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PII:	S0969-8051(21)00531-X
DOI:	https://doi.org/10.1016/j.nucmedbio.2021.12.002
Reference:	NMB 8250
To appear in:	Nuclear Medicine and Biology
Received date:	17 February 2021
Revised date:	4 November 2021
Accepted date:	9 December 2021

Please cite this article as: I. Bastiani, S.J. McMahon, P. Turner, et al., Dose estimation after a mixed field exposure: Radium-223 and intensity modulated radiotherapy, *Nuclear Medicine and Biology* (2021), https://doi.org/10.1016/j.nucmedbio.2021.12.002

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# Dose estimation after a mixed field exposure: Radium-223 and intensity modulated radiotherapy

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#### <u>Abstract</u>

#### Introduction

Radium-223 dichloride ([<sup>223</sup>Ra]RaCl<sub>2</sub>), a radiopharmaceutical that delivers α-particles to regions of bone metastatic disease, has been proven to improve overall survival of men with metastatic castration resistant prostate cancer (mCRPC). mCRPC patients enrolled on the ADRRAD clinical trial are treated with a mixed field exposure comprising radium-223 (<sup>223</sup>Ra) and intensity modulated radiotherapy (IMRT). While absorbed dose estimation is an important step in the characterisation of wider systemic radiation risks in nuclear medicine, uncertainties remain for novel radiopharmaceuticals such as <sup>223</sup>Ra.

#### Methods

24-colour karyotyping was used to quantify the spectrum of chromosome aberrations in peripheral blood lymphocytes of ADRRAD patients at incrementar times during their treatment. Dicentric equivalent frequencies were used in standaro models for estimation of absorbed blood dose. To account for the mixed field nature of the treatment, existing models were used to determine the ratio of the component radiation types. Additionally, a new approach (M-FISH<sub>LET</sub>), based on the ratio of cells containing damage consister which high-LET exposure (complex chromosomal exchanges) and low-LET exposure (simple exchanges), was used as a pseudo ratio for <sup>223</sup>Ra:IMRT dose.

#### Results

Total IMRT estimated doses delivered to the blood after completion of mixed radiotherapy (after 37 IMRT fractions and two [ $^{223}$ Ra]RaCl<sub>2</sub> injections) were in the range of 1.167 ± 0.092 and 2.148 ± 0.096 Gy (dose range across all models applied). By the last treatment cycle analysed in this study (four [ $^{223}$ Ra]RaCl<sub>2</sub> injections), the total absorbed  $^{223}$ Ra dose to the blood was estimated to be between 0.024 ± 0.027 and 0.665 ± 0.080 Gy, depending on the model used. Differences between the models were observed, with the observed dose variance coming from inter-model as opposed to inter-patient differences. The M-FISH<sub>LET</sub> model potentially overestimates the  $^{223}$ Ra absorbed blood dose by accounting for further PBL exposure in the vicinity of metastatic sites.

## Conclusions

The models presented provide initial estimations of cumulative dose received during incremental IMRT fractions and [ $^{223}$ Ra]RaCl<sub>2</sub> injections, which will enable improved understanding of the doses received by individual patients. While the M-FISH<sub>LET</sub> method builds on a well-established technique for external exposures, further consideration is needed to evaluate this method and its use in assessing non-targeted exposure by  $^{223}$ Ra after its localization at bone metastatic sites.

[Insert graphical abstract]

Keywords;

Radium-223, Biodosimetry, Chromosome exchange. Ta geted alpha-particle therapy, Prostate cancer

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#### 1. Introduction

Metastatic castration resistant prostate cancer (mCRPC) is an incurable condition. Prostate cancer has a strong predisposition to forming bone metastases, with upwards of 90% of patients with advanced disease being affected, often with bone as the only site of metastasis (1). As the general standard of care in the UK, patients with mCRPC are offered a number of life-prolonging therapies including chemotherapy, novel anti-hormonals and radium-223 dichloride ([<sup>223</sup>Ra]RaCl<sub>2</sub>). Recent advances in molecular radiotherapy have generated a great deal of interest, particularly the use of  $[^{223}Ra]RaCl_2$ after the landmark phase III trial ALSYMPCA (2-4) showed for the first time an overall survival advantage associated with [<sup>223</sup>Ra]RaCl<sub>2</sub> treatment. Along with survival prolongation, this trial also demonstrated significant improvement in the quality of li<sup>f</sup>, by lelaying the onset of symptomatic skeletal related events and alleviating pain. This led to FDA and EMA approvals and the widespread use of  $[^{223}$ Ra]RaCl<sub>2</sub> for symptomatic mCRPC (5,  $\bigcirc$  <sup>22</sup> Ka has a half-life of 11.43 days, decaying through creation of a succession of short-live. uclides to stable <sup>207</sup>Pb. During its decay chain, <sup>223</sup>Ra emits four  $\alpha$  and two  $\beta$  particles, with 94% c<sup>c</sup> its decay energy released as high LET  $\alpha$ -particles over a short track length of <100  $\mu$ m (7, 8). When fiving tissue is exposed to <sup>223</sup>Ra, this results in the localised induction of clustered D. 'B lesions which are difficult to repair, effectively leading to cell death (9-11).

For symptomatic mCRFC parients, <sup>223</sup>Ra is administered intravenously as [<sup>223</sup>Ra]RaCl<sub>2</sub> under the trade name of Xofigo (5). Once administered, [<sup>223</sup>Ra]RaCl<sub>2</sub> immediately solubilises in the blood resulting in free <sup>223</sup>Ra, which as a calcium mimetic, is cleared from the blood within 24 hours (h) (12) localising to areas of freshly mineralized bone. The uptake of <sup>223</sup>Ra into high-turnover areas of bone is not dependent on a particular malignant signalling process. Thus, [<sup>223</sup>Ra]RaCl<sub>2</sub> could be of benefit in a range of other malignancies which have a predisposition to forming bony metastases such as breast, lung, kidney and myeloma, including those with favourable long-term survival probability. Due to the short range of  $\alpha$ -particles, this effective target cell kill also minimises direct  $\alpha$ -particle exposure to non-target normal cells (7, 13-16). There is also evidence, however, that <sup>223</sup>Ra can lead to high

absorbed doses in sites adjacent to target bone metastases, including osteogenic cells and the red bone marrow (17-20). It has also been reported that additional biological responses via bystander effects may also occur (21).

Currently, the number of studies which seek to understand the biological action of <sup>223</sup>Ra *in vivo* in humans is limited (22), indeed, there remains a lack of scientific rationale to underpin current dosing strategies and there remains a great deal of uncertainty about the heterogeneous distribution of dose at the cellular and tissue levels and the role of direct and indirect effects such as bystander responses. Exposure of non-cancerous cells and tissues surrounding the tumous is of concern in all radiotherapeutic treatments, due to acute toxicity and the potential for delayed late effects, sometimes many years after treatment (23-30). Quantifying the absorbed vose to non-target tissues is an important step in evaluating the potential short- and longer-term secondary radiation risks for patients (31). This is of particular importance if [<sup>223</sup>Ra]Ra(1<sub>2</sub> is to be used earlier in the treatment schedule. A number of dose calculation models have been. A veloped, for example, for calculation of the absorbed dose to blood from radiotherapy (32), however the actual or estimated consequences of these are still being quantified (33).

Cytogenetic analysis of chromo. The aberrations in blood lymphocytes is widely used to estimate the dose of ionising radiation. The aberrations in blood lymphocytes is widely used to estimate the dose of ionising radiation. The divide by an individual following a real or suspected radiation overexposure, to help inform assessment of future health risks (34, 35). The dicentric assay is the most common method of biological dosimetry, not least due to the high radiation specificity and low interindividual variation between yields of dicentric chromosome aberrations (34). In so called criticality situations of mixed field neutron and gamma exposures after nuclear reactor incidents, where individuals are irradiated by both high-LET and low-LET sources, a model for emergency exposure situations has been designed to estimate external doses for the individual exposure components (35). Although originally designed for mixed neutron: gamma exposures, for medical uses of radiation, the ratios of delivered doses from internal and externally applied radiations can be

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readily derived from patient treatment plans. Meaning that this approach can be applied to calculate alpha and X-ray doses in cases of mixed exposures. Complex chromosome aberrations are useful biomarkers of LET (36), and their identification in exposed individuals could provide another approach for estimating the doses from mixed exposures and, thus, to further refine dose estimation methods in mixed exposure scenarios.

