Original Article

Comparison of distal radial access versus standard transradial access in patients with smaller diameter radial Arteries(The distal radial versus transradial access in small transradial ArteriesStudy: D.A.T.A - S.T.A.R study)

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Aims: To evaluate safety and efficacy of distal right radial access (DRRA) compared to right radial access (RRA), for coronary procedures, in patients with smaller diameter radial arteries (SDRA) (radial artery diameter (RAD) <2.1 mm).

Methods and results: This is a retrospective analysis of safety and efficacy of DRRA Vs. RRA in patients undergoing coronary procedures at our cardiac catheterization laboratories over a 10-month period between September 2017 and June, 2018 (first 5 calendar months with RRA- first; next 5 calendar months with DRRA-first). All patients underwent pre-procedure ultrasound of arm arteries. All patients had RAD <2.1 mm (mean RAD 1.63 ± 0.27 mm; RAD ≥1.6 mm in 73.5%). Baseline characteristics were similar between groups. Primary end-point of puncture success was significantly lower in DRRA vs RRA group [79.5% vs 98.5%, p < 0.0001]. Puncture success was also lower in the subgroup of patients with RAD <1.6 mm Vs. ≥ 1.6 mm in the DRRA group (p < 0.0001). The secondary end-point of puncture time was significantly higher (2.1 ± 1.4 min vs. 1.0 ± 0.45 min, p < 0.00001) in the DRRA Vs. RRA group. The occurrence of vascular access site complications (including access site hematomas), radial artery occlusion (RAO) and distal RAO at day 1 and day 30 were similar between RRA and DRRA groups. Nonvascular access-site complication was seen only in the DRRA group.

Conclusion: DRRA is a safe and effective access for coronary procedures; though technically challenging in patients with SDRA (RAD<2.1 mm; mean RAD 1.63 ± 0.27 mm), with lower puncture success and higher puncture time compared to RRA.

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1. Introduction

Transradial access1-2 is associated with decreased mortality and reduced access site bleeding complications in high risk patient groups undergoing percutaneous coronary interventions and is recommended as a class 1 indication in acute coronary syndromes in the European Society of Cardiology guidelines.3 Distal radial artery access (DRRA) at the anatomical snuffbox has been reported4-5...
as a safe and feasible unique alternative to radial artery access at the wrist. The need for patent hemostasis with forearm radial access, to minimize risk of RAO, mandates close vigilance for bleeding to prevent forearm hematoma formation and enable its prompt effective management if it develops, to avert progression to a compartment syndrome. The latter risk may be less with DRRA as not only is the puncture site distal to the forearm, it may be more readily compressible.

Though right arm access is preferred by most operators because of its ergonomic advantage; there is a lack of data on distal right radial access compared to RRA in patients with small diameter radial arteries (SDRA). We aimed to retrospectively analyze procedural performance, outcomes, safety and efficacy of distal right radial access (DRRA) compared to right radial access (RRA), for coronary diagnostic and interventional procedures, in patients with SDRA (RAD < 2.1 mm).

2. Methods

2.1. Study Design

Fig. 1 This is a retrospective analysis of safety and efficacy of DRRA Vs. RRA in patients undergoing coronary procedures at our cardiac catheterization laboratories over a 10-month period between September 2017 and June, 2018. All patients underwent preprocedure ultrasound of arm arteries. All patients had RAD < 2.1 mm. Consecutive patients underwent procedures using RRA first in all cases from September, 2017 to January, 2018 (5 calendar months). Crossover to alternate access due to failure of RRA and use of access other than RRA qualified for exclusion. From February, 2018 till June, 2018 (5 calendar months), DRRA was used first in all cases. Patients who failed DRRA underwent RRA. Crossover to alternate access due to failure of RRA qualified for exclusion. Patients who underwent procedures by ulnar, femoral, brachial or left upper limb access were not included. Baseline characteristics were similar between groups. One operator experienced in transradial (approximately > 15,000 transradial & > 500 prior distal radial procedures) performed all the interventions and another experienced in color Doppler ultrasound and echocardiography performed and read all the color Doppler ultrasounds.

