Prostatic Artery Embolization for Benign Prostatic Hyperplasia: Anatomical Aspects and Radiation Considerations from a Case Series of 210 Patients

Mohamed Shaker 1 Essam Hashem 2 Ahmed Abdelrahman 3 Ahmed Okba 1

1 Department of Diagnostic and Interventional Radiology, Ain Shams University, Cairo, Egypt
2 Department of Diagnostic and Interventional Radiology, Ain Shams University, King’s College Hospital, Cairo, Egypt
3 National Center for Radiation Research and Technology, Cairo, Egypt

Address for correspondence Mohamed Shaker, MD, 13 Mostafa Refaat, Sheraton Al Matar, El Nozha, Cairo Governorate 11799, Egypt (e-mail: mohamed_ghazy@med.asu.edu.eg).

Abstract

Context Prostatic artery embolization (PAE) has been established as a safe and effective treatment option for symptomatic benign prostatic hyperplasia (BPH). Thorough knowledge of detailed prostatic artery (PA) anatomy is essential.

Aims The aim of this study was to provide a pictorial review of PA anatomy and prevalence of related anatomical variants, in addition to other anatomical and radiation dose considerations.

Settings and Design Case series and review of literature.

Materials and Methods We performed PAE for 210 patients from November 2015 to November 2020 under local anesthesia only. Anatomy, procedure duration, fluoroscopy time, radiation dose, technical success, and complications were analyzed.

Statistical Analysis Used Descriptive statistics were analyzed using Microsoft Excel software.

Results A total of 210 patients (420 sides) were analyzed. Double arterial supply on the same side was noted in 12 patients (5.7%). In 10 patients (4.7%), only a unilateral PA was identified. In two patients (0.9%), no PA could be identified. Frequencies of PA origins were calculated. Penile, rectal, and vesical anastomoses were identified with 79 (18.8%), 54 (12.9%), and 41 (9.8%) of PAs, respectively. Median skin radiation dose, procedure time, and fluoroscopy time were 505 mGy, 73 and 38 minutes, respectively. Complications occurred in nine patients (4.3%), none of them was major.

Conclusions Knowledge of PA anatomy is essential when treating BPH by PAE for optimum results. There is no enough evidence to support routine use of preoperative computed tomography angiography and intraoperative cone-beam computed tomography as means of improving safety or efficacy.

Keywords
► prostate
► embolization
► prostatic artery embolization
► angiographic anatomy
► radiation
► fluoroscopy time
► prostatic anastomosis
► nontarget embolization
► cone-beam CT


DOI https://doi.org/10.1055/s-0041-1729134
ISSN 2542-7075

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Published online: 2021-06-22
Introduction

PAE has been established as a safe and effective minimally invasive treatment option for moderate and severe lower urinary tract symptoms (LUTS) due to benign prostatic hyperplasia (BPH).1

Compared with standard urological interventions, PAE is performed as an outpatient day-case procedure,1 suitable for large prostates,2,3 suitable for patients with acute urinary retention,4 and has a lower overall cost.5 PAE has significantly lower risk of severe adverse events compared with standard urological interventions.6-10

PAE is often a challenging procedure requiring certain level of knowledge of the arterial anatomy and set of skills, due to varying degrees of atherosclerosis commonly encountered in this age group.11 We aim to highlight important anatomical data related to PA that are essential for successful PAE as well as radiation exposure considerations, which can be beneficial in such a relatively lengthy procedure.

Subjects and Methods

This retrospective study is based on the angiograms of 210 male patients, who underwent PAE between November 2015 and November 2020, to treat obstructive LUTS due to BPH in patients above the age of 40 with prostate volume more than 40 mL showing no satisfactory response to medical treatment for at least 6 months. Patients with prostate cancer, impaired renal functions, urinary bladder stones, chronic retention, active urinary tract infection, active prostatitis, or uncorrectable coagulopathy were excluded.

