Generating comprehensive comparative evidence on various interventions for penile rehabilitation in patients with erectile dysfunction after radical prostatectomy: a systematic review and network meta-analysis

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Background: We aim to present a comprehensive comparison of various treatments in the management of penile recovery after radical prostatectomy (RP) and provide recommendations for future research.

Methods: Literature search of electronic databases including PubMed, the Cochrane Library, Embase, PsycInfo, and Web of Science, and manual retrieval were conducted from inception through March 2020. “Erectile dysfunction” and “prostatectomy” were used as the Mesh terms. The patients, intervention, comparison, outcome, and study design (PICOS) approach were used to define study eligibility. Two authors independently selected studies, evaluated the methodological quality, and extracted data using Cochrane Collaboration's tools. The data analysis was completed by STATA version 14.2.

Results: A total of 24 studies with 3,500 patients were incorporated in the final analysis after screening 6,131 records. Our findings indicated that vacuum constriction devices (VCD) ranked 1st which meant that patients in VCD group had the best effect regarding mean IIEF scores within 3 months after RP, and no significant difference was observed between VCD and VCD with 20 mg/day tadalafil (V20DT) (MD: 5.44; 95% CI: -0.81 to 11.69). VCD and 50 mg/day sildenafil (VC50DS) showed superiority over 50 mg/day sildenafil (50DS) (MD: 3.75; 95% CI: 2.74–4.76) and intraurethral alprostadil 125–250 μg (MD: 3.05; 95% CI: 0.38 to 5.72), respectively. Moreover, V20DT showed significant superiority over the other interventions for ≥6 months mean International Index Erectile of Function (IIEF) scores after RP. Monotherapy appeared to have similar efficacy in terms of mean IIEF scores and proportion of patients return to baseline, and the effect of phosphodiesterase type 5 inhibitors (PDE5is) did not seem to be affected by the patterns of administration (regular or on demand).

Conclusions: The combination therapy showed certain advantages over monotherapy, and we recommended the combination of VCD and PDE5is to be considered in the clinical management of penile rehabilitation after RP.

Keywords: Penile rehabilitation; erectile dysfunction (ED); radical prostatectomy (RP); network meta-analysis

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Introduction

With the progress of early detection and alleviated health awareness, the morbidity of prostate cancer (PCa) has been growing in the past decades and ranked first among malignancies in men (1,2), among which an increasing number of patients were diagnosed at a younger age (3,4). Though fortunately low is the death rate of PCa, which is reported to be 0.07% to 0.15% (5), prostatectomy related erectile dysfunction (ED) is of great concern for its high occurrence and negative effects on quality of life, especially in the sexually active population. With the improved understanding of cavernous nerves (CNs), the pathophysiological basis underlying radical prostatectomy (RP) induced ED was well studied and the most likely mechanism could be the pathophysiological change of cavernous smooth muscle and tunica albuginea induced by CNs injury in operation (6-8). According to previous studies, the prevalence of RP induced ED was about 55.6% after nerve-sparing laparoscopic radical prostatectomy (nsLRP) and 39.8% after robot-assisted radical prostatectomy (RARP) (9-12). Though supported by current evidence that nerve-sparing robot-assisted radical prostatectomy (nsRARP) could be the most favorable operation for preserving postoperative erectile function (EF) in PCa patients, the postoperative ED prevalence could still be as high as 10–46% (13) and most postoperative patients are unlikely to recover to their baseline EF (14). Concerned the current situation that the mean age patients were when diagnosed with PCa has decreased and sexual activity-related life quality expectancy has grown (15), it is of great importance to establish an effective treatment protocol for postoperative penile rehabilitation. At present, treatment strategies for ED mainly include phosphodiesterase type 5 inhibitors (PDE5is), psychotherapy intervention, pelvic floor muscle training (PFMT), intracorporeal injection of a vasoactive substance, vacuum constriction devices (VCD), and penile prosthesis implant. Other therapeutic approaches including stem cell therapy, nerve transplantation, low-intensity extracorporeal shockwave therapy, erythropoietin, tacrolimus, and hyperbaric oxygenation therapy also show promising efficacy for ED management.

However, it should be noted that most of the prostatectomy-related ED could be resulted from CNs injury and the concept of ED could be categorized into primary impotency and secondary impotency for different pathogenesis. Thus, confusions could be aroused that whether above illustrated therapeutic approaches could be as effective as for RP induced ED and if the guidelines for ED (16,17) could be ideally adaptive for RP induced ED. From the results of the previous meta-analysis, PDE5is, VCD, PFMT, and intracorporeal injection was effective for penile rehabilitation after RP (18-22). Because most of the RCTs conducted in this context were placebo-controlled designed and there is a lack of data from the direct comparison between therapies. It could be difficult for physicians to determine the most preferable treatment strategies for patients when faced with a large variety of options. To make a synthesis of potentially practice-changing evidence and provide a trustworthy evidence-based reference on postoperative penile rehabilitation therapy for physicians, we performed this network meta-analysis. We present the following article in accordance with the PRISMA reporting checklist (available at http://dx.doi.org/10.21037/tau-20-892).

