Re-irradiation in lung disease by SBRT: a retrospective, single institutional study

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Abstract
Background: The loco regional relapse is frequent in the lung disease. The aim of this study was to evaluate the outcomes of re-irradiation by SBRT in terms of Local Control (LC) and toxicities.

Methods: From April 2011 to December 2016, twenty-two patients received a re-irradiation by SBRT. Twenty-seven lesions were treated. The medium BED(10) of re-irradiation was 100.6 Gy (range: 48–151.2 Gy) and the medium EQD2(10) was 93.8 Gy (range: 40–126 Gy). In the previous treatment the medium BED(10) was 97.2 Gy (range: 40–120 Gy), the medium EQD2(10) was 81 Gy (range: 32.5–100 Gy). The median time between the first and the second treatment was 18 months.

Results: Local Control was reached in 18 out of 27 (66%) re-irradiated lesions, with rates of 67 and 54% at 1- year and 2- years respectively. The treatment was well tolerated; the maximum recorded toxicity was Grade 3.

Conclusions: Re-irradiation by SBRT may represent an option for the treatment of lung disease with good results in terms of LC and toxicity.

Keywords: Re-irradiation, SBRT, Lung, Primary or metastatic disease

Background
Lung cancer is one of the most important causes of death for cancer in the world. Patients with early-stage lung cancer have a probability of 4–10% developing a second lung tumor in the first 5 years after treatment [1, 2]. Patients with stage III Non-Small Cell Lung Cancer (NSCLC) treated with concurrent chemo-radiotherapy (CRT) have a risk of 30% for isolated locoregional failure [3]. The lungs are a common site for metastases. A considerable percentage of patients have disease limited to the thorax, which represents an ideal subset for local therapy [4]. A considerable number of patients with primary or metastatic lung lesions need to repeat thorax treatment for recurrent or metachronous disease. For this reason, thoracic re-irradiation starts to be common in clinical practice. Thorax recurrences can be separated in local (lung parenchyma, bronchial stump, or chest wall) and regional (mediastinal lymph nodes). The aim of treatment can be palliative or curative [5]. The improvements of technologies with Stereotactic Body Radiation Therapy (SBRT) have introduced the concept of ablative radiotherapy with curative intent. The SBRT is able to give a high dose of radiation sparing the surrounding normal tissue in 1–5 fractions [6].

The purpose of this study was to evaluate clinical outcomes in terms of Local Control (LC) and toxicity in patients with lung disease (primary or metastatic) which have received re-irradiation by SBRT. Also we have correlated Local Control with tumor histology, status of patient, number of fraction, doses, chemotherapy after treatment, time from the first progression of disease after the first treatment, time between both treatments. Furthermore, we have correlated toxicity with site of treatment, doses and volume.

Methods
From April 2011 to December 2016, twenty-two patients received a re-irradiation by SBRT. Twenty-seven lesions were re-irradiated. There were 15 (68%) male and 7 (32%) female. Median age at time of enrolment was 70 years (range: 47–82 years). This study included patients with comorbidity or advanced...
age or patients with low cardio-respiratory reserve which were unsuitable for surgery or patients refusing surgery. The patients at time of re-irradiation had an Eastern Cooperative Oncology Group (ECOG), PS 0–2, and the intent of treatment was non-palliative. Patients and tumor characteristics are summarized in Table 1. The median time to the progression disease from first treatment was 10 months (range: 0–54 months). The median time between the first and the second treatment was 18 months (range: 6–66 months). For each patient we obtained the disease history and an objective examination. According to these informations we classified for ECOG, PS. The imaging study was performed using TBCT with contrast enhancement or with 18FDG PET-CT to assess the status of systemic disease. The median follow-up time was 13 months (CI: 95%, range = 13–65 months).

**Treatment plan**

All patients underwent computed tomography (CT) simulation and were immobilized in the supine position. A 4DCT system was used to generate an Internal Target Volume (ITV) to evaluate respiratory excursion. The ITV was created on the MIP (Maximum Intensity Projection). When the differentiation between postsurgical/post-radiotherapy scarring and recurrent tumor was not well visible, a fusion of images with 18 FDG PET-CT was done to better define it. The sum of GTV (Gross Tumor Volume) and ITV was expanded by 4 mm in all directions to create the PTV (Planning Tumor Volume). The SBRT plan was created with a conformational 3D technique, using multiple fields (5–7 fields), non-opposing and coplanar. The energy was 6–15 MV. The median prescription dose was 98% (range: 94–99%). The characteristics of lesions are summarized in Table 2. We considered re-irradiation in-field when there was overlap of the two separate dose distributions for the 80% dose level each time [7].