ADDRAD is an approved (NHS REC 15/NI/0074) phase I/ II clinical trial which seeks to address the potential benefit of treating metastatic hormone sensitive prostate cancer patients with androgen deprivation therapy in addition to 6 cycles of [<sup>223</sup>Ra]RaCl<sub>2</sub> and intens ' / modulated radiotherapy (IMRT) to the prostate and pelvic lymph nodes which has recently 'epo ted its first clinical data showing good response and minimal toxicity (37). This patient', rour, therefore presents a unique opportunity to address important radiobiological research que, 'ions pertaining to the doses and cellular and tissue level damage associated with internal <sup>2,3</sup>Ra exposure combined with IMRT. In this study, we describe the use of 24-colour whole chr<sup>4</sup> modulated radiotherapy (multiplex fluorescence *in situ* hybridisation, M-FISH) applied to blood san ple, taken from the ADRRAD cohort of patients, to quantify the patterns of chromosome exchange complexity, with the aim of discriminating damaged cells from each component radiation type ard estimating absorbed radiation dose. In addition, patient specific information is used to cal utate absorbed doses to blood on the basis of three different established or adapted models and the uncertaintie involved in dose estimates are compared, with the overall aim to further understand the uncertaintie involved in dose estimation in this mixed radiation exposure scenario.

#### 2. Methods

## 2.1. Physically derived blood dose estimates

The absorbed blood dose per fraction ( $D_b$ ) was estimated using data from 13 recruited mCRPC patients. [<sup>223</sup>Ra]RaCl<sub>2</sub> was administered over 6 cycles (C1-C6), comprised of one intravenous injection every 4 weeks containing 55 kBq kg<sup>-1</sup> of <sup>223</sup>Ra (38) (Figure 1). As part of the treatment, IMRT is received during the first 7.5 weeks, coinciding with two [<sup>223</sup>Ra]RaCl<sub>2</sub> cycles. The IMRT

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treatment plan for ADRRAD targets the prostate with a dose of 74 Gy, with 60 Gy delivered to lymph nodes. IMRT was delivered in daily fractions of 2 Gy to the prostate and 1.6 Gy to the lymph nodes, the latter as a concomitant boost. Individualised patient treatment schedules are outlined in supplementary Table 1. The absorbed dose to the blood was estimated for <sup>223</sup>Ra based on its pharmacokinetic properties, and the IMRT dose by two simple blood dose models originally derived by Moquet *et al.* (32) with modifications as described.

## [Insert Figure 1]

# 2.1.1. Pharmacokinetic derived absorbed blood dose of <sup>223</sup>R;

In the ADDRAD trial, [<sup>223</sup>Ra]RaCl<sub>2</sub> was administered at *e* h ac 'ivity of 55 kBq kg<sup>-1</sup> of total body weight. Following this, the radionuclide will clear 'recht' e blood by 24 h. Previous studies identified the amount of <sup>223</sup>Ra in the whole blood voluthe *a* varying time points within 24 h of administration (12, 39-41). The median percentage of choulating <sup>223</sup>Ra was averaged at timepoints of 15 min, 4 h and 24 h between all studies reporting median values with upper and lower ranges (39-41). This gave estimates for the percentage of radium in the blood of 22% after 15 min (9-28%), 3% (1.3-4.95%) after 4 h and 0.8 % (0.37-1%) at 24 h. This information was used to estimate the physical absorbed dose to the blood by circul ting <sup>223</sup>Ra.

Firstly, the total activity c<sup> $2^{-3}$ </sup>Ra present in the blood was estimated at each timepoint. This was determined by multiplying the initial injected activity of radium by the estimated percentage of <sup>223</sup>Ra remaining in the blood and the decay constant of <sup>223</sup>Ra, 7.02 × 10<sup>-7</sup> s<sup>-1</sup> (42). The rate of energy deposition was then calculated using the energy release for each decay of <sup>223</sup>Ra and its short-lived daughter products (<sup>219</sup>Rn, <sup>215</sup>Po and <sup>211</sup>Bi) (43), with the total  $\alpha$ -particle energy release estimated to be 4.30 × 10<sup>-12</sup> J. Finally, the dose rate to blood was calculated by dividing this rate of energy deposition by the mass of blood of the patient, which for this work was estimated using 75 mL of blood per kg body weight.

The absorbed dose rates were then plotted as a function of time following injection, and the area under the curve was calculated (GraphPad Prism 9; GraphPad Software; computed using trapezoid rule) to estimate the patient specific dose delivered to the blood for the first 24 hours of each treatment cycle. As less than 1% of the activity remains at 24 h post-irradiation, it was assumed that the additional dose deposited from 24 h to  $t=\infty$  was negligible. The uncertainty on the <sup>223</sup>Ra absorbed blood dose estimates was dominated by the uncertainty in the amount of circulating <sup>223</sup>Ra at each time period, which, as discussed below, could be up to 50% (39-41).

#### 2.1.2.IMRT blood flow model

For this part of the work, data was assessed from 13 ADL.<sup>2</sup> A $\Gamma$  patients, for whom treatment plan and patient specific data were available. The blood flow no iel (BF) enabled estimation of dose within the high dose organ area. The absorbed blood dose per fraction (D<sub>B</sub>) was estimated as follows:

 $D_{B}=D_{f}(V_{95}\div v) \qquad (Equation 1)$ 

Where  $D_B$  was absorbed dose to blood per normalized normalized dose per fraction,  $V_{95}$  represents the high dose volume and  $V_B$  was the total blood volume.

Two variates of the model were used:  $BF_1$ , which was applied as per Moquet *et al* (32) which estimates  $V_B$  by assuming .'5 m of blood per kg and,  $BF_2$  which uses the static volume of blood in the prostate and lymph nodes for  $V_B$ , estimated by calculating a scaling factor between the whole-body volume and the area irradiated (prostate and/or lymph nodes) on the basis of treatment plan information. This scaling factor was then applied to the whole-body blood volume estimates to achieve a static blood volume for prostate and lymph nodes. In both cases,  $D_F$  and  $V_{95}$  were taken from the treatment plans. The uncertainty on  $V_B$  and  $V_{95}$  was estimated to be on the order of 10%, and the uncertainty on  $D_f$  can be estimated on the basis of Moquet and colleagues to be up to 20%. Hence a conservative estimate of uncertainty on  $D_B$  would be approximately +/- 25%.

#### 2.1.3. IMRT CT plan model

For this part of the work, data from 13 ADDRAD patients was assessed, for whom treatment plan and patient specific data were available. The CT planned volume (CTPV) absorbed blood dose model enables dose estimation of both high and low dose areas. The CT volume mapped for each patient was utilized to calculate a scaling factor for each patient relative to whole-body volume, with whole-body volume being calculated based on the average patient weight (kg) over the course of treatment (44). For the following dose models, the whole CT volume was considerer' within the CTPV<sub>1</sub> along with a high dose volume only CTPV<sub>2</sub>. Utilizing the CTPV<sub>1</sub> model, the whole organ dose was estimated, and it was assumed the blood volume had also received to is. Firstly, a scaling factor, S, was estimated as the ratio between the CT plan volume and the whole-body volume. The following was then applied to estimate the absorbed blood dose per fraction:

$$D_{B} = (D_{PB} \div N_{F}) \cdot S \qquad (Equation 2)$$

where  $D_B$  was the absorbed dose to blood per the ction;  $D_{PB}$  was the mean dose in Gy to the body volume covered by the CT scan (specificate each patient, taken from the treatment plans), N<sub>F</sub> was the number of fractions of radiotherapy and S was the patient specific scaling factor. A further estimate, CTPV<sub>2</sub>, was performed in the same manner but in this case firstly calculating the average dose per fraction for prostate and lymph houses, and then calculating a scaling factor based on their volume compared to the whole-bully volume.

The error associated with the plan volume was considered to be on the order of 1-1.5%. The clinical upper acceptable limit in dose delivery to the treatment plan was within 3%, as anything greater than this would trigger re-calibration of dose planned area. The scaling factor was estimated from whole body volume which was estimated by assuming that 1.01 g of human body mass fits within 1 cm<sup>3</sup> and that the tissue density within the target volume was consistent with this for the scaling factor estimation, with an error within 1%. The total error associated with CTPV estimations was considered to be within  $\pm/-6\%$ .

#### 2.2. M-FISH dicentric quantification

#### 2.2.1. Sample collection

Whole blood was received from 5 male patients recruited onto the ADRADD trial (EudraCT 2014-00273-39) with full informed consent (NHS REC 15/NI/0074) at The Belfast Health and Social Care Trust. Blood samples were drawn into lithium heparinised anticoagulant tubes, once every 4 weeks, immediately prior to the first (C1) and then within 24 h prior to each subsequent (C2-C5) [<sup>223</sup>Ra]RaCl<sub>2</sub> administration (Figure 1). The samples were then shipped at room temperature for next day delivery to Brunel University London. Upon arrival, the samples were processed and whole blood stimulated to divide to enable the collection of 1<sup>st</sup> *in vitro* cell division. Anetaphase cells for cytogenetic assessment, as described below.