Methods

Study Selection

In accordance with the statement of preferred reporting items for systematic review and meta-analysis (PRISMA) (23), electronic databases including PubMed, the Cochrane Library, Embase, PsycInfo, and Web of Science were searched to identify RCTs from inception through March 2020 with no limitation to language. “erectile dysfunction” and “prostatectomy” were used as the Mesh terms, and the search strategy used in PubMed was as follows: (((((((((((Prostatectomy[Title/Abstract]) OR Prostatectomies[Title/Abstract])) OR Prostatectomies, Retropubic[Title/Abstract]) OR Prostatectomies, Suprapubic[Title/Abstract]) OR Suprapubic Prostatectomies[Title/Abstract]) OR Prostatectomies[Title/Abstract]) OR Prostatectomies, Retropubic[Title/Abstract]) OR Prostatectomies, Suprapubic[Title/Abstract]) OR Suprapubic Prostatectomies[Title/Abstract])) AND (((Erectile Dysfunction[Title/Abstract]) OR Dysfunction, Erectile[Title/Abstract]) OR Male Sexual Impotence[Title/Abstract]) OR Impotence, Male Sexual[Title/Abstract]) OR Sexual Impotence, Male[Title/Abstract]) OR Male Impotence[Title/Abstract]) OR Impotence, Male[Title/Abstract]) OR Impotence[Title/Abstract])) OR Prostatectomies[Title/Abstract]). The details of search strategy were shown in supplementary material. Besides, we also retrieved reference lists of included studies and related reviews to ensure the comprehensive search.
Four independent reviewers identified potentially eligible studies after removing duplicates and screening titles and abstracts. Full-text articles that met the following inclusion criteria and presented available data were included in the final analysis. Any disagreement was resolved by consensus with a third party. Figure 1 depicts the PRISMA flowchart.

**Selection criteria**

The patients, intervention, comparison, outcome, and study design (PICOS) approach was used to define study eligibility; patients (P): patients with ED after RP; intervention (I): patients treated with any drug, supportive therapy or device; comparison (C): comparisons with different interventions; outcomes (O): EF measured by International Index Erectile Function (IIEF); study design (S): randomized controlled trial (RCTs) with sufficient data for extraction were selected. Meeting abstracts, overlapping population, and duplicated studies were excluded.

**Quality assessment**

Two independent reviewers (DCF, SZL) evaluated the study quality (Figure 2) according to the Cochrane Collaboration’s Risk of Bias (RoB) tool in Review Manager software (https://community.cochrane.org/help/tools-and-software/revman-5). This tool primarily evaluates 7 domains: random sequence generation (selection bias); allocation concealment (selection bias); blinding of participants and personnel (performance bias); blinding of outcome assessment (detection bias); incomplete outcome data (attrition bias); selective reporting (reporting bias); other bias (such as funding sources). Besides, the level of evidence of each included article was rated independently by DCF and SZL using the Oxford Centre for Evidence-Based Medicine criteria (24). This scale graded studies from strongest (level 1) to weakest (level 5) strength of evidence based on study design and data quality. Figure 2 details the RoB summary of the included articles in this study. Overall, included studies had a low risk of random sequence generation (selection bias), incomplete outcome data (attrition bias), and selective reporting (reporting bias). The
Figure 2 Methodological quality of trials included in the meta-analysis and loop inconsistency. (A) risk of bias summary; (B) risk of bias graph; (C) loop inconsistency.
risk of allocation concealment was unclear due to the lack of related descriptions. Besides, 19 of 24 RCTs were rated as 1b and regarded as high quality (Table 1).

Statistical analysis

Data presented as mean and standard deviation (SD) were used. Median and range were used to estimate mean and SD (25). The percentiles, 25th, and 75th percentiles as well as 5th and 95th percentiles, were transformed to SD through the following formula: SD = Norm IQR=(P75-P25)/0.7413 (IQR: inter-quartile range, P75:75th percentile, P25: 25th percentile) (26). Dichotomous data were calculated as risk ratios (RR) and continuous outcomes as mean difference (MD), both with 95% confidence intervals (CIs). The network meta-analysis was performed based on the Bayesian framework model using STATA version 14.2. The random-effects model was used unless the program suggested no heterogeneity. Besides, we performed loop inconsistency with inconsistency factor (IF) and local inconsistency with node-splitting analysis to evaluate the consistency of indirect and direct comparisons. We identified the source of inconsistency through sensitivity analysis and our results would be relatively robust if we cannot explain the inconsistency. Publication bias was assessed by the symmetry of funnel-plots. We ranked the various interventions based on the values of surface under the curve cumulative probabilities (SUCRA). The statistical significance was defined as P<0.05.

