FEASIBILITY OF SIMULTANEOUS INTEGRATED BOOST FOR HIGH-DOSE TREATMENT OF HIGH-RISK PROSTATE CANCER

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Abstract

Background. Radiation therapy for high-risk prostate cancer presents a challenge for cancer radiotherapists. The improvement of treatment outcomes is associated with radiation dose escalation and prophylactic irradiation of lymph nodes, therefore, the development of the new treatment schemes is needed. Simultaneous integrated boost technique based on the volumetric modulated arc therapy is the most efficient treatment option. Material and Methods. The anatomical data of 10 patients with high-risk prostate cancer was used for dosimetry-based treatment planning. Both simultaneous integrated boost and sequential boost technique were considered. The treatment planning goal was to deliver the equivalent dose of 96 Gy at 2 Gy per fraction (\(E_{\text{eq}}D_2=96\) Gy) (\(\alpha/\beta=1.5\) Gy) to the prostate, \(E_{\text{eq}}D_2=62.5\) Gy to the seminal vesicles and \(E_{\text{eq}}D_2=50\) Gy to lymph nodes avoiding damaging the organs at risk, mainly the bladder and rectum. The irradiation was based on volumetric modulated arc therapy with two partially coplanar arcs and two rotations at each arc. The obtained dose distributions were compared with respect to dose-volume histograms and equivalent uniform doses (EUD). Results. In the case of sequential boost, the minimal dose delivered to the prostate was equal to 95.9 \(\pm\) 2.1 Gy, EUD=104.9 \(\pm\) 1.7 Gy. The dose delivered to 2 cm\(^3\) (\(D_{2\text{cc}}\)) bladder was 97.4 \(\pm\) 2.0 Gy. Normal tissue complication probability (NTCP) was 1.64 %. The dose delivered to 2 cm\(^3\) (\(D_{2\text{cc}}\)) rectum was 103.4 \(\pm\) 9.2 Gy and NTCP was 27.4 %. In the case of simultaneous integrated boost, the minimal dose delivered to the prostate was equal to 90.4 \(\pm\) 2.3 Gy, \(E_{\text{UD}}=103.9 \pm 1.3\) Gy. The bladder dose was as high as \(D_{2\text{cc}}=96.1 \pm 5.2\) Gy, NTCP=0.176 \(\pm\) 0.132 %, the rectum dose \(D_{2\text{cc}}=81.1 \pm 6.0\) Gy, NTCP=2.34 \(\pm\) 1.92 %. Conclusion. Volumetric modulated arc therapy along with simultaneous integrated boost have shown the feasibility of simultaneous irradiation of the prostate, seminal vesicles and lymph nodes up to the prescribed dose values without significant over irradiation of the organs at risk (OARs). Dose values in the tumor as high as \(E_{\text{UD}}=103.9 \pm 1.3\) Gy along with prophylactic irradiation of lymph nodes may result in higher tumor control probability value and should be considered for clinical trials.

Key words: prostate cancer, volumetric modulated arc therapy, hypofractionated radiotherapy, simultaneous integrated boost, high risk.
ОСУЩЕСТВИМОСТЬ ОДНОВРЕМЕННОГО ИНТЕГРИРОВАННОГО БУСТА ДЛЯ ВЫСОКОДОЗНОГО ЛЕЧЕНИЯ РАКА ПРЕДСТАТЕЛЬНОЙ ЖЕЛЕЗЫ ВЫСОКОГО РИСКА

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Аннотация

Введение. Лучевая терапия рака предстательной железы высокого риска является важной проблемой клинической онкологии. Повышение эффективности лечения связано с увеличением подводимой дозы и профилактическим облучением лимфатических узлов, поэтому необходима разработка новых схем лечения. Одновременная интегрированная буст-методика на основе ротационной лучевой терапии с модуляцией интенсивности излучения является способом выполнения обоих условий.

