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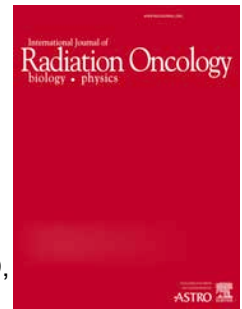
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DIBH (Deep Inspiratory Breath Hold)-based radiotherapy – a clinical review

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**DIBH (Deep Inspiratory Breath Hold)-based radiotherapy – a clinical review*****Running title: DIBH-based radiotherapy – a clinical review***

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**DIBH (Deep Inspiratory Breath Hold)-based radiotherapy – a clinical review**

*Running title:* DIBH-based radiotherapy – a clinical review

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**Abstract:**

Several recent developments in linear-accelerator-based radiotherapy such as fast multileaf collimators, accelerated intensity modulation paradigms like VMAT and flattening filter-free (FFF) high-dose-rate therapy have dramatically shortened the duration of treatment fractions. Deliverable photon dose distributions have approached physical complexity limits as a consequence of precise dose calculation algorithms and online 3D-image-guided patient positioning (Image-Guided RadioTherapy, IGRT).

Simultaneously, beam quality and treatment speed have continuously been improved in particle beam therapy, especially for scanned particle beams.

Applying complex treatment plans with steep dose gradients requires strategies to mitigate/compensate for motion effects in general and particularly for breathing motion. Intrafractional breathing-related motion results in uncertainties in dose delivery and thus in target coverage. As a consequence, generous margins have been used, which, in turn, increases organ-at-risk (OAR) exposure. Particle therapy, particularly with scanned beams, poses additional problems such as interplay effects and range uncertainties. Among advanced strategies to compensate breathing motion such as beam gating and tracking, DIBH-gating is particularly advantageous in several respects, not only for hypofractionated, high single-dose Stereotactic Body RadioTherapy (SBRT) of lung-, liver- and upper abdominal lesions but also for normofractionated treatment of thoracic tumors such as lung cancer, mediastinal lymphomas and breast cancer. This review provides an in-depth discussion of the rationale and technical implementation of DIBH-gating for hypo- and normofractionated radiotherapy of intrathoracic and upper abdominal tumors in photon and proton radiation therapy.

**Key words:** DIBH (Deep Inspiratory Breath Hold), motion management; breathing motion, gating, SBRT (Stereotactic Body Radiotherapy), IGRT (Image-Guided RadioTherapy), Proton therapy, Particle Therapy

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## Introduction

Several recent developments in linear-accelerator-based photon radiotherapy such as intensity modulated radiotherapy (IMRT, [1]) and volumetric modulated arc therapy (VMAT, [2, 3]) allow the application of highly complex treatment plans with steep dose gradients. Photon dose distributions in rigid treatment volumes have approached physically achievable complexity and accuracy limits as a consequence of the introduction of precise dose calculation algorithms (e.g. [4]), daily online soft-tissue based 3-dimensional image-guided patient/target positioning (IGRT, Image-Guided RadioTherapy, [5, 6]) and continuously improved delivery devices with fast collimators [7]. Flattening-filter-free (FFF) high-dose-rate applications [8-15] have dramatically accelerated small-field delivery, particularly for the SBRT (stereotactic body radiotherapy) paradigm while maintaining biological properties of the beam [16, 17]. It has several further advantages such as less scatter from the treatment source, less leaf transmission and head leakage [1, 18].

The combination of all these technical possibilities has refined and accelerated [8] the therapy of both large stationary targets like head and neck cancer [19, 20] as well as smaller mobile targets, resulting in clinical benefits such as excellent local control rates in the treatment of early NSCLC (Non-Small-Cell-Lung-Cancer) or lung/liver metastases with Stereotactic Body Radiation Therapy (SBRT, [21-26]) with very reasonable total treatment times now in the range of 15 min per treatment fraction.

Proton therapy is now applied with increasing frequency, with new treatment facilities being activated on a regular basis. It has made significant technological progress recently with more widespread use of scanned beams and the introduction of 3D-Image-Guidance. Nine times rescanning of a one liter volume within one minute is now technically feasible, bringing into reach treatment deliveries during the time span of one breath hold [27]. An innovative design for image guidance is the integration of a beams-eye view (BEV) imager at Gantry 2 at PSI (Paul Scherrer Institute) in Switzerland, which is a fast parallel beam



scanning proton therapy unit with small spot size and penumbra, allowing X-ray imaging in fluoroscopy mode during treatment delivery [27, 28]. However, target motion implies a much bigger challenge for proton therapy than for photon therapy, especially for a scanned delivery where interplay effects can significantly disturb the planned dose distribution [29]. Furthermore, image guided approaches are much more advanced in photon radiotherapy and online 3D motion monitoring has not been realized for particle therapy to date [30]. Despite constant efforts to mitigate motion effects [31-33] in both advanced photon and proton therapy of body regions that are affected by breathing motion with motion amplitudes of up to 2-3 cm and potentially including hysteresis and deformations [34] there are still methodical improvements needed. Resolving remaining issues may improve the treatment of several disease entities/clinical situations. Among these are:

- Radiotherapy of locally advanced NSCLC, where escalated doses in combination with chemotherapy may improve local control [35, 36], but are limited by normal lung tolerance and methodical imprecisions. Insufficient target coverage prompted by concerns about lung toxicity may have contributed to a lack of efficacy of dose escalation in the treatment of locally advanced lung cancer in the randomized RTOG trial 0617 [37, 38].
- Exposed lung volume also plays a role in considerations regarding secondary malignancy after radiotherapy of all mediastinal tumors [39, 40]. Exposed heart volume after mediastinal or breast radiotherapy is linked to long-term cardiac toxicity [41-46].
- Treatment of non-static targets with passively scattered proton beams, which currently is not unlocking its full potential due to limitations on image guidance.
- Treatment of non-static targets with scanned proton beams, which has only been performed rarely clinically until to date because of concerns regarding interplay-effects [47].

This review describes the different methods and characteristics of available motion management strategies in photon and proton radiation therapy and then outlines how DIBH can be efficiently performed and where it may resolve or mitigate the issues and unmet methodical needs described above. Table 1 provides a synopsis of the dosimetric and clinical characteristics of DIBH treatments and compares them with other currently available motion management strategies regarding advantages and disadvantages.

### 1. Breathing-motion management strategies/methods

Even for short beam-on times now achievable with FFF-treatments, motion management strategies are necessary to compensate for intrafractional breathing motion. Different strategies aim at a reduction of margins between clinical target volume (CTV) and planning target volume (PTV) and/or improved geometrical precision of dose delivery:

- Motion amplitude of free breathing can be reduced by mechanical abdominal compression [48]. Recently however, it has been shown that it is only beneficial for lower lobe tumors and has no or a negative effect for middle/upper lobe tumors [49]. While the *intrafractional* amplitude of tumor motion can be reduced by abdominal compression, *interfraction* motion can be even increased [50]. Mechanical abdominal compression has also been evaluated theoretically [51] and used clinically in particle therapy to reduce intrafractional motion [52].
- One of the most widely used strategies is treatment planning with individual determination of CTV-PTV margins based on a 4D-CT in free breathing [33, 53]. 4D-planning requires appropriately chosen PTV-margins (Internal Target Volume (ITV) concept) considering the end-expiratory and end-inspiratory position of the tumor. Inclusion of all breathing phases during the actual treatment ensures optimal

treatments for small tumors but results in increasing volumes of healthy lung tissue exposed to high doses with increasing CTVs if CTV-PTV-margins are kept constant [54]. For particle therapy, in addition to geometrical considerations, also changes in tissue densities due to motion that affect the particle range, have to be considered when designing margins [55]. Several publications have recently reported uncertainties in the 4DCT-approach regarding breathing pattern [56], motion uncertainties, dosimetry and verification difficulties. Uncertainties have been shown regarding 4DCT-based motion measurements for lung SBRT. Confirmed by MV (MegaVoltage) imaging during beam-on, Zhang et al. [57] have shown that 4DCT may underestimate the overall maximum tumor motion range during lung SBRT. For liver SBRT, a single 4DCT for planning was not always correctly representing the mean motion amplitude (measured by kV (kilovoltage) and MV marker-based imaging) during treatment [58]. A large variation of intra- and interfractional motion patterns for various targets has been also observed [59], especially in antero-posterior direction and in a fraction-duration dependent manner [60, 61]. Measurements of motion of implanted fiducials with daily orthogonal fluoroscopy have shown that 4DCT overestimates daily 3D motion in 39% and underestimated in 53% of the fractions. Breathing pattern varied from breath to breath and from day to day and intrafractional variation of the amplitude was significantly larger than interfractional variation [61]. Free-breathing CBCT potentially underestimates ITV if the respiratory pattern is characterised by a disparate length of time spent in inspiration vs. expiration, potentially leading to misalignments, depending also on tumor size and localisation [62, 63]. 4D-CBCT is the logical continuation of the 4D-concept through the whole treatment chain. It has become available recently and remedies several of the abovementioned issues but trades image quality for time

resolution [64-66]. In particle therapy, the value of 4D-MR imaging has been explored which enables the capturing of motion variations and drift effects [67].

- While 4D-treatment planning results in an individualized choice of PTV margins that may result in an expansion of margins compared to the population mean, real-time target tracking or continuous patient position adjustment with robotic treatment couches with 6 degrees of freedom can minimize PTV margins for all individuals [68, 69]. Several tracking technologies have been clinically established and can, for example be found in the Cyberknife concept [70, 71] or the recently released (and already discontinued) Vero System (Vero SBRT, Brainlab, Feldkirchen, Germany; [72, 73]) with steering of the beam application, or, in an experimental system, with steering of the patient couch [74]. Tracking is typically based on an individual motion model created during treatment planning that is frequently verified by planar EPID-imaging of implanted fiducials or the tumor shadow (when detectable) and/or optical surface tracking (ExacTrac: [75, 76]; Cyberknife: [77]). The clinical introduction of online-4D-Magnetic Resonance Imaging (4DMR) during photon treatment [78, 79] may further advance the concept of instantaneous tumor tracking, but significant developments still have to be made. Tracking seems to be the ideal motion mitigation technique for a steerable particle beam [80]. As it relies on real time 3D imaging information of the patient which is not yet available for particle therapy, it has, however, not been implemented clinically yet.
- Respiratory gating as free-breathing-gating or with voluntary/computer controlled breath hold minimizes PTV margins across a patient cohort, similar to what is achieved by tracking. Free-breathing-gating strategies have typically been used during end-expiration, which occupies the majority of the breathing cycle. This approach therefore allows for the application of large doses during the gating phase. Plan comparison studies, however, demonstrated that IMRT plans for the *inspiration*

phase of the breathing cycle as deep inspiration breath hold (DIBH) resulted in better V10, V20, V40 and mean lung dose when compared to plans for end-expiration also for normofractionated treatments of advanced lung tumors [54, 81-84]. Gating is the most commonly used motion mitigation technique for particle therapy [85].

