ELSEVIER

Contents lists available at ScienceDirect

Critical Reviews in Oncology / Hematology

journal homepage: www.elsevier.com/locate/critrevonc



Modern radiotherapy in cancer treatment during pregnancy



Rosario Mazzola^{a,*}, Stefanie Corradini^b, Markus Eidemüeller^c, Vanessa Figlia^a, Alba Fiorentino^d, Niccolò Giaj-Levra^a, Luca Nicosia^a, Francesco Ricchetti^a, Michele Rigo^a, Mariella Musola^e, Marcello Ceccaroni^e, Stefania Gori^f, Stefano Maria Magrini^g, Filippo Alongi^{a,h}

- a Radiation Oncology, IRCCS Sacro Cuore Don Calabria Hospital, Negrar-Verona, Italy
- ^b Radiation Oncology, University Hospital, LMU Munich, Munich, Germany
- ^c Institute of Radiation Protection, Helmholtz Zentrum München, Neuherberg, Germany
- ^d Radiation Oncology, General Regional Hospital "F. Miulli", Acquaviva delle Fonti-Bari, Italy
- ^e Department of Obstetrics and Gynecology, IRCCS Sacro Cuore Don Calabria Hospital, Negrar-Verona, Italy
- f Medical Oncology, IRCCS Sacro Cuore Don Calabria Hospital, Negrar-Verona, Italy
- g Radiation Oncology, University and Spedali Civili Hospital, Brescia, Italy
- h University of Brescia, Italy

ARTICLE INFO

ABSTRACT

Keywords: Radiotherapy Pregnancy Review Cancer Breast cancer, gynecological malignancies and lymphomas are the most frequently diagnosed tumors in pregnant women. The feasibility of radiotherapy during pregnancy remains a subject of debate and clinicians continue to hesitate on this approach, trying to avoid radiotherapy in most cases. Since the 1990s, several technological advances, including intensity modulated and image guided radiation delivery, have been implemented in radiation oncology to improve the radiation treatment in terms of effectiveness and tolerability. It remains uncertain which short- and long-term health effects the radiation exposure of the fetus may have through advanced radiotherapy techniques. The present systematic literature review aims to summarize the limited current evidences of the feasibility and clinical results of "modern" radiotherapy procedures for the treatment of the most frequently diagnosed tumors in pregnant women.

1. Introduction

The incidence of cancer during pregnancy is a rare event and affects 0.07-0.1% of all pregnancies (Donegan, 1983). Breast cancer followed by gynecological malignancies and lymphomas are the most frequently diagnosed tumors in pregnant women (Donegan, 1983). For each of these oncologic diseases, radiation therapy (RT) is a cornerstone in the multidisciplinary treatment strategy and has a positive impact on longterm survival of non-pregnant women (Early Breast et al., 2011; Fiorentino et al., 2015; Hay et al., 2013; Mazzola et al., 2016; Perez et al., 1998; Corradini et al., 2015; Mazzola et al., 2017). While the treatment of cancer by RT in pregnant patients is aimed at improving the survival of the mother, special considerations are necessary to maximally reduce possible health impairments of the fetus. Since the 1990s, technological and technical improvements in modern RT, such as 3D-conformal RT (3DCRT), intensity modulated RT (IMRT) or volumetric modulated arc therapy (VMAT), have been introduced into clinical practice. These advanced radiation techniques pursue the goal of delivering high doses to the tumor, while sparing the surrounding

tissues or organs at risk in order to improve RT in terms of effectiveness and tolerability (Nutting et al., 2011; Mazzola et al., 2015; GiajLevra et al., 2016; Giaj-Levra et al., 2016). Furthermore, image-guided RT techniques using on-board cone-beam computed tomography (CBCT) have been developed to ensure a precise dose delivery (Dawson and Sharpe, 2006). Modulated-RT does not necessarily result in exposure to a higher dose, because these RT-approaches are designed to limit the high dose to a more restricted volume. On the other hand, the disadvantage of modulated therapies remains the exposure of a larger volume to low doses. Starting from this background, the adoption of advanced RT techniques in pregnant women affected by cancer could increase the probability of short and long-term adverse events for the fetus and, thus, during pregnancy, modulated RT has been cautiously used only in strictly selected cases.

The present systematic literature review aims to summarize the current clinical evidences of the feasibility and clinical results of "modern" radiotherapy procedures for the treatment of the most frequently diagnosed tumors in pregnant women.

E-mail address: rosariomazzola@hotmail.it (R. Mazzola).

^{*} Corresponding author.

2. Fetal exposure and health effects from ionizing radiation during pregnancy

2.1. Fetal adverse events

Gestational age (weeks of amenorrhea) does not correspond to embryonal age (weeks or days from fertilization). Adverse events following fetal irradiation vary according to post-implantation week and dose of irradiation. Specifically, during the organogenesis phase (i.e. weeks 2–7) the main effect is the occurrence of gross malformation and small head size (SHS) without mental retardation. Increased risk of growth retardation and SHS was reported for doses superior to 0.5 Gy (Stovall et al., 1995a; Nakagawa et al., 1997). Normally, the brain develops between weeks 8 and 15 (first trimester of gestation), thus, in this phase, the main potential effects could be SHS and mental retardation. Mental functioning seems not to be impaired for doses below 0.1 Gy, while doses higher than 0.3 Gy might affect its functioning. The incidence of mental retardation for doses between 0.1 and 0.49 Gy is estimated for 6% of the cases (Yonekura et al., 2014).