Материал и методы. Анатомические данные 10 пациентов с раком предстательной железы высокого риска были использованы для дозиметрического планирования лечения. Рассматривались как метод одновременного интегрированного буста, так и метод последовательного буста. Целью планирования было облучение предстательной железы до эквивалентной дозы $EQD_2=96$ Гр ($\alpha/\beta=1,5$ Гр), семенных пузырьков ‒ до $EQD_2=62,5$ Гр и лимфатических узлов – до $EQD_2=50$ Гр без негативного влияния на органы риска, в основном на мочевой пузырь и прямую кишку. Облучение проводилось на основе ротационной лучевой терапии с модуляцией интенсивности излучения с двумя частично компланарными арками и двумя поворотами на арку. Полученные распределения дозы сравнивались с гистограммами «доза-объем» и эквивалентной равномерной дозы (eud).

Результаты. В случае последовательного буста минимальная доза в предстательной железе была равна $95,9 \pm 2,1$ Гр, $EUD=104,9 \pm 1,7$ Гр. Доза в мочевом пузыре в 2 см³ ($D_{2cc}$) составила $97,4 \pm 2,0$ Гр, вероятность осложнений со стороны нормальной ткани – $NTCP=1,64 \%$. Доза в прямой кишке составила $D_{2cc}=103,4 \pm 9,2$ Гр, $NTCP=27.4 \%$. В случае одновременного интегрированного усиления минимальная доза в простате была равна $90,4 \pm 1,3$ Гр, $EUD=103,9 \pm 1,3$ Гр. Доза в мочевом пузыре составила $D_{2cc}=96,1 \pm 5,2$ Гр, $NTCP=0,176 \pm 0,132 \%$, в прямой кишке – $D_{2cc}=81,1 \pm 6,0$ Гр, $NTCP=2.34 \pm 1,92 \%$.

Заключение. Ротационная лучевая терапия с модуляцией интенсивности излучения с одновременным интегрированным бустом показала возможность одновременного облучения предстательной железы, семенных пузырьков и лимфатических узлов до предписанных значений дозы без значительного переоблучения органов риска. Значения дозы в опухоли, такие как $EUD=103,9 \pm 1,3$ Гр, наряду с профилактическим облучением лимфатических узлов, могут привести к более высокому значению вероятности контроля опухоли и должны быть рассмотрены для клинических испытаний.

Ключевые слова: рак предстательной железы, ротационная лучевая терапия, гипофракционированная лучевая терапия, одновременный интегрированный буст, высокий риск.

Introduction

The new millennium is notable for significant achievements in the care of cancer patients. The active introduction of modern diagnostic methods into clinical practice, mainly in screening programs, makes it feasible to identify diseases at an early stage. It fully applies to prostate carcinoma [1].

According to P.A. Herzen Moscow Research Institute Institute’s cancer registry data, 45,763 new cases of prostate cancer were diagnosed, and 13,205
Deaths from prostate cancer were registered in Russia in 2019 [2]. The annual prostate cancer mortality rate increases by about 0.63%. Significant number of deaths is caused by the high-risk prostate that challenges development of new treatment schemes.

External beam radiotherapy (EBRT) is widely used for the prostate cancer treatment. According to the Russian standards, a common treatment scheme prescribes the delivery of total doses in the range of 70–80 Gy with doses per fraction equal to 1.8–2 Gy. Different clinical trials demonstrated that dose increase is the key point to effective treatment. Prostate cancer is characterized by the low \( \alpha/\beta \) ratio equal to 1.5 Gy as stated in Refs. [3, 4]. Further, the \( \alpha/\beta \) ratios of the nearest organs at risk (OARs) are equal to \( \alpha/\beta_{\text{blad}}=3 \) Gy for the bladder and \( \alpha/\beta_{\text{rect}}=3.9 \) Gy for the rectum. This fact has become the key point to start the worldwide development and implementation of hypofractionated treatment schemes. For the last 15 years, the stereotactic body radiation therapy (SBRT) of prostate cancer has been thoroughly investigated because of the possibility to deliver a full treatment course in 4–6 fractions [4]. Several clinical trials were conducted using SBRT for high-risk prostate cancer patients who received 32–37 Gy in 4 or 5 fractions (\( EQD_2=85–108 \) Gy), with the 5-year biochemical recurrence-free survival rates of 69–91% [5].