Preprocedural evaluation included history taking, clinical examination, International Prostate Symptom Score (IPSS) and quality of life (QOL) score, maximum urine flow velocity measurement by urine flowmetry that was done in some cases to be sure of the obstructive nature of the patient complaint, ultrasonographic examination to evaluate the size of the prostate and postvoiding residual urine volume as well as laboratory investigation panel that included prostate-specific antigen (PSA) with free/total PSA ratio, urine analysis, complete blood picture, bleeding profile, and kidney function tests. Transrectal prostatic biopsy was done for patients with suspiciously high PSA levels. Preoperative investigations did not include angiographic study (e.g., CT or magnetic resonance angiography). Cases included in this study were performed on Philips BV Pulsera machine (Philips Healthcare, Andover, MA, United States). Intraoperative CBCT was not used.

Informed consent was obtained. All procedures were performed as day cases under local anesthesia via the right common femoral access. An initial diagnostic internal iliac angiogram was performed using digital subtraction angiography (DSA) by a 5-French Cobra-2 catheter (Boston Scientific, Marlborough, MA, United States) starting by the left side followed by the right side; images were acquired in an ipsilateral anterior oblique of 30 to 50 degrees and a cranial angulation of 10 to 15 degree to determine the origin and number of PAs, followed by selective catheterization of identified PAs using a 2.4 or 2.7 French microcatheter (Progreat, Terumo, Tokyo, Japan). PA angiogram was performed at anteroposterior, oblique, and sometimes lateral views to detect any extra-prostatic supply or anastomosis. CBCT and preoperative CT angiography (CTA) were not utilized, and we depended upon the angiographic anatomy with different projections to avoid nontarget embolization.

Posterior divisions of internal iliac arteries (IIA) and external iliac arteries were only catheterized if additional supply was suspected (e.g., incomplete prostate blush) or PAs could not be identified from the anterior divisions. Embolization was achieved using 300 to 500 µm Embosphere particles (Merit Medical Systems Inc., South Jordan, UT, USA) mixed with contrast and diluted 1:1. The procedure was considered technically successful if at least one PA could be embozilized to stasis.

Postprocedural hemostasis was achieved by manual compression of the femoral artery and patients were discharged on the same day after 4 to 6 hours from achieving hemostasis. Discharge medications included antibiotic prophylaxis, non-steroidal anti-inflammatory medications, and analgesics for 7 to 10 days to control the postembolization symptoms. All patients were evaluated at the clinic 2 to 3 weeks after PAE for the presence of complications and initial response to therapy. Further follow-up at the clinic was performed at 3 and 6 months by ultrasound examination, IPSS, and QOL scores.

Our anatomical review is based upon a simple yet informative classification proposed by Dr Carnevale’s group.11

Anatomy, procedure duration (PD), radiation dose, fluoroscopy time (FT), technical success, and complications were analyzed. Descriptive statistics were analyzed using Microsoft Excel software.

Results

The mean age was 63 years (range: 48–90). Images of the 210 patients (420 sides) were analyzed by the operators (who are also the authors). A total of 418 PAs were angiographically identified. Double arterial supply on the same side was noted in 12 patients (5.7%). All duplicates were observed on the left side except one; internal pudendal artery (IPA) and superior vesical artery (SVA) in six patients; IPA and obturator artery (OA) in 4 patients; two separate arteries arising from the IPA in one patient and two separate arteries arising directly from the right anterior division IIA in another patient (see Fig. 1). In 10 patients (4.7%), only a unilateral PA

Fig. 1 (A) Right internal iliac angiogram showing Duplicated prostatic artery (PA) arising from the anterior division of internal iliac artery. (B) Selective left prostatic artery angiogram showing retrograde filling of a duplicated prostatic artery.
was identified despite thorough search in both internal and external iliac branches; this was due to severe atherosclerotic occlusion; in only two patients (0.9%) no PA could be identified on either side due to severe aortoiliac occlusive disease. Technical success was defined as successful embolization of at least one PA; this was achieved in 208 cases (99%). Bilateral embolization was successfully achieved in 189 cases (90%). Frequencies of PA origins were 167 (39.9%) from SVA (type I), 122 (29.2%) from IPA (type IV), 79 (19%) from OA (type III), 46 (11%) originated directly from anterior division of IIA (type II), and only 4 (0.9%) originated from elsewhere (type V). The unusual origins were one from external iliac, one from superior gluteal artery, and two from inferior gluteal arteries; in the two latter cases, the inferior gluteal arteries were arising separately early from the anterior division rather than being its continuation. Figs. 2-6 show several angiographic images of each anatomical type according to Carnevale’s classification.11 Bilaterally symmetrical PA origins were found in 99 patients (47%). Out of these 99 patients with symmetrical origins, 43 (43.4%) originated from SVA, 28 (28.3%) from IPA, 17 (17.2%) from OA, and 11 (11.1%) directly from the anterior division.

