



# One-Stage vs. Two-Stage Brachio-Basilic Arteriovenous Fistula for Dialysis Access: A Systematic Review and a Meta-Analysis

Khalid Bashar<sup>1</sup>\*, Donagh A. Healy<sup>1</sup>, Sawsan Elsheikh<sup>2</sup>, Leonard D. Browne<sup>3</sup>, Michael T. Walsh<sup>3</sup>, Mary Clarke-Moloney<sup>1</sup>, Paul E. Burke<sup>1</sup>, Eamon G. Kavanagh<sup>1</sup>, Stewart R. Walsh<sup>4</sup>

- 1 Department of Vascular Surgery, University Hospital Limerick, Limerick, Ireland, 2 Department of Acute Medicine, James Connolly Memorial Hospital, Dublin, Ireland, 3 Centre for Applied Biomedical Engineering Research (CABER), Department of Mechanical, Aeronautical & Biomedical Engineering, Materials and Surface Science Institute, University of Limerick, Limerick, Ireland, 4 Department of Surgery, National University of Ireland, Galway, Ireland
- \* khalid@live.ie



# GOPEN ACCESS

Citation: Bashar K, Healy DA, Elsheikh S, Browne LD, Walsh MT, Clarke-Moloney M, et al. (2015) One-Stage vs. Two-Stage Brachio-Basilic Arteriovenous Fistula for Dialysis Access: A Systematic Review and a Meta-Analysis. PLoS ONE 10(3): e0120154. doi:10.1371/journal.pone.0120154

**Academic Editor:** Daniel Schneditz, Medical University of Graz, AUSTRIA

Received: September 9, 2014 Accepted: January 20, 2015 Published: March 9, 2015

Copyright: © 2015 Bashar et al. This is an open access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

**Data Availability Statement:** All relevant data are within the paper and its Supporting Information files.

**Funding:** The authors received no specific funding for this work.

**Competing Interests:** The authors have declared that no competing interests exist.

# **Abstract**

#### Introduction

A brachiobasilic arteriovenous fistula (BB-AVF) can provide access for haemodialysis in patients who are not eligible for a more superficial fistula. However, it is unclear whether one or two-stage BB-AVF is the best option for patients.

#### Aim

To systematically assess the difference between both procedures in terms of access maturation, patency and postoperative complications.

#### Methods

Online search for randomised controlled trials (RCTs) and observational studies that compared the one-stage versus the two-stage technique for creating a BB-AVF.

#### Results

Eight studies were included (849 patients with 859 fistulas), 366 created using a one-stage technique, while 493 in a two-stage approach. There was no statistically significant difference between the two groups in the rate of successful maturation (Pooled risk ratio = 0.95 [0.82, 1.11], P = 0.53). Similarly, the incidence of postoperative haematoma (Pooled risk ratio = 0.73 [0.34, 1.58], P = 0.43), wound infection (Pooled risk ratio = 0.77 [0.35, 1.68], P = 0.51) and steal syndrome (Pooled risk ratio = 0.65 [0.27, 1.53], P = 0.32) were statistically comparable.

#### Conclusion

Although more studies seem to favour the two-stage BVT approach, evidence in the literature is not sufficient to draw a final conclusion as the difference between the one-stage and



the two-stage approaches for creation of a BB-AVF is not statistically significant in terms of the overall maturation rate and postoperative complications. Patency rates (primary, assisted primary and secondary) were comparable in the majority of studies. Large randomised properly conducted trials with superior methodology and adequate sub-group analysis are needed before making a final recommendation.

#### Introduction

The superiority of haemodialysis (HD) access created by means of an Arteriovenous Fistula (AVF) in patients with end stage renal disease (ESRD) has been shown before. Stenosis and thrombosis is less likely to occur in a well-functioning mature AVF when compared to arteriovenous grafts (AVG) and central venous catheters (CVC), resulting in prolonged patency rates for AVFs as has been described previously [1]. Also, AVFs carry a lower risk for infection [2,3]. However, around 20%- 50% of all fistulas fail to mature into a useful HD access [4–7].