SBRT for high-risk prostate cancer does not allow prophylactic whole-pelvis irradiation that is associated with increase in biochemical recurrence-free survival rates [6]. Sequential boost (SEQ) and simultaneous integrated boost (SIB) are two possible ways to increase dose delivered to the prostate and pelvic lymph nodes. The goal of this study was to compare SEQ and SIB schemes for high-risk prostate cancer treatment in order to increase total dose up to \( EQD_2=96 \) Gy.

### Material and Methods

The anatomical data of 10 patients with high-risk prostate cancer (T2cN0M0–T4N0M0 stage) were selected for this study (PSA nadir of 14.4–30.0 ng/ml and Gleason score in the range from 8 to 10). The patients' tomographic data were obtained using a Toshiba Aquilion LB computer tomograph (CT) with 3 mm slice thickness. Before the CT scanning of an abdominal-pelvic region, the patients were immobilized in the treatment supine position using the Combifix frame [7]. The internal organs were fixed following the recommendations in Ref. [8]. The bladder was filled by drinking a fixed amount of water.

The same clinician delineated three clinical target volumes (CTV) for each patient, namely, the prostate (\( CTV_{pr} \)), seminal vesicles (\( CTV_{sv} \)) and lymph nodes (\( CTV_{ln} \)). The planning target volumes (PTVs) included CTVs and small margin (7 mm for \( CTV_{pr} \) and \( CTV_{sv} \), and 10 mm for \( CTV_{ln} \)). The following radiobiological ratios were assumed: \( (\alpha/\beta)_{pr}=1.5 \) Gy, \( (\alpha/\beta)_{sv}=10 \) Gy, \( (\alpha/\beta)_{ln}=10 \) Gy for the prostate, seminal vesicles and lymph nodes, respectively.

The OARs included the rectum, bladder and femoral heads. Table 1 presents the average volumes, standard deviations and confidence intervals (\( p=0.95 \)) of the delineated structures (contours) (Table 1).

<table>
<thead>
<tr>
<th>Structure/Структура</th>
<th>Average/Среднее ± S.D., cm³</th>
<th>CI, cm³</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTV(_{pr} )</td>
<td>44.5 ± 20.0</td>
<td>[30.1–59.0]</td>
</tr>
<tr>
<td>PTV(_{pr} )</td>
<td>146.0 ± 57.0</td>
<td>[106.0–187.0]</td>
</tr>
<tr>
<td>CTV(_{sv} )</td>
<td>8.1 ± 1.4</td>
<td>[6.9–9.3]</td>
</tr>
<tr>
<td>PTV(_{sv} )</td>
<td>210.0 ± 115.0</td>
<td>[128.0–293.0]</td>
</tr>
<tr>
<td>CTV(_{ln} )</td>
<td>504.0 ± 156.0</td>
<td>[392.0–615.0]</td>
</tr>
<tr>
<td>PTV(_{ln} )</td>
<td>1413.0 ± 263.0</td>
<td>[1225.0–1600.0]</td>
</tr>
<tr>
<td>Bladder/Мочевой пузырь</td>
<td>235.0 ± 92.0</td>
<td>[169.0–301.0]</td>
</tr>
<tr>
<td>Rectum/Прямая кишка</td>
<td>75.0 ± 20.3</td>
<td>[61.0–90.0]</td>
</tr>
</tbody>
</table>

Table 1/Таблица 1

The same clinician delineated three clinical target volumes (CTV) for each patient, namely, the prostate (\( CTV_{pr} \)), seminal vesicles (\( CTV_{sv} \)) and lymph nodes (\( CTV_{ln} \)). The planning target volumes (PTVs) included CTVs and small margin (7 mm for \( CTV_{pr} \) and \( CTV_{sv} \), and 10 mm for \( CTV_{ln} \)). The planning target volumes (PTVs) included CTVs and small margin (7 mm for \( CTV_{pr} \) and \( CTV_{sv} \), and 10 mm for \( CTV_{ln} \)). The following radiobiological ratios were assumed: \( (\alpha/\beta)_{pr}=1.5 \) Gy, \( (\alpha/\beta)_{sv}=10 \) Gy, \( (\alpha/\beta)_{ln}=10 \) Gy for the prostate, seminal vesicles and lymph nodes, respectively.