Results

Search results

Six thousand and six hundred twenty-eight records were initially identified from the database, and 60 studies were potentially eligible for further review in full text. Of these studies, 36 were excluded on the basis of inclusion criteria, and 24 studies (27-50) were incorporated in the final analysis.

The 24 articles contained 3,500 patients and were published between 2004 and 2019. In total, 3,500 patients were enrolled in these trials and most of the trials were deemed to be high quality according to the Oxford Centre for Evidence-Based Medicine criteria (24). The 24 studies contained seven interventions, 14 involving PDE5is, 6 involving PFMT, 4 involving vacuum therapy, 1 involving neuromodulatory therapy, 2 involving statin therapy, 1 intracorporeal injection therapy, and 1 involving hyperbaric oxygenation therapy. Of these studies, 8 assessed the combined effects of these treatments. The baseline characteristics of the included studies are described in Table 1. Besides, the network plots of investigated outcomes were presented in Figure 3.

Network meta-analysis of mean IIEF scores

Two parallel-group network meta-analysis were conducted to compare the efficacy of different treatments in terms of short-term results of mean IIEF scores due to available data in the included studies (Figure 4). The first group incorporated 11 studies with 1,235 participants involving 11 interventions (Figure 4E). VCD, VCD and 20 mg/day tadalafil (V20DT), and 200 mg avanafil on demand (200AOD) increased mean IIEF scores within 3 months after RP significantly more than many other treatments (Figure 4E). Most differences between the remaining interventions were small or very uncertain. According to the SUCRA ranking, VCD ranked 1st (99.2) which meant that patients in VCD group had the best effect regarding mean IIEF scores within 3 months after RP (Figure 4A). No significant difference was observed between VCD group and V20DT group (MD: 5.44; 95% CI: -0.81 to 11.69). Patients in 200AOD showed a significant improvement in mean IIEF scores when compared to 100 mg avanafil on demand (100AOD) (MD: 1.60; 95% CI: 1.46 to 1.74). The SUCRA ranking of PFMT has been greatly improved with the assistance of biofeedback (BF) or electrical stimulation (ES). The second group contained 301 patients with four treatments. The network meta-analysis indicated that VCD and 50 mg/day sildenafil VC50DS showed superiority over 50 mg/day sildenafil (50DS) (MD: 3.75; 95% CI: 2.74-4.76) and intraurethral alprostadil 125-250 μg (MD: 3.05; 95% CI: 0.38 to 5.72), respectively (Figure 4F). The SUCRA ranking showed that VC50DS ranked 1st and 50DS ranked the fourth (Figure 4B).

Thirteen studies with 1,052 participants involved 16 interventions and presented available results for ≥6 months mean IIEF scores after RP. Eight of sixteen interventions were associated with significant improvement in mean IIEF scores when compared to placebo (Figure 5). The MDs for interventions related to significant improvement ranged from 1.6 (95% CI: 1.25–1.95) for 80 mg/day atorvastatin (80DA) to 14.22 (95% CI: 9.04–19.40) for V20DT. 10 mg/day atorvastatin and 50 mg sildenafil on demand (10DA50SOD), V20DT, VCD, 20 mg/day tadalafil (20DT),
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Table 1 The main characteristics of the included studies in network meta-analysis

