Selective Bladder Preservation Using Chemo-Radiotherapy in Treatment of Muscle Invasive Bladder Cancer

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Abstract

Background: This study evaluated the outcomes of patients with muscle-invasive bladder cancer (MIBC) stage T2-4a managed by tri-modality therapy by trans-urethral resection (TURB) and concomitant chemo-radiotherapy (CRT).

Aim of Study: We were aiming for preservation of bladder.

Patients and Methods: This study was a prospective randomized clinical trial including 43 patients with MIBC (T2-T4a N0) presented to Clinical Oncology Department, Mansoura University Hospital & Urology center during the period from 5/2008 to 9/2010. Patients were randomized to 2 arms: (I): Included 24 patients, who underwent TURB then concomitant CRT using cisplatin plus paclitaxel, while (II): Included 19 patients received same protocol with cisplatin and 5. Flurouracil. All patients who showed complete response (CR) after induction and consolidation phases were given adjuvant chemotherapy 4 cycles every 3 weeks.

Results: In arm I: 65.2% completed the treatment protocol. In arm II: 63.2% completed the treatment protocol. In arm I: 82.6% showed complete response (CR), 4.3% showed disease progression (DP) & 13% had only partial response (PR). Patients who achieved CR entered the consolidation phase of treatment. In arm II, 78.9% showed CR, 21.1% showed PR. The PFS was 78.26% & 68.42% for both arms respectively. The 3-year OS was slightly better for arm II. The 3-year OS was 60.87% & 68.42% for both arms respectively.

Conclusions: Bladder preservation is a good choice for treatment of MIBC but with good selection of the cases and careful follow up to avoid major toxicities which lead to interruption or stoppage of the preservation protocol.

Key Words: Bladder cancer – Tri-modality treatment – Chemoradiotherapy – Bladder preservation.

Introduction

BLADDER cancer (BC) is the 4th most common malignancy among men in the Western world & accounts for approximately 5-10% of all cancers in Europe and U.S [1]. Bladder cancer (BC) is the ninth most common cancer worldwide. More than 60% of all BC cases and half of deaths occur in less developed regions of the world [2]. Men to women ratio of bladder cancer is 3:1 approximately [3]. According to the results of the National Population-Based Registry Program of Egypt 2008-2011; BC was the 2nd most common cancer in males & the 3rd in both sexes (10.71 & 6.94% respectively) [4].

In general, Muscle-invasive bladder cancer (MIBC) constitutes about 30% of newly diagnosed cases, with about 70% being non-invasive (Ta, Tis, T1). About 20-30% of non-invasive BC cases progress to invasive cancer after transurethral resection of bladder tumor (TURB) [5].

Since the 1970s, radical cystectomy with bilateral pelvic lymph node dissection (PLND) has been the standard treatment of muscle-invasive bladder cancer and this supported by many organizations and guidelines such as National Comprehensive Cancer Network (NCCN) and European Association of Urology (EUA) guidelines [6]. Based on data from multicentric randomized controlled trials (RCTs); the role of neoadjuvant chemotherapy before cystectomy is supported for T2,3,4a lesions with negative lymph node (LN) involvement. Neoadjuvant cisplatin-based chemotherapy increases the median survival and lower the recurrence rate without increased treatment related morbidity or mortality [7].

Meta-analysis of RCT reported an overall survival (OS) and disease-specific survival (DSS) benefit in patients with MIBC receiving adjuvant cisplatin-based chemotherapy after RC. The disease
free survival (DFS) benefit was more obvious in patients with node positive. However the evidence of its use was not strong. These meta-analysis recommended further RCT with adequate sample size [8]. Obviously, bladder preservation is only accepted in the presence of a high cure option with no compromisation of survival expectancy [9]. Many investigations focus on optimizing radiation techniques and incorporating more effective systemic chemotherapy and the proper selection of patients based on molecular markers [10].