The OARs included the rectum, bladder and femoral heads. Table 1 presents the average volumes, standard deviations and confidence intervals (\( p=0.95 \)) of the delineated structures (contours) (Table 1).

The following irradiation schemes were prescribed in order to deliver equivalent dose \( EQD_2=96 \) Gy to the prostate. In the case of SIB, the simultaneous delivery of 75 Gy to the prostate (\( PTV_{pr} \)), 62.5 Gy to seminal vesicles (\( PTV_{sv} \)) and 50 Gy to lymph nodes (\( PTV_{ln} \)) at 25 fractions was prescribed. In the case of SEQ, the three-stage delivery was planned: irradiation of all PTVs up to 50 Gy at 25 fractions, 12.5 Gy boost to \( PTV_{pr} \) and \( PTV_{ln} \) at 5 fractions and final 18.4 Gy boost to \( PTV_{pr} \) at 4 fractions. The prescription for \( CTV_{ln} \) was to deliver no less than 98% of the prescribed physical dose to 98% of the contour volume. For other CTVs and PTVs the limitation was equal to 95% of physical dose to 95% of volume.

All treatment plans were simulated using the Monaco treatment planning system (TPS) v5.51 (Elekta Instrument AB, Stockholm) on the Elekta Synergy linac [9] with photon beam energy equal to 10 MV. A VMAT with two partially coplanar arcs and two rotations at each arc was used. The first arc rotated from 190 to 170 degrees clockwise with a collimator angle...
equal to +45 degrees. The second arc rotated from 170 to 190 degrees counterclockwise with the collimator angle equal to −45 degrees. The plan isocenter was situated in the center of PTV\textsubscript{h} structure. The grid spacing was equal to 3 mm. The beamlet width was equal to 3 mm. The statistical uncertainty per calculation based on the ‘Monte Carlo Photon’ algorithm was equal to 0.8%. The minimal segment width was equal to 0.9 cm. The maximal number of control points per arc was equal to 150.

During the treatment planning in TPS, the combinations of dose-volume and biological cost functions were used for better target irradiation and OARs sparing. For all CTVs, PTV\textsubscript{pr} and PTV\textsubscript{ln}, the ‘Target EUD’ function was combined with ‘Quadratic Overdose’ in order to keep maximal value of physical doses within 110% of the prescribed dose values. The function ‘Target Penalty’ was added for the PTV\textsubscript{ln} to spare OARs. The biological functions ‘Serial’ and ‘Parallel’ as well as their combination were used for the OARs allowing minimisation of normal tissue complication probability (NTCP).

The obtained treatment plans were compared with respect to \(EUD\), dose-volume histograms (DVH), equivalent uniform dose (EUD) and NTCP. EUD and NTCP were calculated using the Niemierko approach [10, 11]:

\[
EUD = \left( \sum_i V_i \left( D_i \left( \frac{a_1}{\alpha_1} + D_i/a_2 \right)^{a_2} \right)^{-a} \right).
\]

Here, \(V\) is the part of the target volume irradiated by a dose \(D_i\), \(V = \sum_i V_i\), \(a\) is the specific parameter equal to \(a = -10\) for prostate cancer, \(n\) is the number of fractions. Differential DVHs for the CTVs and PTVs obtained during the Monaco treatment planning were used for the EUD calculation.