<table>
<thead>
<tr>
<th>Study</th>
<th>Country</th>
<th>Participants (age and number)</th>
<th>GS</th>
<th>PSA* (ng/mL)</th>
<th>DM</th>
<th>TPCDC</th>
<th>SUA</th>
<th>CLS</th>
<th>NES</th>
<th>Intervention</th>
<th>Outcome measures</th>
<th>LoE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aydogdu et al. [2011]</td>
<td>Turkey</td>
<td>N=74 (87.8% completed study); IG = 32; 58.2; CG = 33; 58.1</td>
<td>&lt;8</td>
<td>6.1 [3.6–9.6]</td>
<td>C3G: 4/33; IG: 5/32</td>
<td>At baseline, 3, 6, 12 months</td>
<td>RRP  ≤ 1c</td>
<td>BNES</td>
<td>IG: 3 days/week; Tadalafil 20 mg/day for 6 months following the removal of urethral catheter; CG: no use of tadalafil</td>
<td>IIEF- EF; SEP-2; SEP-3</td>
<td>1b</td>
<td></td>
</tr>
<tr>
<td>Montorsi et al. [2008]</td>
<td>Europe, Canada, South Africa, and USA</td>
<td>N=668 (95.3% completed study); IG1=207; 57.4; IG2=204; 56.8; CG = 206; 57.1</td>
<td>≤ 10</td>
<td>NS</td>
<td>At 0, 1, 3, 6, 9, 10, 11, 12, 13 months</td>
<td>NS</td>
<td>I-II</td>
<td>BNES</td>
<td>IG1=10 months; 10 mg vardenafli nightly [which could be decreased to 5 mg if required]; IG2=9 months flexible dose starting at 10 mg with the option to titrate to 5 mg or 20 mg, on-demand vardenafli; CG = 9 months placebo</td>
<td>IIEF- EF; SEP-2; SEP-3</td>
<td>1b</td>
<td></td>
</tr>
<tr>
<td>Padma-Nathan et al. [2008]</td>
<td>North America, France, Belgium and Australia</td>
<td>N=123 (61.7% completed study); IG1=34; 55 [6]; IG2=28; 55 [9]; CG = 25; 57 [7]</td>
<td>≤ 10</td>
<td>None</td>
<td>At 0, 9 months</td>
<td>ORP; RARP; LRP</td>
<td>Htc-I2c</td>
<td>BNES</td>
<td>IG1=9 months, 5 mg tadalafil once daily; IG2=9 months, 20 mg tadalafil on demand; CG = placebo</td>
<td>IIEF- EF; SEP-3; penile length</td>
<td>1b</td>
<td></td>
</tr>
<tr>
<td>Pavlovich et al. [2013]</td>
<td>USA</td>
<td>N=100 (100% completed the study); IG = 50; 54.3 [42–63]; CG = 50; 53.6 [40–64]</td>
<td>≤ 10</td>
<td>NS</td>
<td>1%</td>
<td>At 1, 3, 6, 9, 12, 13 months</td>
<td>LRP; RARP</td>
<td>Htc-I2a</td>
<td>Yes</td>
<td>IG = nightly 50 mg sildenafil once for 1 year; CG = on demand 50 mg sildenafil once for 1 year [maximum six tablets/month]</td>
<td>IIEF- EF; EPIC</td>
<td>1b</td>
</tr>
<tr>
<td>Barrowsky et al. Germany [2012]</td>
<td>Germany</td>
<td>N=36 (100% completed the study); IG1=11; 56.6; IG2=12; 56.7; CG = 11; 56.6</td>
<td>&lt;7</td>
<td>6.6 [5.2–7.2]</td>
<td>At baseline, 3, 6, and 12 months</td>
<td>RRP I-II</td>
<td>BNES</td>
<td>IG1=12 months, 5 mg/day vardenafli; IG2=12 months, 10 mg/day vardenafli; CG = 12 months, placebo</td>
<td>IIEF-5</td>
<td>1b</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Canat et al. [2015]</td>
<td>Turkey</td>
<td>N=112 (100% completed the study); IG1=38; 62.6 [50–72]; IG2=29.5 [54–72]; CG = 34; 63.52 [52–74]</td>
<td>≤ 10</td>
<td>IG1=6/38; IG2=6/40; CG = 6/64</td>