2.2. Exclusion

1 From September, 2017 to January, 2018 (5 calendar months): Any access other than RRA; including crossover to alternate access were excluded.
2 From February 2018 till June, 2018 (5 calendar months): In case of failed DRRA; RRA was attempted; subsequent failure of which led to cross-over to an alternate access site; selection of which was based on the operator’s judgment (left radial, left or right ulnar, right femoral, or high right radial) and subsequent exclusion from the study. Similarly, in case of puncture failure in the RRA group, access-site was crossed-over and the patients were excluded from the study. Patients who underwent procedures by ulnar, femoral, brachial or left upper limb access were not included.

Keeping in view our objective to assess the safety and efficacy of DRRA compared to standard RRA, in this population with SDRA, not only were those with poorly palpable DRA or RA not excluded; the same puncture and hemostasis techniques were used for both
groups of patients and DRA measurement was not used to pre-select cases because no data exists for its cut-off value, below which DRRA is not recommended. Palpable pulse intensity was graded on a scale of 0–3; 0 indicating no palpable pulse, 1+ indicating faint pulse, 2+ indicating a normal pulse, and +3 indicating a bounding pulse.

Each participant gave written informed consent and the study was approved by the institutional ethics committee.

2.3. Study procedure

2.3.1. Color Doppler Ultrasound Imaging

Each patient underwent an ultrasound examination of bilateral arm arteries (to assess radial artery diameter and brachial artery bifurcation/radial artery anomaly) prior to the cardiac catheterization procedure and at 1 & 30 day follow-up for RAO and distal RAO. The operator was blinded to the results of the forearm ultrasound. Color and pulsed Doppler assessment were undertaken for DRA occlusion in the anatomical snuffbox. Measurement of DRA diameter using ultrasound has been reported to be feasible only in DRAs with >2 mm diameter having straight course.

Further, it is still unclear whether DRA diameter is smaller than or equal to ipsilateral RA, reflecting inconsistency in reproducibility of ultrasonographically assessed DRA diameter. Further, its role in pre-selection of cases for distal radial access is not established, just as with the utility of ultrasound-guided puncture: neither of which is mandatory for DRRA. Hence, neither was utilized.

2.3.2. Distal radial and transradial catheterization procedure

(i) Cardiac Catheterization Procedure: 0.5–1 mL of 2% lidocaine was used as local anesthetic. This dose was adequate for DRRA and RRA. Though there is no data on optimal dose of lidocaine, larger doses made palpation and hence puncture of the DRA or RRA difficult. The same access technique was used in both groups. The superiority of sedliger vs. transfixon using metallic needle vs. sheathed needle with anterior vs. posterior puncture in DRA is not established. All patients received 0.5 mL–1mL Inj Midazolam and 25–50mcg of Inj Fentanyl administered in small aliquots slowly intravaneously for sedation and analgesia; following which the access artery was punctured without ultrasound guidance, using an over the needle cannula system (Jelco TM venous cannula (20-gauge venipuncture catheter needle)) or Terumo sheathed needle, with transfixation or double wall puncture technique by intention, for both groups; as is our usual practice for RA access, even with SDRA (supplementary video-1). Anterior puncture, if it occurred incidentally, was accepted. Puncture of DRA was attempted along the course of DRA downwards from trapezium till the floor of the anatomical snuffbox (Fig. 2A–D) to avoid scraping the periosteum of carpal.

TerumoTM radial sheath was used (5 F or 6 F; per operator discretion) for coronary angiography (CAG) and angioplasty (PCI). Following cannulation, diltiazem (5 mg) and unfractionated heparin (5000 IU) were administered intra-arterially via the radial sheath for diagnostic catheterization (weight–adjusted 70 IU/kg heparin was given for those <50 or >80 kg); for angioplasty, additional dose of 1000–2000 IU of Heparin was given to maintain activated clotting time (ACT) at 250–300sec. Additional diltiazem (5 mg) was administered intra-arterially, with each catheter exchange and prior to sheath removal in all DRA and RA cases. TIG (Terumo TM) was the default catheter for CAG. Guide catheter selection for PCI was as per operator’s discretion. Intra-arterial injection nitroglycerin (50mcg) administration, into access artery was optional at the discretion of the operator.