**Doses**

Seven lesions (26%) were re-irradiated with a single fraction and twenty (74%) lesions were treated with multiple fractions. Table 3 summarized the doses and the number of fractions of first treatment and re-irradiation. At the time of re-irradiation, the choice of single fraction vs. multiple fractions was based on the volume of lesions and on the proximity of the target to critical organs, such as large vessels or bronchial shaft. The doses were measured also in terms of BED (Biological Effective Dose) and EQD2 (Equivalent Dose in 2 Gy), assuming that the α/β ratio is 10 for tumor and acute toxicity [8]. The doses in terms of BED

**Table 1** Characteristics of Patients

| Characteristics                                      | Number | Percent |
|--------------------------------------------------------|--------|---------|
| Total number of patients                               | 22     |         |
| Males                                                  | 15     | 68      |
| Females                                                | 7      | 32      |
| Primitive lung cancer                                  | 13     | 59      |
| NSCLC                                                  | 12     | 54      |
| SCLC                                                   | 1      | 4       |
| PET Positive, No histological report                   | 2      | 9       |
| METASTASIS                                             | 7      | 32      |
| Colon Cancer                                           | 2      | 10      |
| Rectal Cancer                                          | 2      | 10      |
| Breast Cancer                                          | 1      | 4       |
| Carcinoma of the Uterus                                | 1      | 4       |
| Oropharyngeal Cancer                                   | 1      | 4       |
| Histology                                              |        |         |
| Adenocarcinoma                                         | 13     | 59      |
| Squamous Carcinoma                                     | 6      | 27      |
| SCLC                                                   | 1      | 5       |
| PET Positive, No histological report                   | 2      | 9       |
| Performance status                                     |        |         |
| 0                                                      | 12     | 55      |
| 1                                                      | 9      | 41      |
| 2                                                      | 1      | 4       |
| Status of patients                                     |        |         |
| Patients with only one site of disease                  | 4      | 15      |
| Patients with others site of disease, but stable        | 23     | 85      |
| Chemotherapy after re-irradiation                      |        |         |
| Yes                                                    | 8      | 30      |
| No                                                     | 19     | 70      |
| Diagnosis of recurrence disease                        |        |         |
| PET-CT 18FDG                                           | 19     | 70      |
| TBCT                                                   | 8      | 30      |
| Histological Exam                                      | 0      | 0       |
| Localization of lesions                                |        |         |
| Peripheral                                             | 23     | 85      |
| Central                                                | 4      | 15      |

**Table 2** Characteristics of the Lesions

| Characteristics                                      | Number | Range   |
|--------------------------------------------------------|--------|---------|
| Total number of lesions                                | 27     |         |
| Volume                                                |        |         |
| Medium Volume (cc)                                     | 30,8   | 2,7–260,7|
| Diameter                                              | 3,7    | 1,7–7,9 |
| Localization of lesions                                |        |         |
| Out-field                                              | 6      | 22      |
| In-field                                               | 21     | 78      |
Table 3 Number of Fractions and Doses

| Number of fractions of the first treatment (SBRT) | Number | Percent |
|-------------------------------------------------|--------|---------|
| Single Fraction                                 |        |         |
| 30 Gy/ 1 fr                                     | 11     | 42      |
| 23 Gy/1 fr                                      | 3      | 11      |
| Multiple Fractions                              |        |         |
| 45 Gy/ 3fr                                      | 3      | 11      |
| 50 Gy/5 fr                                      | 3      | 11      |
| 60 Gy/ 30 fr                                    | 5      | 18      |
| 30 Gy/ 10 fr                                    | 2      | 7       |

| Number of fractions of the re-irradiation (SBRT) |        |         |
| Single Fraction                                 |        |         |
| 30 Gy/ 1 fr                                     | 4      | 15      |
| 23 Gy/1 fr                                      | 3      | 11      |
| Multiple Fractions                              |        |         |
| 54 Gy/ 3 fr                                     | 3      | 11      |
| 45 Gy/ 3fr                                      | 5      | 18      |
| 50 Gy/ 5fr                                      | 8      | 30      |
| 30Gy/fr 5                                       | 4      | 15      |