<td>At 6 weeks, 12 months</td>
<td>RRP I-II</td>
<td>BNES</td>
<td>IG1=20 mg tadalafil three times per week; IG2=20 mg gadalfili on demand; CG = on treatment; Duration: 12 months</td>
<td>IIEF-6</td>
<td>1b</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Montorsi et al. [2004]</td>
<td>Canada, Germany, Italy, The Netherlands, Spain, UK and USA</td>
<td>N=303 (78.2% completed the study); IG = 201; 59.6 [5.0]; IG = 102; 59.5 [5.2]</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>At baseline, 3 months</td>
<td>RRP  ≤ 11</td>
<td>BNES</td>
<td>IG1=20 mg tadalafil for 12 weeks; CT ≥ 20 mg placebo</td>
<td>IIEF- EF; SEP-2; SEP-3</td>
<td>GAQ</td>
<td>2b</td>
</tr>
<tr>
<td>Muhlall et al. [2013]</td>
<td>USA</td>
<td>N=298 (84.6% completed study); IG1=119; 58.9 [6.8]; IG2=99; 57.5 [6.8]; IG3=58.6 [6.8]</td>
<td>s7</td>
<td>NS</td>
<td>None</td>
<td>At 0, 1, 2, 3 months</td>
<td>RRP; ORP; LRP; RARP ≥ 11</td>
<td>BNES</td>
<td>IG1=100 mg avanafil; IG2=200 mg avanafil; CG = placebo; Duration: 3 months</td>
<td>IIEF- EF; SEP-2; SEP-3</td>
<td>1b</td>
<td></td>
</tr>
<tr>
<td>Raina et al. [2006]</td>
<td>USA</td>
<td>N=109 (87% completed study); IG = 74; 58.2; IG = 35; 58.2</td>
<td>s6</td>
<td>&lt;10</td>
<td>NS</td>
<td>At baseline, 0, 9 months</td>
<td>NS</td>
<td>I-II</td>
<td>BNES</td>
<td>IG: VCD use daily for 9 months; CG: no treatment</td>
<td>IIEF-5; penile length and circumference</td>
<td>1b</td>
</tr>
<tr>
<td>Köhler et al. [2007]</td>
<td>USA</td>
<td>N=28 (100% completed study); IG = 17; 58.2; CG = 11; 60.5</td>
<td>IG1 = 6.7; IG2 = 5.6; IG3 = 6.7; IG4 = 7.5; CG = 5.5</td>
<td>At baseline, 3, 6, 9, 12 months</td>
<td>RRP  ≤ 11</td>
<td>BNES; UNES</td>
<td>IG: early intervention (1 month after RP); IG2=10 minutes/visit VCD for 5 months; CG: control group (6 months after RP)</td>
<td>IIEF-5; penile length and circumference</td>
<td>1b</td>
<td></td>
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<tr>
<td>Engel et al. [2011]</td>
<td>USA</td>
<td>N=23 (87% completed study); IG = 13;</td>
<td>NS</td>
<td>NS</td>
<td>None</td>
<td>At 3, 6, 9, 12 months</td>
<td>RARP</td>
<td>NS</td>
<td>BNES</td>
<td>IG: 20 mg/day tadalafil three times per week plus a VCD, 10 minutes unbanded per day for at least 5 days weekly; IG2=20 mg/day tadalafil three times per week</td>
<td>IIEF-5; penile erection hardness</td>
<td>1b</td>
</tr>
<tr>
<td>Liu et al. [2016]</td>
<td>China</td>
<td>N=64 (100% completed study); IG = 32; 57.6 [4.1]; CG = 32; 56.9 [4.5]</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>At 0, 3 months</td>
<td>ORP</td>
<td>NS</td>
<td>No limitation</td>
<td>IG: 3 months, 50 mg/day sildenafil nightly and VCD; CG: 3 months, 50 mg/day sildenafil nightly</td>
<td>IIEF-5; penile length and circumference; erectile hardness</td>
<td>1b</td>
</tr>
</tbody>
</table>
Table 1 (continued)