- A motion mitigation technique that is unique for scanned particles is rescanning, which refers to repeated irradiations during one treatment fraction to statistically smooth out interplay effects [29]. Rescanning is suggested to be combined with other motion mitigation techniques [86].

The characteristics of DIBH-gating were summarized by a review in the framework of the STIC 2003 (*Soutien aux techniques innovantes couteuses de 2003*, [87]), which confirmed feasibility and good reproducibility of various respiratory-gated radiotherapy (RGRT) systems. Improvement of dosimetric parameters predictive of reduced pulmonary, cardiac and esophageal toxicity by RGRT was described already in this manuscript. Since then, additional data have been published that solidify the rationale for the use of DIBH-gating in various clinical situations and are reviewed in this manuscript.

## 2. Methods for establishing deep inspiration breath hold (DIBH)

DIBH can be achieved by repeated voluntary breath hold or with computer controlled commercially available devices, which can assist DIBH through airway blocking and/or feedback approaches. Breath hold gating signals now automatically trigger treatments across all major treatment device manufacturers.

### a) free DIBH/voluntary breath hold

A fully free (non-computer controlled) breath-hold technique can be used during

radiotherapy for breast cancer aiming at heart, lung and liver dose reduction [88-90]. Voluntary breath hold does not require any additional equipment. To monitor breath-hold, the distance moved by the anterior and lateral skin marks away from room lasers and additional light field verification can be used, therefore voluntary breath hold typically is not completely “uncontrolled” [89]. Despite clear dosimetric benefits (heart and lung) for both 3D tangential and VMAT plans in right- and left sided breast cancer [88, 91, 92] and acceptable precision data even in a randomised setting [92], this method is not yet in widespread use [89], though interest is increasing. The UK HeartSpare study [92] has shown comparable EPI (Electronic Portal Imaging)- and CBCT (cone-beam CT) derived precision data (systematic and random error vector of 3-5mm regarding chest wall position) of voluntary breath hold when compared to computer-controlled breath hold (Active Breathing Coordinator, Elekta). Similar CBCT-based precision data were published by Betgen et al. [93] with good intra-fraction reproducibility of chest-wall position and inter-fraction systematic and random error of 2-5mm and 1.56°. However, in these publications, no position information is provided of OARs (heart/lung) and no intra-fraction EPID verification was performed [92, 93]. Patients and staff preferred voluntary breath hold versus computer-controlled breath hold due to easier workflow and reduced cost [92]. The method seems therefore to be acceptable for breast tangential RT. Given that evaluated patient numbers are low and information on heart/lung position with a 3D soft tissue imaging method (e.g. breath-hold CBCT) was lacking in these studies there is a necessity to further evaluate this issue especially if used in the context of lung/liver RT/SBRT.

#### b) computer-controlled DIBH

##### *Breathing-volume based methods:*

Computer-controlled breath-hold systems aim at creating a static geometrical situation

of the body and the GTV within the body during the planning CT. Breathing volume based methods quantify the inspiration volume with a spirometer. Patient feedback can be established/provided with an open airway audiovisually (a “target zone” is projected on a screen or via video-goggles and the patient is instructed to inhale to reach a certain signal position on the screen [94]), as performed with the SDX System (SpiroDynr’X; France; [87, 95]) or by actually closing the airway for a defined time (as performed with the ABC-System (Elekta AB, Sweden)). Intra- and inter-fractional reproducibility for ABC is 1.7 and 3.7mm [96-103]. Brock et al. [104] measured with repeat breath hold CTs consistent intra-fraction tumor position, but inter-fraction variation of mean (range) values of 5.1 (0-25), 3.6 (0-9.7), and 3.5 (0-16.6) mm in SI, RL and AP directions. However, different breathing maneuvers (thoracic vs. chest breathing) can lead to variations in chest wall position even if inspiration volume is the same, which can lead to uncertainties regarding tumor position [105]. Recently, surface/fiducial tracking methods allow the monitoring of breath hold during one fraction. Data derived from additional optical infrared tracking have shown a mean intrafraction variation vector among breath holds of less than 2.8mm [105]. Uncertainties were observed in the anteroposterior direction (maximal 12mm). This had no influence on target coverage but on OAR doses and therefore optical tracking has been recommended for the surveillance of ABC-based breath-hold [105].

#### *Visual feedback/optical surface detection/tracking*

Breath hold with visual feedback requires optimal patient compliance and has been shown to be accurate for lung lesions with intra-fraction reproducibility of <3mm [106, 107]. Intra- and inter-FGBH (Feedback Guided Voluntary Breath hold) with computer controlled visual feedback (video goggle) resulted in a reproducibility of GTV centroid

positions of  $1.0\pm0.5\text{mm}$ ,  $1.3\pm1.0\text{mm}$ , and  $0.6\pm0.4\text{mm}$  in AP, SI and LR directions, respectively, compared to more than 1cm of tumor motion at free breathing [106].

An indirect approach for breath hold gating is optical surface tracking as it is established with reflectors within the RPM (Varian, Palo Alto, CA, USA [108]), Exac-Trac (Brainlab, Feldkirchen, Germany) or Synchrony (Accuray, Morges, Switzerland) systems or with markerless systems such as alignRT (VisionRT, London, U.K.), or Catalyst (C-RAD, Uppsala, Sweden [109]).

The markerless systems project visible light on the patient and detect the surface and surface movements caused by respiration. This movement detection can be used to verify the tumor position during respiration and to gate the beam during treatment. Several studies with different systems [110-113] compared the agreement of an optical surface tracking system and cone beam computed tomography regarding static targets and found good agreement between both techniques in most situations, indicating the general robustness of this approach. Alderliesten et al. [114] evaluated the accuracy of a 3D surface imaging system compared to CBCT for the guidance of DIBH-RT of left-sided breast cancer and found a good correlation between setup errors detected by both methods. Daily real-time surface monitoring has been shown to ensure accurate inter- and intrafraction repositioning [115, 116], reduced heart dose and acceptable treatment time of left-sided breast cancer patients especially with unfavorable cardiac anatomy [117-120]. Some data indicate that for left breast cancer radiotherapy, surface monitoring systems are superior to spirometer-based systems regarding repositioning of the external surface [121].

### 3. Characteristics and advantages of DIBH

#### a) Possibility to image under DIBH



Serpa et al. [122] have shown that markerless EPID tracking is principally suitable for treatment verification of gated SBRT but marker-based EPID imaging is also being used. For Cyberknife SBRT, breath hold imaging was performed after implantation of 2-4 fiducials directly into the tumor and a maximal tumor vector movement of 3.8mm (detected by kV flat-panel detectors) was reported [123].

Linac-mounted CBCTs that are currently on the market provide the possibility to interrupt imaging/image acquisition with reconstruction after the intended imaging angle has been completed. Such a “stop-and-go” approach allows the acquisition of a complete volume dataset under breathhold [124]. While the acquisition time is longer than that of free-breathing CBCT, image quality is significantly improved over imaging in free-breathing-only, free breathing interlaced with 3-4 breathholds [97, 125] or 4D-CBCT at identical imaging doses. The approach provides superior image quality particularly for middle- and lower-lobe lung tumors (**Fig. 1**) [124] and it also improves soft tissue contrast in upper abdominal lesions (**Fig. 2**). First experiences report feasibility, fastness and better inter-observer variability of DIBH CBCT for lung SBRT [126]. Single-breath-hold CBCT has also been implemented [127] but is not yet broadly used.

A development that is currently undergoing final refinement before clinical testing is combined kV-MV imaging [128, 129] that makes use of both kV and MV imaging devices on a linac in combination with faster gantry movement and dedicated reconstruction algorithms [130, 131]. It offers the possibility to acquire a full 3D-dataset during one breath hold (<15 sec) with acceptable imaging doses and excellent positioning precision [132].

Position of target/surrogate structures in breath-hold for liver/upper abdominal SBRT can be also controlled by stereotactic ultrasound systems [101, 133-135]. Surveillance of breath hold with ultrasound-based tracking is also under development [136, 137]. Breath-hold

imaging can be also completed with MRI-based IGRT systems by matching of intra-treatment orthogonal cine-MRI planes to pre-treatment 3D MRI datasets [78, 138].

#### b) Clinical application and dosimetric features of DIBH

In 1987, the potential for improvement in radiotherapy treatments of mobile targets by reducing respiratory effects has been first reported. An American team noticed that treatment in deep inspiration spared parts of the lungs, and they suggested a need to develop “Radiotherapy Gated to Respiration” [139]. In the following paragraphs we discuss the site-specific advantages of DIBH.

##### i. SBRT of liver lesions

Intra-breath hold liver motion and intra- and inter-fraction reproducibility of liver/diaphragm position relative to vertebral bodies during ABC-based liver SBRT was assessed by kV fluoroscopy as well as MV EPIs and movies [97]. Average maximal diaphragm motion measured by fluoroscopy during a single ABC breath-hold was 1.4 mm, also confirmed by the MV movies. Repeated CT scans in breath hold have shown a mean difference (intrafractional) in the liver surface position of -0.9 mm, -0.5 mm, and 0.2 mm in the CC, AP, and medial-lateral (ML) directions; average absolute interfraction craniocaudal offset in diaphragm position relative to vertebral bodies was 3.7mm [97].

