015;18:100–112.
189. Fefer P, Matetzky S, Gannot S, et al. Predictors and outcomes associated with radial versus femoral access for intervention in patients with acute coronary syndrome in a real-world setting: Results from the Acute Coronary Syndrome Israeli Survey (ACSIS) 2010. *J Invasive Cardiol* 2014;26:398–402.
190. Jao GT, Knovich MA, Savage RW, Sane DC. ST-elevation myocardial infarction and myelodysplastic syndrome with acute myeloid leukemia transformation. *Tex Heart Inst J* 2014;41:234–237.
191. Hamon M, Rasmussen LH, Manoukian SV, et al. Choice of arterial access site and outcomes in patients with acute coronary syndromes managed with an early invasive strategy: The ACU-ITY trial. *EuroIntervention* 2009;5:115–120.
192. Nathan S, Rao SV. Radial versus femoral access for percutaneous coronary intervention: Implications for vascular complications and bleeding. *Curr Cardiol Rep* 2012;14:502–509.
193. Fujii T, Masuda N, Tamiya S, et al. Angiographic evaluation of right upper-limb arterial anomalies: Implications for transradial coronary interventions. *J Invasive Cardiol* 2010;22:536–540.
194. Lo TS, Ratib K, Chong AY, et al. Impact of access site selection and operator expertise on radiation exposure; a controlled prospective study. *Am Heart J* 2012;164:455–461.
195. Yan Z, Zhou Y, Zhao Y, et al. Impact of transradial coronary procedures on radial artery function. *Angiology* 2014;65:104–107.
196. Fleisher LA, Beckman JA, Brown KA, et al. ACC/AHA 2007 Guidelines on perioperative cardiovascular evaluation and care for noncardiac surgery: Executive summary: A report of the American college of cardiology/American heart association task force on practice guidelines (writing committee to revise the 2002 guidelines on perioperative cardiovascular evaluation for noncardiac surgery) developed in collaboration with the American society of echocardiography, American society of nuclear cardiology, heart rhythm society, society of cardiovascular anesthesiologists, society for cardiovascular angiography and interventions, society for vascular medicine and biology, and society for vascular surgery. *J Am Coll Cardiol* 2007;50:1707–1732.
197. Vieira RD, Pereira AC, Lima EG, et al. Cancer-related deaths among different treatment options in chronic coronary artery disease: Results of a 6-year follow-up of the MASS II study. *Coron Artery Dis* 2012;23:79–84.
198. Abbott JD, Ahmed HN, Vlachos HA, Selzer F, Williams DO. Comparison of outcome in patients with ST-elevation versus non-ST-elevation acute myocardial infarction treated with percutaneous coronary intervention (from the National Heart, Lung, and Blood Institute Dynamic Registry). *Am J Cardiol* 2007;100:190–195.
199. Yusuf SW, Daraban N, Abbasi N, et al. Treatment and outcomes of acute coronary syndrome in the cancer population. *Clin Cardiol* 2012;35:443–450.
200. Kurisu S, Iwasaki T, Ishibashi K, et al. Comparison of treatment and outcome of acute myocardial infarction between cancer patients and non-cancer patients. *Int J Cardiol* 2013;167:2335–2337.
201. Goloshchapov-Aksenov RS, Lebedev AV, Mirzonov VA. Primary percutaneous coronary angioplasty in patients with acute coronary syndrome and concomitant cancer. *Vestn Rentgenol Radiol* 2012;(1):17–20.
202. Mauri L, Kereiakes DJ, Yeh RW, et al. Twelve or 30 months of dual antiplatelet therapy after drug-eluting stents. *N Engl J Med* 2014;371:2155–2166.

Catheterization and Cardiovascular Interventions DOI 10.1002/ccd.

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203. Schiano P, Barbou F, Chenilleau MC, Louembe J, Monsegu J. Adjusted weight anticoagulation for radial approach in elective coronarography: The AWARE coronarography study. *EuroIntervention* 2010;6:247–250.