<table>
<thead>
<tr>
<th>Study</th>
<th>Country</th>
<th>Participants (agea and number)</th>
<th>GS</th>
<th>PSA (ng/mL)</th>
<th>DM</th>
<th>TPDC</th>
<th>SUA</th>
<th>CLS</th>
<th>NES</th>
<th>Intervention</th>
<th>Outcome measuresb</th>
<th>LoE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laurienzo et al. [2018]</td>
<td>Brazil</td>
<td>N=112 (93.1% completed the study); IG1=41; 58.5 (5.4); CG = 40; 55.5 (5.9)</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>At baseline, 1, 3, 6 months</td>
<td>NS</td>
<td>I2a-IIIb</td>
<td>NS</td>
<td>IG1: PFMT; IG2: 5/40; 58 (5.4); CG: no treatment</td>
<td>IIEF-5</td>
<td>1b</td>
</tr>
<tr>
<td>Glazner et al. [2011]</td>
<td>UK</td>
<td>N=411 (100% completed the study); IG = 206; 62.4 (5.8); CG = 206; 62.3 (5.6)</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>At 3, 6, 9, 12 months</td>
<td>LRP; abdominal; perineal</td>
<td>NS</td>
<td>NS</td>
<td>IG: therapist-guided PFMT, four one to one sessions, 3 months; CG: no treatment</td>
<td>Number of men unable to achieve any erection 12 months after prostate surgery</td>
<td>1b</td>
</tr>
<tr>
<td>Geraets et al. [2016]</td>
<td>Belgium</td>
<td>N=109 (100% completed the study); IG = 16; 61.1; CG = 17; 61.5</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>At 0, 3 months</td>
<td>ORP; RARP</td>
<td>NS</td>
<td>BNES; UNES</td>
<td>IG: therapist-guided PFMT and ES, during 10 min (biphasic symmetric current (constant voltage), intensity as high as possible, not painful, frequency: 50 Hz and pulse duration: 600 μs); CG: no treatment</td>
<td>IIEF-ES; VAS</td>
<td>1b</td>
</tr>
<tr>
<td>Prota et al. [2012]</td>
<td>Brazil</td>
<td>N=52 (82.5% completed the study); IG = 17; 62.4 (8.6); CG = 16; 64.0 (8.0)</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>At 1.36 and 12 months</td>
<td>RRP</td>
<td>II-III</td>
<td>NS</td>
<td>IG: PFMT and BF, once a week for 12 weeks after catheter removal at postoperatively day 15; CG: no treatment</td>
<td>IIEF-5</td>
<td>2b</td>
</tr>
<tr>
<td>de Lira et al. [2019]</td>
<td>Brazil</td>
<td>N=31 (100% completed the study); IG = 16; 67.3 (6.3); CG = 15; 63.3 (7.62)</td>
<td>No limitation</td>
<td>IG: 9.20 (4.65); CG: 14.1 (11.19)</td>
<td>NS</td>
<td>At baseline, 3 months</td>
<td>RRP</td>
<td>I2c-IIIb</td>
<td>NS</td>
<td>IG: PFMT and BF, two pre-RP physical therapist-guided PFMT sessions, including exercises and electromyographic biofeedback, and verbal and written instructions to continue PFMT until RP, which was then resumed after urethral catheter removal; CG: only usual post-prostatectomy care</td>
<td>IIEF-5; ICIQ-SF</td>
<td>1b</td>
</tr>
<tr>
<td>Oh et al. [2020]</td>
<td>Korea</td>
<td>N=84 (97.6% completed the study); IG = 40; 67.5 (8.9); CG = 42; 65.9 (8.8)</td>
<td>No limitation</td>
<td>IG: 5/40; CG: 9/42</td>
<td>At baseline, 1, 2,3 months</td>
<td>RARP</td>
<td>NS</td>
<td>BNES; UNES</td>
<td>IG: PFMT and BF, 8 times per day; (II) 10 minutes per session of exercise; (III) a minimum of 10 seconds of tension duration with maximal tension intensity; CG: PFMT</td>
<td>IIEF-5</td>
<td>1b</td>
<td></td>
</tr>
<tr>
<td>Chiles et al. [2018]</td>
<td>USA</td>
<td>N=109 (85.3% completed the study); IG = 40;</td>
<td>NS</td>
<td>NS</td>
<td>None</td>
<td>At baseline, 12, 18 months</td>
<td>RARP</td>
<td>NS</td>
<td>BNES</td>
<td>IG: 100% oxygen in a hyperbaric chamber, 10 sessions (90 minutes of 100% oxygen at 2.0 ATA) beginning day 1 after hospital discharge. Sessions were continued daily on Monday through Friday for an additional 9 days. CG: air; Both groups received 50 mg sildenafil daily for 12 months beginning at the completion of hyperbaric treatment at 15 days after surgery</td>
<td>IIEF-EPIC-26</td>
<td>1b</td>
</tr>
<tr>
<td>McCullough et al. [2010]</td>
<td>USA</td>
<td>N=212 (73.5% completed the study); IG = 97; 58.8 (6.4); CG = 95; 55.5 (6.9)</td>
<td>&gt;7</td>
<td>20</td>
<td>NS</td>
<td>At baseline, 0, 1, 3, 6; 9, 10, 11 months</td>
<td>ORP; RARP</td>
<td>NS</td>
<td>BNES</td>
<td>IG: nightly intraurethral alprostadil, initially 125 μg and dose titrated 250 μg at 1 month and maintained for 8 months; CG: 50 μg sildenafil nightly</td>
<td>IIEF-EP</td>
<td>2b</td>
</tr>
<tr>
<td>Mihal et al. [2018]</td>
<td>USA</td>
<td>N=131 (94.6% completed the study); IG = 59; 55.1 (8.2); CG = 58; 54.1 (6.2)</td>
<td>NS</td>
<td>NS</td>
<td>None</td>
<td>At 1 w, 2 w, 4 w, 6 w; 3 m, 6 m, 12 m, 18 m</td>
<td>OPP</td>
<td>II-BNES</td>
<td>IG: tacrolimus 2-3 mg daily for 27 weeks (1 week prior to and 6 months after RP) and followed up for 2 years after RP; CG: placebo</td>
<td>IIEF-5</td>
<td>1b</td>
<td></td>
</tr>
<tr>
<td>Sittari et al. [2019]</td>
<td>Finland/Denmark</td>
<td>N=158 (74.6% completed the study); IG = 62; 61.9 (5.4); CG = 63; 58.8 (6.9)</td>
<td>No limitation</td>
<td>No limitation</td>
<td>IG: 10%; CG: 12%</td>
<td>At baseline, 3, 6, 9, 12 months</td>
<td>NS</td>
<td>I-III</td>
<td>No limitation</td>
<td>IG: 80 mg atorvastatin daily from study inclusion to the day of surgery; CG: placebo</td>
<td>IIEF-5</td>
<td>2b</td>
</tr>
<tr>
<td>Hong et al. [2007]</td>
<td>Korea</td>
<td>N=50 (98% completed the study); IG = 20; 61.3 (4.3); CG = 30; 60.8 (2.3)</td>
<td>IG: 6.2 (0.7); CG: 6.1 (1.2)</td>
<td>IG: 7.97 (2.3); CG: 7.44 (4.5)</td>
<td>NS</td>
<td>At baseline, 0, 6 months</td>
<td>RRP</td>
<td>II-BNES</td>
<td>IG: 10 mg atorvastatin daily from postoperative days 1 to 90 and on demand 50 mg sildenafil; CG: on demand 50 mg sildenafil</td>
<td>IIEF-5</td>
<td>1b</td>
<td></td>
</tr>
</tbody>
</table>

