Phase I/II trial of definitive carbon ion radiotherapy for prostate cancer: evaluation of shortening of treatment period to 3 weeks

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Background: The purpose of this study was to evaluate the feasibility of a new shortened 3-week treatment schedule of carbon ion radiotherapy (CIRT) for prostate cancer.

Methods: Beginning in May 2010, patients with T1b–T3bN0M0, histologically proven prostate adenocarcinoma were enrolled in the phase II trial of CIRT. Patients received 51.6 GyE in 12 fractions over 3 weeks (protocol 1002). The primary end point was defined as the incidence of late adverse events that were evaluated based on the Common Terminology Criteria for Adverse Events version 4.0. Biochemical failure was determined using the Phoenix definition (nadir + 2.0 ng ml⁻¹).

Results: Forty-six patients were enrolled, and all patients were included in the analysis. The number of low-, intermediate-, and high-risk patients was 12 (26%), 9 (20%), and 25 (54%), respectively. The median follow-up period of surviving patients was 32.3 months. Two patients had intercurrent death without recurrence, and the remaining 44 patients were alive at the time of this analysis. In the analysis of late toxicities, grade 1 (G1) rectal haemorrhage was observed in 3 (7%) patients. The incidence of G1 haematuria was observed in 6 (13%) patients, and G1 urinary frequency was observed in 17 (37%) patients. No G2 late toxicities were observed. In the analysis of acute toxicities, 2 (4%) patients showed G2 urinary frequency, and no other G2 acute toxicities were observed.

Conclusions: The new shortened CIRT schedule over 3 weeks was considered as feasible. The analysis of long-term outcome is warranted.

The morbidity in prostate cancer is the second highest in the world, and the mortality in prostate cancer is the fifth highest in the world (Ferlay et al, 2012). There are many treatment modalities for prostate cancer including radical prostatectomy (Roehl et al, 2004), laparoscopic surgery (Touijer et al, 2009), three-dimensional (3D)-conformal radiotherapy (CRT) by photons (Zelefsky et al, 2008), intensity-modulated radiotherapy (IMRT) (Cahlon et al 2008), low-dose rate (LDR) brachytherapy (Peinemann et al, 2011), high-dose rate brachytherapy (Masson et al, 2012), proton beam therapy (Zietman et al, 2010), carbon ion radiotherapy (CIRT) (Ishikawa et al, 2006), hormone therapy (Souhami et al, 2009), and robot-assisted laparoscopic prostatectomy (Menon et al, 2010). A high dose of 78–80 Gy is generally used in 3D-CRT or IMRT for prostate cancer, and definitive radiotherapy using photons or proton beam requires ~ 8 weeks (Zelefsky et al, 2008; Cahlon et al 2008; Zietman et al, 2010). This treatment duration appears to be one of the disadvantages for patients, and some patients choose radical prostatectomy or LDR brachytherapy because of the
treatment duration. Although treatment duration for radiotherapy tends to increase with the increase in the total irradiation dose, because of the biological characteristics of photons and proton beams, it is not easy to shorten the treatment duration by increasing the dose/fraction (Furushawa et al, 2000). Recent studies have reported stereotactic radiotherapy (SRT) for prostate cancer using 35–50 Gy/5 fractions (Boike et al, 2011; King et al, 2013). However, the efficacy and the safety (in higher dose) have not yet been established.

Carbon ion radiotherapy for prostate cancer has been in use since 1995, and an initial clinical trial was initiated using a schedule of 20 fractions over 5 weeks (Akakura et al, 2004). The clinical trial of CIRT for prostate cancer was initiated with a shorter duration than that typically used with photons. The efficacy and the feasibility were established with that schedule (Tsujii et al, 2005; Ishikawa et al, 2006). A new treatment schedule of 16 fractions over 4 weeks was initiated in 2003. This treatment schedule has also shown favourable outcomes and feasibility despite approximately half the duration compared with that of radiotherapy by photons (Okada et al, 2012). The possibility of shortening the treatment duration is one of the advantages of CIRT; however, it is also one of the challenges of CIRT (Tsujii et al, 2004). The purpose of this study (protocol 1002) was to evaluate the feasibility of a clinical trial using a new shortened treatment schedule of 12 fractions over 3 weeks.

