Cardiotoxicity induced by antineoplastic drug Daunorubicin and its amelioration: A review of literature

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Abstract
Daunorubicin (DNR) belongs to anthracycline class of drugs that has been reported with a significant role in treating leukemia. But use of DNR is restricted due to associated cardiotoxicity. In this review, we present the cardiotoxic effects associated with DNR along with some ameliorative agents. DNR induced sub chronic cardiomyopathy, down regulation of cytokines and stem cell markers which may reflect impaired chemotaxis, migration and homing of stem cells; and tissue repair in the heart. DNR may affect the heart by forming hydrophobic interactions and hydrogen bonds with cardiac myosin. DNR cardiotoxicity may change with age of patient as systematic clearance of drug decreases with age. Cardiotoxicity effects may include inhibition of cardioprotective epoxyeicosatrienoic acid. Adenosine, sodium ferulate and pomegranate have been reported as promising cardio-protectants against DNR induced cardiotoxicity. GnRH conjugates containing DNR also showed no associated cytotoxic effects on cardiomyocytes. Future research on amelioration of DNR associated toxicity is highly recommended.

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Introduction
Anthracycline is a type of antibiotic that comes from Streptomyces peucetius bacteria. Each anthracycline drug contains rubicin as suffix in its name, which is derived from Latin word rubidus, that means red. Anthracyclines are aromatic polyketides available in large variety of forms because of structural differences in the aglycone [1]. The generic name anthracycline was given by H. Brockmann to a group of glycosidic compounds whose aglycones, called anthracyclinones, had a basic structure of 7,8,9,10-Tetrahydro 5,12-naphthacene quinone [1]. The common anthracycline drugs include daunorubicin, doxorubicin, epirubicin and idarubicin. Important functional characteristics of the anthracycline antibiotics are (a) the planar anthraquinone ring system (b) quinone groups on the unsaturated rings (c) stereochemistry of the D ring substitution at position 9 and (d) the amino sugar, daunosamine, which provides water solubility and chemical architecture for stabilizing DNA binding [2].

Anthracyclines are used to treat many types of cancer. These are effective against broad range of malignancies including solid and hematological tumors. These are also used against leukemia, lymphoma, breast cancer, bladder cancer, uterine cancer, bowel cancer and other broad-spectrum therapies which make them first-line chemotherapeutic drugs [1]. Anthracyclines are known to inhibit the synthesis of DNA and RNA by intercalation and base modification, thus preventing the replication of rapidly growing cancer cells [3]. These interact with topoisomerase II enzyme which results in disturbance in the replication and transcription processes [3]. The mechanism may involve the production of iron-mediated free oxygen radicals that damage the DNA and cell membranes (Figure 1) [3].

Daunorubicin
The first anthracycline drug identified was DNR from the soil bacterium Streptomyces peucetius. DNR was found effective in treatment of leukemia and lymphomas in 1960’s [4]. It was the first anthracycline drug analog to be characterized structurally and sterochemically [1].

DNR drug is available both in the form of solution and powder which is given intravenously to the patients along with other chemotherapeutic medications [5]. The dose and schedule are determined by the person’s size and the treatment regimen being used. It can be administered alone or in combination with other drugs [6]. DNR is injected once a day, when it is used to treat the acute myeloid leukemia (AML) and once a
week in case of acute lymphocytic leukemia [5]. Urine color of patient turns pink or red due to introduction of DNR. It is not blood, but due to renal clearance of the drug from the patient's body [6]. DNR can act during multiple phases of the cell cycle and is considered cell-cycle specific [7].

DNR is beneficial in the treatment of leukemia, but its use is restricted due to associated cardiotoxic effects. Decrease in cardiac function in AML patient's after treatment with DNR in combination with cytarabine can be determined by drop in left ventricular ejection [8]. DNR may interfere with the pumping action of heart or may lower the body's ability to make blood cells. Thus, patient may need blood transfusion leading to bleeding complications and infections [7]. An early biomarker of DNR-induced cardiotoxicity is troponin and its level may be taken as directly proportional to the amount of cardiac damage. Troponin is an intracellular enzyme which can be detected in the blood when cells have lysed and expelled their contents. In case of DNR toxicity, a rise in troponin may indicate cardiomyocyte apoptosis as well as myofibril degradation [9]. Heart damage may occur during the treatment or after many months or years [5]. A patient can receive only up to a certain amount of DNR during his lifetime. The 'lifetime maximum dose' may be lesser if patient has heart risk factors like old age, radiation to chest and use of other toxic medications [7]. Another side effect of DNR may include leaking out of DNR from administering line into patient's body which can cause damage to the muscles and skin in contact. DNR is a vesicant that may cause extensive tissue damage and blistering if it escapes from the vein. Administered DNR may cause burning sensation through the arm of patient and discoloration around the administering site (Table 1) [7].