#### 2.2.2. Cell culture

For each sample, 0.4 ml of whole blood was vise to inoculate 2.6 ml of freshly prepared media (PBMAX Karyotyping Medium (Therino) isher, Cat. Number 12557021) supplemented with 0.5  $\mu$ g/ml purified phytohaemagglutinin (PHA) (ThermoFisher, Cat. Number R30852801), 10 $\mu$ M 5-bromo-2'-deoxyuridine (BrdU, Sigma-Ad' $\alpha$  h Cat. Number 19-160), 10  $\mu$ l/ml heparin (Sigma-Aldrich Cat. Number 9041-08-1) and cultur. (Tin a humidified incubator at 37°C (95% air/5% CO<sub>2</sub>), at a 45° angle, and with the cap left slighdy open to allow gaseous exchange. Cultures were set up to maximise the yield of 1<sup>st</sup> of the  $\alpha$  vision of peripheral blood lymphocytes (PBLs) and harvested using standard cytogenetic technic ues after a total of 50-60 h. To arrest cells at the metaphase stage of the cell cycle, 50  $\mu$ l/ml of Colcemid KaryoMAX (ThermoFisher, Cat. Number 1521012), a tubulin inhibitor, was added 4 h prior to harvest. After this time, the cultures were centrifuged at 200g for 10 min and the cell pellet re-suspended before the addition of 0.075M KCl hypotonic solution (Fisher Scientific Cat Number10575090) for 8 min at 37°C. Cells were then centrifuged at 200g for 10 min and fixed in 3:1 methanol (Thermo Fisher Catalogue Number 15654570) acetic acid (Thermo Fisher Catalogue Number 1743468) on ice. The fixation process was repeated until the samples appeared clear (~ 5 times), and these were then stored in the freezer at -20°C.

#### 2.2.3. Harlequin stain

Fixed chromosome preparations were dropped onto clean, grease-free slides and assessed for metaphase quality. Harlequin staining was used to assess the number of  $1^{st}$  division metaphase cells. For this, slides were aged on a hot plate for 45 min at 90°C and immersed in Hoescht (Thermo Fisher Scientific Cat Number 62249) for 10 min, then transferred to a flat tray and covered in 2x Saline-Sodium Citrate (SSC) (Thermo Fisher Cat Number 15557036), before being exposed in UV box to  $1.0 \text{ J/cm}^2$  for 1 h. After exposure slides were washed with distilled water twice and air dried. Treated slides were stained in 5% Giemsa (VWR Cat Number 350864) for 4 min. removed and rinsed with distilled water. Once dry, the slides were mounted with coverslips with 4 drops of DPX (Fisher Scientific Cat Number 15538321). Slides were scored using heightfield microscopy with oil immersion at x100 magnification. The fraction of  $1^{st}/2^{nd}/2^{-d}$  division cells was determined based upon the chromatid staining patterns (45, 46). Samples with  $3^{rt}/2^{nd}$  division cells were assayed by M-FISH.

## 2.2.4. Multiplex fluorescence in situ hybridisation (M-FISH)

M-FISH was carried out utilizing 24XCyte staining probe (Metasystems Probe Cat NumD-0125-600-DI) as per manufactu. The protocol. Patient slides were selected from 5 patients according to metaphase spread quality at a thether samples containing  $\leq 5\%$  2<sup>nd</sup> division cells, a minimum of 3 patient slides were painted, there er cycle. In brief, slides were incubated in 2xSSC at 70°C (±1°C) for 30 min. After this time, the cooled slide was transferred into 0.1xSSC at RT for 1 min. Chromosomes were then denatured in 0.07 NaOH at RT for 1 min followed by 1 min incubation in 0.1xSSC, followed by 2xSSC at 4°C, and then dehydrated through immersion in a series of alcohol solutions of ascending strength (70%, 95% and 100%). The 24Xcyte probe was denatured by incubating at 75°C (±1°C) for 5 min, placed on ice briefly and then incubated at 37°C for 30 min. The probe was overlaid on to the slide and left to hybridize in a humidified chamber at 37°C (±1°C) for 2-3 days. Slides were washed in 0.4x SSC preheated to 72°C (±1°C) for 2 min then incubated in 2xSSCT (containing 0.05%)

Tween20) for 30 secs. For counterstaining, the slide was rinsed in double distilled water and left to air dry before application of DAPI/antifade and sealing.

Slides were visualised utilizing 8-position Zeiss Axioplan 2 microscope containing individual filter sets for 24XCyte probe cocktail plus DAPI (FITC, Spectrum Orange, Texas red, Cy5, DEAC and DAPI). Metaphase cells were imaged under x63 oil immersion and captured by Cool Cube driven by Metafer4 version 3.14.191 software. The image files were exported and karyotyped in ISIS version 5.8.11.

#### 2.2.5. Chromosome aberration classification

A cell was classified as being apparently normal if all 46 chr. nosomes were present and contained the appropriate fluorophore combination along their entire tength. Only metaphase cells with good fluorochrome staining were selected for analysis in cell, containing  $\geq 43$  chromosomes. Chromosomal aberrations were identified by colour junction . 'on, the length of each individual chromosome and/or by the presence of chromosomal fragme. 's (Figure 2b). A chromosome interchange involving 2 breaks in 2 chromosomes was categories  $d_{a}$  a simple exchange, and further classified as a reciprocal translocation or dicentric. KI. o chromosomes, which involve 2 breaks in one chromosome were also classed as simple (47) E. changes involving 3 or more breaks in 2 or more chromosomes were classed as complex and assigned the minimal number of breaks, arms and breaks involved (CAB) (48). Chromosome having breaks only, not involving any additional chromosomes, were classed as chromosome breaks. When classifying cells with multiple aberrations, all aberrations were recorded as independent events and the chromosomes involved identified. Where homologous chromosomes were involved, efforts were made to establish whether the homologues were in the same event or in different independent events, mainly by consideration of chromosome length (49). All exchanges were recorded as either complete (all break-ends re-joined), true incomplete (where one or more break-ends fail to find an exchange partner) or one-way (where one or more elements appear to be missing) (50, 51). The potential transmissibility of exchanges was also recorded, where a stable (transmissible) exchange was defined as complete and with no evidence of unstable elements

e.g. dicentric or acentric fragments. Each independent complex event was also determined to be transmissible or non-transmissible and the presence of insertion-type rearrangements were noted (49). Metaphase spreads were categorised as stable only if all the exchange events detected within that spread were classified as stable. Unstable complex chromosomal exchanges containing polycentric chromosomes were broken down in to their dicentric equivalents whereby each additional centromere within a chromosome structure constituted a dicentric equivalent (dicentric equivalent event = n centromere -1) (Figure 2).

#### 2.3. M-FISH dicentric assay dose estimation

## 2.3.1.Mixed field absorbed blood dose ratio

For this part of the work, M-FISH analysis was carried out n-5 individuals with a minimum of n=3patient samples processed for C2-C5. The <sup>223</sup>Ra:IMRT ratio , 'as estimated in two ways. Firstly, by physical dose estimation and secondly, from categorning cells based on the complexity of chromosome exchange observed. For the ph<sup>-</sup> sice dose ratio, the absorbed blood dose was calculated independently for [<sup>223</sup>Ra]RaCl<sub>2</sub> and IMR<sup>T</sup> (section 2.1) for each exposure and was summed over the 4-week time period prior to each sample ten g taken. The average absorbed blood dose per cycle was estimated by averaging the absorbed blood dose across all 13 patients for all treatment cycles. For the dose ratio the overall treatment p. n was used (Figure 1), not the patient specific plan adaptation. Therefore, C2 relates to the rest onse to one injection of [<sup>223</sup>Ra]RaCl<sub>2</sub> and 20 IMRT fractions, C3 relates to two injections of  $1^{223}$ Ra]RaCl<sub>2</sub> and 37 IMRT fractions, with C4 and C5 each including the additional [<sup>223</sup>Ra]RaCl<sub>2</sub> only administrations. Summing these gives the total absorbed blood dose of [<sup>223</sup>Ra]RaCl<sub>2</sub> and IMRT delivered at each time point from C2 to C5. For the M-FISH derived ratio (M- $FISH_{LET}$ ), cells containing at least one complex chromosome exchange were classed as being damaged by the traversal of high-LET  $\alpha$ -particles from the <sup>223</sup>Ra while cells containing only simple chromosome exchanges, as from IMRT (section 2.2.5). The ratio of cells containing at least one complex chromosome exchange to cells containing simple exchanges only was therefore a pseudo ratio for <sup>223</sup>Ra:IMRT absorbed blood dose at each cycle point.

