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Introduction

After radical prostatectomy (RP), 33–40% of patients experience biochemical recurrence (BCR) [1,2]. Improved imaging, particularly positron-emission tomography (PET)/computed tomography (CT) techniques, has enabled early detection of lymph node metastasis [3–5]. In this context, metastasis directed therapy (MDT) either as stereotactic ablative radiotherapy (SABR) or salvage lymph node dissection (sLND), has emerged as an alternative option to standard androgen deprivation therapy (ADT). Several retrospective systematic reviews and studies on the safety and effectiveness of open pelvic- and retroperitoneal sLND, are available [6–9]. Common for many of these reports are the high rate of biochemical recurrence (BCR) after sLND. Recently, Bravi et al. [10] reported results from a long-term follow-up (FU) study in 189 patients operated with sLND between 2002 and 2011. At a median FU of 87 months, 45 patients (24%) had died from PCa. Clinical recurrence-free survival and BCR-free survival at 10 years were 31% and 11%, respectively, and a total of 145 patients (77%) had received ADT. Such data indicate that MDT alone offer limited oncological benefit.

Conclusion: Oncological benefit of sPLND as the only salvage procedure seems to be limited, though almost one third of patients achieved treatment response. Clinical trials are needed to determine if sPLND as part of a multimodal treatment may improve outcome.
In this study we assess our current experience with robotic sPLND. Our aims are to investigate treatment response after robotic sPLND, the time to start of additional treatment and factors associated with these two events. Furthermore, we assessed complications related to this minimally invasive procedure.

Material and methods

Patients

Our study population consists of 69 patients consecutively operated with sPLND at the Oslo University Hospital from June 2014 to December 2018. All patients had post-RP BCR (PSA ≥ 0.2 ng/ml) and lymph node recurrence detected by PET/CT. None of the patients used ADT upon referral for sPLND, and none had distant metastases detected on MRI or bonescan.

PET imaging

All patients underwent PET/CT before sPLND, either with 18 F-fluciclovine (49 patients), 18 F-PSMA-1007 (19 patients) or 18 F-fluorodeoxyglucose(FDG) (1 patient) [17,18]. Imaging was performed on a Siemens Biograph mCT 40 or on a GE Discovery 690 PET/CT-scanner for all but two patients (PET performed at St.Olavs Hospital, Trondheim and in Aarhus, Denmark). Routine PET/CT protocols and reports were applied pre-sPLND.

Surgical procedures

The da Vinci Surgical system was used with a six-port configuration to perform the transperitoneal sPLND procedures (Intuitive Surgical, Sunnyvale, CA, USA). Depending upon previous treatments and findings on imaging, the extent and mode of sPLND was left to the discretion of the surgeon and ranged from targeted removal of the PET positive node(s) to a bilateral or unilateral extended sPLND. Targeted removal of nodes was preferred in patients previously operated with PLND concomitant to RP. Extended sPLND encompassed the internal iliac, obturator, external iliac and common iliac landing areas. Two patients had removal of vesicula seminalis and sPLND due to recurrence in both sites. The length of stay in the hospital after sPLND was routinely one night, the patient arrived at the hospital the same day they were operated and went home the day after without urinary catheter.

Histopathology

Histopathological evaluation of the sPLND specimen assessed the total number of excised lymph nodes, the number of metastatic lymph nodes and the laterality of the excised metastatic nodes. Information of previous RP specimens was retrieved from the medical records (pathological T stage, surgical margins and Gleason ISUP Gleason Grade Groups (GGG)).

Follow up

Postoperative follow-up (FU) consisted of outpatient visits at 6 and 12 weeks at our institution with clinical examination and PSA evaluation. Thereafter the patients were followed by their general practitioner with 3 months intervals, in collaboration with the urological department at the local hospital. PSA, clinical recurrence, postoperative complications and start of the first additional treatment post-sPLND were recorded. Eventual start of ADT was generally offered to patients with rising PSA post-sPLND with or without clinically detected recurrence, but was left to the physician’s discretion and patient preference.