The effects of irradiation in the second trimester (weeks 16–25) are similar to those of the previous trimester. In particular, the main risks include mental and growth retardation, SHS, cataracts, sterility and secondary malignancies. The incidence of mental retardation is 2% for doses below 0.5 Gy (Otake and Schull, 1984). The risk of sterility and neurological diseases is smaller than for irradiations during the previous trimester (Brent, 1983). Finally, for exposures during the third trimester (weeks > 25) the risk of mental and growth retardation and SHS seems low. Nevertheless, evidence for these adverse events were reported for exposures < 0.5 Gy (Miller and Mulvihill, 1976; Stovall et al., 1995b).

Table 1 summarizes the main adverse eventsfollowing fetal irradiation according to the post-conception time.

2.2. Childhood and adult cancer risk after in-utero exposure

No new phantom models studies exists for radiation in pregnancy using new RT-techniques. For this reason, we will focus this section on medical exposures from diagnostic x-rays rather than therapeutic radiation exposures.

Deterministic effects describe a cause and effect relationship between radiation and certain side-effects. They are also called non-stochastic effects and have a threshold, below which the effect does not occur. The stochastic effects may occur by chance without any threshold level of dose, whose probability is proportional to the dose and whose severity is independent of the dose. In the context of radiation protection, the main stochastic effect is cancer induction (Stovall et al., 1995b).

A relationship between childhood malignancy and pre-natal diagnostic X-ray irradiation of the child was first reported by Stewart (Stewart et al., 1956, 1958), in the large Oxford survey of childhood

cancers (OSCC). The dependence of the risk on the number of films exposed was highly significant and adequately described by a linear relationship (Bithell et al., 2018). Wakeford and Little (2003) summarized the efforts to derive the fetal doses and obtained an excess relative risk of around 50 Gy⁻¹ for childhood cancer below the age of 15 years, leading to an excess absolute risk coefficient for incident cases of about 8% Gy⁻¹. They cautioned that the uncertainties related to these estimates were appreciable, and there were reasons to believe that this coefficient could be a systematic overestimate. The great majority of the intrauterine exposures occurred during the third trimester of pregnancy for obstetric reasons. Nevertheless, the relative risk of childhood cancer associated with exposure during the first trimester was found to be a statistically significant ~2.5 times greater than that for the third trimester, implying an increased sensitivity to radiation-induced childhood cancer early in pregnancy. While the OSCC is the largest and most comprehensive study for medical exposures from diagnostic X-rays, many other case-control studies are consistent with an increased risk of childhood cancer and leukemia (Wakeford, 2008; UNSCEAR, 1994).

However, in the atomic bomb survivors of Hiroshima and Nagasaki, by far the largest cohort study of intrauterine exposures, only 2 cases of childhood cancer in the first 14 years of life were observed among 1630 children exposed in utero, and no case of leukemia (Yoshimoto et al., 1988). Since background cancer rates are very small, single cases can have a large influence on risk estimates. (Boice and Miller (1999)) reviewed the arguments for a causal association of intrauterine radiation exposure and subsequent cancer risk. A comparison of the risk estimates between the OSCC and the atomic bomb survivors concluded that the risk estimates of both studies are compatible when taking the uncertainties into account (Wakeford and Little, 2003). In summary, the evidence of increased childhood leukemia and solid cancer risk after intrauterine exposure is convincing, however, the uncertainties about the magnitude of risk are significant.

The same atomic bomb survivors constitute the most significant source of information on adult cancer risk after intrauterine exposure. Delong champ and colleagues (Delongchamp et al., 1997) analyzed solid cancer and leukemia mortality over the age range 17-46 years. Among the 807 in utero survivors with doses over 10 mSv, eight deaths from solid cancers and twocases from leukemia were recorded. The risk for solid cancer was statistically significant with an excess relative risk (ERR) of $2.4\,\mathrm{Sv}^{-1}$ (95%CI: 0.3; 6.7). The magnitude of this excess did not substantially differ from that of those exposed during the first 6 years of life with ERR = $1.4 \, \text{Sv}^{-1}$ (95%CI: 0.4; 3.1). Subsequently, Preston and collaborators analyzed the incidence of solid cancer in the atomic bomb survivors with seven additional years of follow-up for persons with a range of 12-55 years (Preston et al., 2008). Ninety-four cancers were observed among 2452 survivors in utero at the time of bombing. The statistically significant risk was ERR = $1.3 \,\mathrm{Sv}^{-1}$ (95%CI: 0.2; 2.8). Risk for exposure during childhood was somewhat higher with ERR = $2.0 \,\mathrm{Sv}^{-1}$ (95%CI: 1.4; 2.8), however, the difference

Table 1Effects of fetal irradiation according to the postconceptional time.

Post-conceptional week(s)	Embrional period 1	Organogenesis 2–7	First trimester 8–15	Second trimester 16–25	Third trimester > 25
Effect					
Death	+ + +	+	+	_	_
Malformation	_	+++	+	+	_
Growth retardation	-	+++	+ +	+	+
Mental retardation	-	+	+ + +	+	_
Small head size	_	+ +	+++	+	+
Sterility	_	_	+	+	+
Cataracts	_	_	_	+	+
Neurological disease	_	+++	_	+	+
Malignant disease	-	+	+	+	+

⁻ no effect; + demonstrated effect with small incidence; + + demonstrated effect with average incidence; + + + high incidence.

between these estimates was not statistically significant. The analysis suggested a decrease of relative risk with increasing age, both for individuals exposed in utero and during childhood, and the decrease was more marked for those exposed in utero than as children. No variation in the ERR by trimester of exposure was observed for those exposed in utero, and the risk estimates were identical.

