Advances in 4D Treatment Planning for Scanned Particle Beam Therapy – Report of Dedicated Workshops

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We report on recent progress in the field of mobile tumor treatment with scanned particle beams, as discussed in the latest editions of the 4D treatment planning workshop. The workshop series started in 2009, with about 20 people from 4 research institutes involved, all actively working on particle therapy delivery and development. The first workshop resulted in a summary of recommendations for the treatment of mobile targets, along with a list of requirements to apply these guidelines clinically. The increased interest in the treatment of mobile tumors led to a continuously growing number of attendees: the 2012 edition counted more than 60 participants from 20 institutions and commercial vendors. The focus of research discussions among workshop participants progressively moved from 4D treatment planning to complete 4D treatments, aiming at effective and safe treatment delivery. Current research perspectives on 4D treatments include all critical aspects of time resolved delivery, such as in-room imaging, motion detection, beam application, and quality assurance techniques. This was motivated by the start of first clinical treatments of hepato cellular tumors with a scanned particle beam, relying on gating or abdominal compression for motion mitigation. Up to date research activities emphasize significant efforts in investigating advanced motion mitigation techniques, with a specific interest in the development of dedicated tools for experimental validation. Potential improvements will be made possible in the near future through 4D optimized treatment plans that require upgrades of the currently established therapy control systems for time resolved delivery. But since also these novel optimization techniques rely on the validity of the 4DCT, research focusing on alternative 4D imaging technique, such as MRI based 4DCT generation will continue.

Key words: Particle therapy; Proton therapy; Beam scanning; Organ motion; Interplay; 4D.

Introduction

In this report of the 4th 4D treatment planning workshop held in Erlangen, Germany in December 2012 we want to review the recent progress that has been made in the field of mobile tumor treatment with scanned particle beams with an emphasis on the contributions discussed within the workshop.

Abbreviations: 3D: Three Dimensional; 4D: Four Dimensional, Time Resolved; 4DCT(MR): MRI Motion Trace Based 4DCT Generation; CN: Conformity Number; CNAO: Centro Nazionale di Adroterapia Oncologica; CT: Computed Tomography; CTV: Clinical Target Volume; DKFZ: German Cancer Research Center, Heidelberg, Germany; DVH: Dose Volume Histogram; GSI: GSI Helmholtzzentrum für Schwerionenforschung, Darmstadt, Germany; HIT: Heidelberg Ion Beam Therapy Center, Heidelberg, Germany; ICRU: International Commission on Radiation Units & Measurements; IMPT: Intensity Modulated Particle Therapy; ITV: Internal Target Volume; MRI: Magnetic Resonance Imaging; NIRS: National Institute for Radiological Sciences, Chiba, Japan; PCA: Principle Component Analysis; PCR: Phase-controlled Rescanning; PET: Positron Emission Tomography; PSI: Paul Scherrer Institute, Villingen, Switzerland / ETH, Zuerich, Switzerland; SIFT: Scale Invariant Feature Transform; SGSMP: Swiss Society of Radiobiology and Medical Physics.

*Corresponding author: Christoph Bert, Ph.D.
Phone: +49 9131 85 44213
E-mail: christoph.bert@uk-erlangen.de

Christoph Bert, Ph.D.1,2*  
Christian Graeff, Ph.D.2  
Marco Riboldi, Ph.D.3,4  
Simeon Nill, Ph.D.5,6  
Guido Baroni, Ph.D.3,4  
Antje-Christin Knopf, Ph.D.7  