MATERIALS AND METHODS

The primary end point was defined as the incidence of late adverse events. Late adverse events were defined as adverse events occurring in more than 6 months after the start of radiotherapy. Adverse events were evaluated based on the Common Terminology Criteria for Adverse Events version 4.0 (National Cancer Institute, 2009). The secondary end points were (1) biochemical failure-free survival, (2) overall/cause-specific survival, and (3) quality of life. Biochemical failure was determined using the Phoenix definition ( nadir + 2.0 ng ml\(^{-1}\)) (Roach et al, 2006). Biochemical failure-free survival, cause-specific survival, and overall survival were calculated from the day CIRT was started. Risk categories of prostate cancer were defined as follows: low-risk group, initial prostate-specific antigen (PSA) \(< 20 \text{ng ml}^{-1}\) and T1b–T2bN0M0 and Gleason score \(\leq 6\); intermediate-risk group, initial PSA \(< 20 \text{ng ml}^{-1}\) and/or T2cN0M0 and/or Gleason score \(= 7\); and high-risk group, initial PSA \(\geq 20 \text{ng ml}^{-1}\) or T3a/bN0M0 or Gleason score \(\geq 8\). The T stage was determined based on TNM classification 7th edition (Sobin et al, 2009). Clinical stage was determined by digital rectal examination, magnetic resonance imaging, computed tomography, bone scintigraphy, and other diagnostic images. Adverse effects were evaluated by interview of symptoms from patients, urine analysis, stool analysis, cystoscopy, and colonoscopy.

Inclusion and exclusion criteria. Patients who met all the following conditions were included: histologically diagnosed prostate adenocarcinoma, without any previous surgery or radiotherapy for prostate cancer, and T1bN0M0 to T3bN0M0. In addition, the following patients were included in this trial (excluded from another CIRT trial 9904): (1) low-risk patients who underwent hormonal therapy, (2) intermediate/high-risk patients who refused hormonal therapy, and (3) intermediate/high-risk patients who received longer than 6 months of hormonal therapy. Exclusion criteria included the following: (1) having a history of pelvic irradiation; (2) having prognosis for survival of < 6 months; (3) performance status of 3 or 4; (4) receiving other treatments for prostate cancer, with the exception of hormonal therapy; (5) having uncontrolled malignancy(ies) other than prostate cancer; (6) having uncontrolled infectious disease near the irradiation area; (7) medical/psychological/other reasons; and (8) showing PSA close to the lower margin of the PTV. The prescribed dose was 51.2 GyE/12 fr., and >95% of the prescription dose were planned to be irradiated to the CTV. The recommended (not restricted) dose constraints for rectum are V53 GyE (rectal volume to be irradiated 53 GyE) \(\leq 0\%\), V50 GyE \(\leq 7\%\), and V40 Gy E \(\leq 16\%\). Other dose constraints were not defined. The two fields technique (opposing lateral fields) was routinely used for the CIRT planning. Compensators were individually made for each port, and irradiation fields were shaped using multi-leaf collimators. Purgatives or enema was used so that patients have bowel movements at least once a day. All patients were treated with a resinous shell and an image-guided irradiation system under shallow natural breathing, and irradiation was performed 4 days per week. The total irradiation dose was calculated on the basis of the past irradiation schedules. The past protocols of CIRT (63.0 GyE in 20 fractions in protocol 9904[2] and 57.6 GyE in 16 fractions in protocol 9904[3]) showed favourable outcomes with limited toxicity; the total dose of this clinical trial was estimated to be equivalent to the biological effect observed in the past protocols (Okada et al, 2012). Assuming the alpha–beta ratio of prostate cancer is 1.5–3.0 (Akimoto et al, 2005; Fowler, 2005; Khoo, 2005; Pollack et al, 2006), biologically effective dose when \(\alpha/\beta = 1.5\) (BED\(_{1.5}\)) of 63.0 GyE/20 fr. was calculated as 195.3, BED\(_{1.5}\) of 57.6 GyE/16 fr. was 195.8, and BED\(_{3.0}\) of 63.0 GyE/20 fr. was 129.2, and biologically effective dose when \(\alpha/\beta = 3.0\) (BED\(_{3.0}\)) of 57.6 GyE/16 fr. was 126.7. Similarly, BED\(_{1.5}\) and BED\(_{3.0}\) of 51.6 GyE/12 fr. were calculated as 199.5 and 125.6, respectively.