In summary, adult cancer risk was found to increase statistically significant with radiation exposure of the uterus. Risk estimates were similar for intrauterine exposures and for exposures during early childhood. There was no clear evidence for a difference of risk on the trimester of exposure. Compared to childhood cancer risk after intrauterine exposures, the lower relative risk estimates and the decrease of risk with age indicate that adult lifetime cancer risk is likely to be considerably less than projections based on relative risks derived from childhood cancer studies.

3. Clinical Results: fetal exposure and health effects from radiotherapy techniques

3.1. Search strategy and selection criteria for clinical studies

Wesearched PubMed, EMBASEand Cochrane library for articles published in English language between 1 January 1990 and 1 January 2018. Search terms included ("Pregnancy" [MeSH Terms] OR "Pregnancy" [All Fields]) AND ("radiotherapy" [MeSHTerms] OR "radiotherapy" [All Fields] OR ("cancer" [All Fields] AND "radiotherapy" [All Fields]).

We identified additional references by doing a manual search of the references of all included articles. Two independent reviewers (VF and MR) identified potential studies and exported them to an electronic reference management software program (Ref Works, version 2.0). VF and MR determined eligibility by first reviewing the title and abstract and then the full article. Disagreements were solved by consensus; if consensus was not achieved, then a third author (RM) provided an assessment of eligibility.

A study was included if it reported on cancer-related RT for breast cancer, gynecological malignancies and lymphomas in pregnant patients. All studies were analyzed for study design, number of patients, age (mean and range), type of RT, radiation dose, gestational phase and outcomes in terms of adverse events for the fetus. Exclusion criteria were: articles with no detailed information regarding clinical outcomes, review articles, editorials, articles not written in English language. Fig. 1 depicts the study selection approach.

3.2. Breast Cancer

The literature regarding modern breast RT during pregnancy is limited and most experiences come from retrospective case series (see Table 2). The AAPM Task group-36 recommends that a threshold of

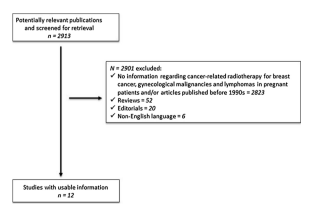


Fig. 1. Search strategy flowchart for the inclusion and exclusion of studies.

Author; Year of publication	Number of patients	Surgery Y/N	Chemotherapy Y/N	Gestational Phase	Gestational Phase RT Technique/ fetal dose	Late toxicity (i.e. mental retardation, growth retardation, second tumors, malformations)
Galimberti et al. (2009)	15 in-vivo measurements in non- pregnant patients	NA	NA	NA	ELIOT, 5-9 MeV 21 Gy in 1Fx /dose in non-pregnant uterus 1.7 mGy (range 0.6-3.2 mGy) with lead shielding	NA
Basta et al. (2015)		¥	Z	17-23 week	Electron beams, 5-8 MeV 54 Gy in 27Fx /fetal dose 53 mGy/15 mGy with lead shielding	Not reported
Antypas et al. (1998), Kouvaris et al. (2000)	. 1	¥	Z	2-6 week	3D-CRT, 6 MV photons 46 Gy in 20Fx /fetal dose 39 mGy	No toxicity at 36 months
Leonardi et al. (2017)	1	¥	Z	15 week	ELIOT, 6 MeV 21 Gy in 1Fx /dose in non-pregnant uterus 0.84 mGy with lead shielding	No toxicity at 60 months
Ngu et al. (1992)	1	Y following delivery	Y following delivery	15-20 week	3D-CRT, 6 MV photons 50 Gy in 25Fx /fetal dose 210 mGy /140-180 mGy with lead shielding	No toxicity at 3 months
Van Calsteren et al. (2010)	2	ND	ND	18-23 week 16-21 week	Thoracic wall 46 Gy Breast 50 Gy	No toxicity at birth
Van der Giessen (1997)	1	QN	ND	24-29 week	3D-CRT, 10 MV photons 50 Gy in 25Fx /fetal dose 160 mGy with lead shielding	No toxicity at 3 months

Patients treated for metastasis or with a palliative dose were excluded. NA: not applicable; ND: no data.

50cGy should always be maintained for the fetal dose of RT during pregnancy (Stovall et al., 1995a). Special external abdominal leaf shields have been evaluated to further reduce the fetal dose up to 50–58% (Stovall et al., 1995a; Luis et al., 2009; Owrangi et al., 2016). Taken together, whole breast irradiation is considered as a relatively safe treatment option during the first two trimesters, as the distance between the radiation fields and the fetus is sufficient and fetal doses do not exceed threshold doses associated with organ malformations.