As the use of combined modality treatment for MIBC has matured, the opportunity for bladder preservation has developed. Preoperative radiation when combined with cisplatin alone, cisplatin and 5-fluorouracil or cisplatin and paclitaxel results in the down-staging to T0 of a significant proportion of patients when TURB, radiation and multi-agent chemotherapy are combined, CR rates of 70% or greater have been achieved. The radiation sensitizing effects of cisplatin have been long recognized, and the opportunity to safely enhance this effect by the simultaneous administration of a second radiation sensitizer such as 5-Fluorouracil or paclitaxel has been a goal of RTOG protocols since 1995 [11].

Patients and Methods

This study is a prospective randomized clinical trial which included 43 patients with muscle invasive bladder cancer (MIBC); T2- T4a N0 M0 who presented to Clinical Oncology & Nuclear Medicine Department, Mansoura University Hospital and Urology Center during the period from May 2008 to September 2010.

Selection criteria:
- Operable patients with muscle invasive bladder cancer, all histotye types.
- AJCC Stages T2-T4a N0 M0.
- ECOG performance scale 0-1.
- Patients must have a transurethral resection of the bladder tumor (TURB) as judged safely as possible.
- Hemoglobin level (Hgb) ≥10mg/dl; white blood cells count (WBC) ≥4000/ml and Platelet count ≥100,000/mm3.
- Serum creatinine <1.5mg%; creatinine clearance of 60ml/min or greater.
- Treatment must begin within 8 weeks following transurethral resection of bladder tumor and endoscopic evaluation.
- Patients must sign study specific informed consent form prior to study entry.

Pre-treatment evaluation:
- History was taken from all patients. They were examined physically including weight, height and surface area. Performance status was assessed.
- Investigations included:
  - Laboratory studies: Complete blood count, liver function tests, serum creatinine, creatinine clearance and serum alkaline phosphatase.
  - Radiological evaluation: Chest X-ray, CT or MRI and bone scan.
  - Endoscopic evaluation by cystoscopic examination, bimanual examination under anesthesia, multiple punch biopsies and complete transurethral resection as thorough as possible.
  - Pathological examination of the biopsy taken by endoscope for pathological type, grade and depth of muscle invasion and urine cytology examination.
- Stage of disease was determined according to AJCC staging (2002).
- Assessment of the performance status was done according to Eastern Co-Operative Oncology Group (ECOG).

Treatment:

For patients with T2-T4a N0 M0 MIBC and who were candidates for radical cystectomy, TURB was done then induction and consolidation CRT were received in the form of paclitaxel, cisplatin, and irradiation (TCI) in Arm I or 5-FU, cisplatin, and irradiation (FCI) in Arm II. The irradiation used was accelerated hyper-fractionation for the tumor with a standard dose schedule for the pelvis.

Surgery:

TURB as much as is judged safely possible then after induction chemo-radiotherapy endoscopic response evaluation was done in week 7 following the completion of the induction CRT by multiple punch biopsies. If there was response in the form of no malignancy (T0), papillary tumor (Ta) or carcinoma in situ (Tcis), the patient would enter consolidation phase. If there was no response in the form of T1 or more, the patient was converted to radical cystectomy at week 9.

Chemo-radiotherapy:

Induction phase: All patients after TURB entered in induction phase of concomitant CRT within 8 weeks from TURB. Patients were randomized to one of two treatment arms for 3 weeks (week 1-3):
Arm I: Patients received concomitant CRT in the form of:
- Paclitaxel 50mg/m² (Day 1, 8, 15).
- Cisplatin 15mg/m² (Day 1 → 3 , 8 → 10, 15 → 17).
- Accelerated hyperfractionated RT, two sessions per day in the form of 1.6Gy to small pelvic fields then 1.5Gy boost to whole bladder 4-6 hours apart (day 1 → 5, 8 → 12, 15 → 17) i.e: bid x13 days (26 fractions with a total dose of 40.3Gy).