Fig. 2 (A) Right internal iliac artery (IIA) DSA showing type I origin of the prostatic artery (PA) arising with the superior vesical artery (SVA) by a common trunk from the anterior division of the internal iliac artery. (B) Left internal iliac artery DSA showing type I origin of the prostatic artery (PA) arising with the superior vesical artery (SVA) by a common trunk from the anterior division of the internal iliac artery (Ant.).

Fig. 3 (A) Right internal iliac artery DSA showing Type II origin of the prostatic artery (arrows) arising by a separate origin from the anterior division of the internal iliac artery (arrowhead). (B) Left internal iliac artery DSA showing Type II origin of the prostatic artery (arrows) arising by a separate origin from the anterior division of the internal iliac artery (Ant.).

Fig. 4 (A) Selective Left obturator angiogram showing Type III origin of the prostatic artery (arrows) arising from the upper third of the obturator artery (Obt). (B) Selective Left obturator angiogram showing Type III origin of the prostatic artery (arrows) arising from the middle third of the obturator artery (Obt).

Fig. 5 (A) Left internal iliac angiogram showing Type IV origin of the prostatic artery (PA) arising from the internal pudendal artery (int. P). (B) Selective internal pudendal angiogram showing Type IV origin of the prostatic artery (PA) arising from the internal pudendal artery (int. P).

Fig. 6 Left hemipelvic angiogram for the same patient shows type V origin of prostatic artery arising from external iliac artery (EIA): (A) Left internal iliac artery (IIA) angiogram revealed absent obturator (Obt) and prostatic arteries; (B) left external iliac artery (EIA) gives rise to the obturator artery; (C) selective angiogram of the obturator artery demonstrating prostatic artery supplying prostatic blush (arrowhead), note accessory pudendal (arrows) artery arising from the left prostatic artery.
Penile, rectal, and vesical anastomoses were angiographically identified with 79 (18.8%), 54 (12.9%), and 41 (9.8%) of PAs, respectively. **Figs. 7-9** show examples of these anastomoses. Accessory pudendal supply arising from PA is also a potential channel for nontarget embolization; this was encountered with 37 PAs (8.8%) (►Fig. 10). In order of decreasing frequency, the utilized protective techniques to overcome the anastomoses were very slow injection of particles, intraprostatic vasodilators (glyceryl trinitrate 200 µg), coils, and Gelfoam. Protective Gelfoam or coil embolization for nontarget anastomoses was used in 10 cases (4.7%). Gelfoam (thin slurry) was used in 2 of the 10 cases; both were vesical anastomoses with no enough landing vessel length or diameter to accommodate the smallest available coil in our inventory (2 mm). Anastomosis between both hemiprostates was detected in 38 patients (18%), manifesting as filling of contralateral PA branches (►Fig. 11). In two cases, the anastomosis was extensive enough to allow full embolization of the entire gland from one artery only.\(^\text{12}\)

**Fig. 7** Selective left prostatic artery angiogram showing intra-prostatic pudendal anastomosis (arrowheads).

**Fig. 8** (A) Selective right prostatic artery angiogram revealing prostatic blush with no angiographically visible anastomosis (B). Intermittent angiography during embolization revealed opening of the rectal anastomosis with retrograde filling of the superior rectal artery (arrowheads) through anastomosis.

**Fig. 9** (A) Selective left prostatic artery angiogram reveals vesical anastomosis (arrows). (B) Selective left prostatic artery angiogram revealed rectal (arrows) and vesical (arrowheads) anastomoses

**Fig. 10** Selective prostatic artery angiogram revealed accessory pudendal artery (arrows), notice the appearance of right hemiprost- tate blush through intra-prostatic anastomosis (arrowheads).

**Fig. 11** selective left prostatic artery angiogram for 2 different patients reveals retrograde filling of the contralateral prostatic artery (arrowheads) through intra-prostatic anastomosis. Notice the accessory pudendal artery (Arrows) arising from the left prostatic artery.