The NTCP can be calculated based on the EUD value as follows [10, 11]:

\[
NTCP = \frac{1}{1 + \left( \frac{TD_{50}}{EUD} \right)^{\gamma_{50}}},
\]

where \(TD_{50}\) is the 50% damage dose, i.e. \(NTCP = EUD \equiv TD_{50} = 50\%\), \(\gamma_{50}\) is a parameter that depends on the steepness of the NTCP curve. During the whole-pelvic irradiation of prostate cancer, the rectum and bladder are irradiated to the highest doses if one does not take into account urethra that is naturally included in PTV\textsubscript{pr}. The following parameter values were used for it: \(\alpha/\beta = \approx 3.9\) Gy, \(TD_{50} = \approx 80\) Gy, \(\gamma_{50} = \approx 4\) and \(\alpha_{rec} = 8.33\) [11–13]; \(\alpha/\beta_{blad} = \approx 3\) Gy, \(TD_{50} = \approx 80\) Gy, \(\gamma_{50} = \approx 4\) and \(\alpha_{blad} = 2\) [12].

**Results**

During the dosimetric simulation in TPS both SIB and SEQ treatment plans and dose distributions were developed for anatomic data of each patient. Fig. 1 shows DVHs averaged over all ten patients for contours of CTV\textsubscript{l} (red curve), PTV\textsubscript{pr} (blue curve), bladder (green curve) and rectum (brown curve). The dashed lines show the standard deviations. In Fig. 1 one can see that developed plans are close to each other with respect to irradiation of the tumour. The DVHs for the bladder and rectum have a larger standard deviation due to anatomical peculiarities of the patients.

Table 2 shows the averaged over all patients minimal \(EQD\) dose in the CTVs and PTVs (\(D_{EQD} = 100\%\)) in the form of a mean value, standard deviation and confidence interval at confidence level of \(p = 95\%\). For the CTV\textsubscript{pr} the maximum dose (\(D_{EQD} = 2\%\)) and \(EUD\) values are also presented in Table 2. In the case of SEQ both minimal and maximum doses are the sums of minimal and maximum doses obtained during each irradiation stage, respectively.

The dosimetric treatment planning results presented in Fig. 1 and in Table 2 demonstrate that developed treatment plans delivered dose to the targets very effectively. In the case of SIB, minimal \(EQD\) isdose in CTV\textsubscript{pr} contour amounted 94%. In the case of SEQ, the result is even better and minimal isdose was as high as 100%. Maximal \(EQD\) dose in CTV\textsubscript{pr} contour was almost equivalent for SIB and SEQ techniques and was within 115% of the prescribed \(EQD\) dose. At the same time, physical dose was within 110% limit for all patients that absolutely corresponded to the prescription. Despite the minimal dose difference, the \(EUDs\) for SIB and SEQ were very close (\(EUD_{SIB} = 103.9 \pm 1.3\) Gy and \(EUD_{SEQ} = 104.9 \pm 1.7\) Gy) that allows to expect almost equivalent treatment efficiency according to Niemierko’s tumor control probability (TCP) model [10, 11]. In the case of PTV\textsubscript{pr} irradiation, minimal SIB \(EQD\) dose delivered to 106% of the contour volume amounted 80.3% that coincides to physical dose 87.3% assuming 25 fractions and \(\alpha/\beta = 1.5\) Gy. SEQ minimal dose was higher resulted in 89.3%.

Table 3 presents the OARs irradiation limits according to RTOG recommendations and treatment planning results. Table 3 shows that the developed SIB plans were within RTOG limitations both for the bladder and the rectum. There were several patients with nominal over irradiation at dose levels \(EQD = 74\) Gy for the rectum and \(EQD = 79\) Gy for the bladder. In the case of SEQ, there was no way to calculate total dose-volume irradiation to compare with RTOG limitations. With respect to dose limitation in 2 cm³ of the contour volume, SIB technique demonstrated obtained dose value \(\approx 6\) Gy higher than the limit both for the rectum and the bladder. In the case of SEQ, dose in the bladder was approx. 7 Gy higher than \(D_{EQD}\) limit, while dose in the rectum significantly exceeded the limit. Table 3 also shows that NTCP values for the bladder irradiation were within 5% for both SIB and SEQ as well as for the rectum in the case of SIB. In the case of SEQ, the rectum NTCP value was as high as \(27.4 \pm 10.1\%\) that demonstrated significant probability of the OAR radiation damage.
Table 2/Таблица 2