Outcome variables

The primary endpoint was treatment response defined as the percentage of patients obtaining PSA < 0.2 ng/ml 6 weeks after sPLND. The type and time of any additional treatment was recorded. Clinical recurrence was based on metastatic findings on MRI or PET/CT imaging post sPLND. Finally, complications were reported post-sPLND according to the Clavien–Dindo classification system [19]. The study was approved by the Regional Ethical Committee (REK 2016/1977), The Data Protection Officer, and the Institutional Review Board.

Statistics

Frequencies and percentages of categorical variables and medians and interquartile ranges (IQR) for continuous variables were calculated. Intergroup differences between targeted sPLND, unilateral extended or bilateral extended sPLND template, were assessed by chi-squared tests for categorical variables and Wilcoxon rank sum test for skewed continuous variables. Multivariable Cox regression was used to predict additional treatment after SLND. Predictors consisted of GGG < 4 vs GGG ≥ 4, time from RP to sPLND, number of positive lesions on PET/CT scan (two and three or more vs one as reference), and PSA level at sPLND. The patients were followed up until December 31, 2019 (end of the study). The Kaplan-Meier estimates displayed survival free of additional treatment. The level of significance was set at 0.05. Data were analysed using the IBM Statistical Package for the Social Sciences (SPSS) Statistics version 25.

Results

Background characteristics

Histology of the RP specimen showed ≥ pT3a in 76% of the patients and 73% with GGG ≥ 3 (Table 1). Thirty patients (43%) had concomitant PLND at RP and of these patients almost one third had metastatic lymph nodes. Thirty-seven patients (54%) had undergone post-RP pelvic radiotherapy before referral for sPLND (Table 1).
**sPLND results**

Median age pre-sPLND was 67 years (Table 2). Sixty patients (85%) had only 1 or 2 positive lesions on PET/CT. Median number of positive lesions was 1 (range 1–20 lesions). The median number of excised lymph nodes was 19 for bilateral extended sPLND (Table 2). No metastatic lymph nodes were described in the sPLND histopathological specimen of five patients (7%). Re-evaluation of the PET images pre-sPLND for these five patients showed that in three patients the PET reported positive pelvic nodes were outside the surgical field (mesorectal nodes in two patients and an inguinal node in one patient). For the other two patients the re-evaluation changed the description of the PET image from positive to equivocal findings. Three patients had a new sPLND (re-sPLND) due to BCR and new findings on PET/CT after the first sPLND (Table 3).

**Oncological outcomes**

Twenty patients (29%) achieved treatment response (Table 3) independently of mode of sPLND (Table 3). At 6 weeks post sPLND more than 50% reduction of PSA value was detected in 38 patients (54%) while 10 patients (14%) had an increase of their pre-sPLND PSA value (Figure 1).

At a median follow-up of 15 months, 14 patients (20%) still had PSA < 0.2 ng/ml (Table 3). Forty-one patients (59%) had started additional treatment at median follow-up 15 months.

Median (IQR) PSA for patients with and without treatment response was 1.1 ng/ml (0.5–2.2) and 1.8 ng/ml (0.9–3.8), respectively (p = 0.03) (Figure 2). In univariable logistic regression none of clinically relevant variables predicted treatment response except for pre-sPLND PSA (odds ratio 0.71, 95% CI 0.50–0.97, p = 0.05). No covariates emerged as statistically significant predictors for treatment response in multivariable logistic regression analysis (Figure 3), but there was a tendency towards association between less likelihood of treatment response and increasing pre-sPLND PSA and increased numbers of PET positive nodes.