Medium biological effective dose (10Gy) (BED10)

| I Treatment                                      | Dose  | Range   |
|-------------------------------------------------|-------|---------|
| 97,2 Gy                                          | 40–120|
| Re-Irradiation                                  | 100,6 Gy | 48–151,2 |
| Medium equivalent dose in 2 Gy (10 Gy) (EQD2)    | DOSES | RANGE   |
| I treatment                                     | 81 Gy | 32,5–100 Gy |
| Re-Irradiation                                  | 93,8 Gy | 40–126 Gy |
| Cumulative medium BED (10)                      | 197,8 Gy | 88–271,2 Gy |
| Cumulative medium EQD2 (10)                     | 164,8 Gy | 72,5–226 Gy |
| Median biological effective dose (10Gy) (BED10) | Doses | Range   |
| I Treatment                                     | 112,5 Gy | 40–120 Gy |
| Re-Irradiation                                  | 100 Gy | 48–151,2 Gy |
| Median equivalent dose in 2 Gy (10 Gy) (EQD2)    | Doses | Range   |
| I treatment                                     | 93,8 Gy | 32,5–100 Gy |
| Re-Irradiation                                  | 83,3 Gy | 40–126 Gy |
| Cumulative median bed (10)                      | 200 Gy | 88–271,2 Gy |
| Cumulative median EQD2 (10)                     | 166,7 Gy | 72,5–226 Gy |

* Treatment done by IMRT technique

** Treatment done by 3D technique

(10) and EQD2 (10) are summarized in Table 3. For each treatment there was the DVH (Dose Volume Histogram) that evaluated the dose received by the organs at risks and by the target. It was not possible to sum the DVH of the first treatment and second treatment because there was not algorithm or image system that permitted it.

Follow up

The first follow-up was at 2 months. We evaluated physical examinations, blood tests, tumor markers and TBCT with contrast enhancement. Then every 3 months for 2 years by first follow-up, after that every 6 months. The first imaging was done by CT scan with contrast enhancement, and then patients had alternatively 18 FDG PET-CT and TBCT. The criteria for radiological response evaluation were extrapolated from the RECIST 1. 1 scale (Solid Tumors Response Evaluation Criteria). Toxicity was classified by the CTCAE scale v.4.

Statistical analysis

The first end point was the Local Control (LC) and the second end point was toxicity. The LC is defined as the
absence of new or progressive lesions within or at the margin of the PTV re-irradiated. In the statistical analysis we described also the Progression Free Survival (PFS), the Disease Specific Survival (DSS) and the Overall Survival (OS). The PFS was calculated from the first day of SBRT (re-irradiation) and was defined absence of any local and distant disease. The DSS was calculated from the first day of treatment (re-irradiation) until death for disease. Patients died for causes not related to disease were not considered as event. The OS was calculated from the first day of re-irradiation by SBRT until death. LC, PFS, DSS and OS were estimated using the Kaplan-Meier method. Differences between survival curves and tumor histology, status of patient, number of fraction, doses, chemotherapy after treatment, time from the first progression of disease after the first treatment and time between both treatments were tested with log-rank test analysis. Statistical analyses were performed using the software package SPSS, version 24 (IBM corporation). A $p \leq 0.05$ was considered statistical significant.

Results

The results in terms of Local Control and survival index are showed in Table 4, those of toxicity in Table 5. Finally, the correlation between the LC, survival index and variables are showed in Table 6.

Local control

Local control was reached in 18 out of 27 (66%) re-irradiated lesions, with rates of 67 and 54% at 1-year and 2-years respectively (Fig. 1). In the univariate analysis of the single fraction vs. multiple fractionation, the median cumulative EQD2(10) of the two treatments $\geq 167$ Gy vs. $< 167$ Gy and the median cumulative BED(10) of the two treatments, $\geq 200$ Gy vs. $< 200$ Gy showed a statistically significant correlation with local control, respectively with a $p$ value of 0.05 (Fig. 2), 0.03 (Fig. 3) and 0.02 (Fig. 4). No other statistically significant correlations were found for the other variables. Considering the heterogeneity of patients was not relevant to consider the other survival index as PFS, DSS and OS.