### 2.3.2.Dicentric absorbed blood dose estimation

The dicentric assay traditionally utilizes Giemsa staining where chromosomes are evenly stained in one colour, (Figure 2a), thus enabling the identification of chromosomes containing more than one centromere along with any associated acentric fragment. In this study, dicentrics were quantified from M-FISH painted metaphase cells (Figure 2b and c).

#### [Insert Figure 2]

Dose estimation was first carried out utilizing the dicentric equivalent frequency as determined by M-FISH. For the IMRT, the <sup>60</sup>Co calibration curve of Lloyd and colleagues, 1986, was used (52) where whole blood was irradiated *in vitro* utilizing a <sup>60</sup>Co s with dose range of 0-5 Gy. The dicentric vield was entered into Dose Estimate V5.2 (5.3) vith the following coefficients  $\alpha = 0.0756 \pm 0.0031$ ,  $\beta$ = 0.0149 ± 0.0060 and C= 0.0004 ± 0.0009 (52). This being a well-established calibration curve, it has been utilized in many exposure scenar  $\sqrt{\gamma}$  for  $\gamma$ -ray and X-ray dose estimation by Public Heath England and was judged to be the most comparable curve in terms of type and energy of radiation exposure. As there is currently  $n^{2}$  Ra calibration data, a calibration curve based on a <sup>239</sup>Pu, which emits  $\alpha$ -particles of a similar e, ergy to <sup>223</sup>Ra (5.16 MeV per  $\alpha$ -particle), was selected. The <sup>239</sup>Pu calibration curve of Purroti  $\gamma t al.$ , 1980, (54) was used, the curve coefficients were:  $\beta = 0.3696 \pm$ 0.0322, C=  $0.0019 \pm 0.0126$  (54). The decay chain of <sup>223</sup>Ra and <sup>239</sup>Pu does differ, however, due to the perceived lack of risk of a radiation accident involving alpha exposures and the complexity of the experiments, alpha curves for biodosimetry are rare. This curve was established by irradiating whole blood with <sup>239</sup>Pu in the range of 0-1.6 Gy. It is important to note, however, that in this study, the likely non-homogeneous, partial body, nature of both the <sup>223</sup>Ra and IMRT exposures beyond the treatment plan details, was not further considered, neither was the microdosimetric heterogeneity of radium in areas of high bone turnover, and thus this represents a key source of unquantified uncertainty. This aspect will need to be incorporated into further development of absorbed blood dose models. To

calculate the absorbed blood dose in this mixed exposure scenario, the "criticality" model was used (35). In brief, all aberrations were firstly assumed to be attributed to <sup>223</sup>Ra and from the dicentric equivalent yield the dose was estimated. The absorbed blood dose ratio (<sup>223</sup>Ra:IMRT) calculated according to section 2.3.1. was then used to estimate the IMRT dose and then the gamma calibration curve used to estimate the dicentric equivalent yield. This IMRT yield was then subtracted from the total yield to give a 'new' <sup>223</sup>Ra dicentric equivalent yield. This iterative process was repeated until self-consistent estimates were obtained.

#### 2.4. Other statistical analysis

Statistical analysis was performed using GraphPad Prism 9 (CraphPad Software). Descriptive statistics are presented as mean ± SE for pooled data. Standard propagation of errors was applied to estimate the uncertainty in the derived dose estimate . A crmality testing indicated ANOVA was appropriate in order to test for differences in floc. ration frequencies between treatment rounds and for differences in estimated doses.

#### 3. Results

## 3.1. Estimating the '10.1 cose from the treatment schedule

Patient data related to the planned treatment for 13 patients was available for physical dose estimation. Table 1 gives the absorbed blood dose per fraction predicted by the BF model, estimated for each patient for the prostate, lymph nodes, and total high dose region combined. The majority of the absorbed blood dose was estimated as being from lymph node exposure, due to the larger volume of irradiation for this tissue (BF<sub>1</sub>). The total prostate-only absorbed blood dose was found to be in the range of 0.880-1.962 Gy by the end of IMRT treatment (30-37 fractions; 37 fractions assumed for dose ratio calculations in section 3.3), comparable to the 0.38-1.92 Gy reported by Moquet *et al.* To consider the differences in lymphatic fluid shift, the static volume was calculated (BF<sub>2</sub>) and, by

combining the BF models ( $BF_{1P} + BF_{2LN}$ ), the cumulative absorbed blood dose was estimated to be between 1.131 - 2.717 Gy.

Table 1. BF model.  $V_B$  blood volume,  $D_B$  blood dose per fraction. BF<sub>1P</sub> from BF<sub>1</sub> prostate dose combined with BF<sub>2</sub> LN dose (BF<sub>2LN</sub>). Patients for whom treatment plan and patient specific data were available n=13. Reported estimated uncertainty of 25% for all BF model variants.

				BF1			BF <sub>2</sub>	$BF_{1P} + BF_{2LN}$
Patient ID	V <sub>B</sub> body (ml)	V <sub>B</sub> prostate (ml)	V <sub>B</sub> LN (ml)	Prostate D <sub>B</sub> (Gy)	LN D <sub>B</sub> (Gy)	Tota' dos D <sub>B</sub> Gy)	LN D <sub>B</sub> (Gy)	Total dose D <sub>B</sub> (Gy)
1	9030	12.9	54.5	0.038	0.132	0.170	0.007	0.045
2	7615	11.7	56.8	0.041	0.164	0.205	0.010	0.050
3	6288	9.5	49.2	0.040	0	0.212	0.009	0.049
4	6560	12.2	57.2	0.060	0`(1	0.291	0.014	0.073
5	6461	13.8	63.9	0.056	v.?09	0.265	0.015	0.071
6	6985	10.8	58.9	0.04.	0.184	0.225	0.011	0.052
7	7650	9.4	52.0	0.033	0.149	0.181	0.008	0.040
8	5700	7.0	32.4	0 0'34	0.124	0.157	0.004	0.037
9	6290	11.5	49.1	0 15,	0.154	0.204	0.007	0.058
10	6629	14.0	52.3	P.U56	0.171	0.227	0.009	0.065
11	5559	9.1	48.4	0.043	0.188	0.231	0.009	0.052
12	7883	7.1	49.2	0.024	0.135	0.159	0.007	0.031
13	6486	10.2	54.9	0.042	0.161	0.203	0.009	0.051

The absorbed blood dose was estimated for the whole CT plan area, including high dose and low dose regions (CTPV<sub>1</sub>) and for high dose regions only (CTPV<sub>2</sub>) to enable comparison with the BF model. Previous CTPV approaches have used a fixed scaling factor of 2.5 with a reported uncertainty of 20%. In this work we instead estimated a patient specific scaling factor between 2.08-2.64, with the largest deviation from the scaling factor for Patient 10 being 16% smaller than the published 2.5 scaling factor. The increased absorbed blood dose CTPV estimates compared to the BF estimates observed here are likely to be reflective of the large lymph node irradiation volume (32). To compare the BF and CTPV models, CTPV was also calculated using the high dose volumes of prostate and lymph

nodes, termed as CTPV<sub>2</sub>. To do so, a scaling factor for each component as detailed for BF<sub>2</sub>, was used. The resulting absorbed blood doses for CTPV<sub>2</sub> were found to be comparable to that of BF<sub>2</sub> within the uncertainty estimates (Table 2). The average absorbed blood dose per fraction of IMRT was estimated as 0.101 Gy for CTPV<sub>1</sub> and 0.016 Gy for CTPV<sub>2</sub> (average n =13 patients), each with an estimated uncertainty on the order of 6%.

BF<sub>1</sub> and CTPV<sub>1</sub> cannot be directly compared as they use different irradiation volumes. The BF<sub>1</sub> model assumes the blood flow within prostate and lymph node regions was dentical and in doing so, it appears to significantly overestimate the lymph node blood volur ie. Up the basis of this and the simple nature of the other model assumptions, this method likel that the largest uncertainty. Accordingly, only  $CTPV_1$  and  $CTPV_2$  are considered further values models represent an estimate of combined absorbed blood dose for high dose regions and low regions (CTPV) and high dose regions only (CTPV<sub>2</sub>). Supplementary Table 1 det ils the patient specific injected activities and CT planning volumes. Table 2 shows the individual absorbed blood dose estimates for each <sup>223</sup>Ra administration, and the IMRT absorbed blood dose estimates for each fraction. The average <sup>223</sup>Ra absorbed blood dose per treatment cycly var  $0.012 \pm 0.002$  Gy. The estimated <sup>223</sup>Ra absorbed blood dose per fraction was not found to be statistically different between patients (P= 0.097). To estimate the <sup>223</sup>Ra absorbed blood dose per reatment cycle, the median values of the activity kinetics reported in the literature were utiliz d. B sed on the highest and lowest limits reported for each timepoint the <sup>223</sup>Ra blood dose estimates r er treatment cycle could vary by up to 50%. This value being reported as a conservative estimate of uncertainty for dose estimates, with more work needed to quantify the variations between patient clearance.