Arm II: Patients received concomitant CRT in the form of:
- 5-FU 400mg/m² (Day 1 → 3, 15 → 17 continuous I.V infusion).
- Cisplatin 15mg/m² (Day 1 → 3, 8 → 10, 15 → 17).

The radiotherapy given was the same as in arm I inclusive.

Consolidation phase: All patients who showed response to induction treatment after inter-assessment entered in a consolidation phase of concomitant CRT at week 8 for 2 weeks. Each patient continued with his original arm for 2 weeks; week 8, 9.

Arm I: Patients received concomitant CRT in the form of:
- Paclitaxel 50 mg/m² (Day 1, 8).
- Cisplatin 15 mg/m² (Day 1 → 3, 8 → 10).
- Accelerated hyperfractionated RT, two sessions per day with a dose of 1.5 Gy to small pelvic fields then 1.5 Gy again to small pelvic fields 4-6 hours apart (day 1 → 5, 8 → 10) i.e: bid x8 days (16 fractions with a total dose of 24Gy).

Arm II: Patients received concomitant CRT in the form of:
- 5-FU 400mg/m² (Day 1 → 3, 8 → 10 continuous I.V infusion).
- Cisplatin 15mg/m² (Day 1 → 3, 8 → 10).

The radiotherapy given was the same as in arm I inclusive.

Adjuvant phase: All patients who ended consolidation phase or those who underwent radical cystectomy entered the adjuvant phase after one month from their last line of treatment whatever it was chemo-radiotherapy or radical cystectomy.

Adjuvant phase included 4 cycles of chemotherapy which were given 3 weeks apart. Chemotherapeutic agents were gemcitabine, paclitaxel and cisplatin according to the following protocol:
- Gemcitabine 1000mg/m² (Day 1, 8).
- Paclitaxel 50mg/m² (Day 1, 8).
- Cisplatin 35mg/m² (Day 1, 8).

Radiotherapy technique:
- Radiotherapy given during induction:
  Treatment schedule: External beam irradiation, 1.6 Gy, was delivered to the small pelvic field in the first RT session followed by an inter fraction period of at least 4-6 hours. During the second RT session, 1.5 Gy was delivered to the whole bladder with a safety margin. The bladder was full before the treatment session to the small pelvic fields and was empty before the treatment session for the whole bladder field.

Target volumes:
- Small pelvic fields:
  The field included the whole bladder, the gross tumor volume, the prostate and the prostatic urethra, and the lymph nodes immediately adjacent to the bladder. The fields were designed using a simulator with the patient having 40 to 50ml air contrast cystogram (20-30ml dye + 20ml air).

  The combination of four shaped anterior, posterior, and lateral fields were used. In the cranial-caudal dimension, the planning target volume (PTV) extended from the lower pole of the obturator foramen to the anterior aspect of the S1-S2 junction. In the anterior and posterior pelvic field, PTV width extended 1.5cm lateral to the bony margin of the pelvis at its widest point. The anterior and posterior fields had been shaped with inferior corner blocks, which shielded the medial border of the femoral heads. For the two parallel-opposed lateral fields, the anterior boundary of the PTV was 1.0cm anterior to the most anterior portion of the bladder mucosa seen on the air contrast cystogram. Posteriorly, the PTV extended at least 1.5cm posterior to the most posterior portion of the bladder or 1.5cm posterior to the bladder tumor mass if it was palpable or identifiable on the pelvic CT scan.

- Whole Bladder fields:
  These fields included the whole bladder plus safety margin 1.5cm all around and were designed during the same simulation with the same air contrast cystogram. Three field techniques were used for the bladder boost in the form of one anterior field and two direct lateral wedged fields or two lateral posterior wedged fields.

- Radiotherapy given during consolidation:
  Consolidation therapy started 7-14 days following a cystoscopic re-evaluation demonstrating a
complete response to the induction therapy. A dose of 1.5 Gy (per fraction) was given to the small pelvic field in two RT sessions per day, with an interfraction period of at least 4-6 hours.