Median skin radiation dose, total procedure time, and FT were 505 mGy, 73 and 38 minutes, respectively.
- Table 1 provides detailed data in this regard. Mean prostate volume, median prostate volume, and range were 125, 115, and 45 to 350 mL, respectively. Postembolization syndrome was dealt with as a normal sequel rather than a complication, as it was encountered in almost all patients with varying degrees and it subsided in all patients within 1 to 2 weeks of conservative management (analgesic, anti-inflammatory, and prophylactic antibiotic).

Nine patients experienced minor adverse events (4.3%): four cases of hematospemria (all subsided within 4 weeks without specific treatment), one case of acute severe pelvic pain on the same night of the operation (successfully treated by intravenous [IV] analgesics on outpatient basis), one case of chronic mild pelvic floor pain that persisted for 3 months, and three cases of urinary tract infection (treated by antibiotic therapy). We encountered no severe adverse events according to standardized incidence ratio (SIR) classification.13 We did not encounter any cases of hematuria, hematochezia, or genital ulcers.

**Discussion**

This study highlights detailed anatomy of the PA and its variants.11 Almost half of the cases had symmetrical origin of PA; this fact should be exploited to spare PD, radiation, and contrast use when searching for the contralateral PA. If the origin is not readily obvious on DSA, one should start by exploring SVA and IPA, because collectively they contribute to about two-thirds of PA origins. Pisco group reported that IPA is the most common origin of PA (35%) followed by SVA (20%).14 Carnevale group reported IPA and SVA origins to be 31.1 and 28.7%, respectively.15 On the other hand, the most common origin in our series was SVA followed by IPA. This agrees with American and German groups reporting that SVA origin as the most common at 35 and 27.5%, respectively.16,17

In our series, anastomoses connecting both hemi-prostates were detected in 18% of cases. The anastomosis was extensive enough to allow complete prostatic embolization through unilateral approach in two cases (0.9%), one of them was previously published as a case report.12 Extensive anastomosis is reported to be 3% in the literature.18 This is extremely useful in cases where prostate supply is identified/accessible on one side only. The true extent of anastomosis may be best detected using the PERFECTED technique described by Carnevale group.19

Protective coils/Gelfoam are extremely useful tools in selected cases, enabling safe embolization of the target arteries. Although coils are most commonly used, Gelfoam use has also been reported in the literature.20 If anastomosis is inaccessible, it is recommended to use intraprostatic vasoocclusion rates combined with very slow injection of particles.21 After such maneuvers, the anastomoses are usually no longer visualized; however, they become more evident when the peripheral resistance increases, after saturation of prostatic vascular bed by particles. Therefore, repeated slow angiograms during the embolization process are important for early detection of saturation, leading to reappearance of flow through the anastomoses into nontarget territory (-Fig. 8 highlights this concept). UK-ROPE study reported a much higher prevalence of extra-prostatic anastomosis, requiring protective coiling in 25.7%. We assume this was due to the power pump injector used during CBCT acquisition (instead of the gentle hand injections that we performed) that might have overestimated relevant anastomosis. Another possible explanation for higher sensitivity of anastomosis detection is their CBCT use. Although they reported mean FT shorter than ours (40.6 vs. 44 minute); complications, PD, skin dose, and percentage of bilateral PAE were all in favor of our series (-Table 2). Other studies reported protective embolization rates ranging from 4.3 to 26.2%.22,23

Scrutiny is essential when searching for possible additional prostatic feeders. In our series, we encountered multiplicity of supply (>2 PAs) in 5.7% of cases. Multiplicity was most encountered as double supply on the left side, which may be because this side is almost always embolized first, with the prostatic bed still well-vascularized, unsaturated with particles yet and with low resistance to blood flow. In the literature, multiplicity of prostatic supply has been described in up to 8.7% of cases.24 Such situation can be misleading due to “satisfaction of search,” leading to suboptimal embolization if one or more of these multiple feeders is missed, which may reduce clinical improvement and increase recurrence due to revascularization.24

Several PAE series used preoperative CTA for patient selection and procedure planning. Sensitivity of CTA for detection of extra-prostatic anastomosis was found to be 57.5% only.25 We did not use CTA because there is no enough evidence to support routine use for the extra IV contrast, radiation, and cost. In our 210 patients, only two cases (<1%) would have been excluded by CTA due to severe aortoiliac occlusive disease. This is not enough to justify the routine use of preoperative CTA for every case. There is insufficient data regarding the proportion of patients excluded from getting PAE based on preoperative CTA findings. This needs further investigation to determine its cost–benefit as a routine investigation for patient selection.