Table 2. Physically derived absorbed blood dose, in Gy. <sup>223</sup>Ra absorbed blood dose was representative of an average dose to the blood during localization period. The absorbed blood dose after all treatment cycles was calculated for n=6 injections with the exception of Patient 5 where n=5. IMRT absorbed blood dose estimated for whole CT plan area CTPV<sub>1</sub> and high dose regions only CTPV<sub>2</sub>. Scaling

factor estimated from patient body volume derived from average weight during treatment. Patient

numbers n=13 patients for whom treatment plan and patient specific data were available.

_	<sup>223</sup> Ra	IMRT		
Patient ID	D <sub>B</sub> (Gy)	$CTPV_1 D_B (Gy)$	$CTPV_2 D_B Dose(Gy)$	
1	0.0116	0.105	0.013	
2	0.0114	0.104	0.015	
3	0.0116	0.101	0.016	
4	0.0118	0.113	0.022	
5	0.0116	0.100	0.020	
6	0.0116	0.112	0.017	
7	0.0117	0.092	0.014	
8	0.0118	ر 20.0	0.012	
9	0.0117	0 )8>	0.015	
10	0.0115	0112	0.017	
11	0.0114	0.1 )8	0.018	
12	0.0116	074	0.012	
13	0.0114	0.088	0.015	

Uncertainties of up to 6% for CTPV models and up to 50% considered for the <sup>223</sup>Ra estimates.

## 3.2. Frequency and type of chrome so a aberration in ADRADD patients

M-FISH analysis was carried out on blood samples received from 5 patients for C1-C5. The control samples of all 5 patients were analysed and following this a minimum of 3 patients were analysed per treatment cycle. The background frequency of chromosomal aberrations was found to be within the expected normal range for individuals in the 50+ age bracket (55, 56). Specifically, frequencies of 0.0,  $0.024 \pm 0.011$  and  $0.020 \pm 0.009$  were observed for simple dicentrics, reciprocal translocations and break-only aberrations, respectively (sample C1; pooled for 5 patients). One cell containing an unstable complex rearrangement was found in Patient 2 (0.004  $\pm$  0.004), for which the origin was not clear.

Simple chromosomal exchanges were observed to significantly increase in frequency between C1 and C2 ( $0.024 \pm 0.011$  to  $0.319 \pm 0.042$  (P < 0.001)) and further at C3 to  $0.484 \pm 0.042$  (P < 0.031) (Supplementary Table 2). No other statistical differences were noted between treatment cycles

(C3-4, P= 0.966 or C4-C5, P= 0.996), suggesting the majority of simple aberrations formed early in the treatment regime persist over the period sampled, and/or the induction of new aberrations was balanced by cell death of unstable (e.g. dicentric) types. The frequency of complex exchanges increased from  $0.058 \pm 0.016$  at C2 to  $0.174 \pm 0.022$  at C3 (P = 0.007), rising further to  $0.265 \pm 0.037$  at C4 and  $0.210 \pm 0.030$  at C5 (Supplementary Table 2).

In terms of classification of complexity, the frequency of damaged PBLs assigned to each exposure type (<sup>223</sup>Ra or IMRT) was reported in Figure 3 and Table 3. This categorisation was based upon the presence or absence of a complex chromosome exchange and shows unit the majority of damaged PBLs sampled, with combined IMRT and <sup>223</sup>Ra contained mostly simply exchanges only at C2 (cells containing simple exchanges  $0.204 \pm 0.025$  and containing at least one complex exchange  $0.050 \pm 0.014$ , P < 0.001) and C3 ( $0.243 \pm 0.021$  and  $0.150 \pm 0.013$ , P = 0.006). After this time, when patients continue to receive <sup>223</sup>Ra only, the fraction of damage  $2^{10}$ BLs with at least one complex exchange increased. In the <sup>223</sup>Ra only cycles, no significant cifference between cells containing a simple aberration and those containing at least one complex exchange was observed (C4 P = 0.992, C5 P = 0.558).

## [Insert Figure 3]

#### 3.3. M-FISH dicentric assay absorbed blood dose estimates

The absorbed blood dose ratios were estimated for C2-C5, this was equivalent to 4 intravenous injections of  $^{223}$ Ra and completed IMRT schedule of 37 fractions by C5. For the physical dose ratio, this was based on the average absorbed blood dose per fraction across all 13 patients for both the IMRT and  $^{223}$ Ra dose. The M-FISH<sub>LET</sub> ratios described in section 2.3.1, was obtained from blood samples received from 5 patients with a minimum of 3 patient samples analysed per cycle. The absorbed blood dose ratios for all models can be seen in Table 3. Applying these blood dose ratios to the dicentric assay, the absorbed blood dose was estimated in three ways (Table 4). The CTPV

methods were derived from the physical absorbed blood dose estimates as per method section 2.1. with  $CTPV_1$  estimating the absorbed blood dose across the planned volume including both high and low dose volumes whilst  $CTPV_2$  including the high dose regions only. The M-FISH<sub>LET</sub> absorbed blood dose was instead estimated from the ratio of cells consistent with high LET exposure (<sup>223</sup>Ra) and low LET exposure (IMRT).

Table 3. Absorbed blood dose ratios. M-FISH derived ratio based on the complexity of cellular damage with <sup>223</sup>Ra assigned cells to include those with at least 1 complex exchange and, IMRT assigned cells defined as containing only simple exchanges. Frequency per total cells scored pooled per cycle, with calculated SE. D<sub>B</sub> per cycle of patient average (1 = 13), complexity of cellular damage from pooled patient data (n=5).

	Cellular	classification		D <sub>B</sub> per cycl	<sup>223</sup> Ra : IMRT			
Cycl e	<sup>223</sup> Ra cells (f)	IMRT cells (f)	<sup>223</sup> R . ∖ີv)	IMRT CTPV <sub>1</sub> (Gy)	IMRT CTPV <sub>2</sub> (Gy)	CTPV 1	<b>CTPV</b> 2	M- FISH <sub>LET</sub>
C2	$0.050 \pm 0.014$	$0.204\pm0.025$	$0.212 \pm 0.002$	2.021 ± 0.050	0.318 ± 0.017	1:174	1:27	1:4
C3	$\begin{array}{c} 0.150 \pm \\ 0.018 \end{array}$	$0.243 \pm 0.02^{1}$	$0.023 \pm 0.003$	$3.739 \pm 0.092$	$\begin{array}{c} 0.587 \pm \\ 0.031 \end{array}$	1:161	1:25	2:3
C4	$0.198 \pm 0.025$	$0.229 \pm 0.02$	$0.035 \pm 0.003$	$3.739 \pm 0.092$	$\begin{array}{c} 0.587 \pm \\ 0.031 \end{array}$	1:108	1:17	1:1
C5	$0.162 \pm 0.021$	0.270 = 9.02 r	$\begin{array}{c} 0.046 \pm \\ 0.004 \end{array}$	$\begin{array}{c} 3.739 \pm \\ 0.092 \end{array}$	$0.587 \pm 0.031$	1:81	1:13	3:4

The absorbed blood dose estimates from the CTPV models suggest the dose after 20 IMRT fractions, measured at C2 (Table 4), was between  $1.327 \pm 0.115$  Gy and  $1.395 \pm 0.115$  Gy. The M-FISH<sub>LET</sub> method was found to be in similar range at  $0.956 \pm 0.114$  Gy at C2. The IMRT absorbed blood dose after the end of fractionation (C3 assuming 37 completed fractions) was estimated for CTPV<sub>1</sub> as 2.148  $\pm 0.096$  Gy and 2.073  $\pm 0.096$  Gy for CTPV<sub>2</sub>, with M-FISH<sub>LET</sub> blood dose estimates of  $1.167 \pm 0.092$  Gy.