Table 4 Index of survival and Local Control

|       | 1 Year | 2 Years |
|-------|--------|---------|
| aLC   | 67%    | 54%     |
| bPFS  | 45%    | 39%     |
| cDSS  | 81%    | 63%     |
| dOS   | 81%    | 63%     |

a Local Control  
b Progression Free Survival  
c Disease Specific Survival  
d Overall Survival

Table 5 Toxicity

|       | aAcute | bLate |
|-------|--------|-------|
| Grade | 1 2 3 4 1 2 3 4 |
| Dyspnea | 4 1 0 0 1 4 1 0 |
| Chest pain | 0 1 0 0 0 1 0 0 |
| Laryngeal hemorrhage | 1 0 0 0 0 0 0 0 |
| Peripheral sensory neuropathy | 0 1 0 0 0 0 0 0 |
| Cough | 1 0 0 0 0 0 0 0 |
| Productive cough | 1 0 0 0 0 0 0 0 |
| Pulmonary fibrosis | 0 0 0 1 1 2 0 0 |
| Rib fracture | 0 0 0 1 0 0 0 0 |
| Pleural effusion | 0 0 0 0 1 0 0 0 |
| Gastro-esophageal reflux | 0 0 0 0 0 1 0 0 |
| Soft tissue necrosis | 0 0 0 0 0 1 0 0 |

The number is the number of case with the toxicity described  
a Within the first 3 months after re-irradiation  
b After 3 month after re-irradiation

Toxicity

The treatment was well tolerated. The maximum recorded toxicity was Grade 3. For 16 lesions (59%) treated we did not find acute toxicity and also for 16 lesions (59%) we did not register late toxicity. Acute and late toxicity are summarized in Table 5. There was one case of dyspnea (4%) G3 as acute toxicity. The same patient (4%) had dyspnea and fibrosis G3 associated with rib fracture G1 as late toxicity. The lesion was in-field. The dose was 50 Gy in 5 fractions, the BED (10) sum was 232.

Table 6 Correlations of Local Control and index of survivals with variables

|       | LC | PFS | DSS | OS |
|-------|----|-----|-----|----|
| Histology | 0.6 | 0.3 | 0.3 | 0.3 |
| Status of patients | 0.7 | 0.5 | 0.3 | 0.3 |
| Number of fractions | 0.05 | 0.09 | 0.06 | 0.06 |
| Median EQD2 (10 Gy) of re-irradiation | 0.4 | 0.4 | 0.7 | 0.7 |
| Median BED2 (10) of re-irradiation | 0.4 | 0.4 | 0.7 | 0.7 |
| Median EQD2 (10 Gy) cumulative | 0.03 | 0.03 | 0.1 | 0.1 |
| Median BED (10) cumulative | 0.02 | 0.05 | 0.08 | 0.08 |
| CHT after re-irradiation | 0.2 | 0.09 | 0.8 | 0.8 |
| Median time between both treatments | 0.4 | 0.5 | 0.6 | 0.6 |
| Median time of first progression of the disease | 0.6 | 0.5 | 0.3 | 0.3 |

Histology: Adc (adenocarcinoma), Scc (Squamous cell carcinoma), Scl (Small cell lung cancer), Pet+, Without Histology  
a Single Lesion vs. Multiple Lesions  
b Single Fraction vs. Multiple Fractions  
c EQD2(10) $\geq 83$ Gy vs. $< 83$ Gy  
d BED(10) $\geq 100$ Gy vs. $< 100$ Gy  
E Cumulative EQD2(10) $\geq 167$ Gy vs. $< 167$ Gy  
F Cumulative BED (10) $\geq 200$ Gy vs. $< 200$ Gy  
G CHT (Chemotherapy) after re-irradiation: Yes vs. No  
H Median time between both treatments $\geq 18$ months Vs. $< 18$ months  
I Median time of the first progression of disease $\geq 10$ months vs. $< 10$ months
5 Gy, the EQD2(10) sum was 193.8 Gy and the volume was 27 cm$^3$. There was one case (4%) of chest pain G3 as late toxicity. It was an in-field lesion. The dose was 45 Gy in 3 fractions, the BED (10) sum was 232.5 Gy, the EQD2 (10) sum was 193.8 Gy, and the volume was 11 cm$^3$. There was one case (4%) with pleural effusion G2 associated with pulmonary fibrosis G3, as late toxicity. It was an out-field lesion. The dose was 50 Gy in 5 fractions, the BED (10) sum was 172 Gy, the EQD2(10) sum was 148.3 Gy. The volume was 46.6cm$^3$. All the lesions had a peripheral localization.