Literature is scarce, but successful treatments with good maternal and fetal outcome using modern RT techniques have been reported during pregnancy. Specifically, Antypaset al. (Antypas et al., 1998) treated a patient in the first trimester of pregnancy using 6 MV photon beam 3D-conformal RT to a total dose of 46 Gv in 20 fractions. The fetal dose was estimated to 39 mGy using in-vivo and phantom measurements. A report from New South Wales (Australia) described a RT treatment in the second trimester gestation (Ngu et al., 1992). The patient underwent neoadjuvant RT of the breast and lymphatic pathways with 50 Gy in 25 fractions with 6 MV photons. With shielding using lead covering and blocks, the fetal dose measured with thermoluminescent dosimeters was 140-180 mGy. Similarly, van der Giessen (Van Der Giessen, 1997) reported on a case treated during 24-29 weeks of gestation with a fetal dose of 160 mGy during breast RT with 50 Gy and proper lead shielding. In contrast, Antolaket al. (Antolak and Strom, 1998) reported a significantly lower fetal dose of 15 mGy with abdominal shielding. Of note, this patient suffered from recurrent breast cancer in the area of the mastectomy scar and received chest wall irradiation using electron beams (5-8 MeV). In the analysis of cancer during pregnancy by Van Calsteren et al. (2010), two breast cancer patients treated with RT in the second trimester are included. They were treated with 46 Gy and 50 Gy to the chest wall and breast, respectively. Unfortunately, no fetal doses are reported. In terms of fetal outcome of these examples, all pregnancies were carried to term. There was no elective termination of pregnancy or in utero death, and all children were in good health at birth without congenital abnormalities. However, long-term follow-up is missing in most cases. Two children had no evidence of neuro-developmental impairments at 36 and 60 months follow-up, respectively (Leonardi et al., 2017; Kouvaris et al., 2000).

To minimize risk-exposure, a possibleRT strategy is a localized dose escalation using an intraoperative boost to the tumor bed (Sedlmayer et al., 2017). The European Institute of Oncology, in Italy, evaluated the feasibility of electron beam intraoperative RT (ELIOT) as an anticipated boost using in-vivo dosimetry in fifteen non-pregnant patients (Galimberti et al., 2009). All cases received 21 Gy intraoperative RT with electrons using a mobile linear accelerator. Dosimetric measurements showed a mean dose in utero of 1.7 mGy (range 0.6–3.2 mGy) with shielding. In 2011 the first pregnant patient was treated at 15th week of gestation. The estimated dose to the fetus was 0.84 mGy. Therefore, ELIOT can be considered as a treatment option for anticipated boost therapy during the first or second trimester and postpone whole breast RT after the childbirth. One throwback is the longer time

interval between boost delivery and whole breast irradiation, which could increase the risk of local recurrence. Nevertheless, in cases were chemotherapy is administered, this time interval stays within regular time lines (Corradini et al., 2014).

In summary, from the experiences here reported, using the previously described specific precautions, 3DCRT or IORT approaches could be considered in selected cases for breast cancer irradiation during pregnancy. Due to the lacking of clinical data, to date, intensity modulated RT and other modern techniques seem to be not recommended.

3.3. Gynecological tumors

Pelvic irradiation during pregnancy is a major challenge because, regardless of the utilized technique, fetal exposuredose is always significant and leads to serious adverse effects for the fetus. With regard to the tumor site, cervical cancer is the most common gynecological cancer during pregnancy, while vulvar, endometrial and ovarian cancer are extremely rare and usually managed with upfront surgery. In contrast, invasive cervical cancer can also be treated with definitive RT or RT-chemotherapy in non-pregnant patients.

Concerning pregnant-patients affected by gynecological cancers, when pelvic RT is performed with thefetus in utero, spontaneous abortion will invariably occur, usually within 3–6 weeks (Gustaffson and Kottmeier, 1962; Prem et al., 1966). If cervical cancer is diagnosed after the 20th week of gestation, a treatment delay can be considered in the interest of the fetuswithout significantly affecting the prognosis (Hunter et al., 2008).

According to the inclusion criteria of the present review, a single study was found. Specifically, Sood et al. (1997) performed a case-control analysis of 26 women with cervical carcinoma who were diagnosed during pregnancy and treated primarily with RT. Patients were treated with external beam 3D-CRT (mean dose, 46.7 Gy) followed by brachytherapy. Two patients with Stage IA2 disease were treated in the third trimester; in these last cases, infants had an uncomplicated neonatal course. On the other hand, seven patients underwent radical hysterectomy due to positive pelvic nodes; thus, abortion of the fetuswas performed. Finally, three patients diagnosed during the first trimester were treated with radiation with the fetus in situ, and all had spontaneous abortions 20–24 days after the start of radiation (after a mean dose of 34 Gy). Astatistical analysis revealed no differencesin terms of overall survival between the pregnant group and the control arm.

In summary, from the few experiences here reported, RT in pregnant patients affected by gynecological malignancies remains contraindicated due to the high risk of abortion and fetal damage.

3.4. Hodgkin and Non-Hodgkin lymphoma

A limited number of experiences have evaluated the role of RT in lymphomas in pregnant women, as described in Table 3. Woo et al.

Table 3

Available articles regarding Hodgkin and Non-Hodgkin Lymphoma irradiation during pregnancy.