*年龄报告为均数和SD unless specified; **仅测量与勃起功能相关的措施；***PSA before treatment; IG: intervention group; CG: control group; BF: biofeedback; EF: erectile function; ES: electrical stimulation; GS: Gleason score; RP: radical prostatectomy; PDE5is, phosphodiesterase type 5 inhibitors; IIEF: International Index Erectile of Function; PSA, prostate specific antigen; DM, diabetes mellitus; TPDC, time points of data collection; NES, nerve sparing; bilateral NES, unilateral NES, UNES; SUA, Surgical approach; CLS, Clinical stage; LoE, level of evidence; RRP, radical retropubic prostatectomy; NS, not specified; EF: erectile function; SEF: sexual encounter profile; ORP, open RP; LRP, laparoscopic RP; RARP, robot-assisted RP; EPIC, Expanded Prostate Cancer Index Composite; GAO, global assessment question; VAS, visual analog scale; PFMT, pelvic floor muscle training.

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Figure 3 Network plots and funnel plots. Mean IIEF scores within 3 months after surgery: (A,E) and (B,F). Mean IIEF scores ≥6 months after surgery: (C,G). The proportion of mean IIEF scores ≥22: (D,H).
Figure 4 Surface under the curve cumulative probabilities (SUCRA) ranking and network results of mean IIEF scores within 3 months after surgery. Mean IIEF scores within 3 months after surgery: (A,E) and (B,F). Mean IIEF scores ≥6 months after surgery: (C). The proportion of mean IIEF scores ≥22: (D).
Figure 5: Network meta-analysis results of mean IIEF scores ≥6 months after surgery.

10 mg/day vardenafil (10DV), and 5 mg/day vardenafil (5DV) improved mean IIEF scores significantly more than many other interventions (Figure 5). Moreover, V20DT showed significant superiority over the other interventions, which was consistent with the SUCRA ranking (Figure 4C).

**Network meta-analysis of the proportion of IIEF scores ≥22**

Ten studies with 1,723 patients involved 13 interventions, and presented usable results for the proportion of IIEF scores ≥22 greater than or equal to 6 months after RP. Of these interventions, only PFMT and BF (PFBF) were conducted as a combination therapy. Six of thirteen treatments had higher proportions of IIEF scores ≥22 than tacrolimus, which was consistent with the SUCRA ranking (Figure 4D). Despite 50 mg sildenafil on demand (50SOD) ranking 1st, most differences among these interventions were not significant (Figure 6). Notably, the results of network meta-analysis between the proportion of IIEF scores ≥22 and mean IIEF scores ≥6 months after RP were approximately congruent in terms of monotherapy.

**Consistency and convergence analysis**

No significant inconsistency was observed among the various interventions through node-splitting analysis, which meant the consistency model was reliable. No obvious loop inconsistency was detected by the test of loop inconsistency and the IF was relatively small (Figure 2). Besides, no significant publication bias was observed in the outcomes according to the funnel plot (Figure 3).

**Discussion**

Different from the other secondary impotency, post-prostatectomy ED is mainly caused by neurovascular bundle injury. The erectile nerve plays an important role in the process of erection initiating and maintenance by activating the release of nitric oxides (NOs) and substantially increasing the prostaglandin E-1 (PGE-1) level (51-53). It has been demonstrated that nerve terminal released NOs have relaxant effects on the penis by increasing oxygenated blood flow to the erectile cavernous tissue and relaxing the smooth muscle fibers of the arteries and arterioles of the erectile tissue (54-58). This effect could be altered when the integrity of the nerve was damaged. Originated from the pelvic plexus, CN walks along the anterolateral side of the...
**Figure 6** Network meta-analysis results of the proportion of mean IIEF scores ≥22. 20DT, 20 mg/day tadalafil; 20TOD, 20 mg tadalafil on demand; 100AOD, 100 mg avanafil on demand; 200AOD, 200 mg avanafil on demand; PFMT, pelvic floor muscle training; PFES, PFMT+ electrical stimulation (ES); PFBB, PFMT and biofeedback (BF); TAC, tacrolimus; VCD, vacuum constriction devices; V20DT, VCD+20DT; 80DA, 80 mg/day atorvastatin; 5DV, 5 mg/day vardenafil; 10DA50SOD, 10 mg/day atorvastatin+50SOD; 10DV, 10 mg vardenafil on demand; 5DT, 5 mg/day tadalafil.
prostate (59). As the anatomical structure of the CN and prostate could be so tight, CNs injury is common during operation. Though more mini-invasive and precise surgical approaches like nsRARP have been widely recommended for its advantage in providing better visualization of the operative field for surgeons hopefully for protecting CN and a lower occurrence of RP induced ED was achieved, the morbidity of post-prostatectomy ED could still be high (13,60). As is believed that not only by cutting, coagulation, traction, and compression of the pelvic tissues during operation could CNs injury be caused, postoperative local hypoxia and postoperative neuropraxia-related fibrotic and apoptotic changes in the erectile tissue were also responsible for post-prostatectomy ED (61,62). Despite the prominent understanding of the mechanism of post-prostatectomy ED, CNs injury is inevitable, even with the progress of surgical techniques. Thus, ED is a common sequela in patients after RP for clinically localized PCa, with a negative impact on the quality of life and intimate relationship of patients and partners. Besides, the persistent lack of ED after neuropraxia can itself cause deleterious chain reactions that can negatively affect EF. Ligation of the accessory internal pudendal arteries and neuropraxia results in hypoxia and a lack of nocturnal erections. This results in corpus cavernosum fibrosis and the transformation of trabecular smooth muscle by collagen, which itself results in the loss of the venous occlusion mechanism required to maintain an erection (63). Nerve damage coupled with reduced arterial inflow may exacerbate hypoxia and ultimately lead to apoptosis (63). Therefore, the focus of improving ED after RP is to increase the amount of oxygen in the cavernous body and reduce tissue fibrosis and apoptosis.