Statistical analysis. Time to event was calculated from the first day of CIRT to the date of the event. The follow-up period was calculated from the first date of CIRT to the date of last follow-up. The Kaplan–Meier method was used for survival analyses.

RESULTS

The phase I/II clinical trial of CIRT in 12 fractions for prostate cancer (protocol 1002) was approved by the research ethics review board of our institution on May 2010, and then the clinical trial was open for enrollment. Although the scheduled registration period was 2 years, registration was closed on October 2011 because the number of patients reached 46. Between May 2010 and October 2011, 282 prostate cancer patients were enrolled in CIRT protocols. Of the 282 patients, 197 patients were eligible for the protocol 9904[3] phase II clinical trial. In all, 46 of 85 patients who did not meet the eligibility criteria for protocol 9904[3], but wished
to participate in clinical trial (protocol 1002) were enrolled in this clinical trial. The remaining 39 patients who did not meet the eligibility criteria for protocol 9904(3) and who did not wish to participate in the protocol 1002 trial were treated in the protocol 0507 G trial with the same dose fractionation as 9904(3) (Figure 1).

The characteristics of 46 patients are shown in Table 1. According to the risk classification, patients with T1c–T2b, T2c, and T3a/b were 25, 6, and 15, respectively. The number of patients with initial PSA < 20 ng ml⁻¹ and ≥ 20 ng ml⁻¹ were 34 and 12, respectively. The number of patients with Gleason score ≤ 6, ≥ 7, and ≥ 8 was 12, 15, and 19, respectively. Finally, 12 patients (26%) were classified as low risk, 9 patients (20%) were classified as intermediate risk, and 25 (54%) patients were classified as high risk. Forty-five patients underwent hormonal therapy, and one of the forty-five patients underwent castration due to liver function failure after LH-RH analogue injection. One patient who was intermediate risk did not undergo hormonal therapy because he refused hormonal therapy.

All 46 enrolled patients completed the scheduled irradiation. The median overall treatment time (OTT) of CIRT was 20 days (range, 16–21 days), and there were no treatment interruptions or treatment delays. The median follow-up period of surviving patients from the start of CIRT was 32.3 months (range, 23.5–38.9 months), and the median follow-up period of surviving patients from the start of treatment including hormonal therapy was 40.0 months. Two deaths were observed during the follow-up period. There were no deaths due to disease and no treatment-related deaths. The two deaths were due to intercurrent disease. One of the intercurrent deaths was in a 64-year-old patient who had hypertension, hyperlipidaemia, diabetes mellitus, hyperuricaemia, cardiovascular disease, and an abdominal aortic aneurysm (AAA). This patient suddenly died from AAA rupture 4 months after CIRT without elevation of PSA. No relationships were found among CIRT, hormonal therapy, and the death; therefore, this case was determined to be intercurrent death without recurrence. A 78-year-old patient who had been diagnosed with interstitial pneumonitis (IP) before treatment for prostate cancer also died. Although IP was stable during the treatment for prostate cancer, the patient died from exacerbation of IP and pneumothorax 24 months after the start of hormonal therapy and 17 months after CIRT, without elevation of PSA. No relationships were found between hormonal therapy, CIRT, and exacerbation of IP; therefore, this case was also determined as intercurrent death without recurrence.