<table>
<thead>
<tr>
<th>Contour/Контур</th>
<th>SIB, EQD₂ Gy</th>
<th>SEQ, EQD₂ Gy</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTVₚ,ₚ, min dose/CTVₚ,ₚ, мин. доза</td>
<td>90.4 ± 2.3 [88.8, 92.0]</td>
<td>95.9 ± 2.1 [94.4, 97.4]</td>
</tr>
<tr>
<td>CTVₚ,ₚ, max dose/CTVₚ,ₚ, макс. доза</td>
<td>110.6 ± 1.4 [109.6, 111.6]</td>
<td>110.5 ± 2.0 [109.0, 111.9]</td>
</tr>
<tr>
<td>CTVₛ, EUD</td>
<td>103.9 ± 1.3 [103.0, 104.8]</td>
<td>104.9 ± 1.7 [103.7, 106.1]</td>
</tr>
<tr>
<td>PTVₚ,ₚ, min dose/PTVₚ,ₚ, мин. доза</td>
<td>77.1 ± 2.5 [75.3, 78.9]</td>
<td>85.8 ± 3.7 [83.2, 88.5]</td>
</tr>
<tr>
<td>CTVₛ, min dose/CTVₛ, макс. доза</td>
<td>64.9 ± 0.9 [64.3, 65.6]</td>
<td>74.2 ± 6.2 [69.7, 78.6]</td>
</tr>
<tr>
<td>PTVₛ,ₚ, min dose/PTVₛ,ₚ, мин. доза</td>
<td>60.5 ± 2.5 [58.7, 62.2]</td>
<td>60.9 ± 4.1 [58.0, 63.9]</td>
</tr>
<tr>
<td>CTVₛ, min dose/CTVₛ, макс. доза</td>
<td>47.8 ± 0.9 [47.1, 48.4]</td>
<td>48.9 ± 1.0 [48.2, 49.6]</td>
</tr>
<tr>
<td>PTVₛ,ₚ, min dose/PTVₛ,ₚ, мин. доза</td>
<td>44.7 ± 0.8 [44.1, 45.3]</td>
<td>45.0 ± 1.3 [44.1, 46.0]</td>
</tr>
</tbody>
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Table 3/Таблица 3

<table>
<thead>
<tr>
<th>Parameter/Параметр</th>
<th>Bladder/Мочевой пузырь</th>
<th>Rectum/Прямая кишка</th>
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<tr>
<td>EQD₂=59 Gy</td>
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<td>–</td>
</tr>
<tr>
<td>EQD₂=64 Gy</td>
<td>V&lt;50%</td>
<td>22.9 ± 4.8 %</td>
</tr>
<tr>
<td>EQD₂=69 Gy</td>
<td>V&lt;35%</td>
<td>19.9 ± 4.2 %</td>
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<tr>
<td>EQD₂=74 Gy</td>
<td>V&lt;25%</td>
<td>16.8 ± 3.7 %</td>
</tr>
<tr>
<td>EQD₂=79 Gy</td>
<td>V&lt;15%</td>
<td>14.1 ± 3.2 %</td>
</tr>
<tr>
<td>Dₚ,ₚ, Gy</td>
<td>&lt;90Gy</td>
<td>96.1 ± 5.2 Gy</td>
</tr>
<tr>
<td>NTCP %</td>
<td>&lt;5 %</td>
<td>0.2 ± 0.1 %</td>
</tr>
</tbody>
</table>

Note: Dₚ,ₚ – Dose in 2 cm³ of the contour volume.
Примечание: Dₚ,ₚ – доза в 2 см³ объема контура.