The previously simulated small pelvic fields were treated during the consolidation phase.

Radiation dose specification:

The induction radiotherapy course delivered 20.8 Gy to the small pelvic fields and 40.3 Gy to the bladder with the tumor and safety margin. The radiation given during the consolidation treatment was 24 Gy to the pelvis and the whole bladder. Radiation given in both phases resulted in a total dose to the bladder with the tumor volume and safety margin of 64.3 Gy over 8 weeks in 42 fractions and a total dose of 44.8 Gy to the pelvic lymph nodes.

Patients follow-up:

• During treatment:

Patients were followed up during radiotherapy weekly for acute radiation toxicity according to World Health Organization (WHO) for small bowel and urinary tract toxicity. Early effects were recorded weekly during treatment and after 4 weeks.

Patients were followed up during chemotherapy for symptoms & sign of toxicity either hematological or non-hematological toxicity using WHO cancer toxicity criteria for grades of toxicity.

• After the end of treatment:

Patients were followed up every 1-2 month by clinical examination, every 3 months by abdomi- pnal CT or MRI and endoscopy after end of treatment when needed.

Treatment regimens were compared together. Study primary end point was to evaluate response rate after induction phase of treatment. The secondary endpoints were to evaluate the overall survival & progression free survival for three years, failure rates (local and distant) and treatment toxicities.

Statistical analysis:

Data entry and analyses were performed using SPSS statistical package version 10 (SPSS, Inc., Chicago, IL, USA). The quantitative data were presented as a mean, standard deviation, median and range. Student t-test was conducted to compare the mean of continuous variable for two different groups of individuals. The qualitative data were presented as number and percentage. The chi-square ($\chi^2$) was used to find the association between variables of qualitative data. Kaplan-Meier Survival Analysis was used to find out overall survival and progression free survival. Relative risk and 95% confidence intervals were calculated for factors affecting response. The p-value of $\leq 0.05$ and $< 0.001$ indicate significant and highly significant results respectively at confidence interval 95%.

Results

Out of the 43 eligible patients who entered the study, one patient died before completion of the induction phase due to unrelated cause and was excluded from the study.

Patient characteristics: Characteristics of the patients and tumors are summarized in the table below (Table 1). Almost, the base line characteristics of the patients were well balanced between the two treatment groups.

<table>
<thead>
<tr>
<th>Patient characteristic</th>
<th>Arm I (n=23)</th>
<th>Arm II (n=19)</th>
<th>p-value</th>
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<tbody>
<tr>
<td>No.</td>
<td>%</td>
<td>No.</td>
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<tr>
<td>Age (Years)</td>
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<td></td>
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<td>&lt;60</td>
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<td>17.4</td>
<td>2</td>
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<tr>
<td>≥60</td>
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<td>82.6</td>
<td>17</td>
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<td>Sex</td>
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<td>M</td>
<td>23</td>
<td>100</td>
<td>16</td>
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<tr>
<td>F</td>
<td>-</td>
<td>-</td>
<td>3</td>
</tr>
<tr>
<td>Tumor stage</td>
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<td></td>
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<tr>
<td>T2</td>
<td>9</td>
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<td>6</td>
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<tr>
<td>T3a</td>
<td>-</td>
<td>-</td>
<td>4.3</td>
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<tr>
<td>T3b</td>
<td>13</td>
<td>56.5</td>
<td>11</td>
</tr>
<tr>
<td>T4a</td>
<td>-</td>
<td>-</td>
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<td>Tumor grade</td>
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<td></td>
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</tr>
<tr>
<td>G I</td>
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<td>4.3</td>
<td>2</td>
</tr>
<tr>
<td>G II</td>
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</tr>
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<td>13</td>
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<tr>
<td>G I</td>
<td>1</td>
<td>4.3</td>
<td>6</td>
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</table>