The acute and late toxicities observed with CIRT are shown in Table 2. The most frequent acute toxicity, grade 1 (G1) urinary frequency was observed in 34 out of 46 (74%) cases. Grade 2 (G2) urinary frequency was observed in 2 out of 46 (4%) cases. One of the patients with G2 urinary frequency had been diagnosed with benign prostatic hypertrophy before being diagnosed with prostate cancer, and the patient had pollakiuria before CIRT. The urine sample of the other patient tested positive for bacteria; therefore, G2 urinary frequency was considered to be due to CIRT and a complication of urinary infection. No grade ≥ 3 toxicities were observed. The most frequent late toxicity observed was G1 urinary frequency in 17 out of 46 (37%) cases. The second most frequent late toxicity was G1 haematuria observed in 6 out of 46 (13%) cases. The median time to occurrence of G1 haematuria was 20.7 months (range, 3.5–34.4 months). Grade 1 rectal haemorrhage was observed in 3 (7%) patients and G1 proctitis was observed in 2 (4%) patients.

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**Figure 1. Chart of the trial. The patients who did not meet the inclusion criteria of 9904(3) (CIRT over 4 weeks) and who wished to participate were enrolled in the study (1002).**

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**Table 1. The characteristics of 46 patients**

| Enrolled patients | 46 |
|-------------------|----|
| Gender            | Male 46 (100%) |
| Age (years)       | Median 66 (range 54–83) |
| Stage (UICC2009)  | T1c 17 (37%)  |
|                   | T2a 5, T2b 3, T2c 6 (30%) |
|                   | T3a 13, T3b 2 (33%) |
| Initial PSA (ng ml⁻¹) | Median 11.1 (range 3.559–242.7) |
| Gleason score     | 3 + 3 = 6 (12%) |
|                   | 3 + 4 = 7 (6%) |
|                   | 3 + 5 = 8 (9%) |
|                   | 4 + 4 = 8 (17%) |
|                   | 4 + 5 = 9 (24%) |
| Risk              | Low 12 (26%)  |
|                   | Intermediate 9 (20%) |
|                   | High 25 (54%) |
| ADT               | With ADT 45 (98%) |
|                   | Castration 1 (2%) |
|                   | Without ADT 1 (denied) (2%) |
| Median (range) duration of ADT by risk (months) | Low risk 7.7 (5.5–25.2) |
|                   | Intermediate risk 10.0 (7.3–28.3) |
|                   | High risk 26.7 (9.7–52.9)* |

*Abbreviations: ADT = androgen deprivation therapy; PSA = prostate-specific antigen.

*One patient who underwent castration was excluded (duration of ADT 3.5 months).*
Follow-up period of surviving patients | Median (months) (range) | 32.3 (23.5–38.9)
---|---|---
Survival | Alive | 44 (96%)
| Dead | 2 (4%)
Cause of death | Intercurrent death | 2 (100%)
| Death from primary disease | 0 (0%)
| Treatment-related death | 0 (0%)

### Grade

#### Acute toxicity by risk

| Toxicity | Low | Intermediate | High |
|---|---|---|---|
| Skin (radiation dermatitis) | 46 (100%) | 0 (0%) | 0 (0%) |
| Rectum (haemorrhage) | 43 (94%) | 3 (7%) | 0 (0%) |
| Rectum (proctitis) | 45 (98%) | 1 (2%) | 0 (0%) |
| GU (haematuria) | 40 (87%) | 6 (13%) | 0 (0%) |
| GU (urinary frequency) | 29 (63%) | 17 (37%) | 0 (0%) |
| GU (urinary stricture) | 46 (100%) | 0 (0%) | 0 (0%) |

#### Late toxicity by risk

| Toxicity | Low | Intermediate | High |
|---|---|---|---|
| Skin (radiation dermatitis) | 46 (100%) | 0 (0%) | 0 (0%) |
| Rectum (haemorrhage) | 43 (94%) | 3 (7%) | 0 (0%) |
| Rectum (proctitis) | 45 (98%) | 1 (2%) | 0 (0%) |
| GU (haematuria) | 40 (87%) | 6 (13%) | 0 (0%) |
| GU (urinary frequency) | 29 (63%) | 17 (37%) | 0 (0%) |
| GU (urinary stricture) | 46 (100%) | 0 (0%) | 0 (0%) |

### Abbreviation: GU – genitourinary. Acute grade 2 urinary frequency was seen in two patients (4%). There were no grade 2 or more toxicities in the late toxicity.