(ii) Hemostasis: Though various techniques, from compression device to bandage to...

Even manual compression have been used for hemostasis by different operators, the superiority of any one technique over the other is not established. Gauze ball/bandage roll (2.5 cm × 5 cm) and elastic bandage (5 cm × 15 cm) without patent hemostasis was used in both groups (Fig. 3A–F) per our routine practice with RA access, on account of its overall simplicity and to maintain homogeneity between groups especially since the feasibility and relevance of patent hemostasis is not established in DRRA.

The bandage was loosened 1 h post-procedure followed by palpation of hand and forearm proximal to bandage every 3 min for 1 h for early detection of hematoma. Per our usual practice, patients were discharged 3 h after a coronary angiogram; with advice to remove the bandage 12 h later.

Patient demographics including left ventricular ejection fraction and hemodynamic instability; pre-procedural characteristics, palpability of DRA and RA (poorly- or well-palpable), ultrasound data (radial diameter and anomalies), patient anxiety (on a scale of...
3: mild to severe), procedural characteristics, ≥ grade 2/3 pain during puncture (scale 1–3), anterior vs posterior punctures, peri-procedural radial artery spasm (≥ grade 3/4 of spasm grading by Chugh et al); and study end-points were recorded (Table I-II).

Radial artery occlusion (RAO) and distal RAO were also recorded on days 1 and 30 post-procedure, clinically by palpating the radial artery & distal radial artery, with simultaneous compression of ipsilateral ulnar artery & by color Doppler ultrasound. All clinical assessments were performed by an experienced reader.

2.4. Study end-points

2.4.1. Primary end-points

Puncture Success was defined as successful insertion of the radial sheath. Reason for puncture failure was recorded (inability to obtain bleed back in the needle hub; inability to insert length of sheath wire into the DRA after achieving bleed back into the needle hub; inability to insert sheath successfully into DRA).

2.4.2. Secondary end-points

Puncture Time (Intra-procedural) was defined as the time taken from first contact of puncture needle with skin to successful cannulation of the access artery with uninterrupted smooth passage of length of sheath wire. Puncture was discontinued per operator discretion or after 5 min in primary angioplasty cases. Reason for prolonged puncture time was noted (delay in obtaining bleed back in the needle hub from puncture of the DRA; delay in cannulating the DRA with wire and/or sheath).

Radial Artery Occlusion (RAO) on Color Doppler Ultrasound (Day 1 & 30) Defined as slow flow or no flow on 2D color Doppler, or low velocity signal on pulsed doppler or monophasic flow in a previously blocked radial suggesting collateral flow.

Distal Radial Artery Occlusion on Color Doppler Ultrasound (Day 1 & 30) Defined as flow reversal on color Doppler ultrasound in the radial artery in anatomical snuffbox.

Vascular Access Site Complications (VASC): Included forearm hematomas (≥ 5 cm) and hematomas localized to hand (due to bleeding and extravasation); compartment syndrome, pseudo-aneurysm formation, dissection and AV fistulas. To prevent progression to compartment syndrome, any patient whose forearm hematoma was ≥ 5 cm, was closely monitored, together with compression all along the radial artery in the forearm using gauze and bandage; in addition inflation of a sphygmomanometer cuff on the arm to above systolic blood pressure (with intermittent deflations). For ‘hand’ hematomas, additional gauze and bandages were applied on palmar and dorsal aspect of the hand, at the DRA site. A plethysmograph probe on the index finger or thumb of the affected hand was used to monitor for hand-ischemia.