1University Clinic Erlangen, Radiation Oncology, Universitätstraβe 27, 91054 Erlangen, Germany  
2GSI Helmholtzzentrum für Schwerionenforschung, Abteilung Biophysik, Planckstraße 1, 64291 Darmstadt, Germany  
3Dipartimento di Elettronica, Informazione e Bioingegneria, Politecnico Milano, P.za Leonardo da Vinci 32, 20133 Milano, Italy  
4Bioengineering Unit, Centro Nazionale di Adroterapia Oncologica, Strada Campeggi 53, 27100 Pavia, Italy  
5German Cancer Research Center, Im Neuenheimer Feld 280, 69120 Heidelberg, Germany  
6Joint Department of Physics, The Institute of Cancer Research and The Royal Marsden NHS Foundation Trust, Downs Road, Sutton, Surrey SM2 5PT, United Kingdom  
7Paul Scherrer Institute, 5232, Villingen, Switzerland / ETH, Zuerich, Switzerland
The workshop series started in 2009 at the Paul Scherrer Institute (PSI), Villigen, Switzerland. About 20 people attended from 4 institutes (GSI Helmholtz Centre for Heavy Ion Research (GSI), Darmstadt, Germany, German Cancer Research Center (DKFZ), Heidelberg, Germany, University of Tübingen, Tübingen, Germany, PSI). A special report was published as a result of the discussions in the plenary sessions of the workshop (1). It summarizes recommendations for the treatment of mobile targets with actively scanned particles and a list of requirements to elaborate and apply these guidelines. The paper was awarded with the Swiss Society of Radiobiology and Medical Physics (SGSMP)/Varian recognition award 2010. In 2010, the 2nd edition of the workshop was carried out at GSI with about 30 participants; the 3rd edition 2011 at the CNAO facility, Pavia, Italy already attracted more than 40 participants. To restore the original aim of the workshop, i.e., an informal platform to discuss current approaches, challenges and future research directions in 4D treatment planning, the participants agreed at the 4th edition of the workshop in 2012 at the University of Erlangen-Nuremberg, Germany (about 60 participants from about 20 institution including representative of commercial vendors), that in the future, the number of participants per institute will be limited to two persons. To stimulate discussions on still confidential research projects and to enable an open controversy about failures, representatives of commercial vendors will be excluded. In return, an annual report will be established to summarize all novelty. The next edition of the workshop is planned in November 2013 at PSI, Switzerland.

In the last years, the research on beam scanning for treatment of mobile tumors has finally led to first clinical applications, e.g., for hepato cellular tumors that are subject to respiratory motion (2). These clinical approaches can still be considered as first steps, i.e., research work has to continue, mainly in the fields outlined already in the previous report of Knopf et al. In that context, the scope changed from 4D treatment planning to complete 4D treatments, including appropriate imaging, motion detection, beam application, and quality assurance techniques suitable for the interplay prone irradiation of mobile tumors with a scanned beam.

The structure of the report is geared to the outlook chapter of the report summarizing the 1st workshop (1). We will focus on progress in 4D treatment planning options and in 4D treatment application techniques with an emphasis on contributions from participants of the workshop. For a broader overview on the field of the treatment of mobile tumors with scanned particle beam therapy please consult one of the following recent review papers (3-5). Even broader knowledge on cross-cutting aspects such as motion monitoring for treatment of mobile tumors exists in the photon treatment community; for details we refer the reader to review articles (6-8).

**Progress in 4D Treatment Planning**

**4D Dose Calculation**

4D dose calculation is an essential part of 4D treatment and has been first reported for proton therapy almost a decade ago (9). Also for scanned particle beams research codes capable of 4D dose calculations have been established at several centers in the last years (10-14), providing the foundation for more advanced research as mentioned below. The codes include handling of proton as well as particle beams and apart from analytical codes also Monte Carlo based solutions have been reported (15-17). The advanced codes enable 4D dose calculations taking into account different delivery parameters as well as numerous realistic patient motion scenarios (11, 18). Thus, simulation of different application techniques (e.g., active vs. passive energy modulation, rescanning vs. beam tracking) as well as the patient specific reconstruction of treatment deliveries is possible (see section 5.4).

4D dose calculation is not limited to intra-fractionally moving tumors but also inter-fractionally changing geometries such as prostate cancer was studied and presented at the workshops. The motion phases are then from multiple days and, e.g., represent different rectum filling. Another intermediate approach between 3D and full 4D treatment planning is the dose calculation on repeated breath hold CTs. The robustness of proton treatment plans for lung cancer patients against interfractional variations during voluntary breath hold was recently studied in a master project carried out at PSI (19).