Primary end point: The response to induction phase was evaluated in week 7 after completion of induction chemo-radiotherapy. Inter-assessment was done by cystoscopic examination and multiple punch biopsies. In arm I: One patient (4.3%) showed disease progression (DP), 3 patients (13%) had only partial response (PR) and 19 patients (82.6%) showed complete response (CR). Patients who achieved CR entered the consolidation phase of treatment protocol. In arm II, 4 patients (21.1%) showed PR and 15 patients (78.9%) showed CR, so they entered the consolidation phase as shown in Fig. (1). There was no significant difference between both arms regarding the response to induction phase of treatment protocol ($p=0.537$).
All patients (10 patients in arm I & 6 patients in arm II) with early stage tumor T2+T3a showed CR (100%) in both arms of the study. On the other hand, patients with advanced tumors T3b+T4a showed CR in 69.2% of patients (9 patients in arm I & 9 patients in arm II). There was no statistically significant difference between both groups regarding response ($p=0.1$ & 0.25 in both arms respectively).

Secondary end points:

**PFS & OS rates:**

The median duration of follow-up was 16 months (range: 2-35 months) with a mean duration of $15.26 \pm 9.88$ months.

Arm I demonstrated a superior PFS at 3-years compared with arm II. The PFS was 78.26$\%$ & 68.42$\%$ for both arms respectively. The median PFS was 21 & 18 months in both arms respectively. Similarly, PFS at 6 months, 1-year and 2-years were better in arm I of the study. The corresponding PFS values were 87$\%$, 78.26$\%$ and 78.26$\%$ for arm I. On the other hand, PFS values for arm II were 85$\%$, 73.68$\%$ and 68.42$\%$ respectively. However there were no statistically significant differences in PFS between both arms ($p=0.485$) as shown in Fig. (2).

Overall survival at 3-years was slightly better for arm II. The 3-year OS was 60.87$\%$ & 68.42$\%$ for both arms respectively. The median OS was 25 & 23 months for both arms respectively ($p=0.683$). There were no statistically significant differences between both arms regarding OS at 3-years. OS at 6 months, 1-year and 2-years were 77$\%$, 69.57$\%$ and 65.22$\%$ respectively for arm I. On the other hand, the corresponding figures for arm II were 79$\%$, 78.95$\%$ and 68.42$\%$ respectively as shown in Fig. (3).

Univariate analysis was done to study the impact of various prognostic factors on PFS & OS and we found that tumor stage is the only factor that affected PFS with statistical significance but other factors were non-significant.

**Failure rates:**

Loco-regional failure was found in one out of 23 patients (4.3$\%$) in arm I. Local recurrence occurred in the bladder and extended outside the wall to perivesical fat. On the other hand in arm II, 2 out of 19 patients (10.5$\%$) developed local recurrence, one case extended to pelvic lymph nodes and the other case developed recurrence at the bladder and extended throughout the wall to perivesical fat and surrounding pelvic organs; the rectum. There were no statistically significant differences between both arms ($p=0.581$).
Distant metastasis occurred in 3 out of 23 patients (13%) in arm I. Disease metastasized to bone in one case and in the second two cases to lung and bone. In arm II, 3 out of 19 patients (15.8%) had distant metastases. One case developed lung, bone and liver metastases and the other two cases developed only bone metastases. There was no statistically significant difference between both arms regarding occurrence of distant metastasis.

Treatment toxicity:

All patients were evaluated for toxicities according to WHO grading system. The most frequently occurring adverse effects were dysuria, frequency, diarrhea and anemia. Only one patient in arm I developed grade IV neutropenia that was successfully treated. Otherwise no other major hematologic toxicity was encountered in any of the two arms of the study. Dysuria & frequency were the most frequently observed non-hematologic toxicities followed by diarrhea. Gastrointestinal complications were more or less equivalent in both arms of the study. Side effects were tolerable and manageable. No treatment-related deaths was encountered in any of the two study groups.