Non-Vascular Access Site Complications: any access-site related permanent or temporary, motor or sensory nerve injury including localized transient numbness and paresthesia at 1 and 6 weeks corroborated by a neurological evaluation.
Composite Secondary End Point: including hematoma, non vascular access site complications, RAO, and DRAO at day 1.

2.5. Statistical analysis

Descriptive statistics were used for initial analysis. Continuous data are presented as mean with SD. Dichotomous data and categorical data was presented as percentages. Difference between groups of continuous data were examined using the Fischer exact or Student t test and the chi-square test was used to compare dichotomous and categorical data. Statistical significance was assumed at a value of \( p < 0.05 \). All statistical analyses were performed with STATA for Windows. The sample size calculation was performed assuming primary endpoint of puncture success as 98.7%7 in RRA group and 88.6% in the DRRA group4. Overall, 120 patients per group (total \( = 240 \)) were deemed adequate to achieve 90% power considering an alpha error of 0.05. We identified potential variables that may influence our primary end-point with a univariate logistic regression model. A multi-variate logistic regression model with all clinically relevant variables was then established to estimate odds ratios (ORs) with 95% confidence bounds.

3. Results

After exclusions, as per Figure 1; 545 patients were included in the study. Pre-procedural (Table 1) and procedural characteristics (Table II) were similar between the groups. Pain \([> grade 2]/3\) was not reported during DRRA. Finally, 328 people underwent successful RRA and 209 underwent successful DRRA (Fig. 1). In three cases, DRRA was used for coronary angiogram; following which the sheath was removed and gauze/bandage applied. The patients underwent coronary angioplasty a few hours later through the RRA with gauze/bandage still on at the DRRA site.

| Table 1 | Patient Demographics and Pre-procedural ultrasound characteristics. |
|---------|--------------------------------------------------------|
|         | Clinical Characteristics | Right Radial Group (n = 282) | Distal Right Radial Group (n = 263) | p Value |
|         | Age, y | 53.8 ± 12.9 | 55.1 ± 11.9 | 0.21 |
|         | Females/Male | 84/198 (29.7%/70.3%) | 79/184 (30%/70%) | 0.95 |
|         | Weight (kg) | 65.6 ± 9.7 | 64.0 ± 9.0 | 0.94 |
|         | Height (cm) | 169.3 ± 7.0 | 169.4 ± 7.5 | 0.59 |
|         | BMI | 22.8 ± 3.6 | 22.3 ± 3.1 | 0.66 |
|         | Tobacco Smoker/Cheewer | 200 (70.9%) | 156 (59.3%) | 0.004 |
|         | COPD | 52 (18.4%) | 36 (13.6%) | 0.132 |
|         | Hypertension | 34 (12%) | 32 (12.1%) | 0.968 |
|         | Diabetes Mellitus | 37 (13.1%) | 37 (14%) | 0.747 |
|         | Heart Failure with EF < 40% | 44 (15.6%) | 49 (18.6%) | 0.35 |
|         | Hemodynamic Instability | 45 (15.9%) | 61 (23.1%) | 0.06 |
|         | Previous procedure using same access | 70 (24.8%) | 61 (23.1%) | 0.56 |
|         | Forearm Ultrasound | RA size (mm) (mean ± SD) | 1.59 ± 0.27 | 1.62 ± 0.26 | 0.948 |
|         | | RA size, Males | 1.63 ± 0.27 | 1.66 ± 0.27 | 0.26 |
|         | | RA size, Females | 1.46 ± 0.22 | 1.51 ± 0.21 | 0.17 |
|         | Vascular Anomalies | Intimal Thickness | 0 (0%) | 2 (0.8%) | 0.18 |
|         | | Parallel Radial Artery | 6 (2.1%) | 13 (4.9%) | 0.97 |
|         | | Radial Artery Loop | 1 (0.4%) | 1 (0.4%) | 0.97 |
|         | | Accessory Radial Artery | 1 (0.4%) | 0 (0%) | 0.35 |
|         | | Occluded Artery | 1 (0.4%) | 3 (1.1%) | 0.15 |