204. Chan AO, Jim MH, Lam KF, et al. Prevalence of colorectal neoplasm among patients with newly diagnosed coronary artery disease. *Jama* 2007;298:1412–1419.
205. Neugut AI, Lebowitz B. Is the prevalence of colorectal neoplasm higher in patients with coronary artery disease? *Nat Clin Pract Oncol* 2008;5:248–249.
206. Shivaraju A, Patel V, Fonarow GC, et al. Temporal trends in gastrointestinal bleeding associated with percutaneous coronary intervention: Analysis of the 1998–2006 Nationwide Inpatient Sample (NIS) database. *Am Heart J* 2011;162:1062–1068.
207. Topol EJ, Nissen SE. Our preoccupation with coronary luminology. The dissociation between clinical and angiographic findings in ischemic heart disease. *Circulation* 1995;92:2333–2342.
208. Nissen SE, Gurley JC. Assessment of the functional significance of coronary stenoses. Is digital angiography the answer? *Circulation* 1990;81:1431–1435.
209. Kern MJ, Lerman A, Bech JW, et al. Physiological assessment of coronary artery disease in the cardiac catheterization laboratory: A scientific statement from the American Heart Association Committee on Diagnostic and Interventional Cardiac Catheterization, Council on Clinical Cardiology. *Circulation* 2006;114:1321–1341.
210. Mintz GS, Popma JJ, Pichard AD, et al. Limitations of angiography in the assessment of plaque distribution in coronary artery disease: A systematic study of target lesion eccentricity in 1446 lesions. *Circulation* 1996;93:924–931.
211. De Bruyne B, Hersbach F, Pijls NH, et al. Abnormal epicardial coronary resistance in patients with diffuse atherosclerosis but “Normal” coronary angiography. *Circulation* 2001;104:2401–2406.
212. Cheneau E, Leborgne L, Mintz GS, et al. Predictors of subacute stent thrombosis: Results of a systematic intravascular ultrasound study. *Circulation* 2003;108:43–47.
213. Prati F, Kodama T, Romagnoli E, et al. Suboptimal stent deployment is associated with subacute stent thrombosis: Optical coherence tomography insights from a multicenter matched study. From the CLI Foundation investigators: The CLI-THRO study. *Am Heart J* 2015;169:249–256.
214. Nilsson G, Holmberg L, Garmo H, et al. Distribution of coronary artery stenosis after radiation for breast cancer. *J Clin Oncol* 2012;30:380–386.
215. Lipshultz SE, Sallan SE. Cardiovascular abnormalities in long-term survivors of childhood malignancy. *J Clin Oncol* 1993;11:1199–1203.
216. La Francesca S, Frazier OH, Radovancevic B, et al. Concomitant cardiac and pulmonary operations for lung cancer. *Tex Heart Inst J* 1995;22:296–300.
217. Saxena P, Tam RK. Combined off-pump coronary artery bypass surgery and pulmonary resection. *Ann Thorac Surg* 2004;78:498–501.
218. Tsuji Y, Morimoto N, Tanaka H, et al. Surgery for gastric cancer combined with cardiac and aortic surgery. *Arch Surg* 2005;140:1109–1114.
219. Ozsoyler I, Yilic L, Bozok S, et al. Off-pump coronary artery bypass surgery in patients with coronary artery disease and malign neoplasia: Results of ten patients and review of the literature. *Heart Vessels* 2006;21:365–367.
220. Schoenmakers MC, van Boven WJ, van den Bosch J, van Swieten HA. Comparison of on-pump or off-pump coronary artery revascularization with lung resection. *Ann Thorac Surg* 2007;84:504–509.
221. Samuels LE, Kaufman MS, Morris RJ, Styler M, Brockman SK. Open heart surgery in patients with chronic lymphocytic leukemia. *Leuk Res* 1999;23:71–75.