Several studies reported FT, PD and radiation higher than our series. -Table 2 summarizes these findings, in addition to...
to other technical and clinical parameters. Some studies reported FTs comparable to ours; however, PDs were significantly longer. This can be attributed to the arrangements required to perform intraoperative CBCT (e.g., repositioning the patient/table, preparing the injector, resuming the procedure with a new run).

In a prospective single-operator study specifically investigating radiation exposure during PAE, CBCT contributed to ~9% of the total dose-area product. FT from the same study was not significantly reduced to justify the extra-radiation exposure. Moreover, the study did not comment on postprocedure complications. 26 Performing CBCT requires arrangements that may prolong total PD. Another study specifically investigating the utility of CBCT reported PD longer than ours. Moreover, bilateral embolization was performed in 83% of cases only. They reported reasons for unsuccessful embolization were as follows: severe arteriosclerosis/stenosis (8 patients had unilateral PA stenosis and 1 patient had bilateral stenosis), nontarget anastomoses inaccessible by coil embolization (3 patients), vasospasms of PA (2 patients), and inability to identify a PA (3 patients). Interestingly, the study did not comment on postprocedure complications. 27 Several other studies that utilized CBCT did not comment on adverse events either (►Table 2). Bagla et al reported 16% hematospермia despite utilizing CBCT. 28 Throughout literature, routine use of preoperative CTA and intraoperative CBCT did not consistently reduce FT, PD, radiation dose or increase bilateral embolization rate. 22,29 Studies that did not use CTA/CBCT did not report significantly higher complication rates. In our series, there was a low complication rate, and we did not encounter any serious adverse events. There are scarce comparative data highlighting any statistically significant benefit of CBCT, in terms of the previously mentioned parameters. 30 This area needs more

Table 2 Preoperative CTA, intraoperative CBCT, FT, PD, DAP, skin dose, bilateral PAE, and serious complications. Studies are ordered ascendingly according to FT because it is the most consistently reported item. The definition of serious adverse events/serious complications was variable across studies. Numbers represent mean value unless specified otherwise.