Author; Year of publication [Ref]	Number of patients	Chemotherapy Y/N	Gestational Phase	RT Technique/ fetal dose	Late toxicity (i.e. mental retardation, growth retardation, second tumors, malformations)	Abortion Y/N
Woo et al. (1992)	16	Y	II and III	3D-CRT 14 to 55 mGy/ Cobalt 100 to 136 mGy	None	N
Lishner et al. (1992)	22	Y	I and II	N.A.	1	Y
Evens et al. (2013)	90	Y	II and III	N.A.	None	N
Nisce et al. (1986)	7	N	II and III	N.A./20-500mGy	None	N
Peccatori et al. (2009)	1	N	III	3D-CRT 36 Gy/18 fractions to the 90% isodose	None	N

(Lishner et al., 1992) reported on 16 women diagnosed with early stage sclero-nodular HL who received a supra-diaphragmatic radiation treatment of the mediastinal, axillar, cervical lymph node levels or a mantle-field irradiation with a RT dose between 35 and 40 Gy. Four to five half-value layers of lead were used to shield the uterus during RT and the dose to the fetus was estimated in the range from 14 to 55 mGy, for treatments with 6 MV photons. All pregnancies were carried to termand all children were physically and mentally normal.

Similar results have been published in an investigation focusing on fifty pregnant women diagnosed with HL. Of the fifty pregnancies, there were forty deliveries (two of which were stillbirths), five miscarriages and four therapeutic abortions. Clinical data were collected from 22 cases of 38 live-births. Of the 22 babies who were exposed to multimodal therapy in utero, one was exposed to chemotherapy alone, two babies were exposed to a combination of chemotherapy and RT during and after the first trimester six cases received RT during the first trimester of pregnancy. Finally, seven cases received the treatment after the first trimester. The authors found no differences between the babies born to women with HL when compared to the mother-risk matched controls in terms of birth weight, mean gestational age or method of delivery. In a single case, a malformation of hydrocephaly was observed in a patient who was diagnosed with HL before conception and treated with chemotherapy alone in the first trimester (Lishner et al., 1992).

More recently, Evens et al. (2013) investigated the effectsof chemotherapy, RT or a combination of both in HL and NHL in a series of ninety pregnant women. In this population of study, RT was administrated in four cases with a diagnosis of stage I and IIA with a dose prescription of 25–30 Gy. No spontaneous abortions, neonatal intensive care unit admission or malformations were reported.

In summary, from the experiences here reported, RT in pregnant patients affected by HL and NHL seems to be feasible even if no specific use of modern RT-techniques has been explored.

4. Discussion

Oncologic treatment of pregnant women is always an interdisciplinary challenge and should be managed by an expert team of radiation-oncologists, gynecologists, neonatologists, medical oncologists, psychologists and others professional figures with the aim to find an individual treatment plan in consensus with the patient/couple. It is well known, that pregnant patients should be treated similarly to nonpregnant cancer patients and that a comparable survival can be achieved (Amant et al., 2013). Treatments in hospitals with obstetric high-care units are strongly advocated (de Haan et al., 2018). It is recognized that an elective pregnancy interruption has not always a beneficial effect on survival (King et al., 1985). In breast cancer, for example, an international consensus recommendations support a gestation stage-based treatment approach, where the treatment can be postponed until post-partum in near-term patients at more than 37 weeks of gestation (Amant et al., 2010).

The main aim of the current literature review was to assess the role of RT in the modern era concerning the management of the most frequently diagnosed tumors in pregnant women (Donegan, 1983). Clinical data are very limited and the main experiences since the 1990s derive from retrospective series. Thus, several concerns remain unsolved regarding the potential use of RT in the modern era for cancer treatment during pregnancy. In terms of dose reduction to healthy tissue (and to fetus) advanced techniques seem to be rather contraproductive because of low dose bath by IMRT or RT-rotational approaches. Additionally, image-guided RT does not help to reduce dose to the pregnancy in utero. On board CBCT itself may increase radiation burden. In addition, significant dose contribution for fetus in utero could arise from scattering, for which no valid model exists.

Thus, several clinical considerations need to be evaluated. More deeply, concerning the management of breast cancer in pregnancy, modified radical mastectomy or breast conserving surgery with removal

of axillary lymph nodes or sentinel node biopsy is safely applicable during all trimesters of pregnancy (Gentilini et al., 2010). The scenario seems to be more complicated if breast cancer is diagnosed during the first trimester. Besides the option of an induced abortion, surgery and RT are the only treatment options, as chemotherapy is contraindicated. One rationale for mastectomy is to avoid postoperative RT after breast conserving surgery. In cases of breast-conserving surgery, it remains an open question if RT should be administered during pregnancy or should be delayed to after delivery.

In this last clinical scenario, the use of hypo-fractionated breast RT during pregnancy could be a potential option to reduce the overall treatment time. Unfortunately, no clinical data are available in literature

Looking at the limited current evidences of the feasibility and clinical results of "modern" radiotherapy, the need of RT during pregnancy might be limited to the few patients that will not receive adjuvant chemotherapy. For all the others, chemotherapy during pregnancy allows a safe postponement of RT after delivery.

Some studies focused on the estimation of fetal dose exposure during breast RT using computed dose estimations or phantom measurements. Fetal dose during breast irradiation using a tangential field technique is mainly influenced by patient scatter and out-of-field doses of direct radiation leakage and collimator scattering of the head of the linear accelerator (Diallo et al., 1996; Van der Giessen, 1997).

The use of physical wedges in tangential breast irradiation can significantly increase scattered radiation and should therefore be avoided. Physical wedges are commonly used to improve dose uniformity to the target volume (Stovall et al., 1995b).