Based on the activity of the last years, codes for 4D dose calculation are no longer the bottleneck for precise studies assessing the dosimetric outcome of treatment techniques or even delivered treatments. The challenge in the next years will be the transition from research codes into a commercially available product and the development of robust deformable image registration codes since vector field are an essential input to 4D dose reconstruction algorithms.

**Motion Modeling for 4D Planning**

As in 3D treatments, many of the 4D treatment planning approaches currently rely on the 4DCT acquired during treatment planning. This dataset represents a snapshot of the patient’s motion and many studies showed that the motion parameters change during the course of treatment and even within a few minutes (20-22). It is thus essential to exactly quantify the expected changes and their dosimetric influence and to develop solutions that overcome that clinical situation. In addition, the community needs to strive for better image guidance options which are still less advanced in particle therapy centers than in the photon treatment community despite the fact that particle treatments can physically
be delivered more precisely. Then, e.g., 4D Cone-Beam CTs could be acquired of an immobilized patient and ideally be used in an online, adaptive treatment option (23).

The issue of accurate motion modeling in 4D treatment planning has been addressed by investigating the effects of breathing irregularities in 4DCT (24). Higher correlation between external surrogates and internal lung motion was found for regular vs. irregular breathers, with prediction errors mostly dependent on the peak to peak range of motion. A novel 4D CT resorting technique based on the use of multiple surrogates has been also investigated: results show that monitoring of multiple surrogates can handle the task of breathing phase detection more accurately, resulting in reduced image artifacts in presence of limited breathing irregularities (25).

To overcome the restriction of snap shot 4DCTs and to reduce imaging dose, PSI investigated, if 4D magnetic resonance imaging (MRI) data can be the basis for 4DCT generation, named 4DCT(MRI) (12). 4DMRI can be acquired over several minutes since no ionizing radiation is applied. Boye et al. extracted motion vectors from the 4DMRI and wrapped them to stationary 3DCT to derive 4DCT(MRI)s with the help of motion modeling. 4DCT(MRI)s obtained in this way represent motion behavior over several breathing cycles including variations in amplitude and breathing frequency as well as baseline drifts. Thus, no assumptions on breathing regularity have to be made during 4D treatment planning. This enables the investigation of treatment plans against motion variations. Furthermore, motion variations can be considered in treatment plan optimization in order to obtain 4D optimized plans that are robust against expected variation such as phase shifts between motion surrogate and the actual tumor motion. 4DCT(MRI)s can further be used to support motion monitoring, as described in section 5.1.

**Evaluation of Different Deformable Registration Methods**

4D treatment planning heavily relies on deformable image registration. Numerous codes are available from vendors and within the research community. Since the application of those codes varies, also the research on their validation does. In a cross-center study, Brock et al. studied various codes for typical radiotherapy sites with respect to accuracy and reproducibility (26). They report large discrepancies in the reported shifts with a majority of the codes performing at voxel size level of the underlying dataset.

Among the workshop participants, an automatic feature detection was proposed relying on the Scale Invariant Feature Transform (SIFT) method for validation of deformable image registration (27). The automated detection results are comparable to expert-user-based detection, and is applicable to both 4D CT and 4DMRI datasets (28). The method was studied in adaptive radiotherapy and showed that SIFT based metrics are correlated to detected anatomical changes over the course of treatment. Also the use of regularization methods in deformable image registration was analyzed, as a way to increase the physiological consistency of the quantified deformation field (29). This has been applied to head & neck treatments and requires extension to mobile sites, in order to establish optimal regularization parameters.

A detailed study with respect to 4D dose calculation in scanned proton beams has been reported by Zhang et al. (18). For single-field treatments, where no motion mitigation was used a maximum (mean) dose difference (averaged over three cases) of 32.8% (2.9%) was observed with regards to the use of different deformable registration algorithms to extract motion information from 4D images. This registration ambiguity-induced uncertainty indicate the necessity to interpret 4D dose distributions for scanned proton therapy as approximation, inevitably bonded with error bars. Quantification and presentation of deviations in 4D treatments is an essential topic in itself. Not only the registration quality but also many of the other parameters such as internal-external correlation of motion monitoring devices or assumptions in the treatment application technique are prone to uncertainties. Hild et al. proposed different quantification options on the basis of 4D treatment plans for lung tumors (30).