<table>
<thead>
<tr>
<th>Study</th>
<th>CTA</th>
<th>CBCT</th>
<th>FT (min)</th>
<th>PD (min)</th>
<th>Dose area product (Gy.cm²)</th>
<th>Skin dose (mGy)</th>
<th>Bilateral PAE</th>
<th>Serious complications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pisco et al31</td>
<td>Yes</td>
<td>No</td>
<td>19.5</td>
<td>77</td>
<td>241.5</td>
<td>N/A</td>
<td>98.1% (618/630)</td>
<td>0.3% (2/630)</td>
</tr>
<tr>
<td>Garzón el al32</td>
<td>No</td>
<td>Yes (2/5)</td>
<td>29.1</td>
<td>N/A</td>
<td>523.9</td>
<td>2674.2</td>
<td>N/A</td>
<td>N/A</td>
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<td>Wang et al33</td>
<td>Yes</td>
<td>Yes</td>
<td>30</td>
<td>105</td>
<td>N/A</td>
<td>N/A</td>
<td>86.3% (101/117)</td>
<td>0% (0/117)</td>
</tr>
<tr>
<td>Bagla et al34</td>
<td>No</td>
<td>Yes</td>
<td>30.2</td>
<td>72</td>
<td>559.2</td>
<td>N/A</td>
<td>95% (18/19)</td>
<td>0% (0/19)</td>
</tr>
<tr>
<td>Andrade et al36</td>
<td>No</td>
<td>Yes (7/25)</td>
<td>30.9</td>
<td>N/A</td>
<td>450.7</td>
<td>2420.3</td>
<td>100% (25/25)</td>
<td>N/A</td>
</tr>
<tr>
<td>Schott et al27</td>
<td>No</td>
<td>Yes</td>
<td>30.9</td>
<td>89.4</td>
<td>134</td>
<td>N/A</td>
<td>83% (83/100)</td>
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<tr>
<td>du Pisanie et al16</td>
<td>Yes</td>
<td>No</td>
<td>31</td>
<td>72.9</td>
<td>59.2</td>
<td>N/A</td>
<td>98.5% (66/67)</td>
<td>N/A</td>
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<td>Gao et al9</td>
<td>No</td>
<td>No</td>
<td>33.2</td>
<td>89.7</td>
<td>113</td>
<td>N/A</td>
<td>94.7% (54/57)</td>
<td>1.7% (1/57)</td>
</tr>
<tr>
<td>Enderlein et al17</td>
<td>No</td>
<td>Yes</td>
<td>35</td>
<td>156 (median)</td>
<td>432.1 (median)</td>
<td>N/A</td>
<td>94.2% (98/104)</td>
<td>N/A</td>
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<tr>
<td>du Pisanie et al16</td>
<td>Yes (86/150)</td>
<td>Yes (64/150)</td>
<td>39.1</td>
<td>119.3</td>
<td>197.5</td>
<td>N/A</td>
<td>96.6% (145/150)</td>
<td>N/A</td>
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<td>Hacking et al25</td>
<td>Yes</td>
<td>Yes</td>
<td>40.6</td>
<td>145.2</td>
<td>221</td>
<td>2072.8</td>
<td>88% (165/187)</td>
<td>0.9% (2/216)</td>
</tr>
<tr>
<td>Our study</td>
<td>No</td>
<td>No</td>
<td>44</td>
<td>77</td>
<td>N/A</td>
<td>578</td>
<td>90% (189/210)</td>
<td>0% (0/210)</td>
</tr>
<tr>
<td>Carnevale et al24</td>
<td>No (MRI)</td>
<td>Yes</td>
<td>45.8–49.2</td>
<td>144.8–147.5</td>
<td>N/A</td>
<td>N/A</td>
<td>100%</td>
<td>0% (0/30)</td>
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<tr>
<td>Chiaradia et al35</td>
<td>No</td>
<td>Yes</td>
<td>47</td>
<td>N/A</td>
<td>454</td>
<td>N/A</td>
<td>66.7% (4/6)</td>
<td>0% (0/6)</td>
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<tr>
<td>Abt et al7</td>
<td>No</td>
<td>Yes</td>
<td>50.8</td>
<td>122.2</td>
<td>176.5</td>
<td>N/A</td>
<td>75% (36/48)</td>
<td>4.2% (2/48)</td>
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<tr>
<td>Amouyal et al19</td>
<td>No (MRI)</td>
<td>Yes (25/32)</td>
<td>54</td>
<td>139</td>
<td>364.7</td>
<td>3,065</td>
<td>97% (31/32)</td>
<td>0% (0/32)</td>
</tr>
<tr>
<td>de Assis et al3</td>
<td>No (MRI)</td>
<td>No</td>
<td>55.4</td>
<td>158</td>
<td>N/A</td>
<td>N/A</td>
<td>94.3% (33/35)</td>
<td>2.9% (1/35)</td>
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<td>Grosso et al36</td>
<td>Yes</td>
<td>No</td>
<td>69</td>
<td>165</td>
<td>N/A</td>
<td>N/A</td>
<td>69.2% (9/13)</td>
<td>0% (0/13)</td>
</tr>
</tbody>
</table>

Abbreviations: CBCT, cone-beam computed tomography; CTA, computed tomography; DAP, dose-area product; FT, fluoroscopy time; MRI, magnetic resonance imaging; N/A, not available; PAE, prostatic artery embolization; PD, procedure duration.
comparative data, preferably randomized controlled trials. Finally, a recent large American study found no significant difference between CTA and CBCT regarding FT, PD, radiation dose, or contrast volume.\textsuperscript{16} – Table 2 summarizes and compares data from literature regarding technical, radiation, and clinical outcomes (including our study).

Limitations of the Study

Our study has no intraoperative CBCT control group.

Conclusion

Detailed knowledge of PA anatomy is essential for treating BPH by PAE to enhance technical success, reduce complications due to nontarget embolization, shorten the procedure, and reduce radiation exposure. There is no enough evidence to support routine use of preoperative CTA and intraoperative CBCT as means of improving safety or efficacy. Randomized controlled studies are required in this regard.

Conflict of Interest

None.

References