Similarly, the use of intensity-modulated techniques showed a five-fold increase of fetal dose as compared to 3D-CRT, because higher monitor units (MUs) are generally used. MUs represent a measure of the dose delivered by a single beam of ionizing radiations (Öğretici et al., 2016). Finally, during breast irradiation, the most important factors determining the fetal dose are the field size and distance from the radiation field. The absorbed dose by the fetus increases with gestational stage, as the distance from the radiation field edge to the uterine fundus is narrowing by around a centimeter per week. Thus, it is of great interest to determine the expected change during the course of RT to give an estimate of the fetal dose (Mazonakis et al., 2003). The height of the fundus scan be estimated with the help of weekly ultrasound measurement or clinical examinations (Luis et al., 2009).

In contrast, it is a major challenge to irradiate the pelvic region for gynecological tumors in pregnant women, as preservation of fetal life is not really possible. In this last clinical scenario, it has to be decidedwhether the treatment should be postponed until after delivery or whether elective termination of pregnancy should be chosen.

Finally, Hodgkin and Non-Hodgkin Lymphomas are very common haematological diseases in young women. Over the last decades, chemotherapy cycles have been shortened and RT volumes decreased, in order to reduce the risk of late side effects. This last strategy allowed decreasing the possible fetal exposure to ionizing radiation. In fact, the reported risk of mental retardation, growth retardation, secondary tumors and organ malformations in clinical practice seems to be very low. Obviously, the irradiation of supra-diaphragmatic disease represents the best clinical situation with regard to a possible exposure of the fetus to ionizing radiation.

Possible statements, within the present literature review, could derive from limited series (i.e. 22 patients diagnosed with breast cancer in pregnancy and 118 with lymphoma); therefore, there is a quite small fundament for any clinical recommendation.

Apart from the crucial role of high-expertise centers in the management of pregnant patients, many efforts are needed to ensure safe RT options. New technologies such as heavy particle therapy could also become a treatment option for pregnant women affected by cancer. Nevertheless, the uncertainties regarding the production of neutrons by heavy ions and the subsequent fetus exposure remain unclear (Brenner

and Hall, 2008). The available clinical experiences are scarse (Münter et al., 2010). Undoubtedly, future investigations or prospective clinical trials will have to encompass important issues.

In summary, even if technological advances have been introduced into clinical practice, the role of RT during pregnancy in modern era needs specific precautions and it could be still considered only in selected cases of breast cancer and lymphomas. Due to the lacking of clinical data, the potential role of intensity modulated RT and other modern techniques remain uncertain.

In the absence of valuable data, a case-by-case assessment is needed to address concerns about potential risks for the fetus and the impact on the oncological outcome for the patient.

Conflict of interest statement

All the Authors declare no Conflict of Interest.

References

- Amant, F., Deckers, S., Van Calsteren, K., et al., 2010. Breast cancer in pregnancy: recommendations of an international consensus meeting. Eur. J. Cancer 46 (18), 3158-3168
- Amant, F., von Minckwitz, G., Han, S.N., et al., 2013. Prognosis of women with primary breast cancer diagnosed during pregnancy: results from an international collaborative study. J. Clin. Oncol. 31 (20), 2532–2539.
- Antolak, J.A., Strom, E.A., 1998. Fetal dose estimates for electron-beam treatment to the chest wall of a pregnant patient. Med. Phys. 25 (12), 2388–2391.
- Antypas, C., Sandilos, P., Kouvaris, J., et al., 1998. Fetal dose evaluation during breast cancer radiotherapy. Int. J. Radiat. Oncol. Biol. Phys. 1;40 (4), 995–999.
- Basta, P., Bak, A., Roszkowski, K., 2015. Cancer treatment in pregnant women. Contemp. Oncol. (Pozn.) 19 (5), 354–360.
- Bithell, J.F., Draper, G.J., Sorahan, T., et al., 2018. Childhood cancer research in Oxford I: the oxford survey of childhood cancers. Br. J. Cancer 119 (6), 756–762.
- Boice Jr, J.D., Miller, R.W., 1999. Childhood and adult cancer after intrauterine exposure to ionizing radiation. Teratology 59 (4), 227e33.
- Brenner, D.J., Hall, E.J., 2008. Secondary neutrons in clinical proton radiotherapy: a charged issue. Radiother. Oncol. 86, 165–170.
- Brent, R.L., 1983. The effects of embryonic and fetal exposure to X-ray, microwaves, and Ultrasound. Clin. Obstet. Gynecol. 26, 484–510.
- Corradini, S., Niemoeller, O.M., Niyazi, M., et al., 2014. Timing of radiotherapy following breast-conserving surgery: outcome of 1393 patients at a single institution. Strahlenther. Onkol. 190 (4), 352–357.
- Corradini, S., Niyazi, M., Niemoeller, O.M., et al., 2015. Adjuvant radiotherapy after breast conserving surgery – a comparative effectiveness research study. Radiother. Oncol. 114 (1), 28–34.
- Dawson, L.A., Sharpe, M.B., 2006. Image-guided radiotherapy: rationale, benefits, and limitations. Lancet Oncol. 7 (10), 848–858.
- de Haan, J., Verheecke, M., Van Calsteren, K., et al., 2018. Oncological management and obstetric and neonatal outcomes for women diagnosed with cancer during pregnancy: a 20-year international cohort study of 1170 patients. Lancet Oncol. 19 (3), 337–346.
- Delongchamp, R.R., Mabuchi, K., Yoshimoto, Y., et al., 1997. Cancer mortality among atomic bomb survivors exposed in utero or as young children, October 1950-May 1992. Radiat. Res. 147 (3), 385–395.
- Diallo, I., Lamon, A., Shamsaldin, A., et al., 1996. Estimation of the radiation dose delivered to any point outside the target volume per patient treated with external beam radiotherapy. Radiother. Oncol. 38 (3), 269–271.
- Donegan, W.L., 1983. Cancer and pregnancy. CA Cancer J. Clin. 33, 194–214.
- Early Breast, Cancer, Trialists' Collaborative Group (EBCTCG), Darby, S., McGale, P., Correa, C., et al., 2011. Effect of radiotherapy after breast-conserving surgery on 10year recurrence and 15-year breast cancer death: meta-analysis of individual patient data for 10,801 women in 17 randomised trials. Lancet 378, 1707–1716.
- Evens, A.M., Advani, R., Press, O.W., et al., 2013. Lymphoma occurring during pregnancy: antenatal therapy, complications, and maternal survival in a multicenter analysis. J. Clin. Oncol. 31 (32), 4132–4139.
- Fiorentino, A., Mazzola, R., Ricchetti, F., et al., 2015. Personalized–not omitted–radiation oncology for breast cancer. J. Clin. Oncol. 33 (36), 4313–4314.
- Galimberti, V., Ciocca, M., Leonardi, M.C., et al., 2009. Is electron beam intraoperative radiotherapy (ELIOT) safe in pregnant women with early breast cancer? In vivo dosimetry to assess fetal dose. Ann. Surg. Oncol. 16 (1), 100–105.
- Gentilini, O., Cremonesi, M., Toesca, A., et al., 2010. Sentinel lymph node biopsy in pregnant patients with breast cancer. Eur. J. Nucl. Med. Mol. Imaging 37 (1), 78–83.
- GiajLevra, N., Sicignano, G., Fiorentino, A., et al., 2016. Whole brain radiotherapy with hippocampal avoidance and simultaneous integrated boost for brain metastases: a dosimetric volumetric-modulated arc therapy study. Radiol. Med. 121 (1), 60–69.
- Giaj-Levra, N., Sciascia, S., Fiorentino, A., et al., 2016. Radiotherapy in patients with connective tissue diseases. Lancet Oncol. 17 (3), e109–17.
- Gustaffson, D.C., Kottmeier, H.L., 1962. Carcinoma of the cervix associated with pregnancy. A study of the Radiumhemmet's series of invasive carcinoma during the period 1932-1956. Acta Obstet. Gynecol. Scand. 41, 1–21.
- Hay, A.E., Klimm, B., Chen, B.E., et al., 2013. An individual patient-data comparison of