Lüchtenborg used different registration options in a treatment planning study assessing beam tracking (31). They report, that rigid registrations should be used for calculation of beam tracking parameters since distances are preserved which is essential if over/under-doses due to changes in spacing of Bragg-peaks should be avoided.

**4D Treatment Plan Optimization**

4D treatment plan optimization can be classified as the explicit incorporation of motion data into the cost function of the optimization. It has been reported especially in the photon community since several years (32) also within the context of our workshop series, which included the studies of Suh et al. (33, 34).

The potential of 4D optimized treatment plans are one of the results of a recent study of Knopf et al. (35). They studied the consequences of different beam weight distributions when treating mobile targets. For static targets, beam weights were optimized in order to achieve best target dose conformity. This usually results in many low weighted spots and a few high weighted spots at the distal edge of the target. For the treatment of mobile tumors target dose conformity is compromised by blurring, thus different beam weight optimization objectives might be considered. In their preliminary study it was shown that treatment plans with a “smooth”
beam weight distributions are significantly more robust when treating moving targets with scanned proton beams (36). By designing appropriate robustness constraints, such smooth distribution could be the outcome of a treatment plan optimized in 4D.

Eley et al. investigated the possibilities of 4D optimization in beam tracking with a scanned carbon beam (37). They investigated for an artificial geometry as well as for a lung tumor patient, if 4D optimized treatment plans could reduce the dose to nearby organs at risk. The GSI in-house treatment planning system was thus extended to fully incorporate 4DCTs and deformation maps in the optimization process. They found comparable target coverage for both cases and a reduced maximal dose to the heart as organ at risk.

The work is complemented by Graeff et al., who developed a more general framework for 4D optimization based on subdividing the target volume to ease the technical demands in the optimization. This is promising or even essential, since memory and calculation demands might be too high, if all (typically ∼10) motion states are incorporated without further constraints. The sub-sections of the target volume are chosen with respect to the delivery constraints. The authors showed for nine lung cancer patients that highly conformal target dose distributions can be achieved without detrimental inverse interplay patterns in the entrance channel which are typically observed in beam tracking plans (38). An alternative approach is reported by Graeff et al. in this issue.

As in 4D dose calculation, also 4D optimization still relies on precise 4DCT and deformation vector field data. Appropriate procedures and/or techniques have to be established and assessed prior clinical use (see also section 4.2).

**Experimental Validation of 4D Treatments**

Experimental validation of implemented 4D treatment options is essential since the technical demands of the treatment delivery systems are often much higher than for 3D treatments and can thus be the bottleneck of potential solutions. Validation requires adequate motion phantoms with detection systems. A list of phantom features was reported in the previous workshop report (1).

Each group performing experiments focusing on moving targets uses motion platforms, either commercial solutions or in-house built. These devices allow typically a one-dimensional (translational or rotational) motion which is not representing a patient geometry but often sufficient for an initial study of a new technique. It is then the preferred technique due to easy handling but investigators have to keep in mind that study results might not be transferable to patient geometries. The platforms can carry radiographic films (40), scintillation detectors (41), biological probes (42), probes for positron emission tomography (PET) measurements (43), or water phantoms with ionization chambers (44).

Recently, developments focused on complex phantoms that should mimic the patient geometry. Zakova et al. together with the Centre Suisse d'Electronique et de Microtechnique (CSEM) currently develop an anthropomorphic thorax phantom which is completely metal-free and CT as well as MRI compatible (45). Motion can be controlled by air inflation into an air-tight lung, which is surrounded by a realistically expanding rib cage. The dose distribution can be recorded with an ionization chamber or Gafchromic films placed directly into a tumor moving within the lung compartment. First irradiation tests with a scanned proton beam showed that the phantom allows extensive dosimetric studies under realistic circumstances (45). Steidl et al. report about a robot based thorax phantom which is used for validation studies in scanned carbon beams (46). A robotic arm is used to move a lucite block mimicking the tumor and equipped with radiographic films and 20 ionizations chambers. The tumor can move in 6D and motion is in correlation with an independent thorax phantom that is based on a plastic skeleton covered by rubber representing the skin. Initial tests with and without beam have been successful (46) and the phantom has since been used in a number of 4D validation studies (11, 47).