Fig. 1. DVHs for developed SIB and SEQ plans. Solid curves show average values, dashed curves show standard deviations.
Рис. 1. DVHs для разработанных планов SIB и SEQ. Сплошные кривые показывают средние значения, пунктирные кривые – стандартные отклонения. Красная кривая – CTVₚ; синяя кривая – PTVₚ; зеленая кривая – мочевой пузырь; коричневая кривая – прямая кишка. Предписанное значение дозы составило EQD₂=96 Гр
Discussion

Dosimetric comparison of SIB and SEQ treatment plans showed that both techniques allowed to deliver the high-value prescribed doses to all targets that included prostate, seminal vesicles and lymph nodes. In the case of SEQ, the minimal dose in the solid tumour $CTV_{\mu}$ and the dose in the $PTV_{\mu}$ were significantly higher than the doses delivered during SIB irradiation ($EQD_{SSW}=95.9\,\text{Gy}$ vs $EQD_{SSW}=90.4\,\text{Gy}$). The payback was the increased level of the OARs irradiation in the case of SEQ. Rectum irradiation was unfavourable in the case of SEQ resulted in $D_{2cc}=103.4 \pm 9.2\,\text{Gy}$ and average NTCP level as high as 27.4 %. SIB irradiation also resulted in nominal over irradiation of the OARs. However, the NTCP values were within 5 %. According to Sumida et al., NTCP values that are lower than 5 % are clinically acceptable following the TD5/5 concept, which is defined as the tolerated NTCP of 5 % within five years after radiation therapy [15]. Despite the difference in minimal doses in $CTV_{\mu}$ contour, the EUD values for SIB and SEQ were almost equal (EUD104 Gy).

Increase in the treatment efficiency of localized high-risk prostate cancer is associated with increase in equivalent dose $EDQ_{\alpha}$ delivered to the solid tumor. The prostate cancer radiobiological ratio is equal to $\alpha/\beta=1.5\,\text{Gy}$ [2, 3]. Such value allows $EDQ_{\alpha}$ dose increase due to hypofractionated radiation that could be delivered in different ways. The first option is to use moderate hypofractionation with doses up to 3 Gy per fraction delivered by EBRT (IMRT or VMAT). Both SIB and SEQ techniques may be used. The second option is the high-dose-rate brachytherapy boost after EBRT. Both first and second options allow irradiation of solid tumor and prophylactic irradiation of lymph nodes. The third option is based on SBRT with high fraction dose that could be used for solid tumor irradiation, only, excluding seminal vesicles and lymph nodes.

Arcangeli et al. presented results of comparison of conventionally-fractionated and hypofractionated EBRT treatment of high-risk prostate cancer [16]. In 2002–2007, 168 patients were treated in randomized trial either conventionally (80 Gy, 40 fraction) or in the hypofractionated mode (62 Gy, 20 fractions – $EQD_{2}=81.5\,\text{Gy}$ at $\alpha/\beta=1.5\,\text{Gy}$). The 10-year biochemical recurrence-free survival rates were 72 and 65 % in hypofractionated and conventional groups, respectively. The overall survival was also higher in the hypofractionated group, being 75 vs 64 %. Thus, a long follow-up period showed that moderate hypofractionation resulted in better prognoses both due to shorter overall treatment time and slightly higher $EQD_{2}$ dose value. Moderately-hypofractionated therapy also showed no significant increase of toxicity. The difference was observed only for Grade 1 long-term GU toxicity [16].

Tamihardja et al. reported long-term outcome of moderately hypofractionated radiotherapy of high-risk prostate cancer based on image-guided IMRT or VMAT [17]. The dose was delivered in the SEQ mode. The doses delivered to the high-dose PTV that included prostate and proximal 2 cm of the seminal vesicles amounted 76.2 Gy in 33 fractions ($EQD_{2}=83\,\text{Gy}$ at $\alpha/\beta=1.5\,\text{Gy}$). Low-dose PTV that included base of seminal vesicles was irradiated up to 60 Gy (1.82 Gy per fraction). Pelvic lymph nodes of the patients were irradiated up to 45.5 Gy (1.82 Gy per fraction) [18]. The 5-year biochemical relapse-free survival was 79.4 % for high-risk disease. The 5-year prostate cancer-specific survival was 89.3 %. Cumulative 5-year late GU toxicity and late GI toxicity grade ≥2 was observed in 26.3 and 12.1 % of the patients, respectively. Cumulative 5-year late grade 3 GU/GI toxicity occurred in 4.0/1.2 % [17].