**Discussion**

Primary and metastatic lung diseases often relapse, creating difficulties in the type of therapeutic approach to be use. The relapses may occur in different sites: in the same site or another site of lung or in the lymph node. It can relapse outside of lungs. Generally the first approach for lung disease recurrence is surgery or chemotherapy. Actually progression in technologies, especially SBRT, gives a new treatment approach, not only as palliative intent [9, 10]. There are few data about the efficacy, the doses, the number of fraction and the ideal time between the two treatments. Usually, the target of
the first treatment is larger than the target of retreatment. More attention is focused about toxicity, involving more reasonable choice of doses and volume of retreated lesions [11]. The aim of this study was to evaluate the efficacy and tolerability of SBRT re-irradiations treatment of curative intent. We had a LC of 66%, with rates of 67% at 1-year and 54% at 2-years. The others studies showed rates of LC from 59 to 95% at 1-year and from 50 to 92% at 2-years [12]. Liu et al. at MD Anderson in a retrospective study have analysed 386 patients with primitive or metastatic lung disease treated by SBRT for a single lung recurrence, with a maximum diameter ≤ 4 cm. The dose was 50 Gy in 4 fractions, the local control was 95% at 1-year, the median follow-up was 16 months [13]. Kilburn et al. studied a group of 33 patients with primitive and metastatic lung disease treated by SBRT for in-field and out-field lesions. The dose was 50 Gy in 10 fractions. The local control at 2 years was 67%. The median follow-up was 17 months. They found greater efficacy in terms of local control with multiple fractions vs. single fractionation (Log Rank, \( p = 0.006 \)) [14]. As know, the tumor radio resistance depends on the degree of hypoxia. Therefore, if we have more fractions we have more time to reduce the degree of hypoxia.
and improve the tumor response [12]. In our study, we observed a statistically significant correlation of single fraction vs. multiple fractions ($p = 0.05$), median cumulative EQD2(10) of the two treatments $\geq$ 167 Gy vs. $< 167$ Gy ($p = 0.03$) and median cumulative BED (10) of the two treatments $\geq$ 200 Gy vs. $< 200$ Gy ($p = 0.02$). Comparing these data with those of other studies, we observed discordance about local control results and number of fraction. A possible reason is that the lesions treated with single fraction are smaller than lesions treated with multiple fractions. We did not find any statistical correlation with volume of lesions. Regarding the dose, in terms of BED(10) and EQD2(10), our data confirm the data of Reyngold et al. They found a statistical correlation between LC and BED(10) $\geq$ 100Gy for the re-irradiation ($p = 0.04$) [4]. Trakul et al. showed good results on correlation between LC and BED when the time between the two treatments was greater than 16 months ($p = 0.042$) [15]. Reyngold et al. confirmed these good data with an interval greater than 36 months (HR = 0.25, $p = 0.05$). In addition, the same study showed good results on LC for a PTV volume $< 75$ cm$^3$ ($p = 0.03$) and a KPS $\geq$ 80 ($p = 0.03$) [4]. Kilburn et al. found a correlation between PTV and LC ($p = 0.03$) [14]. The lesion volume is very important to define the radio resistance, because in a bigger volume there were a lot of hypoxic cells [7]. We did not find the same correlations in our study. In our study no major toxicity findings or fatal events were identified. The tolerance was good, with 59% of cases with no acute toxicity and the same percentage with no late toxicity. The maximum acute toxicity was one case of dyspnea G3. The maximum late toxicities were a case of dyspnea G3, a case of chest pain G3 and two case of pulmonary fibrosis G3. There were no correlation with dose, localization and volume of lesions. Kelly et al. showed a correlation between out-field lesions and Grade 3 pneumonitis ($p = 0.03$). However, they did not find correlation with volume, lesion localization (central or peripheral), dose or with history of lung disease. Patients treated for in-field lesions had more frequently chest wall pain and low rates of pneumonitis, while patients treated for out-field lesions had a higher frequency of pneumonitis and less probability of chest pain. Radiation pneumonitis is an inflammatory process. One hypothesis is that previously irradiated areas can be fibrotic, and less susceptible to the inflammatory process caused by a new radiotherapy. No clear explanation can be found for chest pain. One hypothesis is that the nervous damage, fractures and myositis were caused by the re-irradiation [16]. Trovò et al. found an association between the dose received from the heart and the incidence of radiation pneumonitis. Patients who developed grade 3 to 5 pneumonitis had a higher heart Dmax (mean Dmax = 27 Gy), $p < 0.05$ [17]. Liu et al. described pre-SBRT parameters that could influence the incidence of Grade 3–5: ECOG PS 2–3 ($p = 0.008$), a pre-SABR FEV1 $\leq$ 65% ($p = 0.012$), a V20 $\geq$ 30% in the composite plan ($p = 0.020$), and previous bilateral mediastinal PTV ($p = 0.024$) [13]. The limits of our study were a small number of patients, a lack of homogeneity in the first treatment and re-irradiation, and a missing algorithm able to sum the doses for OARs and target between the two treatments. We have not reported the results about the index of survival, the correlation with variables because patients and tumor characteristics were too heterogeneous and the study sample was too small to find statistically significant results.