On the experimental side, the feasibility and efficacy of the advanced scanning techniques was tested systematically in numerous studies. Rescanning (41), gating (48), beam tracking (44, 49) as the main motion mitigation techniques, but also the validity of 4D treatment planning systems (11, 38, 50)
have been studied. More details will be reported in section 5 covering the mitigation techniques itself.

Validation further includes in vivo dosimetry. An established method for ion beam therapy is the in-beam and offline use of PET. Carbon-11 and C-10 that result of fragmentation of the primary C-12 beam serve as positron emitters (51). In the last years the Helmholtz-Centre Dresden Rossendorf (HZDR), GSI and HIT worked on 4D extensions for in-beam PET within the EU funded projects ULICE and ENVISION. Laube et al. reported on simulations, reconstruction methods and experimental results for 4D in-beam PET (43, 52). The proposed methods are suitable to judge the relevant parameters for treatments of intra-fractionally moving tumors with a scanned carbon beam. Initial steps towards routine implementation at the HIT facility have been successful (17).

Activities based on 4D PET further focused on the use of 4D CT motion models to optimize 4D PET imaging (53, 54). Such an optimization can either be applied to treatment planning 4D PET/CT studies or to post-irradiation PET imaging (PET-based dosimetry), in order to make the most of the reduced count statistics induced by particle irradiation.

Despite all efforts, validations will always lack the clinical scenario and typically focus on one special technique (e.g., motion phantoms to validate motion monitoring systems that do not allow dosimetric quantification or assessment of deformable image registration). Thus, careful introduction of new techniques into clinical application has to follow despite potential shortcomings accompanied by, e.g., the proposed in vivo dosimetry techniques and stringent follow-up of the patients.

**Progress in Beam Application Techniques**

Treatment of intra-fractionally moving organs will require dedicated means if scanned beams are chosen as treatment technique. In the context of this manuscript we will refer to all procedures as techniques and report the current status of the involved groups. Many of them require precise motion monitoring and often also margins forming the internal target volume (ITV) from the clinical target volume (CTV).

**Motion Monitoring and Motion Prediction**

Volumetric methods such as 4DCT or 4DMRI are used as part of motion modeling in treatment planning to describe the anatomy for dose calculation and treatment plan optimization. During treatment delivery these methods are not (yet) available, despite first approaches to combine photon linacs and MRI exactly for this purpose (55). Thus, 1D motion signals or motion surrogates are frequently used for the purpose of detecting the 4DCT state or the 3D position of a marker in real-time during treatment delivery (3, 56). Especially for precise treatment techniques like beam tracking (section 5.5), surrogates are not ideal due to potential miscorrelation to the internal targets. Vice versa, purely fluoroscopic based detection of radio-opaque fiducials offers precise motion information, but results in additional x-ray doses (57) even though these can be at dose levels comparable to other image guidance options (58). Thus, alternative techniques have been studied.

In order to precisely track tumor motion online it is essential to obtain information on the 3D motion vector throughout the region of interest. Any sparsely acquired surrogate motion is generally not sufficient to describe the deformable behavior in three dimensions. In a recent study, it has been shown that 3D deformable motions can be estimated from surrogate motions obtained from either BEV or dual X-ray imaging systems for treatments in the liver (59, 60). The method requires motion sampled over a number of breathing cycles for each patient before treatment using some form of 4D imaging, for example 4DMRI. On the base of this motion library a Principle Component Analysis (PCA) can be applied to build subject specific motion models. Motion models based on markerless surface detection have also been explored, relying of deformable surface registration to achieve accurate motion monitoring of specific anatomical landmarks (61).