Median survival time without additional treatment was 17 months (95% CI 6–28) (Figure 4). The pre-sPLND PSA value, number of excised positive lymphnodes and time

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**Table 1.** Patient characteristics, n = 69.

| Age at radical prostatectomy (RP), years, median (IQR) | 62 (56–66) |
| Pre-RP PSA value (ng/ml), median (IQR) | 13.0 (11.0–18.5) |
| Undetectable PSA after RP, n (%) | Yes 39 (56), No 29 (44), Unknown 1 (2) |
| Pathological T stage RP specimen, n (%) | pT2 17 (25), pT3a 35 (51), pT3b 17 (25) |
| Gleason score RP specimen, n (%) | ISUP grade group 1 10 (0), ISUP grade group 2 18 (26), ISUP grade group 3 35 (51), ISUP grade group 4 23 (33) |
| Pathological nodal status, n (%) | pNx 39 (57), pN0 21 (30), pN1 9 (13) |
| Surgical margins, n (%) | Positive 17 (25), Negative 52 (75) |
| Radiotherapy post –RP, n (%) | 38 (54) |

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**Table 2.** Pre-, peri- and postoperative sPLND data stratified on mode of sPLND.

<table>
<thead>
<tr>
<th>sPLND overall n = 69</th>
<th>Targeted sPLND n = 30</th>
<th>Unilateral extended n = 11</th>
<th>Bilateral extended n = 28</th>
<th>Intergroup differences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age at sPLND, years, median (IQR)</td>
<td>67 (62–71)</td>
<td>67 (62–71)</td>
<td>68 (62–72)</td>
<td>67 (62–71)</td>
</tr>
<tr>
<td>Months from RP, median (IQR)</td>
<td>49 (24–79)</td>
<td>36 (21–59)</td>
<td>79 (54–93)</td>
<td>53 (25–83)</td>
</tr>
<tr>
<td>PSA at sPLND</td>
<td>1.4 (0.8–3.4)</td>
<td>1.4 (0.7–3.2)</td>
<td>1.6 (0.9–3.0)</td>
<td>1.9 (0.9–4.3)</td>
</tr>
<tr>
<td>Number of positive PET/CT lesions per patient, n (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>40 (57.7)</td>
<td>20 (66.7)</td>
<td>7 (58.3)</td>
<td>14 (50.0)</td>
</tr>
<tr>
<td>2</td>
<td>19 (26.8)</td>
<td>5 (16.7)</td>
<td>3 (25.0)</td>
<td>11 (39.3)</td>
</tr>
<tr>
<td>≥3</td>
<td>10 (14.1)</td>
<td>5 (16.7)</td>
<td>2 (16.6)</td>
<td>3 (10.8)</td>
</tr>
<tr>
<td>Patients with malignant histology in sPLND specimen</td>
<td>63 (90)</td>
<td>26 (86.7)</td>
<td>9 (75.0)</td>
<td>28 (100)</td>
</tr>
<tr>
<td>Number of excised nodes, median (IQR)</td>
<td>12 (7–19)</td>
<td>8 (3–12)</td>
<td>12 (7–18)</td>
<td>19 (12–25)</td>
</tr>
<tr>
<td>Number of excised metastatic nodes, median (IQR)</td>
<td>2 (1–4)</td>
<td>1 (1–3)</td>
<td>3 (1–5)</td>
<td>3 (1–5)</td>
</tr>
<tr>
<td>Clavien Dindo grade of complications, n (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Grade 1*</td>
<td>11</td>
<td>5</td>
<td>1</td>
<td>7</td>
</tr>
<tr>
<td>Grade 2**</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Grade 3a***</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Grade 3b***</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

*8 patients with temporary lymphedema and 3 patients with neurapraxia.

*1 patient with deep venous thrombosis.

*1 patient with drainage of abscess.

*Injury of ureter.
between RP and sPLND significantly predicted the time to additional treatment post-sPLND (Supplementary Table 1).

At the end of study 2 patients had died, one of them due to prostate cancer.

**Postoperative complications**

There were no perioperative complications. Post-sPLND complications occurred in 14 patients (20%) (Table 2). In eleven of these patients, the complications were classified as Clavien-Dindo grade 1 (Table 2). Complications occurred in 9 of 38 patients who had undergone postoperative radiation therapy after primary RP, compared to 5 complications in 32 patients without postoperative radiation (Chi-squared \( p = 0.5 \)).