While SBRT of lung and liver lesions was initiated with stereotactic body frames including devices to limit liver excursion during treatment with the sole objective to improve dose delivery accuracy and thus reduce PTV-margins, DIBH has been introduced soon after the clinical introduction of SBRT to immobilize the diaphragm

movement less invasively. Intrafraction precision was excellent when using fiducial markers and EPID imaging (maximal craniocaudal offset 1,7mm; [97]). Clinical results were comparable to those reported for body-frame fixation. Meanwhile, based on DIBH, a minimally invasive frameless workflow could be established together with ultrasound [101, 134] or CBCT. Results have also been excellent for hepatocellular carcinoma [140, 141], where radiotherapy as a bridging treatment before transplantation or as definitive therapy has seen renewed interest [142].

Particle radiotherapy has seen an increasing role in the treatment of hepatocellular carcinoma due to the potential of increased normal-liver sparing [143]. Often hypofractionated regimens are applied [52, 144, 145]. The combination of high motion sensitivity of particle treatments with the unforgiving character of hypofractionation (little statistical smoothing of interplay effects, sensitivity towards drift effects due to increased fraction duration) makes the application of motion mitigation techniques essential. Clinically, abdominal pressure plates and gating is most commonly used to mitigate motion effects. Especially for scanned proton therapy the combination of rescanning with other motion mitigation techniques like gating or breath-hold have been suggested [28, 47].

## ii. SBRT of lung lesions

Theoretical advantages of radiotherapy for lung cancer in deep inspiration breathhold (DIBH) have been already published in 2005 by Underberg et al. [146]: a maximally expanded healthy lung tissue allows minimizing lung dose; complete immobilization of the PTV allows reduction of PTV margins which again reduces lung dose [147, 148], **Fig. 3**. This approach has since increasingly been used for simple and reliable tumor immobilization, reduction of lung exposure [149] and heart

protection [51].

Scotti et al. [150] investigated the impact of ABC-based DIBH on PTV margins and OAR sparing for 3DCRT and SBRT for lung cancer. In comparison to free-breathing CT, PTV margins could be reduced and all dosimetric lung parameters (V20, MLD) were significantly improved using DIBH-gating.

Corresponding with the dosimetric data, clinical results of DIBH-based SBRT are promising (and comparable with results of 4DCT based/mixed SBRT cohorts [24]) for both primary lung tumors and for metastases. Actuarial 1-yr local control rates are between 90-95% (3yr-LC 82-88%;) with very low toxicity [131, 151-154]. Results seem to depend on applied dose and size of PTV [151] and the method seems to be suitable even in the re-irradiation situation [155].

By creating a static situation during treatment, DIBH prevents interplay effects. While these are likely of minor importance in modulated photon radiotherapy (with some exceptions) [156], they can significantly disturb particle treatment plans [157, 158]. Especially for lung indications, non-rigid deformations which relocate high- (ribs) and low-density (soft tissue) regions, can result in severe over- or under-shoots. Therefore, methods to restrict motion or to mitigate motion effects are highly desired.

Georg et al. evaluated passively scattered proton treatments and intensity modulated proton (IMPT) plans for shallow breathing with abdominal compression and DIBH [51]. Irrespective of treatment modality they found that DVH were improved with the DIBH technique. However, the differences between shallow breathing and DIBH did not reach statistical significance. They state that although respiration controlled proton and ion beam therapy with gating and tracking approaches is technically

feasible, shallow breathing with abdominal compression or DIBH are probably more practical for the delivery of high fractional doses. Stuschke et al. showed the robustness of single field uniform dose proton plans and IMPT plans for lung patients in a breath hold scenario [159]. As this was only a planning study, concerns about the feasibility to deliver the dose for one treatment field entirely during one breath hold were raised. As inter-breath hold positional variations during the same fraction tend to be larger than intra-breath hold variations a scan across the whole target volume during one breath hold would be required to ensure robustness. Lin et al. estimated that energy switching times/average spot delivery times of 1s/5ms are required to deliver treatment fields in about 74% of lung SBRT cases within one breath hold [160]. Current commercial systems are mainly slower than that. A system that fulfils these requirements is the Gantry 2 at the Paul Scherrer Institute (PSI) in Switzerland, which is a fast parallel beam scanning proton therapy unit with small spot size and penumbra, which was optimized for the treatment of moving targets [27, 161].

### iii. Normofractionated treatments of advanced lung tumors

As discussed above, PTV-margin reduction is essential in radiotherapy of locally advanced NSCLC to maximally exploit normal tissue tolerance in order to escalate tumor doses. Given the fact that methodical insufficiencies may have invalidated the results of RTOG 0617 [37, 38, 162], breathing management, potentially in combination with adaptive strategies as now tested in RTOG 1106 [162]. Potentially particle therapy will be mandatory for any further attempts to improve survival based on better local control.

Hanley et al., [163] as well as Rosenzweig et al. [164] published planning studies comparing dosimetric parameters of FB vs. DIBH reporting the advantages of DIBH

as early as fifteen years ago. Hanley et al. also provided proof of tolerability of breathing maneuvers by the patient [163]. Mah et al. [165] expanded on this, reporting their initial experience of a feasibility study with DIBH for NSCLC.

Since then, dosimetric advantages with reduced lung and cardiac dose have been repeatedly demonstrated for DIBH-RT in the setting of advanced lung cancer treatments [87, 148]. In plan comparison studies, IMRT plans in inspiration were significantly favorable regarding V10, V20, V40 and mean lung dose if compared to expiration plans also for normofractionated treatments of advanced lung tumors [54, 81-84].

Clinical outcome regarding toxicity and economic aspects has also been analyzed by Giraud et al [87] in the framework of the STIC project between 2004 and 2008 in 20 French centres. The reported dosimetric benefits were correlated clinically with a significant reduction of pulmonary acute toxicity, and pulmonary, cardiac, and esophageal late toxicities [87].

*iv. DIBH to reduce cardiac and pulmonary toxicity after adjuvant radiotherapy of breast cancer*

Cardiac damage has been the main concern in whole-breast radiotherapy. While improved RT-techniques seem to have measurably reduced cardiac toxicity [166], every measure should be taken to minimize cardiac exposure to doses in excess of 30Gy [41]. A very recent review summarizes the advantages of DIBH in breast cancer radiotherapy [167].

Data for DIBH-RT of left-sided breast cancer confirmed good reproducibility [168] and dosimetric advantages such as reduced lung and cardiac dose [169-171] in comparison with free-breathing planning. Sung et al. have shown significant

reduction in irradiated heart volume and V25 using DIBH if compared to plans in free breathing [170]. Verhoeven et al. [172] compared plans of supine FB (free breathing), supine DIBH and prone FB. While target coverage was similar with all modalities, doses to the heart, LAD (left anterior descending coronary artery) and contralateral breast could be most effectively reduced by supine DIBH planning. A prospective trial has shown that ABC-based breath hold can reduce the mean heart dose by 20% and dose to the lung [173]. Reduced cardiopulmonary dose by DIBH was reported by several other groups [46, 174-178] even for nodal irradiation [179]. A possible drawback of the method is the potentially higher dose to the contralateral breast [180], however, second cancer risk estimation was the same for FB and DIBH plans [181].

While the dosimetric benefits of DIBH treatments for breast cancer are striking, a recent manuscript reporting functional imaging results after DIBH- or conventional RT did not find a difference in cardiac muscle perfusion at 6 months after treatment [182]. While the correlation between these imaging changes and clinical late effects is by no means established [41] these results may be explained by too high sensitivity of the chosen imaging method or heart volumes exposed to high doses in this series that were still too large even with DIBH despite low mean heart doses. At this stage there is therefore no clinical proof of DIBH benefits.

A comparative study for whole breast irradiations between IMRT and IMPT by Mast et al. states significant dose reduction to the heart and LAD-region for IMPT even without breath hold [183]. The results showed that a breath-hold technique had no added value when using IMPT. However, using breath hold may improve the robustness of the IMPT technique, since the tissue shift will be less in breath hold.

#### v. *Hodgkin's Disease*

In addition to the potential reduction of functional damage to normal tissue, in

patients with Hodgkin's Lymphoma, a supremely curable disease frequently encountered in younger patients, reduction of irradiated tissue may reduce second cancer risk [184], adding a further motivation to perform breath hold treatments in these patients. Involved node radiotherapy in DIBH has been shown to be safe and effective [185]. Dosimetric advantages with reduced lung and cardiac/coronary dose have been demonstrated for supradiaphragmic Hodgkin lymphoma [185] also in a prospective phase II study [186] and especially for tumors of the upper mediastinum [185] and in combination with IMRT [187]. Long-term toxicity data with functional imaging are missing yet.

Protons have been pointed out to theoretically provide both excellent high-dose conformality and reduced integral dose [184]. In combination with breath-hold they could enable superior treatments for involved-field and involved-node treatment of mediastinal Hodgkin lymphoma. Clinical evidence is, however, not yet available.

vi. Other tumor entities:

Dosimetric advantages with reduced lung and cardiac dose have been also demonstrated for thoracic esophageal cancer [188, 189].

The non-invasive ablation of kidney tumors has become an intriguing concept, now that evidence regarding abscopal effects of large radiation doses is mounting [190]. It is already being explored within the framework of clinical studies (NCT02334709: Phase I-II, SBRT+tyrosine kinase inhibitors, Ghent). Both online image guidance with ultrasound [136, 137] and online MR-imaging [78, 138] now provide the technical basis for these treatments that will benefit dramatically from breath hold strategies.

So far only limited experience of particle therapy treatments in combination with breath hold can be found in the literature. Studies are restricted to the above



mentioned indications. The reason for that is that moving targets present a special challenge for particles and are not commonly clinically treated yet. A clinical trial for lung cancer, breast cancer, gastrointestinal indications and lymphomatous malignancies has recently been completed at the Abramson Cancer Centre of the University of Pennsylvania [191]. Outcomes will give more evidence on the benefit of DIBH treatment in the context for proton radiotherapy.

#### 4. Recent developments that have facilitated the use of DIBH and outlook

Quality assurance and workflow for breath hold application is fast and easy [8, 87]. Frequently voiced concerns regarding DIBH have concentrated on the necessity for optimal patient collaboration/compliance with the procedure, sufficient pulmonary reserve and the longer treatment time in comparison to non-gated or tracked treatments [192]. With the advent of fast MLCs, VMAT and particularly the FFF-technology, the prolongation of treatment time of a gated over a non-gated treatment has been dramatically reduced [8]. Patient collaboration is excellent under these conditions if assisted breath hold is used and a minimum of training is provided. DIBH has been shown to be safe and effective [147] and to have positive effects in fractionated therapy of various thoracic and upper abdominal tumor entities (table 1).