- combined modality therapy and ABVD alone for patients with limited-stage Hodgkin lymphoma. Ann. Oncol. 24, 3065–3069.
- Hunter, M.I., Tewari, K., Monk, B.J., 2008. Cervical neoplasia in pregnancy. Part 2: current treatment of invasive disease. Am. J. Obstet. Gynecol. 199, 10–18.
- King, R.M., Welch, J.S., Jr, Martin, et al., 1985. Carcinoma of the breast associated with pregnancy. Surg. Gynecol. Obstet. 160 (3), 228–232.
- Kouvaris, J.R., Antypas, C.E., Sandilos, P.H., et al., 2000. Postoperative tailored radiotherapy for locally advanced breast carcinoma during pregnancy: a therapeutic dilemma. Am. J. Obstet. Gynecol. 183 (2), 498–499.
- Leonardi, M., Cecconi, A., Luraschi, R., et al., 2017. Electron beam intraoperative radiotherapy (ELIOT) in pregnant women with breast Cancer: from in vivo dosimetry to clinical practice. Breast Care (Basel) 12 (6), 396–400.
- Lishner, M., Zemlickis, D., Degendorfer, P., et al., 1992. Maternal and foetal outcome following Hodgkin's disease in pregnancy. Br. J. Cancer 65 (1), 114–117.
- Luis, S., Christie, D., Kaminski, A., et al., 2009. Pregnancy and radiotherapy: management options for minimising risk, case series and comprehensive literature review. J. Med. Imaging Radiat. Oncol. 53 (6), 559–568.
- Mazonakis, M., Varveris, H., Damilakis, J., et al., 2003. Radiation dose to conceptus resulting from tangential breast irradiation. Int. J. Radiat. Oncol. Biol. Phys. 55 (2), 386–391.
- Mazzola, R., Ferrera, G., Alongi, F., et al., 2015. Organ sparing and clinical outcome with step-and-shoot IMRT for head and neck cancer: a mono-institutional experience. Radiol. Med. 120 (8), 753–758.
- Mazzola, R., GiajLevra, N., Alongi, F., 2016. Radiation dose-response relationship for risk of coronary heart disease in survivors of hodgkin lymphoma. J. Clin. Oncol. 34 (24), 2940–2941.
- Mazzola, R., Ricchetti, F., Fiorentino, A., et al., 2017. Weekly cisplatin and volumetric-modulated arc therapy with simultaneous integrated boost for radical treatment of advanced cervical Cancer in elderly patients: feasibility and clinical preliminary results. Technol. Cancer Res. Treat. 16 (3), 310–315.
- Miller, R.W., Mulvihill, J.J., 1976. Small head size after atomic irradiation. Teratology 14, 355–358.
- Münter, M.W., Wengenroth, M., Fehrenbacher, G., et al., 2010. Heavy ion radiotherapy during pregnancy. Fertil. Steril. 94 (6), 2329 e5-7.
- Nakagawa, K., Aoki, Y., Kusama, T., 1997. Radiotherapy during pregnancy: effects on fetuses and neonates. Clin. Ther. 19, 770–777.
- Ngu, S.L., Duval, P., Collins, C., 1992. Foetal radiation dose in radiotherapy for breast cancer. Aust. Radiol. 36 (4), 321–322.
- Nisce, L.Z., Tome, M.A., He, S., Lee 3rd, B.J., Kutcher, G.J., 1986. Management of coexisting Hodgkin's disease and pregnancy. Am. J. Clin. Oncol. 9 (2), 146–151.
- Nutting, C.M., Morden, J.P., Harrington, K.J., et al., 2011. Parotid-sparing intensity modulated versus conventional radiotherapy in head and neck cancer (PARSPORT): a phase 3 multicentrerandomised controlled trial. Lancet Oncol. 12 (2), 127–136.
- Öğretici, A., Akbaş, U., Köksal, C., et al., 2016. Investigation of conformal and intensity modulated radiation therapy techniques to determine the absorbed fetal dose in pregnant patients with breast cancer. Med. Dosim. 41 (2), 95–99.
- Otake, M., Schull, W.J., 1984. In utero exposure to A-bomb radiation and mental retardation: a reassessment. Br. J. Radiol. 57, 409–414.
- Owrangi, A.M., Roberts, D.A., Covington, E.L., et al., 2016. Revisiting fetal dose during radiation therapy: evaluating treatment techniques and a custom shield. J. Appl. Clin. Med. Phys. 17 (5), 34–46.
- Peccatori, F.A., Azim, H.A., Jr, Pruneri, G., Piperno, G., Raviele, P.R., Preda, L., et al., 2009. Management of anaplastic large-cell lymphoma during pregnancy. J. Clin. Oncol. 27 (25), e75–7.
- Perez, C.A., Grigsby, P.W., Chao, K.S., et al., 1998. Tumor size, irradiation dose, and long-term outcome of carcinoma of uterine cervix. Int. J. Radiat. Oncol. Biol. Phys. 41 (2), 307–317.
- Prem, K.A., Makowski, E.L., McKelvey, J.L., 1966. Carcinoma of the cervix associated with pregnancy. Am. J. Obstet. Gynecol. 95 (1), 99–108.
- Preston, D.L., Cullings, H., Suyama, A., et al., 2008. Solid cancer incidence in atomic bomb survivors exposed in utero or as young children. J. Natl. Cancer Inst. 100 (6), 428–436.
- Sedlmayer, F., Reitsamer, R., Wenz, F., et al., 2017. Intraoperative radiotherapy (IORT) as boost in breast cancer. Radiat. Oncol. 12 (1), 23.
- Sood, A.K., Sorosky, J.I., Mayr, N., et al., 1997. Radiotherapeutic management of cervical carcinoma that complicates pregnancy. Cancer 80 (6), 1073–1078.
- Stewart, A., Webb, J., Giles, D., et al., 1956. Malignant disease in childhood and diagnostic radiation in utero. Lancet 1;271 (6940), 447.
- Stewart, A., Webb, J., Hewitt, D., 1958. A survey of childhood malignancies. Br. Med. J. 1 (5086), 1495–1508.
- Stovall, M., Blackwell, C.R., Cundiff, J., et al., 1995a. Fetal dose from radiotherapy with photon beams: report of AAPM Radiation Therapy Committee Task Group No. 36. Med. Phys. 22 (1), 63–82.
- Stovall, M., Blackwell, C.R., Cundiff, J., et al., 1995b. Fetal dose from radiotherapy with photon beams: report of AAPM Radiation Therapy Committee Task Group No. 36. Med. Phys. 22 (1), 63e82.
- UNSCEAR, 1994. United Nations Scientific Committee on the Effects of Atomic Radiation.