Yamazaki et al. investigated the dose increase efficiency in the case of very-high risk prostate cancer with the worst prognosis including clinical stage T3b–T4, primary Gleason pattern 5, or more than four biopsy cores with Gleason score 8–10 [19]. Yamazaki compared EBRT with prescribed dose values equal to $EQD_{2}=72\,\text{Gy}$ (group A) with different boost schemes that included either EBRT boost up to $EQD_{2}=80\,\text{Gy}$ or high-dose brachytherapy boost up to $EQD_{2}=129\,\text{Gy}$ (group B). Group B showed superior 5-year biochemical disease-free survival rate (81.2 % vs. 66.5 % at group A). Accumulated late toxicities in gastrointestinal and genitourinary tracts with Grade ≥2 were similar among groups [19]. Yamazaki concluded that both high-dose EBRT and high-dose-rate brachytherapy boosts could be good options for improving the biochemical disease-free survival in T3-T4 stages of localized prostate cancer.

Due to low value of radiobiological ratio $\alpha/\beta=1.5\,\text{Gy}$ for prostate cancer, the possible way to increase dose values in the high-risk tumors is associated with increase of the fractional dose up to 10 Gy per fraction during SBRT treatment. There were several clinical trials that studied SBRT efficiency in the case of high-risk prostate cancer.

Katz and Kang in Ref. [20] presented the data on 65 % rate of disease-free survival for a 8-year follow-up period (7-year disease-free survival amounted 68.2 %) for 35 Gy or 36.25 Gy delivered in 5 fractions ($EQD_{2}=85 – 90.6\,\text{Gy}$). Kang et al. in Ref. [21] showed results of image-guided SBRT of high-risk prostate cancer with the doses equal to 32 – 36 Gy delivered in 4 fractions ($EQD_{2}=87 – 108\,\text{Gy}$). The reported 5-year freedom from biochemical relapse rate amounted 90.9 %. Bernettich et al. reported on 86.7 % of 5-year freedom from biochemical relapse rate for the patients treated with 35 – 37.5 Gy delivered in 5 fractions ($EQD_{2}=85 – 96.4\,\text{Gy}$) [22].

Royce et al. analyzed the prostate cancer SBRT trials and modeled a dose-response tumor control probability for the endpoint of freedom from biochemical relapse [5]. In the case of high-risk cancer, authors analyzed five-year freedom from biochemical relapse.
of 85 patients reported in 3 previously mentioned studies [5]. Royce used Poisson TCP model:  

\[ TCP = 2^{-\exp \left[ \gamma \left( 1 - \frac{EQD_2}{D_{50}} \right) \right]} \]

where the following parameters were found by the approximation of trials data: \( \gamma = 4.50 \) (CI 2.82 – 6.53) and \( D_{50} = 84.2 \) (CI 81.4–86.8) [5]. Fig. 2 shows Poisson TCP model with Royce parameters that is compared with results of clinical trials. The expected efficiencies of the calculated SIB treatment plans are also shown with results of clinical trials. The expected TCP model with Royce parameters that is compared (Katz and Kang [20], Kang [green line] [21] and Bermetich (blue curve) [22].

Blue point shows expected TCP for the prescribed dose value \( EQD_2 = 96 \) Gy. Blue point with rectangular shows the expected TCP value and uncertainty region for the simulated SIB irradiation taking into account \( EUD = 103.9 \pm 1.3 \) Gy

Fig. 2. Comparison of Royce TCP model (black dashed curve) with clinical trials by Katz and Kang (red line) [20], Kang (green line) [21] and Bermetich (blue curve) [22].

Blue point shows expected TCP for the prescribed dose value \( EQD_2 = 96 \) Gy. Blue point with rectangular shows the expected TCP value and uncertainty region for the simulated SIB irradiation taking into account \( EUD = 103.9 \pm 1.3 \) Gy.


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Funding

This work has been partly supported by the Tomsk Polytechnic University Development Programme “Priority 2030”.

SIBERIAN JOURNAL OF ONCOLOGY. 2023; 22(3): 57–65
Conflict of interests
The authors declare that they have no conflict of interest.