Discussion

Our patients showed an age range from 48-78 years with a median age of 64 years in arm I and 70 years in arm II, this is consisted with the worldwide reported median age. Khosravi-Shahi and Cabezón-Gutiérrez reported that median age at diagnosis was 65 years, and 70% of patients with bladder cancer were >60 years of age [12]. Our study showed male predominance with male to female ratio 5.3:1. This ratio is higher than the reported world-wide ratio which is about 3:1 [12]. This may be explained by the small number of the sample to be presented with the world wide ratio.

Arm I showed higher CR than arm II but the difference did not reach the statistical significance. In our study, we used cisplatin and paclitaxel plus RT in arm I but in arm II we used cisplatin and 5-fluorouracil also plus RT. Our results are comparable with the studies conducted by Sabaa et al. [13], Khader et al. [14] & Efstathiou et al. [15]. Sabaa et al. studied 104 patients with MIBC that were treated by complete TURB followed by CRT in the form of gemcitabine and cisplatin and conventional radiotherapy after the maximum resection of their tumors. Complete response was shown by 78.8% [13] & this typically is equal to the results of arm II. The slightly higher CR rate observed in arm I may be due to the use of paclitaxel which may be more potent than 5-fluorouracil and gemcitabine. Khader and his colleagues studied 14 patients with MIBC. Initial therapy consisted of TURB followed by induction chemotherapy, then irradiation with concurrent platinum-based agents. The bladder and pelvic lymphatics were treated via a four-field box technique to a total dose of 4500cGy (180cGy daily fractions in 5 consecutive days). Additional therapy with irradiation (up to 6400 cGy) was delivered to the bladder with safety margin to complete responding patients [14]. They achieved 73% CR and this is much near to CR obtained in our results. Efstathiou et al., reported the MGH experience with selective bladder preservation in the treatment of 348 patients with MIBC (T2-T4a). Patients underwent concurrent cisplatin-based chemotherapy and RT after maximal TURB plus neoadjuvant or adjuvant chemotherapy. Repeat biopsy was performed after 40Gy, with initial tumor response guiding subsequent therapy. Those patients showing CR received boost chemotherapy and RT. CR was achieved in 72% of patients [15]. This agrees with our results as they used multidrug regimen with platinum-based agents concomitant with radiotherapy in addition to using chemotherapy as adjuvant or neoadjuvant in the treatment protocol that may contribute to the higher CR obtained. Other clinical researches using cisplatin in combination with RT in patients with MIBC have demonstrated objective tumor responses reaching up to 80%, with acceptable toxicity. These were observed in the studies conducted by Zouhair et al. [16], Rodel et al. [17], Chen et al. [18] and Hagan et al. [19]. Similarly, other studies also used cisplatin plus radiotherapy after TURB and obtained results comparable to those demonstrated in our study: Weiss and his colleagues had CR that reached up to 90.3% [20], Perdona et al. showed 88.4% CR [21] and Joung et al. showed 75% CR [22]. On the other hand, the results obtained in our series are better than that reported by Gamal El Deen et al. [10], Ibrahim et al. [23] and Nowak-sadzikowska et al. [24] studies. CR was achieved by Gamal El Deen in 67.3% of 55 patients with MIBC [10]. Our results were much higher in both arms and this may be owing to the use of combination chemotherapy in induction and consolidation phases and also the use of unconventional fractionation in the form of accelerated hyperfractionation. Ibrahim et al., found that 60% of their patients achieved CR [23]. Similarly, this study showed lower results than ours which may also be attributed to the use of multidrug regimens and unconventional fractionation schedules of radiotherapy in both arms of our study.
Statistical analysis to study the impact of different prognostic factors on response revealed that tumor stage and tumor grade were the most predictive factors for this initial response.