Several studies have been retrieved which demonstrated different effects of DNR on heart. Different models including rat and human have been used. In a study, male Wistar rats were treated with DNR to induce acute cardiomyopathy (6 × 3 mg/kg, i.p., every 48 hr, DAU-A) or subchronic cardiomyopathy (15 mg/kg, i.v., DAU-C). In this experiment, reduced body weight, decrease in left ventricular weight and elevated Nppa, Nppb and Myh7 (isomyosins) levels were observed in both groups, but Myh6 decreased only in DAU-C group. Up-regulated gp91phox and down-regulated Abcb8 were also found only in DAU-C group. Experiment also showed decreased expressions of Scf and Vegf (cytokines), as well as of stem cell markers. All these effects caused down-regulation of cytokines and stem cell markers which reflected impaired chemotaxis, migration and homing of stem cells; and tissue repair in the heart in sub-chronic but not acute model of DAU cardiomyopathy [10]. In another study, the heart samples were collected from donors with and without Down syndrome. These samples were used to investigate the determinants for anthracycline-related cardiotoxicity including cardiac daunorubicin reductase activity (DA), carbonyl reductase (CR)/ aldo-keto reductase (AKRs) protein expression, mitochondrial DNA content (mtDNA), and AKR7A2 DNA methylation status. Clinical evidences suggested that patients with leukemia and Down syndrome were at increased risk for anthracycline-related cardiotoxicity. CR and AKRs catalyze the reduction of DNR and doxorubicin into cardiotoxic C-13 alcohol metabolites. Studies revealed that cardiac mtDNA content, mtDNA (4977) deletion frequency, and AKR7A2 protein content are the most important variables in determining DNR reductase activity [11,12].

Introduction of chemotherapeutic drug targeting techniques potentially increased the tumor selectivity of drugs and decreased their cardiotoxicity. Increased expression of gonadotropin-releasing hormone (GnRH) receptors on the surface of tumor cells has been reported. Sixteen different GnRH-conjugates containing doxorubicin, DNR and methotrexate were developed and investigated. Their cytotoxicity was determined on primary human cardiac myocytes (HCM) and human umbilical vein endothelial cells (HUVEC). As a result, anticancer drug-GnRH-based conjugates with no cytotoxic effect on cardiomyocytes were found. In the future, these compounds could provide a better targeted antitumor therapy without any cardiotoxic adverse effects [13]. Myocardial cells were used to investigate the dose and time-dependent cellular enzyme release induced by either Adriamycin or DNR primary cultures in rat. Myocardial cells with exposure to DNR or Adriamycin (24 hour) at concentration of 1.8 or 18

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μM produced significant release of creatine, phosphokinase and lactic dehydrogenase without a detectable decrease in cell viability. With pre-incubation of the myocardial cells with varying concentrations of adenosine (10 μM to 1 mM) for 24 hr before the addition of anthracycline decreased or prevented drug-induced enzyme release, thus presenting adenosine as an effective myocardial protectant. It had no significant effect on cellular uptake of DNR, nor did adenosine adversely affect the oncotic activity of DNR against L1210 leukemia cells. On the contrary, myocardial protectants including N-acetyl-L-cysteine, alpha-tocopherol and carnitine were found to be ineffective in preventing anthracycline-induced enzyme release [14].