3D real-time data without ionizing radiation can be measured by ultrasound as shown by several groups for interfractional motion assessment (organ positioning) (56). For intra-fractional motion monitoring, Jenne et al. reported the use of ultrasound at the 4th workshop in Erlangen. By means of dedicated transducers a 2D plane of the patient can be scanned. Either the tumor is visualized directly, or internal surrogates such as the diaphragm are used. If combined in two directions, pseudo-3D data are achieved. The feasibility of ultrasound based beam tracking has been reported by Prall et al. (62). They showed that delay compensation is possible via neural networks and present experimental data indicating the feasibility of ultrasound based compensation without tracking parameters from treatment planning.

At DKFZ motion monitoring using the Calypso-System (Medical Systems Inc., Palo Alto, CA) for internal prostate motion and more recently also for lung tumor motion was studied (63). The system uses field generating coils to be implanted inside or close to the tumor whose position is detected by an electromagnetic detector array. Thus no additional radiation dose is applied to the patient and real time tumor motion tracking is feasible (64, 65). Recent attempts are trying to use the system also for proton therapy (66). In case of tumor tracking accurate and real-time tumor motion detection is required. To overcome the problem of system latencies additional emphasis has been put on the evaluation
of motion prediction algorithms for real-time tumor tracking. The study of Krauss et al. emphasized the relative importance of adequate model parameter optimization compared to the actual prediction model selection (67).

A combination of internal motion detection and surrogates can be achieved by dedicated correlation models. The group at Politecnico Milano studied several options, focusing on the accuracy of different correlation models by retrospective clinical data analysis (68-70). The issue of adaptive modeling and robustness of the correlation function of controlled breathing irregularities has been quantified (69, 70). To show feasibility within the scanned particle therapy framework, model based motion detection has been used for beam tracking at GSI (47) (see also section 5.5 and Fattori et al. in this issue).

An alternative technique could be particle radiography which has been proposed for different purposes for decades (71, 72). With current technology, radiography as well as tomography is possible (73, 74), but so far not used for intra-fractional target motion detection. Preliminary work has been carried out to study the potential of particle radiography to monitor soft tissue motion (75, 76). In these studies, prior knowledge represented by the treatment planning CT is used to enhance the soft tissue contrast, so that particle radiography can be optimized for motion detection in soft tissue targets, avoiding the use of implanted surrogates to reach adequate accuracy.

**Margin-based Approach**

Intra-fractional motion is typically dealt with by using margins surrounding the CTV. ICRU report 62 (77) advises that variations in size, shape and position of CTVs relative to anatomic reference points can be considered for ITVs. In addition to geometrical margin adaption, changes of water equivalent path length have to be considered for particle therapy, as already mentioned in the proton report of the ICRU (78). These considerations are applied since several years in passively shaped particle therapy but are often based on, e.g., overwriting of CT-numbers in the planning CT (79).

A number of years ago, Engelsman et al. (80) proposed to use 4DCT as basis of ITV definition in scattered proton beam therapy. The work has been implemented for scanned beams as well (81, 82) but the original implementation is limited to single-field uniformal dose approaches because of field-specific mapping of motion induced changes in particle range and uses out-dated dose calculation models. A full consideration that is also applicable to intensity modulated particle therapy (IMPT) has been reported by Graeff et al. (83). They transform the geometrical ITV into a field-specific water-equivalent path length ITV and use several motion phases to model the motion depended shape of the range-adapted ITV. The proposed method has been tested successfully on the data of a lung cancer patient.

A recent study by Knopf et al. shows that CTVs significantly differ in size from geometrical ITVs and range adapted ITVs (35). Furthermore, range adapted ITVs and geometrical ITVs differ significantly in size and are spatially displaced, particularly for lung patients. Range-adapted ITVs show a strong field dependency in shape.

**Rescanning**

One way to overcome interplay effects are multiple irradiations per treatment fraction, referred to as rescanning (84). For acceptable treatment times this approach requires high scan speeds and thus also fast beam monitoring systems. In case of treatment plan application times within minutes, also (multiple) breath-hold based approaches are then feasible. There exist many possible rescanning strategies and naming schemes (scaled/slice-by-slice/level vs. iso-layered vs. volumetric/uniform, (85, 86)). In simulations by Zenklusen et al. (86) it was shown that continuous line scanning seems to be the most elegant solution: it provides higher repainting rates and produces superior results but is probably more difficult to realize. Recently the effectiveness of volumetric and slice-by-slice rescanning was investigated in relation to slow and fast beam delivery systems by Bernatowicz et al. (87). Similar effect to rescanning can be achieved with multiple fields per plan (10) or by using dedicated fractionation schemes (88).