| Table 2 | Procedural characteristics and Secondary end-points in RRA vs. DRRA groups. |
|---------|----------------------------------------------------------|
|         | Procedural Characteristics | Right Radial (RRA) Group (n = 328) | Distal Right Radial (DRRA) Group (n = 209) | p Value |
|         | Sheath Size | 5 F | 156 | 107 | 0.41 |
|         | | 6 F | 172 | 102 | 0.95 |
|         | Procedure Type | Coronary Angiogram | 236 | 152 | 0.01 |
|         | | PCI (Single vessel) | 79 | 45 | 0.02 |
|         | | PCI (Two vessel) | 2 | 1 | 0.05 |
|         | | Primary PCI | 11 | 11 | 0.08 |
|         | | Radial Artery Spasm | 3 (0.9%) | 3 (1.4%) | 0.31 |
|         | Puncture Time (min) | 1.17 ± 0.8 | 2.08 ± 0.9 | 0.00001 |
|         | Vascular Access Site Complications | 2 (0.6%) | 4 (1.9%) | 0.16 |
|         | Non Vascular Access Site Complications | 0 (0%) | 3 (1.4%) | 0.031 |
|         | Radial Artery Occlusion | Day 1 | 8 (2.4%) | 5 (2.4%) | 0.97 |
|         | | Day 30 | 7 (2.1%) | 6 (2.9%) | 0.58 |
|         | Distal Radial Artery Occlusion | Day 1 | 8 (2.4%) | 5 (2.4%) | 0.93 |
|         | | Day 30 | 5 (1.5%) | 6 (2.8%) | 0.28 |

*PCI: Percutaneous Coronary Intervention.
3.1. Primary endpoint

Of the 263 patients in DRRA group, 209 had puncture success, and 54 had puncture failure. Puncture success was significantly lower in the DRRA compared to RRA group (79.5% vs 98.5%, \( p < 0.0001 \)). After excluding the patients with poorly palpable DRA \((n = 31)\), puncture success was 88.7% vs 98.5% for DRRA Vs. RRA, \( p < 0.0001 \). The reasons for puncture failure were: no bleed back in the needle hub \((n = 13)\); inability to insert sheath wire into DRA \((n = 36)\); and inability to insert sheath into DRA in \((n = 5)\) patients. All patients who had puncture success had procedure success; though in 8 cases, the procedure was completed using alternative access and these patients were therefore excluded from the study \((\text{Fig. 1})\).

We performed multivariate logistic regression analysis using the following clinical variables: age, sex, BMI, hemodynamic status \((\text{stable vs. unstable})\), radial artery diameter \(<1.6 \text{ vs. } \geq 1.6 \text{ mm})\,13\) patient anxiety. Although RAD was unrelated to puncture success or time, there were more puncture failures in patients with RAD \(<1.6 \text{ mm vs. } \geq 1.6 \text{ mm})\,14\,14\) in the both groups. \((\text{Table III})\).

3.2. Secondary endpoints

3.2.1. Puncture time

Secondary end-point of puncture time was significantly longer in patients with DRRA compared with RRA \((2.1 \pm 1.4 \text{ min vs. } 1.0 \pm 0.45 \text{ min, } p < 0.00001)\). Prolonged puncture time occurred because of delay in obtaining bleed back in the needle from puncture of the distal radial artery in 37%; in 63% it occurred from delay in cannulating the distal radial artery with wire and/or sheath.

3.2.2. Radial Artery Occlusion

Rates of RAO at Day 1 and 30 \((2.4\% \text{ vs. } 2.4\%, p = 0.97 \text{ and } 2.1\% \text{ vs. } 2.9\%, p = 0.58 \text{ respectively})\) were similar in both groups \((\text{DRRA vs. RRA})\).

3.2.3. Distal radial artery occlusion

Distal RAO on Day 1 & 30 were 2.4\% vs 2.5\% \((p = 0.93)\) and 2.8\% vs 1.5\% \((p = 0.28) \text{ respectively} \((\text{DRRA vs RRA})\).