1 (2%) patient. No G2 late toxicities were observed during the follow-up period.

Treatment outcomes are shown in Table 3. There were no biochemical failures (PSA nadir + 2.0) during the follow-up period. Almost all patients showed a good response to the hormonal therapy, and the PSA nadir was <0.1 ng ml⁻¹ in 43 out of 46 (94%) patients. Two (4%) patients had a PSA nadir in the range of 0.1–1.0 ng ml⁻¹, and 1 patient who refused to undergo hormonal therapy had a PSA nadir of 1.91 ng ml⁻¹. At the last follow-up date, 26 (57%) patients had rising PSA (≥0.1 ng ml⁻¹) after cessation of hormonal therapy, and the remaining 20 patients, including 5 patients undergoing hormonal therapy. One patient who underwent castration did not show rising PSA. None of the patients showed rising PSA during the hormonal therapy.

### DISCUSSION

**Gastrointestinal adverse events.** In definitive radiotherapy for prostate cancer, late adverse effects tend to be more problematic than acute adverse effects. The rate of ≥G2 rectal toxicity after radiotherapy has been reported as 5.1% in the analysis of patients who enrolled in the RTOG 75-06 and 77-06 trials (Pilepich et al, 1984). However, a lower total dose of radiation of ~65 Gy was used in those trials, so their results cannot be compared with recent data. The results of a randomised controlled trial by Pollack et al (2002) suggested that the rates of ≥G2 rectal toxicity were 12% in patients who received 70 Gy/35 fr. and 26% in patients who received 78 Gy/39 fr., and the rate of rectal toxicity was significantly higher in the high-dose group. Those results may be higher than those of recent studies because the trial was performed from 1993 to 1998, before image-guided radiotherapy (IGRT) or IMRT had been developed. Vargas et al (2005) analysed 331 patients treated with a median dose of 79.7 Gy by 3D-CRT and reported that the rate of ≥G2 rectal toxicity was 20% and the rate of G3 rectal toxicity was 4% at 3 years. Rectal toxicity has improved compared with the report of Pollack et al, despite using a higher total dose of radiation, seemingly due to development of radiotherapy techniques. In the analysis of 743 patients treated with 75.6–81.0 Gy by 3D-CRT at Memorial Sloan-Kettering Cancer Center, it was reported that the incidence of ≥G2 rectal toxicity was 15% (Zelefsky et al, 1999). Although the incidence of rectal toxicity is approximately twice as high as that in the lower dose group (64.8–70.2 Gy), the rate of rectal toxicity has been improving. In a subsequent trial at the same institution that compared 3D-CRT and IMRT using a total dose of 81 Gy, the rate of rectal toxicity of IMRT was significantly lower than that of 3D-CRT (2% vs 14%) (Zelefsky et al, 2001). A similar result has been reported in the study on GI toxicity comparing 79.2 Gy by IMRT and 79.2 Gy by 3D-CRT (≥G2 GI toxicity at 3 years; 15% vs 22%, P = 0.039) (Michalski et al, 2013). Although dose escalation is inevitable from the viewpoint of treatment outcomes for prostate cancer, IMRT or IGRT allows for a high dose of radiation with feasible toxicity.
In the present phase II/III study of hypofractionated CIRT, the incidence of G1 late rectal toxicity was 7% (3 out of 46) and that of G2 late rectal toxicity was 0%. There were no acute rectal toxicities. The dose distribution of CIRT using Bragg-peak is better than that of photons, and the dose distribution of CIRT with less beam scattering is also better than that of protons. The ability to easily decrease the irradiated rectal volume is considered to be one of the reasons for lower rectal toxicity.