Predictive factors for post-prostatectomy ED seem to be associated with age, preoperative baseline function, comorbidities index, an extension of the nerve-sparing procedure, surgery type (intrafascial, interfascial or extrafascial), surgical techniques (open, laparoscopic, RARP), operator’s surgical experience, and the use of cautery-free dissection or the use of pinpointed low-energy cauteryization (6,13). Although some patients might gradually recover 2 years or longer after surgery, only a few patients can return to their baseline EF (14). The concepts of penile rehabilitation were proposed to facilitate recovery of EF after RP as much as possible with any drug or device. Based on the current RCTs in our study, treatments currently available include PDE5is, PFMT, vacuum therapy, neuromodulatory therapy, statin therapy, intracorporeal injection therapy, hyperbaric oxygenation therapy, and combination therapies of these interventions. This network meta-analysis indicated that V20DT might have a better effect than other interventions within 1 year after surgery. However, the effectiveness of VCD may be affected by the equipment itself and the patients’ experience. That’s why VCD ranked 1st within 3 months after RP. PFMT might perform better with the guidance of BF and BS. Monotherapy appeared to have similar efficacy in terms of mean IIEF scores and proportion of patients return to baseline, and the effect of PDE5is did not seem to be affected by the patterns of administration (regular or on demand).

The goal of penile rehabilitation includes promoting early recovery of EF, reducing the loss of penis length and circumference, and improving overall sexual intercourse satisfaction. There is no definite conclusion regarding the timing of penile rehabilitation. Mulhall et al. (64) found patients in starting interventions within 6 months after surgery had a significantly higher mean IIEF-5 scores than those who starting interventions 6 months or longer after RP (22 vs. 16, P<0.001). Besides, stem cell therapy currently shows some promising results (65-67).

Despite ED after RP is a neurovascular injury event, psychological trauma may be a factor that cannot be ignored. Diagnosis of PCa is a life-changing event that can cause considerable psychological stress on patients and their sexual partners, and patients’ anxiety can lead to ED before treatment after cancer diagnosis and before pathological results after prostate biopsy (68-70). The psychological effects before RP may have a substantial impact on preoperative and postoperative EF, exacerbating the adverse effects of surgery on EF (70).

The current trend in disease management is individualized treatment, and penile rehabilitation after RP is no exception. To our knowledge, this is the first study that synthetically commenced evaluating the efficacy of various interventions on penile rehabilitation for patients suffering from ED after RP. Despite comprehensive analysis being performed to make the summary results accurate and convincing, our study does have the following limitations. Firstly, limited RCTs might result in publication bias, and prevent us from reaching sufficient conclusions. Secondly, although considerable progress has been made in the study of EF rehabilitation after RP, there are still no conclusions in this field. Many studies lack the necessary data, such as using patient self-reported questionnaires, and there is significant heterogeneity, such as the definition of population, study design, and outcome measures. Besides,
biological data such as penis scans or Doppler ultrasound are scarce. Finally, we were unlikely to further evaluate the long-term results of various treatments due to the lack of sufficient data in the included studies, which might make our analysis defective.

Conclusions

The combination therapy shows certain advantages over monotherapy, and we recommended the combination of VCD and PDE5is to be considered in the clinical management of penile rehabilitation after RP. High-quality meta-analysis is still warranted based on well-designed RCTs, and other emerging therapies with promising results are also worth further exploring.

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Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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