3.2.4. Access-site complications

Vascular access-site complications were similar in both DRRA \((\text{Fig. 4A-B})\) and RRA groups \((1.9\% \text{ vs. } 0.35\%, p = 0.12)\). Forearm hematoma occurred only with RRA and hematomas in the DRRA group were localized only to the hand and were managed successfully.13 Non-vascular complications were limited to localized transient paresthesia, with mild sensory impairment \((\text{at 1 week, resolved in all patients by } 6 \text{-week follow-up})\) over the dorsum of the hand in the region of anatomical snuffbox: seen in 3/209 DRRA cases \((1.4\%) \text{ vs. } 0\% \text{ in RRA group} \((p = 0.031)\).

Subgroup of patients with radial artery diameters \(<1.6 \text{ mm and } \geq 1.6 \text{ mm and their outcomes} \((\text{Table III})\). Majority \((73.5\%)\) patients had RAD \(<1.6 \text{ mm}. The distribution of patients with RAD \(<1.6 \text{ mm was } 239/328 \text{ and } 148/209 \text{ patients in the RRA Vs. DRRA groups respectively} \((\text{Table III})\).

3.3. DRRA group

Numerically higher rates of RAO and DRAO at days 1 and 30 as well as higher puncture failures were seen in patients with RAD \(<1.6 \text{ mm} \text{ V patients with RAD } \geq 1.6 \text{ mm} \,(p < 0.0001); however puncture times were similar in both groups. Further, higher rates of a composite of hematomas, non-vascular access-site complications, RAO & DRAO at Day 1 were seen \((p < 0.001)\) in the subgroup with RAD \(<1.6 \text{ mm.}

3.4. RRA group

Patients with radial artery diameters \(<1.6 \text{ mm})\) had higher puncture times \((p < 0.05)\) and lower puncture success. Vascular access site complications, RAO and DRAO on ultrasound at Day 1 and Day 30, were also higher in these patients though not statistically significant. However, higher rates of a composite of hematomas, non-vascular access site complications, RAO & DRAO at Day 1 were seen \((p < 0.001)\).

4. Discussion

The use of DRA for coronary procedures, has been shown to be feasible15 in small observational studies. To the best of our knowledge, unlike most current studies on distal left radial access,5,6,7 this is the first study comparing puncture success, puncture time and vascular and non-vascular access site complications of DRRA with RRA in patients with SDRA.

Unlike previous studies4,5,9,10 in which poorly-palpable DRA, reported in up to 41\% patients, were excluded; our study included all-comers without pre-selection, including those with a weak and poorly-palpable DRA pulse, with the aim to assess DRA as a safe and effective access compared to RRA. The puncture success was therefore lower \((79.5\%)\) and puncture time was prolonged in our study. Also, after excluding only those with poorly-palpable DRA, our puncture success rate in DRRA group was 88.7\%; with a failure rate of 11.3\% compared to 10\% reported in other studies which carefully selected study patients and excluded those unsuitable for DRRA.5,9,10 This failure rate in DRRA reported across studies is attributable to the unique anatomical challenges of DRA including (i) angulation and tortuosities in the DRA in its course over the trapezius to the floor of anatomical snuffbox, and (ii) presence of branches of the deep palmar arch, of which the DRA too is a branch. However, since the RA in the forearm does not have any of these unique challenges; puncture failures and puncture time were considerably less in the RRA group.