In the future, DIBH will likely facilitate the development of new treatment paradigms and the refinement of existing ones.

Therapy with scanned particle beams will likely be more robust and more mobile targets will therefore be accessible to this treatment paradigm.

Online MR-based IGRT will provide the possibility for instant replanning on a daily basis.

DIBH in this context increases the similarity of target/body geometry from treatment day to treatment day and thus may facilitate instant replanning using previous knowledge.

## Conclusion

DIBH-gating is a precise, reliable technique that is applicable to most patients and, with the advent of fast delivery techniques, no longer results in excessive treatment times (**Fig. 4**). It facilitates the application of complex treatment plans with steep dose gradients to moving targets for both photon and particle therapy by widening the therapeutic window and improving dosimetric accuracy.

### Legends to tables and figures:

**Table 1.** Characteristics of DIBH treatments within the framework of advantages and disadvantages of currently available motion management strategies

**Fig.1.** Comparison of imaging paradigms for lung lesions: Upper row: CBCT under repeat breath hold, including free breathing phases into the reconstruction. Note the blurring at the tumor surface and diaphragm. Lower row: CBCT stop-and-go (same number of frames in reconstruction but all frames acquired under breath-hold conditions). Note the improved image quality and reduction of blurring.

**Fig.2.** A-C: Helical treatment planning CT for comparison, D-F: Excellent CBCT-image quality in the upper abdomen with stop-and-go acquisition (all frames acquired under breath hold).

**Fig.3.** Treatment planning for lung SBRT: A) comparison of PTV and lung DVHs in FB vs. DIBH, coronal and sagittal matched planning CTs in DIBH and FB. B) treatment plan without breathing management (predominantly end-expiration). C) treatment plan in DIBH. Note expanded lung tissue and smaller PTV margins.

**Fig. 4.** Hallmarks of DIBH workflow. Left, breath curve and patient with Catalyst; right, breathing curve and patient with ABC.