Studies have been found which were conducted to compare the efficacy and toxicity of DNR with idarubicin. In one of the studies, 1809 participants met the eligibility criteria and were included in the meta-analysis. The patients treated with idarubicin presented a significantly greater complete response rate after the first course of induction therapy compared with those treated with high dose of DNR. Moreover, a significantly lower rate of refractory acute AML was observed in patients receiving idarubicin as compared to DNR [15]. The interaction between DNR and cardiac myosin can be one of the modes of cardiotoxicity. To investigate this fact, forty groups of mice were used in which mice were treated with DNR orally, and three DNR-treated groups in which mice were injected intraperitoneally with DNR at seven bolus doses of 2.0, 4.0, and 6.0 mg/kg body weight, respectively. Results revealed that weakly acidic environment (pH 4.0-6.0) or higher temperature (30-37 °C) promoted the interaction between DNR and cardiac myosin, causing variations in conformation and normal physiological functions of cardiac myosin. Thermodynamic studies showed that the binding of DNR to cardiac myosin was a spontaneous process driven by entropy. It also indicated that hydrophobic interaction and hydrogen bonds may play essential roles in the combination of DNR with cardiac myosin. In addition, 4.0-6.0 mg/kg DNR-treated mice exhibited obvious increase in myocardial enzyme level, and reductions in blood cell count [16]. Oxidative stress and free radical formation are also involved in DNR cardiotoxicity. Pomegranate may play a significant role in scavenging free radical activity. To find the effect of pomegranate on DNR induced cardiotoxicity, 21 male Sprague rats were divided into three groups out of which Group A received distilled water, Group B was treated with DNR (20 mg/kg via intraperitoneal injection daily for 12 days for total cumulative dose of 240 mg/kg) and Group C was the pre-treatment group with pomegranate (25 mg/kg for 6 days orally, then DNR 20 mg/kg administrated concomitantly for the next 6 days with a cumulative dose of 120 mg/kg). Cardiac troponin I ([cTn I] pg/ml), malondialdehyde (MDA) (ng/ml), interleukin 17 (IL-17 pg/ml), and cardiac lactate dehydrogenase (LDH) (pm/ml), were used as biomarkers to measure the severity of cardiotoxicity. In group B, DNR induced lipid peroxidation and pro-inflammatory changes. In Group C, pomegranate pre-treatment demonstrated a significant cardio-protection in DNR-induced cardiotoxicity through reduction in oxidative stress, lipid peroxidation, pro-inflammatory, and cardiac injury biomarkers [17].

Significant age-related changes were noticed in DNR and daunorubicin kinetics in the rats that may alter susceptibility to acute systemic toxicity and chronic cardiotoxicity. Systemic clearance of DNR was decreased in older rats in comparison to younger rats. Moreover, concentrations of DNR in plasma and heart were found to be higher in older as compared to younger rats [18]. DNR-induced cardiotoxicity was mediated through the induction of cardiotoxic hydroxyeicosatetraenoic acids and/or the inhibition of cardioprotective epoxyeicosatrienoic acids (EETs). To investigate into it, Sprague-Dawley rats were treated with DNR (5 mg/kg i.p.) for 24 h, whereas human ventricular cardiomyocytes (RL-14) cells were exposed to DNR in the presence and absence of 4-[[trans-4-[(tricyclo[3.3.1.13,7]dec-1-ylamino)carbonyl]amino]cyclohexyl]oxy]-benzoic acid (tAUCB), a soluble epoxide hydrolase (sEH) inhibitor. This study provided the evidence that DNR induces cardiotoxicity through a sEH-mediated EETs degradation-dependent mechanism [19]. Cardiotoxicity induced by DNR could be protected by sodium ferulate (SF). Forty juvenile Sprague Dawley (SD) rats were divided into four groups. 1st group was taken as control whereas 2nd, 3rd and 4th groups were treated with DNR, DNR+SF, SF respectively. Juvenile rats were intraperitoneally treated with DNR (2.5 mg/kg every week for a cumulative dose of 10 mg/kg) preparation immature myocardial injury model in presence with SF (60 mg/kg) oral treatment for 25 days. It was found that SF could inhibit the decrease in heart rate induced by DNR (p < 0.05). SF treatment also resulted in an increase in the left ventricular end diastolic pressure, heart rate, the maximal left ventricular systolic speed and the maximal left ventricular diastolic speed responding to isoproterenol stimulation (p < 0.01); SF improved the myocardial ultrastructural injuries and inhibited the decline in cardiac Troponin I expression caused by DNR (p < 0.05) [12].

Conclusion

Daunorubicin (DNR) has been reported with a significant role in treating leukemia but its use is restricted due to associated cardiotoxicity. DNR has been reported to induce sub-chronic cardiomyopathy as well as down regulation of cytokines and stem cell markers. DNR may affect the heart by forming hydrophobic interactions and hydrogen bonds with cardiac myosin. Other cardiotoxic effects may include inhibition of cardio-protective epoxyeicosatrienoic acid. In the field of amelioration, adenosine, sodium ferulate and pomegranate have been reported as capable cardio-protectants against DNR induced cardiotoxicity. Future research on amelioration of DNR associated toxicity is highly recommended so as to minimize the associated risk.

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