Further, though debatable, DRA diameter is generally believed to be smaller3,8,10 than the RA. This may help explain the higher puncture failure of DRA Vs RRA when used as default access. Again, because the mean RAD \((1.63 \pm 0.27 \text{ mm})\) in our study population was smaller than the reported mean RAD \((>2.1 \text{ mm})\) in other studies3,9,10; puncturing a proportionately smaller DRA was even more challenging with more puncture failure and longer puncture time. Not only does this explain an even higher puncture failure rate in the DRRA group in our study compared to that in other studies3,9,10; it also explains a higher DRRA puncture success rate after exclusion of patients with poorly-palpable DRA from the DRRA group. Again, although the higher puncture failure in DRRA group may be because ultrasound-guided puncture was not used, there is no data to support this and even data on ultrasound-guided RA puncture has limitations.15 Further, because a majority of our study patients had a radial artery diameter of \(<1.6 \text{ mm, such higher failure rate with DRRA is not unexpected. South East Asians are known to have smaller radial arteries, increasing their risk of procedural complications and puncture failure compared to their Caucasians.15 Also, prior adequate experience of the operator with coronary procedures via RRA and DRRA rules out the possibility of learning curve being responsible for higher puncture failures in DRRA group in the study. A trend for a higher hematoma rate, localized to the hand only was observed in the DRRA group \((\text{Fig. 3}), and is attributed to a greater number of failed puncture attempts over the DRA, with resultant trauma to the artery and its small branches in the snuff box. None of these patients had remote forearm bleed from puncture of a perforator or branch by wire in the study. Though the higher incidence of 'hand' hematoma may have been because of use
Fig. 4. (A,B) (left to right) Complication of distal right radial access: Hand hematoma following DRRA, localized to hand (A) Dorsal (B) palmar aspect of hand.
of gauze and bandage for DRRA hemostasis; there is no published data to support this possibility. On the contrary, because of gauze and bandage use in both DRRA and RRA, there was homogeneity in hemostasis technique allowing for meaningful comparison between groups. Forearm hematomas occurred only in the RRA group and with our prompt, aggressive management\(^1\) of forearm hematomas \(>5 \text{ cm}\); none of our patients developed compartment syndrome.\(^1\)

Transient numbness over the region of the snuffbox (corroborated on neurological assessment), only reported in the DRRA group, was likely due to inadvertent trauma of sensory branches of the superficial radial nerve in the proximity during puncture. All patients that developed this complication had a radial artery diameter \(<1.6 \text{ mm}\), which may have contributed to increased puncture attempts and in-turn neurological injury to the snuff box.

Though not powered for small differences in RAO; RAO was similar (\(p = NS\)) in both groups with similar baseline characteristics (Tables. I-II), with use of same technique for puncture and hemostasis, with same sheath length, as well as artery/sheath ratio based on usage of comparable 5 F and 6 F sheaths; comparable radial artery spasm, and other procedural characteristics.

Contrary to the lower RAO reported with DRA in earlier studies which were also not powered for conclusion,\(^ {25} \) our study found similar rates of RAO and DRAO between groups, and the reasons were: (i) direct (puncture site) and indirect (presence of \( > 1 \text{ mm} \) flow between sheath and radial artery related to RA: sheath mismatch\(^ {24} \) especially in SDRA; all substrates for thrombosis and precursors of RAO;\(^ {25} \). Though it is unclear whether RAO would have been different between groups, had patent hemostasis been used; it seems extremely unlikely, because the RAO rate in our study with the gauze-ball and bandage technique, but without patent hemostasis was comparable to that reported with patent hemostasis in PROPHET II.\(^ {25} \)

Unlike in the PROPHET trial\(^ {5,18} \) which used only plethysmography to assess radial artery patency; in our study, RAO (and DRAO) were diagnosed with greater accuracy using doppler ultrasound.\(^ {18} \)

Patent hemostasis has been shown to reduce the rate of RAO\(^ {5} \) while larger sheaths have been found to be associated with higher RAO\(^ {15} \); yet, with use of bigger sheaths (6 F Vs 5 F, in 52.4% Vs, 47.6%) and without patent hemostasis, RAO was 2.9% in the standard radial group in our study, compared to 3% RAO reported in PROPHET II study\(^ {25} \) which used patent hemostasis, smaller (5 F) sheaths in 100% and did not include any PCI cases in the study. One definite advantage of using DRRA as default is that the RA in forearm may be available for repeat procedures in situations where DRA is occluded or inaccessible because of hematoma or tender-ness from a recent prior procedure.\(^ {25} \) Our study has some important limitations; (i) Like many other initial publications on the subject,\(^ {4,5,10} \) this too was a non-randomized, single center, single operator retrospective analysis (ii) The study was not powered to detect a small difference in RAO between groups (iii) As per our usual practice, patent hemostasis was not used; though this helped meaningful comparison by maintaining homogeneity between groups (v) The findings may not be applicable to other ethnic groups with larger RA diameters (vi) Ultrasound guided puncture was not used (vii) DRA diameter was not measured.