Rescanning is a strong approach to mitigate interplay effects, but it does not address dose blurring due to motion. Therefore, it is believed that best results can be obtained by combining rescanning with other motion mitigation techniques and/or margins. At NIRS, most likely rescanning will be combined with gating and referred to as phase-controlled rescanning (PCR) (89). They reported several technical and simulation studies in the last years and are close to treat the first patient. Also at PSI rescanning alone is only seen as first level of motion mitigation. As a next step it is foreseen to use gated rescanning or slow tracking (35, 60). The second refers to a treatment during repeated breath holds with an adaption of the beam position for possible variation between different breath holds. As ultimate solutions 4D optimized rescanning or beam tracking and retracking (90) is envisioned.

**Gating**

Gating has been the method of choice for lung and liver treatments with a scattered carbon beam at NIRS for >10 years (91). Based on motion monitoring data (section 5.1), the beam is only turned on in a defined part of the breathing cycle, typically at end-exhale. Thus, the effective motion
amplitude is reduced. Also for scanned beam delivery this approach will reduce the interplay effects, but gating alone will not result in homogeneous CTV coverage since also reduced amplitudes induce interplay effect. NIRS thus combines gating with rescanning (PCR, see previous section). An alternative seems to be irradiation with increased beam overlap, e.g., larger beam spot sizes at identical raster grid spacing as for stationary tumors (48). Based on numerous simulations and experimental studies of GSI and HIT (92, 93) that included assessment of the sensor delays (94) and dosimetric verification (95), gating was recently introduced clinically for the treatment of hepato cellular cancer at HIT (2). Apart from gating, mainly abdominal compression was used to control the motion extent of the patient, but the required mitigation approaches are similar since in each case the interference effect of residual motion amplitudes needs to be compensated. Richter et al. recorded the motion surrogate trace and the scanner progress data during treatment delivery of each fraction and used the GSI in-house 4D treatment planning system to reconstruct the delivered 4D dose distribution for each fraction and for the complete treatment. Even though the focus of that study was feasibility of the proposed method, they could show appropriate CTV coverage for a number of patients (96).

**Beam Tracking**

Beam tracking uses data from motion monitoring and potentially 4D treatment planning to compensate target motion by adaptation of the scanned pencil beam (49, 97). Previous studies concentrated on simple phantom geometries and had the goal of feasibility checks since the application and therapy control needs are challenging. With respect to system performance (98) but also dosimetrically (49, 99, 100), beam tracking works precisely. In case of non-translational motion, e.g., rotations, adjustment of the pencil beam position is insufficient since the pre-irradiation mainly of proximal parts of the target by distal iso-energy layers changes. Luchtenborg et al. studied this influence and proposed an adaption of deposited particle numbers combined with real-time calculations as solution (44).

In 2012, experiments were conducted and reported at the 4th 4D treatment planning workshop that extended from simple phantoms to complex, patient-like geometry. Scenarios based on the robotic thorax phantom presented in section 4.6 were studied using correlation model based motion monitoring solutions from the Politecnico Milano group (see section 5.1). The experiments included baseline drifts of the tumor and correlation mismatch between thorax surface and tumor which will cause dose deviations if beam tracking delivery is fully based on the optimized 4D treatment plan. Thus, in the lateral plane, compensation offsets were directly determined in the motion monitoring system which can therefore compensate changes with respect to the treatment planning 4DCT scan such as baseline drifts. Integration of the system (see Fattori et al., this issue), motion correlation models (47), and also the dosimetric outcome worked as expected. Beam tracking is thus on the verge to commercial implementation or research based clinical assessment.