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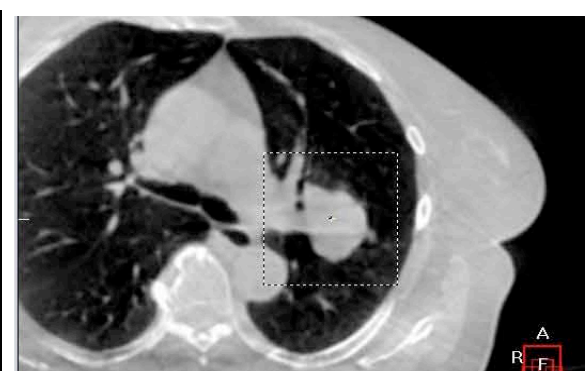
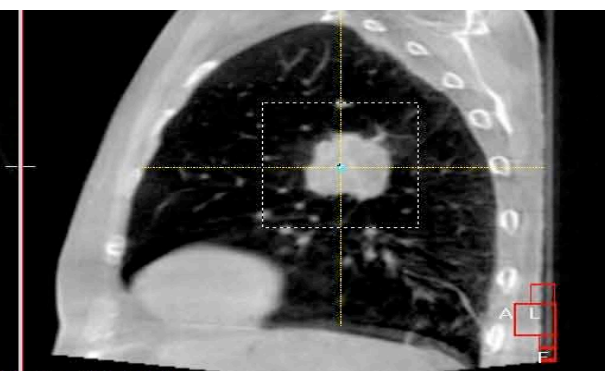
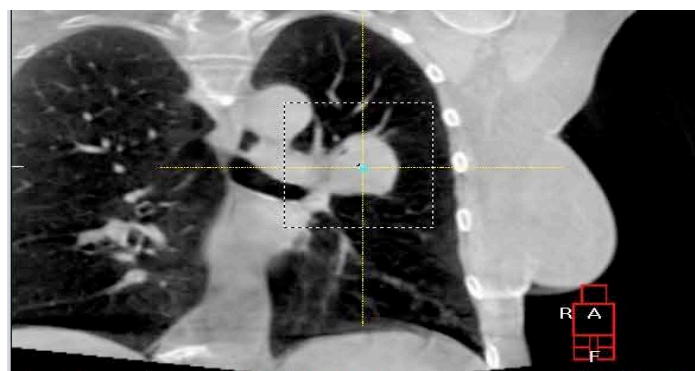
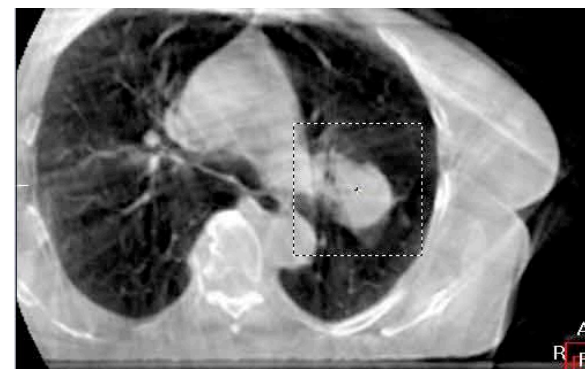
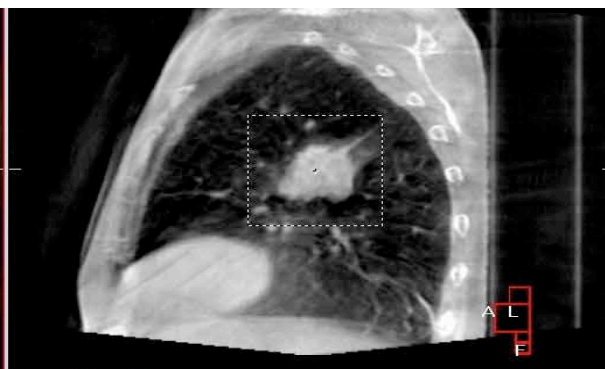
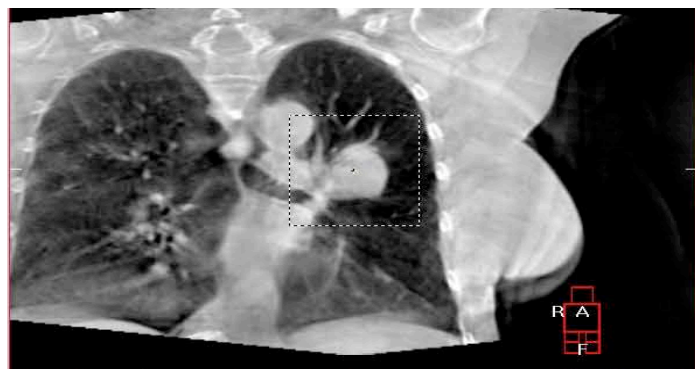
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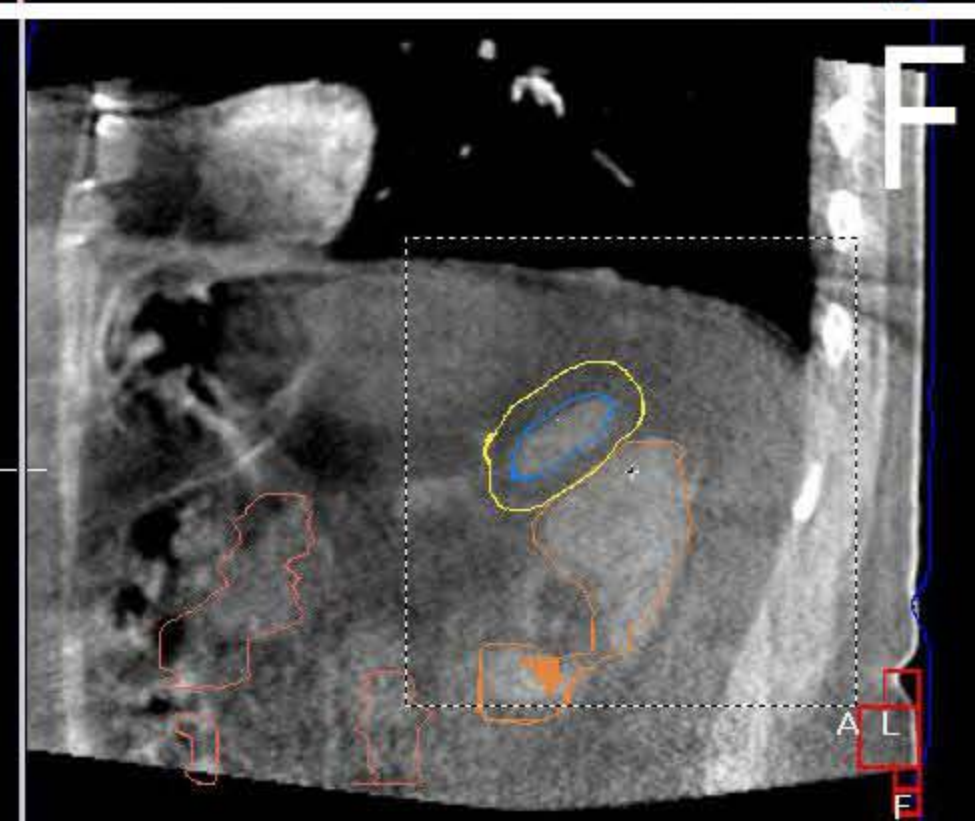
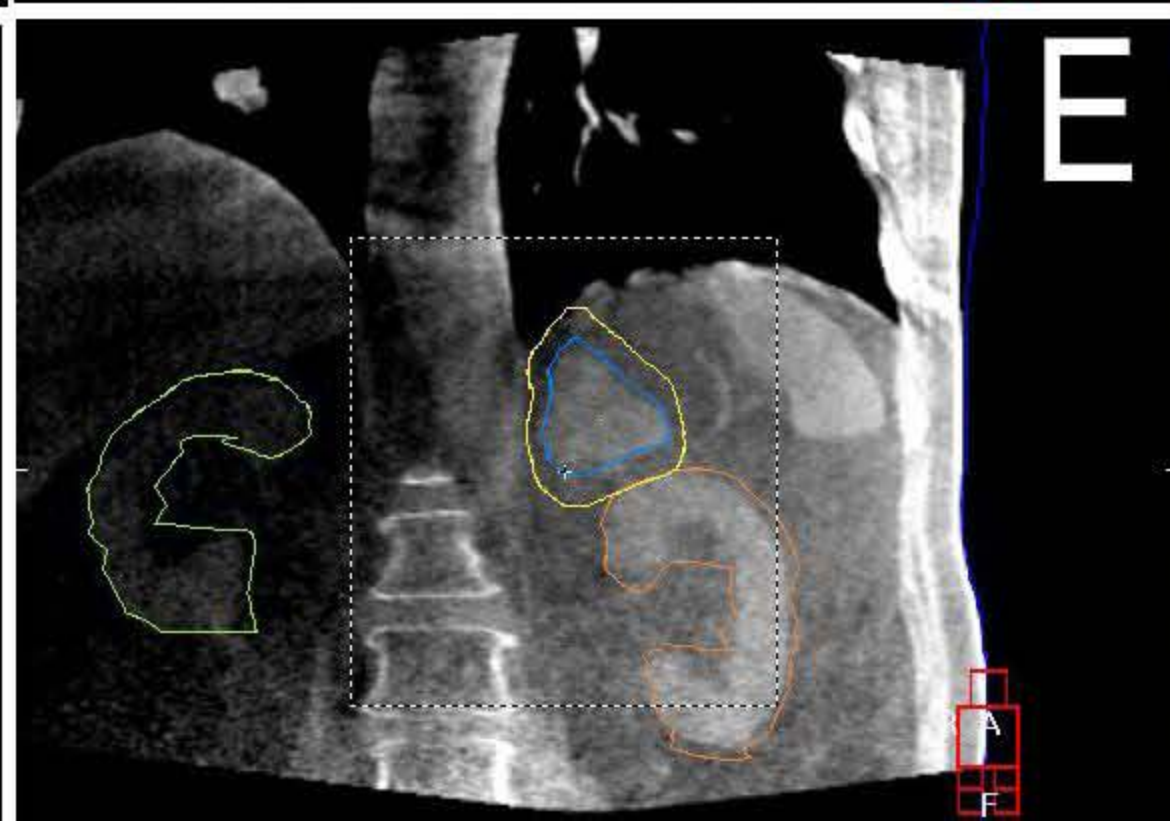
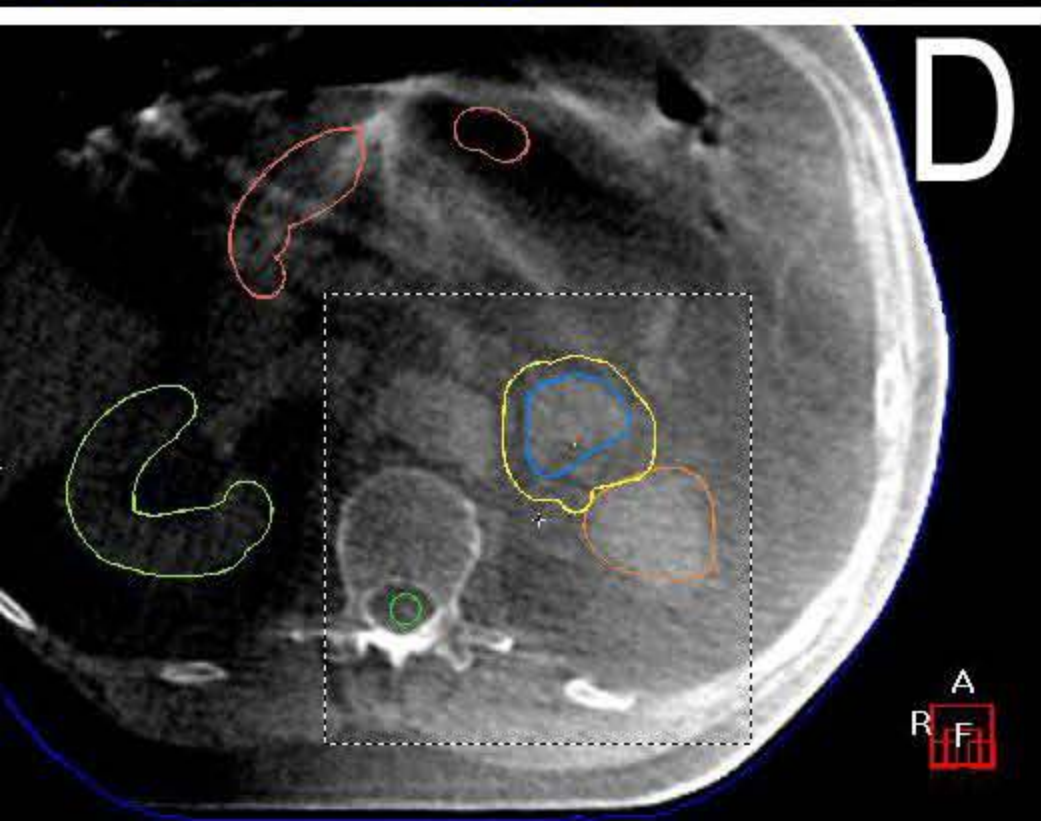
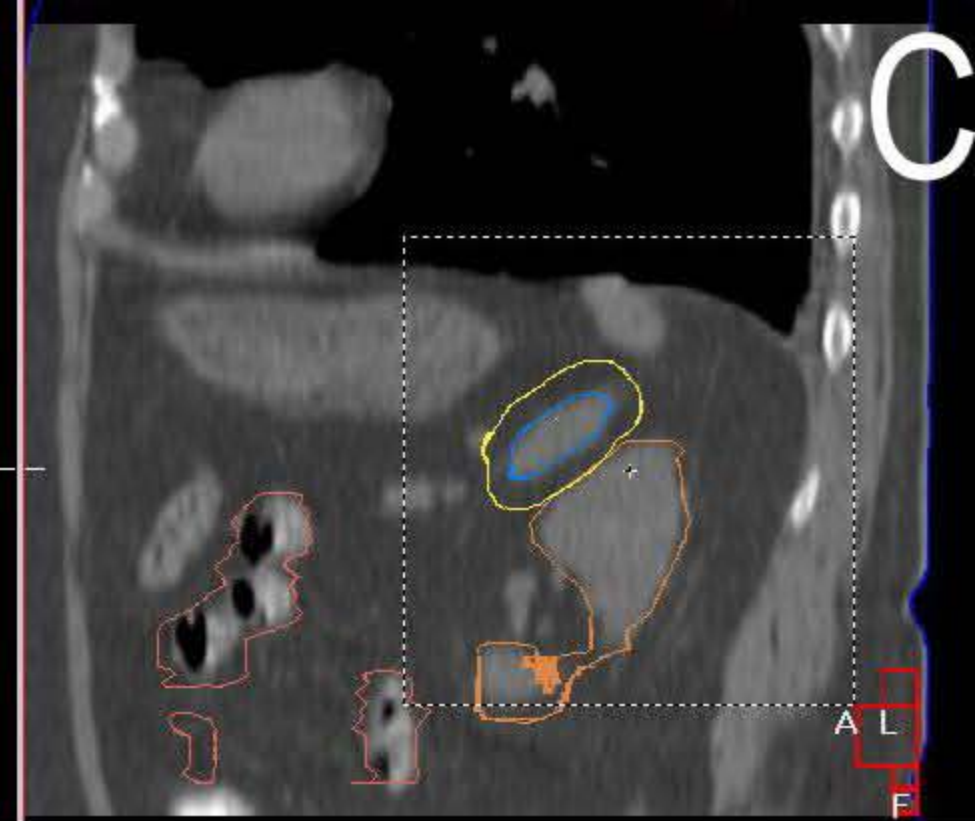
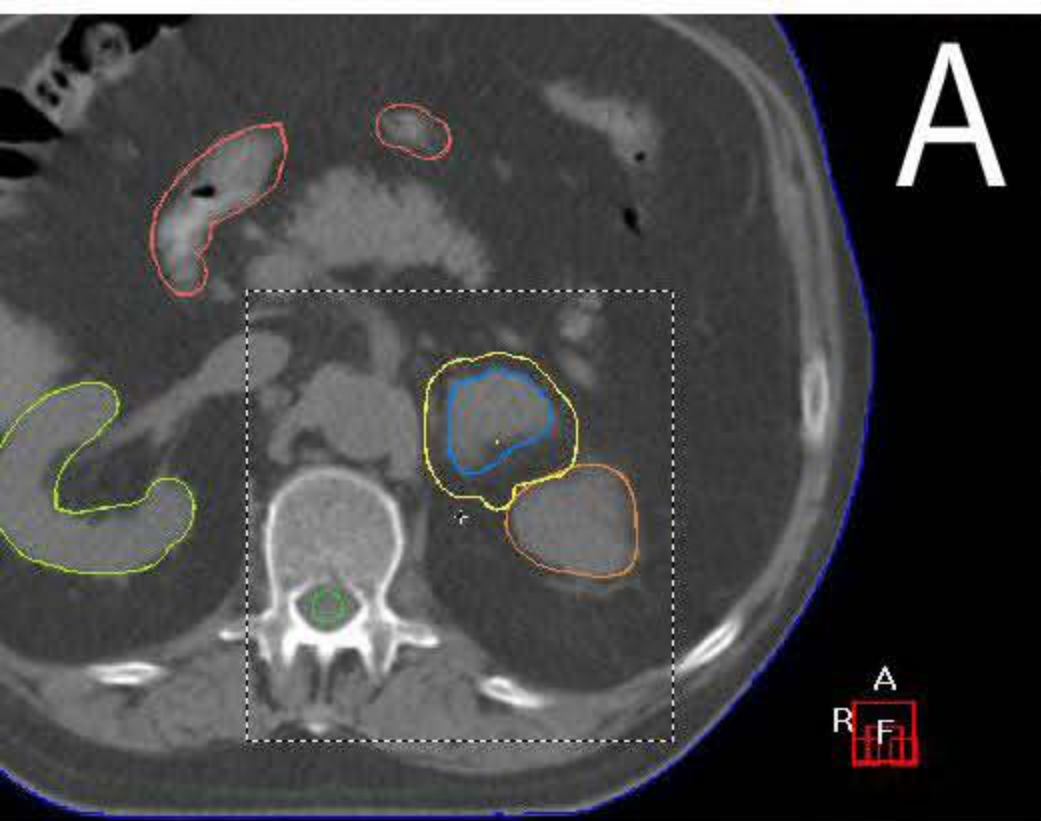