### 5. Conclusion

DRRA is a safe and effective access for coronary procedures; though technically challenging in smaller diameter radial arteries, with lower puncture success and higher puncture time compared to RRA. The rates of vascular complications, RAO and distal RAO are similar between RRA and DRRA groups. Larger randomized trials are needed to further evaluate advantages of DRRA over RRA.

### Conflict of interests

None to report for all co-authors.

### Funding disclosures

No disclosure for all co-authors.

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**Table III**

Outcomes in patients with radial artery diameters \(<1.6 \text{ mm}\) vs \(\geq 1.6 \text{ mm}\).

|                               | Radial Artery Diameter \(<1.6 \text{ mm}\) | Radial Artery Diameter \(\geq 1.6 \text{ mm}\) | \(p\) Value |
|-------------------------------|------------------------------------------|-----------------------------------------------|------------|
| **DRRA Group**               |                                          |                                               |            |
| **Puncture Success (%)**      | (\(n = 148\)) 77.5%                     | (\(n = 61\)) 89.8%                           | 0.0001     |
| **Puncture Time (min)**       | 2.02 + 1.43                             | 2.19 + 1.43                                 | 0.48       |
| **Vascular Access Site Complications** | 4 (1.7%)                              | 0 (0%)                                       | 0.21       |
| **Non Vascular Access Site Complications** | 3 (1.4%)                              | 0 (0%)                                       | 0.21       |
| **Radial Artery Occlusion**   |                                          |                                               |            |
| Day 1                         | 4 (2.7%)                                 | 1 (1.6%)                                     | 0.34       |
| Day 30                        | 6 (4.1%)                                 | 0 (0%)                                       | 0.11       |
| **Distal Radial Artery Occlusion** |                                    |                                               |            |
| Day 1                         | 5 (2.1%)                                 | 0 (0%)                                       | 0.16       |
| Day 30                        | 5 (2.1%)                                 | 1 (1.6%)                                     | 0.71       |
| **Composite Secondary Endpoint * **|                                  |                                               | 0.001      |
| **RRA Group**                | (\(n = 239\)) 16 (10.8%)                | (\(n = 89\)) 1 (1.6%)                        |            |
| **Puncture Success (%)**      | 96%                                      | 100%                                         | 0.05       |
| **Puncture Time (min)**       | 1.20 + 0.82                              | 0.76 + 0.34                                 | 0.05       |
| **Vascular Access Site Complications** | 1 (0.7%)                              | 1 (1.6%)                                     | 0.51       |
| **Non Vascular Access Site Complications** | 0 (0%)                                | 0 (0%)                                       | 0.42       |
| **Radial Artery Occlusion**   |                                          |                                               |            |
| Day 1                         | 7 (2.9%)                                 | 1 (1.1%)                                     | 0.64       |
| Day 30                        | 6 (2.5%)                                 | 1 (1.1%)                                     | 0.44       |
| **Distal Radial Artery Occlusion** |                                    |                                               |            |
| Day 1                         | 5 (2.1%)                                 | 1 (1.1%)                                     | 0.56       |
| Day 30                        | 4 (1.7%)                                 | 1 (1.1%)                                     | 0.49       |
| **Composite Secondary Endpoint * **|                                  |                                               | 0.001      |

\* Composite of access site hematoma, non-vascular access site complications, radial artery occlusion day 1, and distal radial artery occlusion day 1.
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