**Genitourinary adverse events.** Frequent GU adverse events include haematuria, urinary stricture, and urinary frequency. The incidence of ≥G2 late GU toxicity was ~5% in the RTOG 75-06 and 77-06 trials (Pilepich et al., 1984). In the aforementioned phase III clinical trial reported by Pollack et al. (2002), the rates of G2 and G3 late GU toxicities in the 151 patients in the 78 Gy group were 10% and 3%, respectively, whereas the rates of G2 and G3 late GU toxicities in the 150 patients in 70 Gy group were 7% and 1%, respectively. The GU toxicity tended to increase according to the irradiation dose, but there were no significant differences. The study at the Memorial Sloan-Kettering Cancer Center reported that the rates of ≥G2 late GU toxicity of the high-dose group (75.6–81.0 Gy) at 3 and 5 years were 10% and 15%, respectively (Zelefsky et al., 1999). In the following study comparing IMRT and 3D-CRT in 1100 patients, the rates of G2 and G3 late GU toxicities were 13% and 1.5%, respectively, in the group who received ≥75.6 Gy (Zelefsky et al., 2001). Unlike the rates of rectal toxicity, there was no significant difference between 3D-CRT group and IMRT group in the late GU toxicity (P = 0.32). In a recent study, ≥G2 GU toxicity at 2 years of 3D-CRT group (76 Gy) and IMRT group (78 Gy) were 40% and 30%, respectively (P = 0.011) (Sweitzer et al., 2014). Although significant difference was shown in GU toxicity in that study, the difference is smaller than that of rectal toxicity. In another study on SRT for prostate cancer, a total of two patients (7%) show grade 3 genitourinary (GU) toxicity in the arm of 47.5 and 50 Gy in five fractions (Boike et al., 2011). Further, because the median follow-up time of 47.5-Gy group and 50-Gy group is 18 and 12 months, respectively, the incidence of late adverse effect may increase in the future. It is believed that sufficient data are needed when performing the short-term irradiation by X-rays.

In the present study of CIRT, the rate of ≥G2 late GU toxicity was 0% (Table 2). Grade 1 haematuria was observed in 6 (13%) patients and G1 urinary frequency was observed in 17 (37%) patients. None of these patients required any treatment, and symptoms were improved with observation. No other ≥G2 late toxicities were found. However, GU late toxicity sometimes occurs ≥36 months after radiotherapy; therefore, the follow-up period of the present study was not sufficient to observe all late GU adverse events (Zelefsky et al., 1999). However, because the long-term incidence of late GU toxicity can be predicted from the incidence at 2–3 years (Zelefsky et al., 1999), the rate of late GU toxicity of this phase II/III CIRT study is highly favourable.

**Biochemical failure and metastases.** There were no biochemical failures or distant metastases as of the last follow-up date. The PSA nadir of almost all patients (94%) showed good response to the treatment (<0.1 ng ml⁻¹), which may be due to the fact that all but one patient received hormonal therapy. Hormonal therapy to the low-risk patients may be overtreatment, but this trial was designed to be able to include these patients. Twenty-six of forty-six (57%) patients showed rising PSA ≥0.1 ng ml⁻¹ after cessation of hormonal therapy, but there were no patients with rising PSA >2.0 ng ml⁻¹ above the PSA nadir. Several studies have reported on PSA bouncing after radiotherapy or PSA rising after hormonal therapy (Crook et al., 2007; Pinkawa et al., 2007). It has been reported that long-term PSA rising after hormonal therapy is a concern for the recovery of serum testosterone level (Pickles et al., 2002). After completion of CIRT, the PSA level was not so different from that in patients treated with radiotherapy by photons and hormonal therapy. Long-term outcomes will be reported in the future.