222. Christiansen S, Schmid C, Loher A, Scheld HH. Impact of malignant hematological disorders on cardiac surgery. *Cardiovasc Surg* 2000;8:149–152.
223. Fecher AM, Birdas TJ, Haybron D, et al. Cardiac operations in patients with hematologic malignancies. *Eur J Cardiothorac Surg* 2004;25:537–540.
224. Potapov EV, Zurbrugg HR, Herzke C, et al. Impact of cardiac surgery using cardiopulmonary bypass on course of chronic lymphatic leukemia: A case-control study. *Ann Thorac Surg* 2002;74:384–389.
225. Patel MR, Dehmer GJ, Coronary Revascularization Writing G, et al. ACCF/SCAI/STS/AATS/AHA/ASNC/HFSA/SCCT 2012 appropriate use criteria for coronary revascularization focused update: A report of the American College of Cardiology Foundation Appropriate Use Criteria Task Force, Society for Cardiovascular Angiography and Interventions, Society of Thoracic Surgeons, American Association for Thoracic Surgery, American Heart Association, American Society of Nuclear Cardiology, and the Society of Cardiovascular Computed Tomography. *J Thorac Cardiovasc Surg* 2012;143:780–803.
226. Yeh ET, Tong AT, Lenihan DJ, et al. Cardiovascular complications of cancer therapy: Diagnosis, pathogenesis, and management. *Circulation* 2004;109:3122–3131.
227. Jaworski C, Mariani JA, Wheeler G, Kaye DM. Cardiac complications of thoracic irradiation. *J Am Coll Cardiol* 2013;61:2319–2328.
228. Williams PD, Palmer S, Judkins C, et al. Right and left heart catheterization via an antecubital fossa vein and the radial artery—a prospective study. *J Invasive Cardiol* 2014;26:669–673.
229. Cooper LT, Baughman KL, Feldman AM, et al. The role of endomyocardial biopsy in the management of cardiovascular disease: A scientific statement from the American Heart Association, the American College of Cardiology, and the European Society of Cardiology. Endorsed by the Heart Failure Society of America and the Heart Failure Association of the European Society of Cardiology. *J Am Coll Cardiol* 2007;50:1914–1931.
230. Tsang TS, Enriquez-Sarano M, Freeman WK, et al. Consecutive 1127 therapeutic echocardiographically guided pericardiocenteses: Clinical profile, practice patterns, and outcomes spanning 21 years. *Mayo Clin Proc* 2002;77:429–436.
231. Silvestry FE, Kerber RE, Brook MM, et al. Echocardiography-guided interventions. *J Am Soc Echocardiogr* 2009;22:213–231. quiz 316–217.
232. Tsang TS, Seward JB, Barnes ME, et al. Outcomes of primary and secondary treatment of pericardial effusion in patients with malignancy. *Mayo Clin Proc* 2000;75:248–253.
233. Virk SA, Chandrakumar D, Villanueva C, Wolfenden H, Liou K, Cao C. Systematic review of percutaneous interventions for malignant pericardial effusion. *Heart* 2015;101:1619–1626.
234. Swanson N, Mirza I, Wijesinghe N, Devlin G. Primary percutaneous balloon pericardiotomy for malignant pericardial effusion. *Catheter Cardiovasc Interv* 2008;71:504–507.
235. Bavaria JE, Szeto WY, Roche LA, et al. The progression of a transcatheter aortic valve program: A decision analysis of more than 680 patient referrals. *Ann Thorac Surg* 2011;92:2072–2076. discussion 2076–2077.
236. Kogoj P, Devjak R, Bunc M. Balloon aortic valvuloplasty (BAV) as a bridge to aortic valve replacement in cancer patients who require urgent non-cardiac surgery. *Radiol Oncol* 2014;48:62–66.

Catheterization and Cardiovascular Interventions DOI 10.1002/ccd.

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