Ibrahim et al., reported that local failure was recorded in 40% of patients and distant metastasis was reported in 25% [23]. Kaufman et al., reported loco-regional failure rate as 27.5% and distant failure rate of 31.25% [9]. Sabaa et al. found that local failure rate in their study was 16.2% out of the evaluable 74 patients and distant failure rate was 24.3% [13]. Zapatero et al., reported the pattern of failure for all patients and according to the treatment protocol. 24.5% experienced local bladder relapse and 15% developed distant metastasis [25]. However, this failure rates are more or less far from our results. In our current study, after a median follow-up of 16 months the local failure rate for the whole studied patients was 3 out of 42 patients (7.1%). One out of 23 patients (4.3%) in arm I and 2 out of 19 patients (10.5%) in arm II developed local failure. This may be explained by the shorter follow-up duration. On the other hand, distant failure rate for the whole group was 6 out of 42 patients (14.3%). Three out of 23 patients (13%) in arm I and 3 out of 19 patients (15.8%) in arm II developed distant failure and this much more comparable with Zapatero et al., results [25]. The relative difference between treatment failure rates of our study and other series could be explained by the relatively shorter follow-up period in our study compared to other series that had larger sample size and longer follow-up periods.

Gamal El Deen et al., reported treatment toxicity observed among 37 patients who ended chemotherapy and radiotherapy protocol [10]. They are different from our results regarding genitourinary (GU) and gastrointestinal (GI) complications. This may be explained by the use of accelerated hyper-fractionation in our series, as it is well known that this altered fractionation increases acute side effects to the irradiated area. Hematological complications more or less are equivalent to our results. Hematological complications were recorded in 27% of their cases, GI complications in 24.3% of their cases and GU complications in 10.8% of their cases. Nowak-Sadzikowska et al., treated 27 patients with CRT. Anaemia mainly of grade I (GI) occurred in 11%, leucopenia of GI in 37% and thrombocytopenia mainly of G III in 22%. GI and GU complications were mainly of GI and GII. Gastrointestinal complications were occurred in 44% of patients with G I and G II in 15% of patients [24]. These results are much more comparable with our results.

At a median follow-up period of 16 months (range: 2-35 months) with a mean duration of 15.26±9.88 months, our protocol reported higher PFS at 3-years in arm I than arm II. They were 78.26% & 68.42% for both arms respectively. On the contrary, OS at 3-years was slightly better for arm II. The 3-year OS was 60.87% & 68.42% for both arms respectively. Both rates PFS and OS did not show statistically significant difference between both arms (P= 0.485 & 0.683 respectively). Gamal El Deen reported that OS at 3-years was 79.23% [10]. The 3-year OS for the patients without hydronephrosis that are similar to our patient's selection criteria was 59%. This compares favorably with the prior RTOG protocol 89-03 studied by Shipley et al. at [26] and this is also comparable with our results in both arms. Other studies by Zietman et al. at [27] and Kaufman et al. [28] reported 3-year OS of 83% with more or less similar radiation fractionation and similar chemotherapy schedules as in arm II of our study. However, our results are a little a bit lower for both arms. This may be due to the smaller number of our sample in comparison to the two previous studies. Houssett and his colleagues reported 63% OS [29] and this is comparable with the results of both arms of our study. Ibrahim and his colleagues recorded that the 2-year survival and progression free survival (PFS) rates were 67% and 58%, respectively [23]. The OS of this study is mostly similar to our result and DFS is more or less near to our results in arm II but far from the results of arm I. The results obtained by Joung and his colleagues were that DFS and OS rates for all 20 patients were 51.1% and 38.6% at 5 yr, respectively [22] and this was markedly different from our results. Of the 80 patients studied by Kaufman et al., 44 survived, with an OS rate of 67% at 36 months and 56% at 60 months. DFS rate was 73% at 36 months and 71% at 60 months [9]. This study design is the most similar to our study and their OS and DFS at 3-years are much more similar to our results in both arms.

Conclusions:

Bladder preservation protocol is a good choice for treatment of bladder cancer but with good selection of the cases and careful follow-up of them to avoid major toxicities that may lead to interruption or even stoppage of the preservation protocol. Many studies should be continued to study more the effect of this protocol on PFS and OS and patient quality of life.
References