**Treatment period.** The standard total dose of definitive radiotherapy for prostate cancer by photons is considered to be ≥70 Gy. In the EORTC (European Organisation for Research and Treatment of Cancer) 22863 trial, 70 Gy/35 fr. over 7 weeks was used (Bolla et al., 2010). After which, the total dose was increased to 75.6–81.0 Gy for intermediate- or high-risk patients. These treatments require 8.5–9 weeks to complete the irradiation (Zelefsky et al., 2001). In proton therapy as well as photon radiotherapy, 7.5 weeks are required for the completion of the treatment of 74 GyE, and ≥8 weeks are required for the dose escalated treatment using 82 GyE (Slater et al., 1998; Goen et al., 2011). Although there are few studies that evaluated OTT of radiotherapy and prognosis in the treatment of prostate cancer, a significant relationship between OTT of radiotherapy and recurrence rate has been reported in a recent study (D’Ambrosio et al., 2008). Although there were no significant differences between the short-OTT group and the long-OTT group in the patients who received >74 Gy, patients in the long-OTT group tended to have a higher rate of biochemical failure than the patients in the short-OTT group (Liu et al. and Liu and Liu, 2001). From these studies, it can be said that dose escalation contributes to the improvement of local control; however, extension of the treatment period due to the increase in dose is considered to be unfavourable for local control. Recently, SRT for prostate cancer has been suggested as short-term external body radiotherapy (Madsen et al., 2007). A study concluded that SRT could be completed with a schedule of five fractions with tolerable toxicity. However, the biological treatment effects and adverse effects can be different from conventional dose/fraction because dose/fraction has a large impact on biological effect in radiotherapy by photons. But the long-term follow-up results of SRT have not been reported.

Although the median follow-up period was 31 months, the results of the present study demonstrated feasibility and local control at this time point. More than 1600 prostate cancer patients

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**Table 3. Biochemical failure and the changes of serum PSA level**

| Biochemical failure | Failure (−) | 46 (100%) |
|---------------------|-------------|-----------|
| PSA nadir + 2.0     | Failure (+) | 0 (0%)    |
| PSA nadir           | Median      | 0.01 ng ml⁻¹ |
| <0.01               | 20 (44%)    |
| <0.01 to <0.1       | 23 (50%)    |
| <0.1 to <1.0        | 2 (4%)      |
| <1.0 to <2.0        | 1 (2%)      |
| <2.0                | 0 (0%)      |
| PSA value at latest follow-up date | <0.01 | 7 (15%) |
| | <0.01 to <0.1 | 12 (26%) |
| | <0.1 to <1.0  | 23 (50%) |
| | <1.0 to <2.0  | 4 (9%)     |
| | <2.0          | 0 (0%)     |
| PSA rise ≥0.1       | +           | 26 (57%)  |
| | −             | 20 (43%)   |
| Local recurrence    | 0 (0%)      |
| Distant metastasis  | 0 (0%)      |

Abbreviation: PSA = prostate-specific antigen. Although almost all patients show PSA rise after termination of ADT, none of the patients showed biochemical failure.
have been treated with CIRT in the > 18 years it has been available. It has been reported that the long-term outcome of CIRT was very favourable even in high-risk prostate cancer (Tsujii et al., 2005). Compared with other EBRTs, with the exception of SRT, CIRT for prostate cancer has steadily shortened the treatment time while maintaining good local controllability. The biological characteristics of CIRT are considered to be one of the advantages of this modality. The biological effect of the carbon ion beam is less susceptible to dose/fraction than that of photons (Kanai et al., 1999). However, ~ 2 months of hospital visits and admissions for prostate cancer patients is one of the disadvantages. Shortening the treatment period is favourable not only for treatment outcome but also for quality of life of the patients.

CONCLUSIONS

The results of the phase I/II clinical trial shortening the treatment period of CIRT for prostate cancer have been reported. The treatment period of irradiation has been shortened from 4 weeks to 3 weeks without occurring ≥ G2 adverse events with a median follow-up of 32 months. Long-term outcomes (e.g., biochemical failure-free survival and local control rate) of the 3-week treatment period are required.

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