 Sources and Effects of Ionizing Radiation. No. E.94.IX.11. New York: United Nations
- Van Calsteren, K., Heyns, L., De Smet, F., et al., 2010. Cancer during pregnancy: an analysis of 215 patients emphasizing the obstetrical and the Neonatal outcomes. J. Clin. Oncol. 28 (4), 683–689.
- Van der Giessen, P.H., 1997. Measurement of the peripheral dose for the tangential breast treatment technique with Co-60 gamma radiation and high energy X-rays. Radiother. Oncol. 42 (3), 257–264.
- Wakeford, R., 2008. Childhood leukaemia following medical diagnostic exposure to

- ionizing radiation in utero or after birth. Radiat. Prot. Dosimetry 132 (2), 166–174. Wakeford, R., Little, M.P., 2003. Risk coefficients for childhood cancer after intrauterine irradiation: a review. Int. J. Radiat. Biol. 79 (5), 293–309.
- Woo, S.Y., Fuller, L.M., Cundiff, J.H., et al., 1992. Radiotherapy during pregnancy for clinical stages IA-IIA Hodgkin's disease. Int. J. Radiat. Oncol. Biol. Phys. 23 (2), 407–412
- Yonekura, Y., Tsujii, H., Hopewell, J.W., et al., 2014. International commission on radiological ProtectionICRP publication 127: radiological protection in ion beam radiotherapy. Ann. ICRP 43 (4), 5–113.
- Yoshimoto, Y., Kato, H., Schull, W.J., 1988. Risk of cancer among children exposed in utero to A-bomb radiations, 1950–84. Lancet 2 (8612), 665–669.