For the <sup>223</sup>Ra absorbed blood dose at C2, the CTPV<sub>1</sub> and CTPV<sub>2</sub> were found to be  $0.008 \pm 0.025$  and  $0.048 \pm 0.032$ , both lower than the M-FISH<sub>LET</sub> estimate of  $0.150 \pm 0.046$  Gy. For C3 the <sup>223</sup>Ra absorbed blood dose estimated from CTPV methods was between  $0.013 \pm 0.025$  and  $0.082 \pm 0.033$  Gy (CTPV<sub>1</sub> and CTPV<sub>2</sub> respectively) with the M-FISH<sub>LET</sub> absorbed blood dose estimated at  $0.719 \pm 0.073$  Gy. By the final sample point studied here (C5), the absorbed blood dose from <sup>223</sup>Ra was estimated as  $0.024 \pm 0.027$  Gy by CTPV<sub>1</sub> and  $0.141 \pm 0.042$  Gy for CTPV<sub>2</sub> with the M-FISH<sub>LET</sub> absorbed blood dose estimate at  $0.665 \pm 0.080$  Gy (see Figure 4).

[Insert Figure 4]

					$CTPV_1$		$CTPV_2$		M-FISH <sub>LET</sub>	
Cyc le	Patie nt	Cells score	E eq	Dicentric uivalents	<sup>223</sup> Ra	IMRT	<sup>223</sup> Ra	IMRT	<sup>223</sup> Ra	IMRT
	ID	a	(yı	eld $\pm$ SE)						
2	1	55	8	$(0.145 \pm 0.066)$	$\begin{array}{c} 0.007 \pm \\ 0.033 \end{array}$	$1.268 \pm 0.246$	$0.044 \pm 0.054$	$1.201 \pm 0.246$	$\begin{array}{c} 0.207 \pm \\ 0.105 \end{array}$	$0.845 \pm 0.245$
	3	101	16	$(0.158 \pm 0.067)$	$\begin{array}{c} 0.008 \pm \\ 0.029 \end{array}$	$\begin{array}{c} 1.329 \pm \\ 0.182 \end{array}$	$\begin{array}{c} 0.046 \pm \\ 0.043 \end{array}$	1.261 ± 0.182	$\begin{array}{c} 0.221 \pm \\ 0.081 \end{array}$	$\begin{array}{c} 0.899 \pm \\ 0.181 \end{array}$
	4	104	21	$(0.202 \pm 0.048)$	$0.009 \pm 0.029$	$\begin{array}{c} 1.516 \pm \\ 0.180 \end{array}$	$0.053 \pm 0.045$	$1.448 \pm 0.180$	$\begin{array}{c} 0.263 \pm \\ 0.086 \end{array}$	$1.070 \pm 0.179$
	Tota 1	260	45	$(0.173 \pm 0.035)$	$\begin{array}{c} 0.008 \pm \\ 0.025 \end{array}$	$1.395 \pm 0.115$	$0.048 \pm 0.032$	1.327 ± 0.115	$\begin{array}{c} 0.235 \pm \\ 0.055 \end{array}$	$0.959 \pm 0.114$
	1	100	30	$(0.30 \pm 0.0820)$	$\begin{array}{c} 0.012 \pm \\ 0.031 \end{array}$	$1.874 \pm 0.184$	$\begin{array}{c} 0.071 \pm \\ 0.051 \end{array}$	1.799 ± 0.184	$\begin{array}{c} 0.584 \pm \\ 0.128 \end{array}$	$\begin{array}{c} 0.947 \pm \\ 0.182 \end{array}$
	2	101	38	$(0.376 \pm 0.076)$	$\begin{array}{c} 0.013 \pm \\ 0.031 \end{array}$	$2.113 \pm 0.184$	$0.080 \pm 0.053$	$2.038 \pm 0.184$	$0.701 \pm 0.139$	$1.138 \pm 0.182$
3	4	101	45	$(0.446 \pm 0.111)$	$0.014 \pm 0.032$	$\begin{array}{c} 2.310 \pm \\ 0.185 \end{array}$	$0.088 \pm 0.055$	2.234 ± 0.184	$0.801 \pm 0.149$	$\begin{array}{c} 1.300 \pm \\ 0.182 \end{array}$
	5	105	45	$(0.429 \pm 0.065)$	$0.014 \pm 0.031$	$2.264 \pm 0.181$	0.086 <del>-</del> 0.053	$2.188 \pm 0.181$	$0.777 \pm 0.144$	1.261 ± 0.179
	Tota 1	407	158	$(0.388 \pm 0.042)$	$0.013 \pm 0.025$	$2.148 \pm 0.096$	0.032 - 0.733	$2.073 \pm 0.096$	$0.719 \pm 0.073$	$1.167 \pm 0.092$
4	1	48	15	$(0.313 \pm 0.099)$	$0.018 \pm 0.042$	$1.908 \pm 0.264$	0.100 ± 0.^82	$1.797 \pm 0.264$	$\begin{array}{c} 0.680 \pm \\ 0.198 \end{array}$	$0.789 \pm 0.262$
	2	102	53	$(0.520 \pm 0.062)$	$0.023 \pm 0.035$	2.497 ± 0.184	6 141 ±	$2.384 \pm 0.184$	$1.048 \pm 0.169$	$1.216 \pm 0.181$
	3	103	32	(0.311 ± 0.116)	$0.018 \pm 0.033$	1.902 ± 0 18?	$0.106 \pm 0.058$	$1.791 \pm 0.182$	0.677 ± 0.136	0.785 ± 0.179
	Tota 1	253	100	$(0.395 \pm 0.052)$	$0.020 \pm 0.028$	0.1 <sup>-</sup> ∠± 0.119	$0.121 \pm 0.043$	$2.050 \pm 0.119$	$0.833 \pm 0.097$	$0.966 \pm 0.116$
5	1	106	47	$(0.443 \pm 0.083)$	0.028 ± 0.037	$2.2 > 0 \pm 0.180$	$0.169 \pm 0.070$	$2.142 \pm 0.180$	$0.864 \pm 0.151$	$1.175 \pm 0.178$
	4	101	35	$(0.347 \pm 0.084)$	0.025 ± 0.236	$2.009 \pm 0.184$	$0.147 \pm 0.068$	$1.862 \pm 0.183$	$0.705 \pm 0.140$	$0.959 \pm 0.182$
	5	102	18	$(0.176 \pm 0.043)$	0.017 <u>-</u> ).( <i>s</i> _	$1.394 \pm 0.182$	$0.099 \pm 0.057$	$1.254 \pm 0.181$	$0.392 \pm 0.105$	$0.533 \pm 0.180$
	Tota 1	309	100	$(0.324 \pm 0.042)$	0.0°4 ≞ € 927	$1.937 \pm 0.108$	$0.141 \pm 0.042$	1.791 ± 0.108	$0.665 \pm 0.080$	$0.905 \pm 0.105$

Table 4. Absorbed blood doses <sup>4</sup>ur.ng treatment (Gy) calculated on the basis of  $CTPV_1$ ,  $CTPV_2$  and M-FISH<sub>LET</sub> models. Patients completed during treatment cycles were coded and identifier number (ID) assigned, n=5 patients were assayed by M-FISH to determine the dicentric equivalent yield. The absorbed blood dose was calculated with use of dicentric assay for each patient with the dose ratios estimated from CT plan and M-FISH<sub>LET</sub> derived models. The SE was reported for the absorbed blood dose estimation, this being the largest uncertainty for M-FISH<sub>LET</sub> dose estimation.  $CTPV_1$  and  $CTPV_2$  reported SE of absorbed blood dose estimation with a propagated conservative uncertainty estimated up to 50% not included.

## 4. Discussion

In the future,  $\alpha$ -particle emitters, such as <sup>223</sup>Ra, alone or in combination, are likely to be adopted in the treatment of a wide range of cancers, contributing to more targeted, personalised medicine with the potential to impact a large number of patients (17, 57, 58). To date, however, there are only limited studies using biological endpoints which seek to understand the biological action of <sup>223</sup>Ra *in vivo* in humans, to improve the optimal delivery of these and, to minimise the risk of adverse effects through radiation exposure of normal tissues (59-63). Quantifying the absorb d blood dose delivered from the treatment is an important step in maximising clinical efficacy and  $\epsilon$  valu, ting radiation risks, estimates of which are currently based on population studies (31). In this tudy, whole blood was sampled from patients recruited onto the ADRRAD trial (37). The treatman, includes a planned mixed field exposure from daily IMRT fractions, 37 x 2 Gy to prostate while concomitant boost (37 x 1.6 Gy) to lymph nodes, over a period of 7.5 weeks, together  $w_{1,1} \in$  intravenous injections of  $[^{223}Ra]RaCl_2$  over a period of 20 weeks. Dose estimation for nor tar eted tissues in mixed exposure scenarios is challenging. To account for the mixed field nature of the treatment, existing models were used to determine the ratio of the component radiation types. Additionally, a new approach (M-FISH<sub>LET</sub>), based on the ratio of cells containing <sup>4</sup>amage consistent with high-LET exposure (complex chromosomal exchanges) and low-' ET exposure (simple exchanges), was used as a pseudo ratio for <sup>223</sup>Ra:IMRT absorbed bloo 1 dos e.