**Delivery of 4D Optimized Treatment Plans**

4D optimization incorporates the different phases of the 4DCT data in treatment plan optimization. Apart from dedicated ITV concepts, these algorithms can also result in a 4D treatment plan, i.e., parameter files that are dependent on the motion phase of the patient. Such plans require a dedicated treatment control system which is more complex than for 3D treatments and of similar but different complexity to particle number compensated beam tracking. Such a control system has been implemented at GSI and its feasibility has been shown for different flavors of 4D optimization.

For both 4D optimization methods described above, the treatment plan consists of a set of 3D plans to be delivered to specific motion phases. The treatment control system thus switches between these plans according to a motion monitoring signal. Depending on target geometry and the irradiation sequence determined by the target motion, some motion phases do not or no longer require irradiation. In this case, the beam is gated during this motion phase. Smooth delivery thus requires fast, multiple gating during a spill. Delivery can be greatly facilitated by intensity control and flexible spill timing.

It should be noted that for these methods, the sequence of beam spots is determined online by the actual measured motion. A specific scanpath as for 3D delivery can thus not be preplanned. For both methods, the feasibility of delivery was shown in a film experiment (38).

**Concluding Remarks**

Scanned particle beam application to intra-fractionally moving tumors are still challenging, but the developments reported within the scope of the four 4D treatment planning workshops and on other occasions show that methods and techniques have been developed that will allow safe treatment deliveries at several centers with a strong research focus in the near future. First patients have already been treated at HIT with a scanned carbon beam and gating or abdominal compression as motion mitigation technique and at Rinneacker Proton Therapy Center in Munich, Germany using apnea. PSI will rely on rescanning and potentially gating, NIRS plans to mimic scattered beam applications by rescanning used in combination with beam gating as in the current clinical practice.
The achievements reported above are predominantly reported by and implemented in centers with a strong research focus. One of the main challenges will be the transition of those ideas into medical products that vendors can offer to all facilities of the meanwhile fairly broad particle therapy community using beam scanning. While many of the new machines, e.g., can provide fast beam gating, there is typically no 4D treatment planning functionality available that would support parameter choices based on assessment of the expected 4D dose distribution. One potential reason for that slow transition into products could be the zoo of options reported from the research community with no clear component portfolio visible yet. For each of the required technologies (motion monitoring, motion mitigation, treatment planning, …) there are several options available, some of them even in clinical use, such as correlation model based motion monitoring, as in the Cyberknife Synchrony or the Brainlab/Mitsubishi Vero system, but the required or at least beneficial integration into a particle therapy center is not yet available. Many of those options were developed for photon radiotherapy, another indication that the particle therapy community still applies other standards than the modern photon therapy techniques. Similar for 4D verification systems and a quality assurance workflow, which was not covered in sufficient detail in this workshop series but which will require new ideas due to the random and thus not really predictable manner of interplay effects. That integration might speed up in case of standardization, i.e. defined protocols for, e.g., the beam delivery sequence or the motion monitoring signal such that different 4D treatment planning systems could import the data for calculation of the delivered 4D dose distribution.

From a research perspective there is still the need for precise and three dimensional motion monitoring, ideally including the range of the particle beam. Potential solutions, such as 4DCT(MRI) in combination with a surrogate and a PCA based model have been proposed recently (12, 60) and also 4DPET or prompt-γ imaging might allow real-time options at some point. Ideally, such precise imaging options have to be combined with (real-time) treatment plan adaptation algorithms to allow precise treatments of moving organs even in hypofractionated or even single-fraction treatment schedules (stereotactic body particle radiosurgery). The management of variable uncertainties including interplay effects in fewer fractions becomes even more challenging since the fractionation itself will not lead to mitigation of dose inhomogeneities (10, 101, 102). Still, such fractionation schemes proved to be effective in clinical studies at NIRS (103) and thus wide-spread use can be anticipated not only because they are preferred by many patients.

For the still pretty young workshop series and future generation of researchers these challenges are motivating and the clinical results of the first patients treated with the advanced mitigation techniques mentioned within the scope of this report will certainly trigger new ideas. The next option to exchange experiences and details within that field will be the 5th workshop, taking place at PSI in Villigen, Switzerland on November 28/29 2013. Interested colleagues – especially radio-oncologists – can contact one of the authors.

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