**Table 1.** Characteristics of DIBH treatments within the framework of advantages and disadvantages of currently available motion management strategies

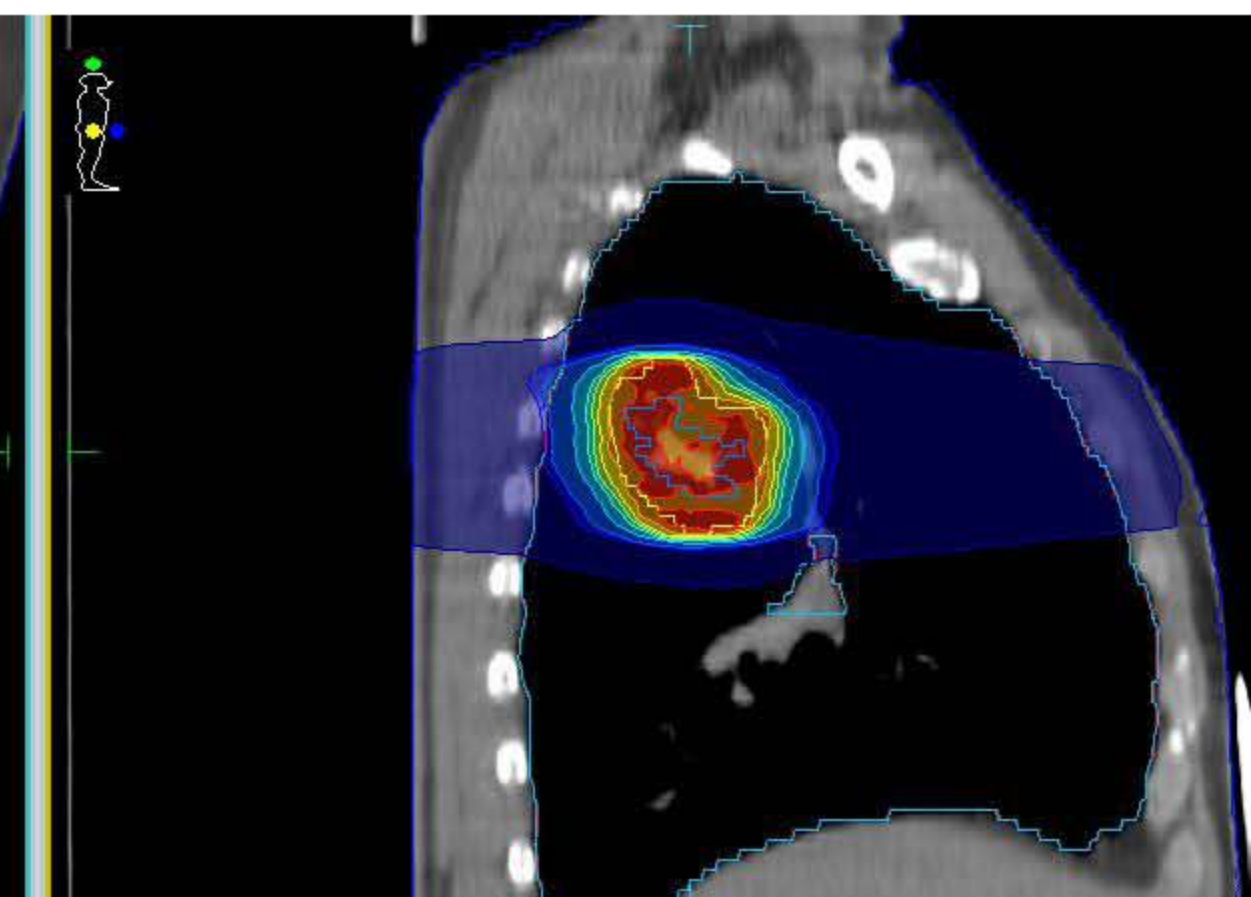
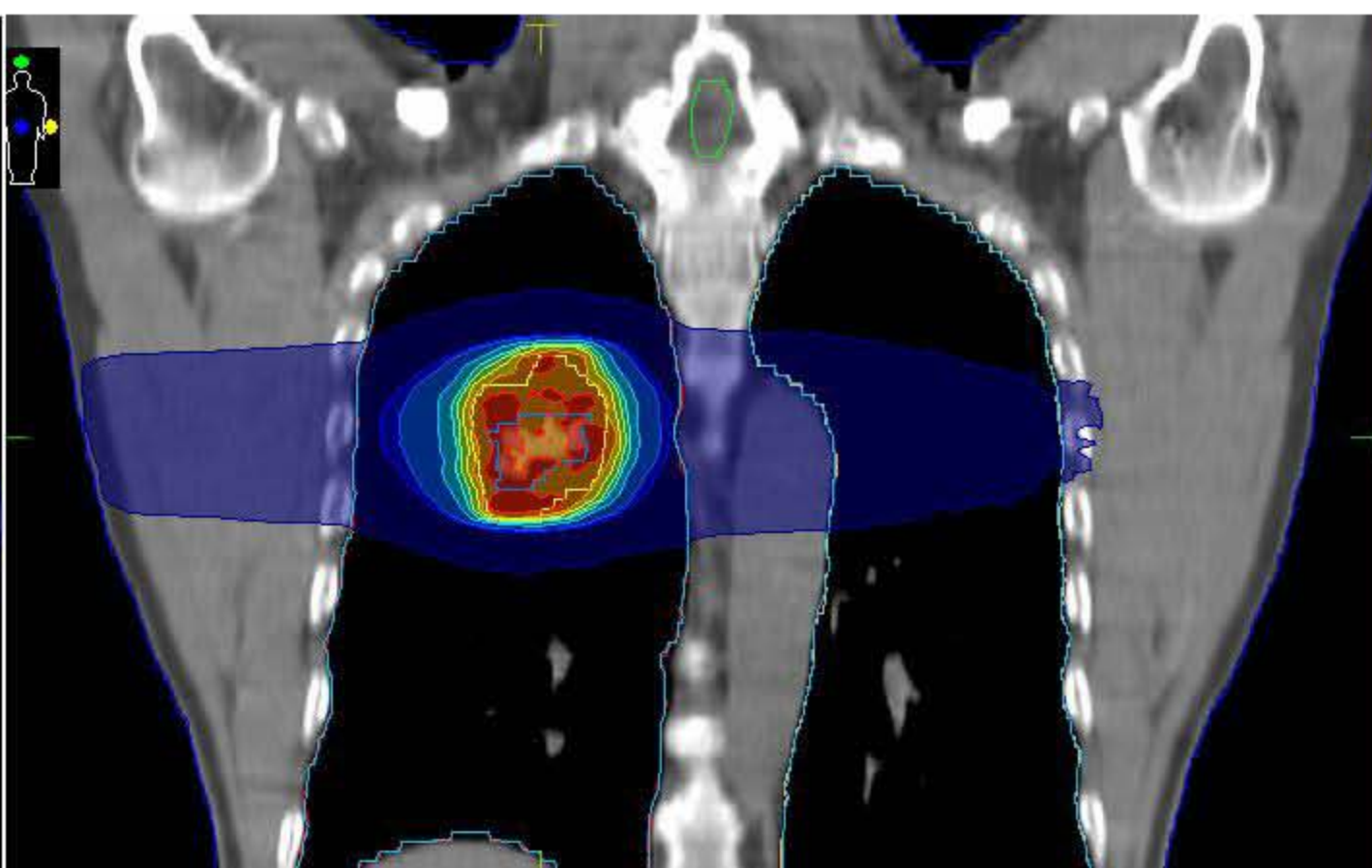
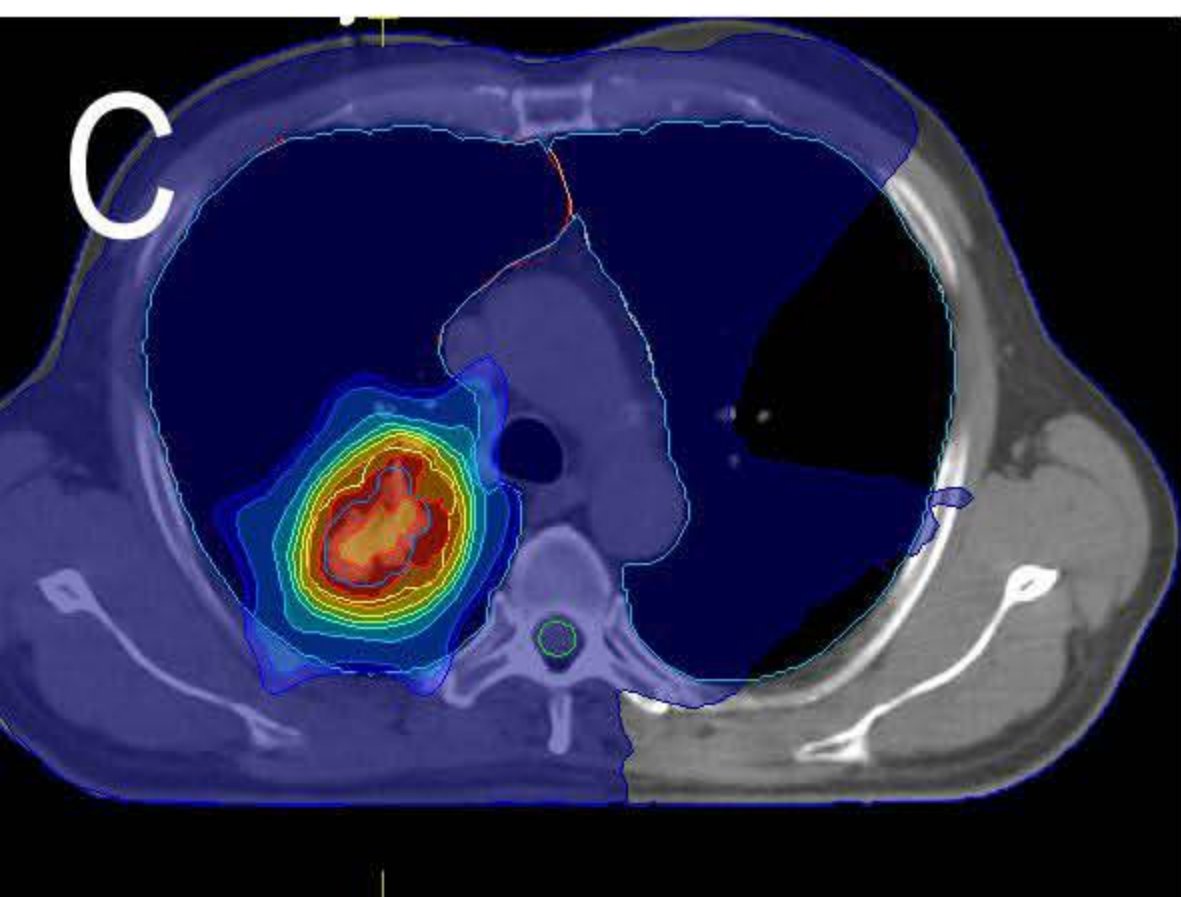
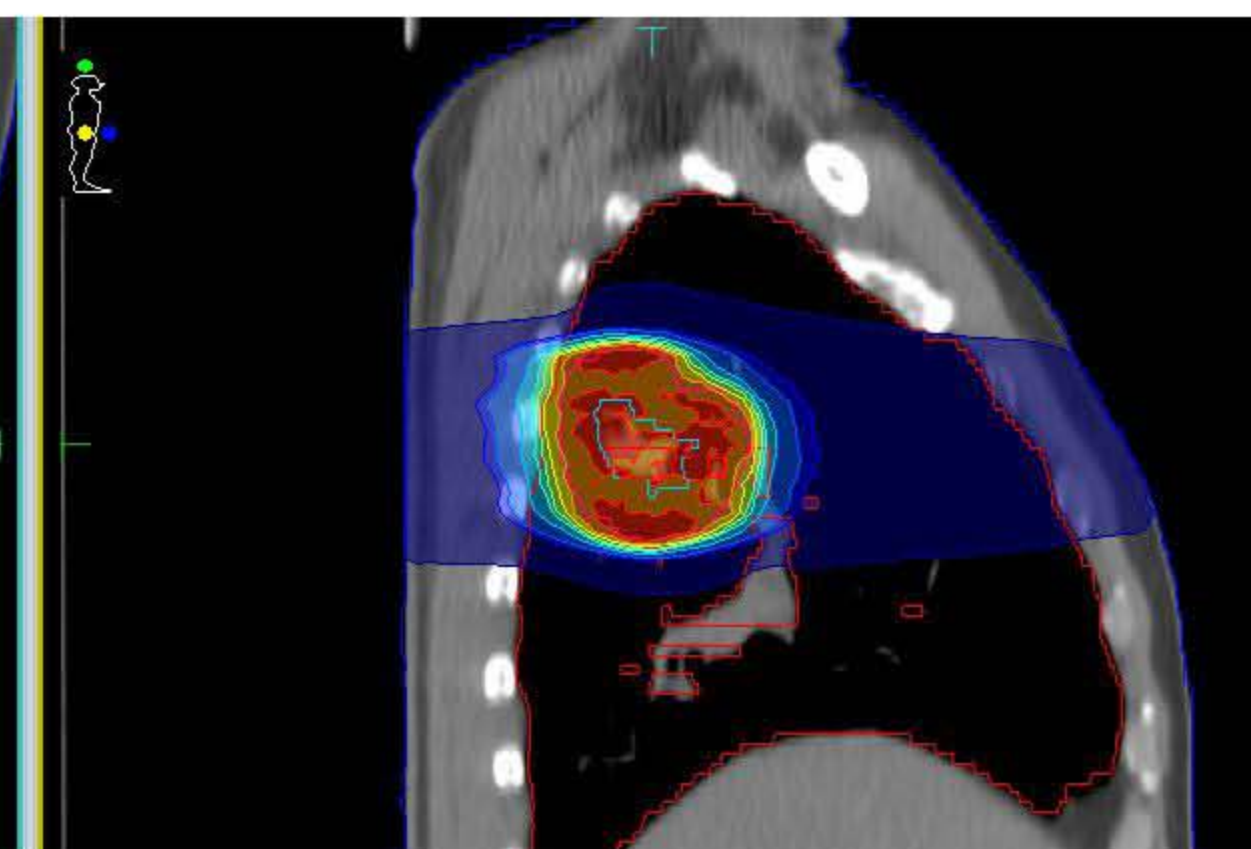
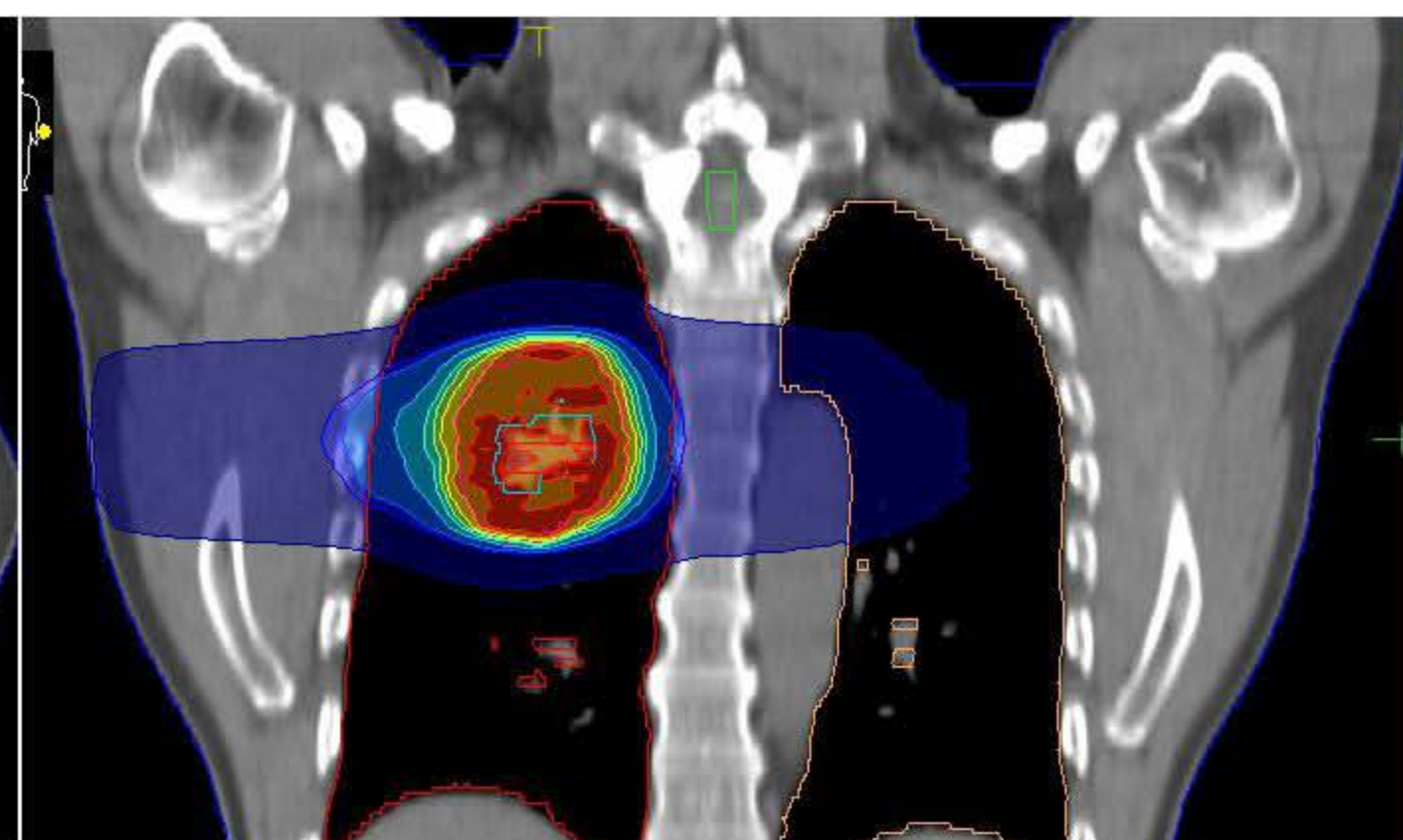
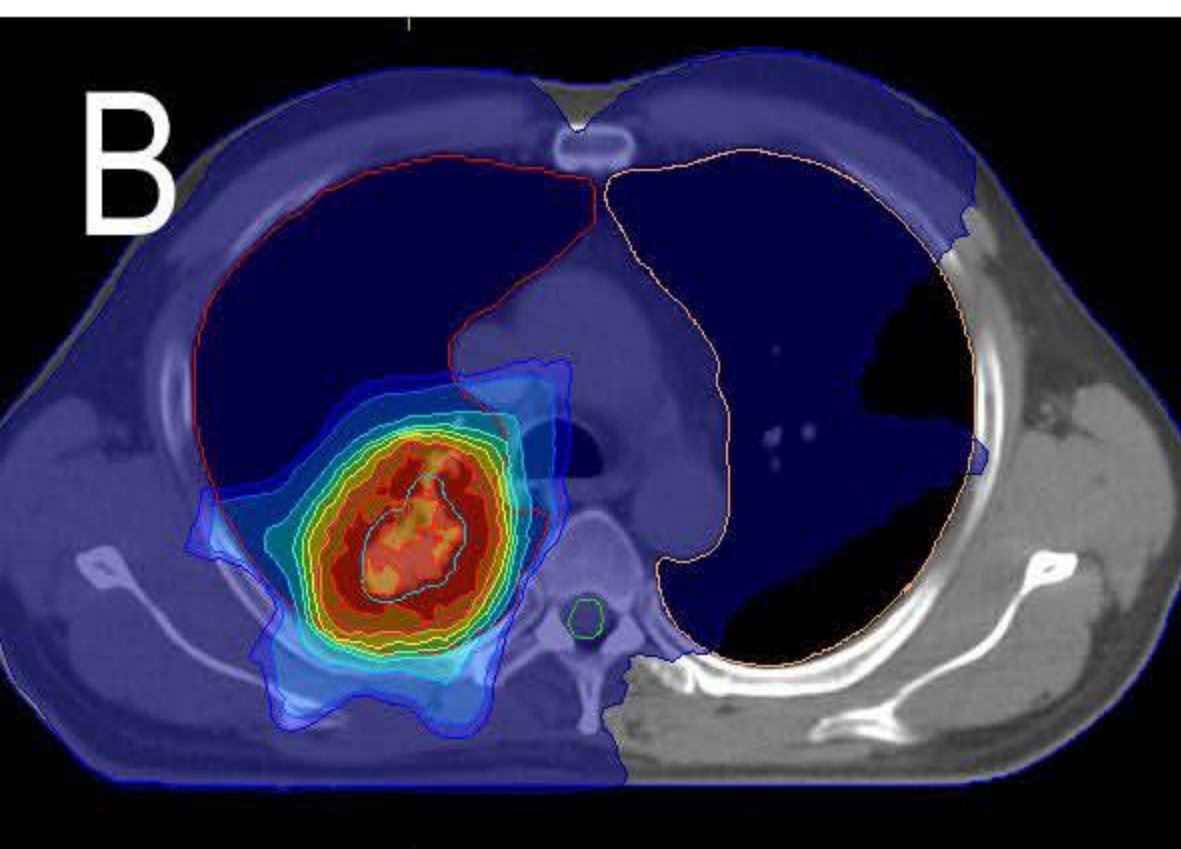
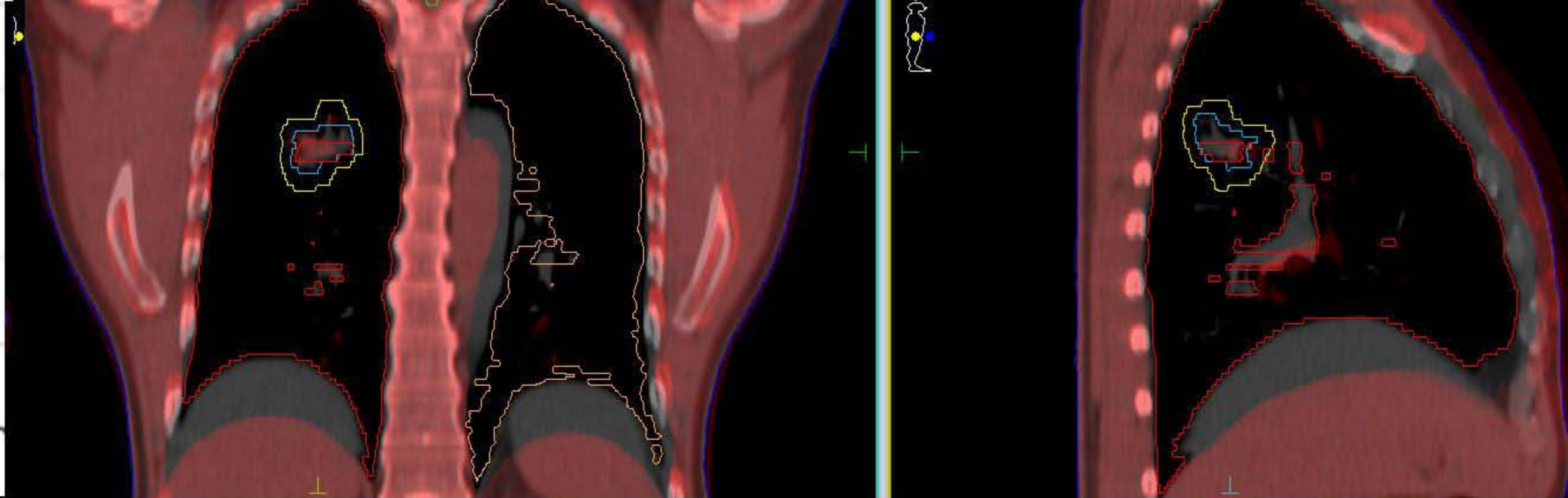
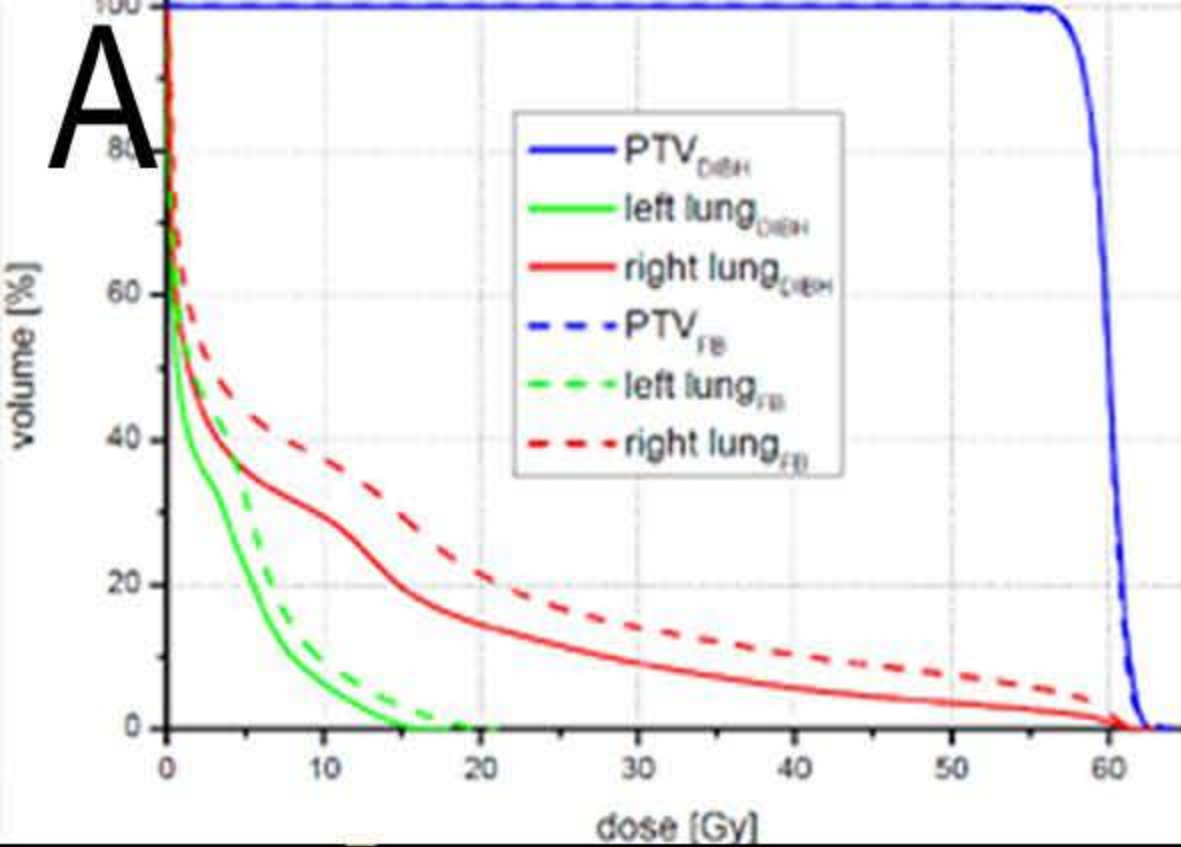
<b><u>Motion compensation method</u></b>	<b><u>DIBH</u></b>	<b><u>Spontaneous breathing gating</u></b>	<b><u>Real-time Tracking</u></b>	<b><u>ITV/individualized margins</u></b>
<b>Available techniques</b>	- free DIBH - computer-controlled DIBH (spirometry, surface tracking with markers or markerless)	Spirometry, surface tracking with markers or markerless	- couch tracking - steering of beam	Treatment planning with 4DCT, potentially with abdominal compression
<b>Imaging</b>	All imaging under DIBH: planning-CT, CBCT, ultrasound surveillance in breath hold, simultaneous VMAT-CT during treatment	- Dynamic planar or ultrasound imaging - 4DCBCT