<sup>223</sup>Ra exposure was dictated by its unique pharmacokinetic properties and its ability to target calciumdependent bone turnover. Once intravenously administered, <sup>223</sup>Ra is rapidly cleared through the gastrointestinal tract and the remainder through the kidneys (39, 41). Previous studies on rodents found minimal uptake of <sup>223</sup>Ra in non-targeted areas such as kidneys and the spleen (13, 64) with the highest absorbed doses being observed in humans in neighbouring sites of target bone metastases, including osteogenic cells and the red bone marrow (17, 19, 20). Here, the <sup>223</sup>Ra absorbed blood dose was estimated from the injected activity using an existing clearance model, based on the quantity of <sup>223</sup>Ra in circulation at three time points (15 min, 4 h and 24 h post intravenous administration). The

uncertainty for the <sup>223</sup>Ra physical absorbed blood dose estimations per cycle was based on the errors associated with the injected activity, patient weight and clearance models used. The injected activity per patient was estimated from the known activity in the syringe prior to administration and the remaining activity after injection, the error associated with this was considered negligible. Patient individual weights were found to fluctuate during treatment by 8% (up to 6.2 kg lost by C6). Therefore, the <sup>223</sup>Ra absorbed blood dose was estimated for each chosen timepoint from the data available at the specific treatment cycle rather than an average across treatment along with the median values of <sup>223</sup>Ra in the blood at each timepoint (averaged between rep<sup>-</sup>r<sup>t</sup>ed studies). The average absorbed blood dose per cycle was estimated as  $0.012 \pm 0.002$  Gy (aver, ge of Table 2 as per methods 2.3.1). In order to estimate the uncertainty for the clearance model, the reported ranges in terms of upper and lower range limits were taken from the literature (29-51) with a variation from the median of up to 50%. The average uncertainty on the delivered  $^{223}$ Ra  $_{\circ}$  tivity was thus estimated to be as high as 50%. The additional absorbed blood dose delivered after 24 h period was not considered in this model. Although the additional absorbed ble of cose is likely to be within the 50% uncertainty described, it may not be implicitly negligible and does form an additional contribution to the uncertainty not quantified here.

The physical absorbed blood dosc reported here are higher than those reported by Stephan *et al* using <sup>224</sup>Ra and in a similar range to bose reported by Schumann *et al* for <sup>223</sup>Ra (65,66). The decay kinetics and energies of <sup>224</sup>Ra are breadly comparable to those of <sup>223</sup>Ra, however, as Stephan and colleagues report, there are a number or limitations associated with the application of the ICRP 67 model (67), including the lack of information regarding the local distribution of the activity. The aim of the <sup>223</sup>Ra clearance calculations in this work was not to test the ICRP model, rather, to provide a very simple kinetic method for validation of the newly proposed biological absorbed blood dose estimation methods presented. Nevertheless, further work is required to assess the most appropriate means of estimating dose, and this will include reassessment of more detailed kinetic models including those from ICRP.

IMRT utilizes conformal beams to accurately target a designated area minimising exposure to nontarget organs/tissues. Unlike <sup>223</sup>Ra, this treatment is tailored to the disease burden of each patient with the number of fractions, dose and area covered dependent upon patient specific information (Supplementary Table 1). Using this, estimates were carried out employing two models previously described by Moquet and colleagues, which were designed to provide a relatively simple assessment of RT doses (32). The first model, the blood flow model (BF), was based on the time taken for blood to flow through the high dose planned area. Moquet *et al.*, assumed that 6 L of whole blood flows through the high dose region within a 1-min IMRT exposure time (3). For this study, individual patient weights were available, therefore blood volume was more a cur, tely estimated by assuming each kg of body weight contains 75 ml of blood (68),  $(BF_1)$ . The absorbed blood dose per fraction was then estimated to range between 0.159-0.291 Gy and this war equivalent to a cumulative dose of 5.872-8.715 Gy at IMRT treatment completion (see suppleme. tary table 1). Although lymph nodes are in proximity of vascularized tissue, they do not character blood, instead they circulate lymph fluid.  $BF_1$  was therefore likely to overestimate the vm<sup>-</sup> h node absorbed dose. The fluid shift within lymph nodes is approximated to  $\sim 4$  L per day (6.) which may be negligible during treatment. To model the lymph node exposure, the static absorb d ol od dose was calculated (BF<sub>2</sub>) using similar principles, subsequently scaled down to lymph. de planned volume. Lymph node absorbed blood dose per fraction represents 0.004-0.015 G, which was equivalent to 0.152-0.508 Gy by IMRT completion (patient individualised sch dule . This BF2 model was likely to be a more accurate representation of static PBL irradiation in this tissue that may then be filtered to the circulatory system. By combining the two models, the cumulative absorbed blood dose to high dose volume likely was between 1.131-2.470 Gy by completion of individualised IMRT schedule. Neither of these models consider the low dose regions that are also exposed during treatment; therefore, so both will underestimate the total blood dose. The original work of Moquet and colleagues (32) was relatively simple and requires further validation to ascertain exposure circumstances in which such models can be applied. In this study, patient specific data was utilised to refine these models, with limited success. The uncertainties in CTPV absorbed blood dose were derived by propagation of the uncertainties associated with factors in equations 1 and 2. For CTPV physical absorbed blood dose these were the

planned volume (considered minimal at 1-1.5% variation), the dose to planned volume and the estimated scaling factors. The dose to the plan region was expected to have an uncertainty of less than 3%, based on published estimates of the upper bound of inter-treatment dosimetric uncertainty (70) and greater errors would be detected by treatment QA and trigger re-planning and re-validation. To estimate the whole-body volume, it was assumed that 1.01 g of human body mass fits within 1 cm<sup>3</sup> and that the tissue density within the target volume was consistent with this for the scaling factor estimation, with an error within 1%. The total error associated with CTPV estimations was considered to be 6%. It is important to note, however, that the estimates of uncertainty are themselves uncertain and further work is needed here to better understand how patient specific data can contribute to such estimates. In addition, more detail with respect to beam-on times are required to improve the BF dose model, in particular, for larger dose volumes, therefore,  $de_{2}e^{\frac{1}{12}}e^{-\frac{1}{12}}$  use previously, this model was not considered further.

To enable the absorbed blood dose estimation by Coentric assay in this mixed (<sup>223</sup>Ra and IMRT) exposure scenario, the absorbed blood dose ra.<sup>1</sup> between each source was estimated per treatment cycle. This then facilitated use of the "criticality" dose estimation technique, originally designed to separate and quantify neutron and gamma exposures following a nuclear accident or incident (35), but here used to assess the IMRT and topad cicle absorbed doses, on the basis of either the treatment planning information (CTPV 1, tios), or the categorisation of complexity of aberrations observed using M-FISH. The dicentric cosage was applied to two separate models, the first utilizing the ratio derived from physical models, <sup>223</sup>ra clearance and IMRT CTPV models, and the second utilizing the M-FISH<sub>LET</sub> ratio based on the categorisation of high LET to low LET exposed cells. The absorbed blood dose estimations were carried out for 5 patients for which blood samples were drawn and the dicentric equivalent frequency estimated. The CTPV derived ratios assume that IMRT induced aberrations accumulated in the circulatory blood pool by C3 are not cleared from the peripheral pool in the following C4 and C5 treatment cycles and similarly for the <sup>223</sup>Ra, it was assumed the aberrations accumulate through treatment with no clearance. This results in a plateau of IMRT dose for CTPV<sub>1</sub> as the dose ratio was based on a large IMRT component with a small <sup>223</sup>Ra dose. For CTPV<sub>2</sub>, as the

model was based on high dose regions areas, the IMRT estimated dose was lower than CTPV<sub>1</sub>, therefore when expressed as a ratio with <sup>223</sup>Ra, this assumes a higher proportion of absorbed blood dose to be attributed to <sup>223</sup>Ra, which decreases the estimated dose for C4 and C5. The dicentric assay was estimated from the ratio of physical <sup>223</sup>Ra and CTPV dose estimates with a conservative error of 50% being attributed to the <sup>223</sup>Ra dose, this was considered to be the largest uncertainty. Future studies will aim to reduce this uncertainty by increasing the number of patients in the study and by including later timepoints.