**Discussion**

To our knowledge, this is the largest single institution study of robotic sPLND. Twenty patients (29%) achieved treatment response. At a median follow-up of 15 months 14 of these patients (20%) still had PSA < 0.2 ng/ml. Decreasing levels of pre-sPLND PSA predicted treatment response. Finally, 20% of patients experienced postoperative complications, mostly Clavien-Dindo grade 1.

Other studies have found similar discouraging rates of treatment response rates. Siriwardana et al. [14] reported 31% treatment response in a series of 35 patients. In another study Linxweiler et al. [15] found treatment response in 33% of patients in a series of 36 patients. The distribution of percentage PSA decline in our series was similar to a recent study with 68 patients [20]. Siriwardana et al. [14] and Linxweiler et al. [15] found that 50% of patients had started additional treatment after a median FU of 12 months, which is in accordance with our results which showed that 41 patients (59%) had received additional treatment at median follow-up 15 months.

The rate of postoperative complications is comparable to Siriwardana et al. [14] who reported complications in 23% of 35 patients, all Clavien–Dindo grade \( \leq 2 \). In a study by Devos et al. [12] comparing open sLND with robotic sLND, postoperative complications within 30 days after surgery were more prevalent in the open sLND group compared to the robotic group (41.6% vs. 20%, \( p = 0.04 \)). The robotic approach might be favorable due to more meticulous dissection and better control of lymphatic vessels because of higher magnification and in depth tissue visualization.

As in the paper by Siriwardana et al. [14], we did not find that GGG in RP specimen was a predictor of treatment response. Similarly, Bravi et al. [10] found that GGG was not a predictor for cancer-specific mortality. We found that pre-sPLND PSA value, number of excised positive lymphnodes and time between RP and sPLND significantly predicted the time to additional treatment post-sPLND. However, the initiation of such treatment after sPLND represents a rather
uncertain endpoint because it is often based on vague criteria and left to the discretion of the patient and physician.

One argument often used in favour of sPLND is that this intervention may delay start of ADT. However, without comparison to a control group it is impossible to adequately assess the oncologic benefit of sPLND. This is illustrated by the remarkable observation in the only randomized study of MDT [11], where 35% of patients undergoing surveillance experienced spontaneous PSA decline without receiving any therapy.

One may also question if delaying ADT is of any benefit. A report last year suggests that 30% of improvement in prostate cancer specific survival during the recent years is due to adjuvant ADT [21]. Messing et al. [22] showed already in 1999 in a randomized study that in RP patients with metastatic lymph nodes immediate ADT has beneficial effect on survival and progression compared to deferred ADT.

The median yield of lymph nodes in bilateral extended sPLND was 19 nodes (IQR 11–24) which is similar to a systematic review by Ploussard et al. [7]. In our study most of the clinical recurrence after sPLND was due to new lymph node metastasis detected on PET/CT which implies that sPLND did not remove all metastatic lesions.

In their long-term outcome study Bravi et al. [10] found a significant survival benefit associated with the administration of ADT after sLND, suggesting that sLND as a local intervention should be considered part of a multimodal approach. In the ongoing STORM study [23] the aim is to investigate if additional whole pelvic radiotherapy after MDT plus adjuvant ADT may more effectively treat micro metastasis and thereby improve oncological outcomes.

Although 20% of patients still had PSA < 0.2 ng/ml after median 15 months, we failed to find any significant predictor for treatment response; hence an important clinical question is how to better select patients for MDT. A promising field of research in this respect is liquid biopsy, such as miRNA derived from oligometastatic cancers which may identify patients who most likely not will benefit from MDT due to high numbers of circulating tumor cells [24–25] and thereby be spared aggressive local treatment.

There are some important limitations to our study. The most important ones are the retrospective nature of the study, a relatively small sample size yet large for a robotic sPLND series, short median FU and the lack of a control group. Another weakness is the heterogenic diagnostic work-up pre-sPLND with different PET/CT tracers and that five different surgeons alternated in performing the sPLND procedures.

**Conclusion**

Oncological benefit of robotic sPLND as the only salvage procedure seems to be limited, though almost one third of patients achieved treatment response. Clinical trials are
needed to determine if sPLND as part of a multimodal treatment may improve outcome.

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