immediately before treatment - simultaneous VMAT-CT during treatment	-Dynamic planar or ultrasound imaging and VMAT-CT possible during treatment, depending on platform used	- 4DCT, 4DMR for treatment planning - 4DCBCT immediately before treatment - simultaneous VMAT-CT during treatment
<b>PTV margins</b>	Small (residual motion after breath hold)	Small (residual motion in gating window)	Small (tracking inaccuracy)	Large (end-expiratory-to-end-inspiratory position)
<b>Characteristics of achievable dose distribution</b>	- Reduced lung dose due to lung expansion and smaller PTV - Typically reduced cardiac dose and dose to most other OAR	depends on gating phase (inspiration: favourable; expiration: unfavorable)	-typically high exposure of lung and other OAR because treatment is performed during all breathing phases. Dose ideally has to be accumulated on a dynamic model.	typically high exposure of lung and other OAR due to treatment in all breathing phases and large margins. Dose ideally has to be accumulated on a dynamic model
<b>QA</b>	Standard treatment and imaging QA	Standard treatment and imaging QA	QA of the dynamic treatment process in addition to standard QA	Standard treatment and imaging QA
<b>Patient convenience</b>	- Optimal patient collaboration/compliance needed - Sufficient pulmonary reserve needed	Patient collaboration and regular breathing pattern needed	Patient collaboration and a sufficiently slow breathing pattern needed	frequently abdominal compression needed to reduce target motion
<b>Treatment time</b>	Longer treatment time	Longer treatment time	Short treatment time	Short treatment time
<b>Scanned Particle Therapy</b>	Minimal risk of interplay effects	Small risk of interplay effects	Small risk of interplay effects	Higher risk of interplay effects
<b>Toxicity</b>	For small lesions low for all techniques, for larger lesions no comparative data available, theoretical benefits for DIBH			













- Planning CT (DIBH, no frame, no rigid fixation, no abdominal pressure)  
↓
- Planning on single phase DIBH dataset.  
No need to contour on multiple breathing phases  
↓
- IGRT with CBCT acquired in repetitive DIBH  
↓
- Fast Delivery (DIBH, FFF, fast MLC) with static anatomical geometry identical to planning CT