The M-FISH<sub>LET</sub> model was based on the chromosomal aberration spectrum in PBL sampled 4 weeks after each [<sup>223</sup>Ra]RaCl<sub>2</sub> administration. Following IMRT completio. th - dose was estimated at 1.167  $\pm 0.092$  Gy. This was based on the assumption that all IMRT indicated aberrations would be of simple type while <sup>223</sup>Ra aberrations were of complex type. As a lerger proportion of cells containing complex aberrations than simple chromosomal aberrations was chserved, this was reflected in the dose ratio. Accordingly, the dose attributed to <sup>223</sup>Ra was proportionally larger than that attributed to IMRT. The resulting IMRT absorbed blood dose estimate.' by M-FISHLET was therefore lower than both CTPV dose estimates. The physically derived <sup>223</sup>Ra <sup>3</sup>stimates were representative of the period taken for <sup>223</sup>Ra to clear from the blood. Due to the compling schedule being every 4 weeks, it cannot be excluded that absorbed dose fron.<sup>223</sup>Ra was also received by circulating PBLs in the vicinity of metastatic sites, especially as nota tatic sites tend to be highly vascularized (71-73). The M-FISH<sub>LET</sub> absorbed blood dose estimates and better account for this, with the <sup>223</sup>Ra dose by C5 estimated to be  $0.665 \pm 0.080$  Gy. This estimate was significantly larger than CTPV<sub>1</sub>  $0.024 \pm 0.027$  Gy and CTPV<sub>2</sub>  $0.141 \pm 0.042$  Gy. The largest uncertainty in the M-FISH<sub>LET</sub> absorbed blood dose estimation was found to be also in C2 whereby the variation in the frequency of cells consistent with IMRT exposure (cells containing simple aberrations only) was up to 13% and for those consistent with <sup>223</sup>Ra exposure of up to 28%. The error from the calibration curve used was estimated to be 12 and 23% for IMRT and <sup>223</sup>Ra absorbed blood dose estimates, respectively. The total propagated error on absorbed dose was within 30% for the M-FISH<sub>LET</sub> derived estimates.

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In this study, the complexity of chromosome exchange observed in PBLs was used as a biomarker of radiation quality (36) from which to make estimates of absorbed blood dose ratio, termed the M-FISH<sub>LET</sub>. Based upon principles of radiation track structure and PBL cell geometry (36, 74), all cells which contained at least one complex chromosome exchange (75, 76) were categorised as having been traversed by high LET  $\alpha$ -particles emitted from the <sup>223</sup>Ra, while damaged cells containing only simple chromosome exchanges (reciprocal translocation, dicentrics and rings) were categorised as being exposed to low LET radiation from IMRT. Although in vitro studies do show the majority of high-LET induced damage to result in complex chromosome aberrations largely independent of dose (77-81) it is also the case that the simple exchanges can be directly induired after  $\alpha$ -particles of lower incident LET (74) and, exposure to low-LET radiation will resu t in he formation of complex exchanges, in a manner strongly dependent upon dose (50 02 82). For instance, an increasing fraction of complex exchanges of up to 20-40% have previous. V been attributed to exposure of a large target field in IMRT treated prostate cancer patien' 3 (4 35). Therefore, it is likely that IMRT absorbed blood dose may be underestimated usir g the M-FISH<sub>LET</sub> reported here. Given the potential usefulness of this ratio in cases of unknown exposures where physical information is not available, further work to determine frequencies of complex exchange occurrence in IMRT only and <sup>223</sup>Ra only treated patients is required.

As the M-FISH<sub>LET</sub> absorbed do. er: tio and the dicentric quantification was based on cytogenetic observations of sampleu <sup>D</sup>b<sup>1</sup> the resulting doses estimated will be directly affected by haematopoietic cell death and repopulation dynamics. The IMRT dose estimates were found to plateau between C3-C5 suggesting cells containing unstable aberrations remain over the time course studied. However, IMRT has been shown to significantly decrease the number of PBLs in circulation, and therefore the clearance and repopulation dynamics should also be taken into consideration for the CTPV models (25, 86, 87). White blood cell counts have been found to increase within 6-8 weeks of therapy completion with a significant increase in lymphocyte population after 3 months (26, 88). The 4-week period between C3-C4 (after end of IMRT) could provide a sufficient break for haematopoietic cells to boost PBL repopulation, which if the case, would have the effect of diluting

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the frequency of persisting unstable chromosomal events. Newly induced aberrations by subsequent <sup>223</sup>Ra treatment cycles would then add to this aberrant cell pool. There is also the potential of bystander responses playing a role both in cell turnover and aberration formation (89). Further work on blood samples representative of later [<sup>223</sup>Ra]RaCl<sub>2</sub> administrations (C6) and follow up samples (up to 1 year post start of treatment), together with patient haematological counts, will help elucidate the cellular dynamics of damaged PBL. An assessment of the occurrence and type of stable chromosome exchange from these samples will also offer the potential to make estimates of absorbed dose delivered to the bone marrow.

In conclusion, in this study we have evaluated a number of absorbed ble of dose methods for mixed field exposure, in a unique population of patients receiving external beam photons and a systemically delivered  $\alpha$ -emitter <sup>223</sup>Ra. We highlight key observations and have number to establish an approach from which we can make dose assessments to better understand mixed field exposures. The models presented provide an initial estimation of cumulative absorbed dose received to the blood during incremental IMRT fractions and [<sup>223</sup>Ra]RaCl<sub>2</sub> injections, all of which move towards assessing patient specific dose information for mixed field treatment to help optimise treatment outcomes and minimise patient risk in the future.

## 5. Acknowledgements

This project received funding from Public Health England, as part of the PHE PhD studentship scheme. This work was supported by the Movember Prostate Cancer UK Centre of Excellence (CEO13\_2-004) and the Research and Development Division of the Public Health Agency of NI (COM/4965/14).

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#### **Figure Legends:**

**Pictorial abstract.** Blood doses were estimated using models of <sup>223</sup>Ra blood clearance and clinical IMRT treatment plans, as well as through a novel application of the M-FISH dicentric assay based on markers of high and low LET radiation exposure.

Figure 1. Treatment timeline. IMRT daily fractions for 7.5 weeks. [<sup>223</sup>Ra]RaCl<sub>2</sub> was administered every 4 weeks for a total of 6 injections. Blood samples were collect d at each cycle, with a control sample collected at week 0 prior to treatment start (C1). Subsequer . blood samples were taken every 4 weeks just prior to the next [<sup>223</sup>Ra]RaCl<sub>2</sub> administration, up to C6. Hence blood samples from C2 and C3 are representative of mixed field exposure while C4-C6 une<sup>-23</sup>Ra only. For this study only samples up to C5 were analysed.

Figure 2. Dicentric equivalent scoring. a) 1<sup>st</sup> division Gremsa stained metaphase imaged under bright field microscopy x100 magnification under on Chromosome with two centromeres highlighted along with acentric fragment b) Pseudo colour processed image painted by M-FISH, captured at x63 magnification under oil. Complex chromosome heat exchange between three chromosomes, dicentric chromosome highlighted by red arrow along with other components in yellow. c) DAPI channel of same cell (b) highlighting same aberration as a simple dicentric exchange.

Figure 3. Frequency of ce.'s containing either simple ( $\bullet$ ) or at least one complex ( $\blacktriangle$ ) chromosome exchange. Data pooled from 5 patients total, n=3 patients/cycle (C2,3 and 5) and n=4 for C3. Error bars represent standard error of the mean for pooled data.

Figure 4. Comparison of <sup>223</sup>Ra dose estimates. M-FISH<sub>LET</sub> ratio was implemented in the dicentric assay estimates of blood dose for n=5 patients (patient ID 1-5), the estimates for these patients were compared across all other models. CTPV<sub>1</sub> and CTPV<sub>2</sub> calculated from the dicentric assay utilizing the physical blood dose estimates by <sup>223</sup>Ra and respective IMRT models. Simple linear regression plotted

for CTPV<sub>1</sub> Y = 0.019\*X + 0.004, R<sup>2</sup> = 0.559 and for CTPV<sub>2</sub> Y = 0.115\*X + 0.025, R<sup>2</sup> = 0.600. Physical <sup>223</sup>Ra dose estimated through clearance of <sup>223</sup>Ra from circulation 24 h post administration. Simple linear regression plotted for Y = 0.028\*X + 0.012, R<sup>2</sup> = 0.301. The error associated with the physical <sup>223</sup>Ra was conservatively estimated to 50% and propagated to CTPV<sub>1</sub>, CTPV<sub>2</sub>. The error for

M-FISH<sub>LET</sub> dose was propagated as 30% for C2 and 14% for C3-C5.



**Graphics Abstract** 

#### **Treatment schedule**



**Blood sampling** 









Figure 3



Figure 4