**Conclusions**

In this study we showed that SBRT may represent a new chance of treatment of curative intent in the recurrence lung disease. Toxicity was acceptable. However, more knowledge and studies with a higher number of patients, and greater homogeneity of treatment and longer follow-up could help to understand better who is the best candidate to obtain optimal results in terms of LC and toxicity.

**Abbreviations**

18FDG PET-CT: PET 18-F-Fluorodeoxyglucose (FDG); Positron Emission Tomography integrated with Computed Tomography; 3D: 3 Dimensional; 4DCT: 4Dimensional Computed Tomography; BED (10): Biological Effective Dose ($\alpha/\beta = 10$ Gy); cc: Centimeter Cubic; CI: Confidence Interval; CRT: Chemo-Radio-Therapy; CTCAE: Common Terminology Criteria for Adverse Events; Dmax: Dose Maximum; DSS: Disease Specific Survival; DVH: Dose Volume Histogram; ECOG: Eastern Cooperative Oncology Group; EQD2 (10): Equivalent Dose in 2 Gy ($\alpha/\beta = 10$ Gy); FEV1: Forced Expiratory Volume in one second; G: Grade; GTV: Gross Tumor Volume; HR: Hazard Ratio; ITV: Internal Target Volume; KPS: Karnofsky Performance Status Scale; LC: Local Control; MIP: Maximum Intensity Projection; MV: Mega Volt; NSCLC: Non-Small Cell Lung cancer; OARs: Organs At Risk; OS: Overall Survival; PFS: Progression Free Survival; PS: Performance Status; PTV: Planning Target Volume; RECIST: Solid Tumors Response Evaluation Criteria; SBRT: Stereotactic Body Radiation Therapy; TBCST: Total Body Computed Tomography; V20 $\geq$ 30%: Percent volume of lung exposed to at least 20Gy $\geq$ 30%

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**Availability of data and materials**

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

**Authors’ contributions**

DC, MV and MFO participated in design and coordination of study. DC, SDM, PB contributed to data acquisition and contributed to the statistical analysis. GM, VDS, PB contributed to interpretation of data. DC and SD drafted the article. IR, GM and MFO critically reviewed/revised the article. All authors read and approved the final manuscript.
Ethics approval and consent to participate
Under a S. Andrea Hospital IRB-approved protocol the data were prospectively collected and retrospectively analysed to evaluate the efficacy and the tolerance. Written informed consent was obtained by all patients.

Consent for publication
The data was anonymized. The analysis was performed retrospectively.

Competing interests
The authors declare that they have no competing interests.

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