The preferred location for placing an AVF for the first time is distally at the radius, thus making it possible to place a second fistula proximally if the first one failed to mature. The order of preference for creating an AVF [8-10]:

- 1. Distal Radio-Cephalic
- 2. Proximal Radio-Cephalic
- 3. Brachio-Cephalic
- 4. Brachio-Basilic (transposed Basilic vein)

This order is in agreement with the National Kidney Foundation Kidney Disease Outcomes Quality Initiative (NKF KDOQI) guidelines [11]. However, fistulas created distally at the wrist are less likely to mature compared to proximal AVF, at the same time proximal AVF require less intervention and are likely to last longer [12]. The decision of where to create the AVF can be helped by preoperative vascular mapping using ultrasound imaging which is expected to improve chances of creating an AVF that will likely mature into a useful dialysis access [13,14]. Placement of a primary forearm fistula is feasible in 40% to 50%, with an upper arm fistula possible in an additional 25% to 35% of patients [15]. An AVF prevalence of  $\geq$  65% has been recommended in the KDOQI guidelines for patients undergoing HD [11], this prevalence is currently higher in Europe (67%- 91%) compared to the US (24%- 47%) [15–18]; however, the prevalence of AVFs in the US varies significantly among different dialysis units [15,19].

Dagher was the first to describe the use of basilic vein to create an AVF in the upper arm between the end of basilic vein and the side of the brachial artery to act as access for long term haemodialysis [20]. Since then, the procedure has seen several changes and modifications. Superficialisation of a brachiobasilic fistula to make it more susceptive to cannulation can be achieved either by an elevation technique without mobilisation to bring the vein superficial to the surgically reconstructed deep fascia and subcutaneous tissue in the anatomic location of the basilic vein [21], or by a transposition technique by mobilising the entire length of the basilic vein to position the vein anterolaterally through a subcutaneous flap [22].

Some of the debate surrounding brachiobasilic arteriovenous fistulas (BB-AVF) has been focused on the decision to choose between one-stage vs the two-stage techniques. The one-stage procedure aims to create a fistula between the basilic vein and the brachial artery in the upper arm in one procedure. This would require a long incision to gain access and mobilise the basilic



vein making sure the anastomosis is not placed under tension and no obvious stenosis is present proximally. The main advantage of this technique is the shorter waiting time required to cannulate the fistula. Also the one-stage will prevent the patient from having to undergo another procedure and is more cost effective as hospital resources will be used only once. One of the main disadvantages of this technique is the long incision which will require a longer time to heal and also carries a higher risk for wound-related complications. Also the procedure takes longer and is more demanding [23–25]. Moreover, in a study by Anaya-Ayala et al assessing the anatomy of basilic vein found that only 66% of patients are expected to have a "normal" basilic vein entering one of two paired brachial veins close to the axilla, while up to 34% will have an "abnormal" variant that would negatively influence the newly created fistula maturation [26].

The two-stage procedure allows the basilic vein to become arterialised and as such, more resistant to torque and will become easier to mobilise in the second procedure as it gets transformed into a bigger and stronger structure. The hope is that operative difficulty and complications would be reduced with improved patency rates [27].

This review was designed to systematically assess the difference between both procedures in terms of access maturation and survival, as well as complications and interventions required to maintain patency for haemodialysis.

#### Methods

This systematic review and meta-analysis were conducted according to the Preferred Reporting Items for Systematic Review and Meta-Analysis (PRISMA) guidelines [28]. No published protocol exists for this review.

# Eligibility criteria

We searched for randomised controlled trials (RCTs) and observational studies that compared the one-stage technique with the two-stage technique for creating a brachiobasilic arteriovenous fistula (BBAVF) for haemodialysis access. Case series and review articles were excluded from this review.

# Search strategy

A search of the literature for relevant studies was conducted in August 2014 using the following terms: (["Basilic Vein" OR "Basilic"] AND ["Fistula" OR "Arteriovenous" OR "Access"] AND "dialysis"). We searched the online databases of: Medline, CINAHL, EMBASE, the Cochrane library and Google Scholar. We did not restrict our search by publication date or status, however, we only included studies published in English language and those conducted on humans. We also searched the bibliographies of included trials for additional studies. A summary of the study selection process can be found in the PRISMA flow diagram below [Fig. 1]. Studies were not restricted based on the duration of follow-up.

The main outcome measures for this review were successful maturation and development of postoperative complications, namely wound haematoma, wound infection and steal syndrome. Secondary outcomes were primary and secondary patency rates. Definitions for "maturation", "primary patency" and "secondary patency" were those specified in individual studies.

#### Data collection

KB and DH independently extracted the data from included studies on a Microsoft Excel spreadsheet. Any differences in recording the outcomes of interest were discussed between two



### Fig 1: PRISMA 2009 Flow Diagram

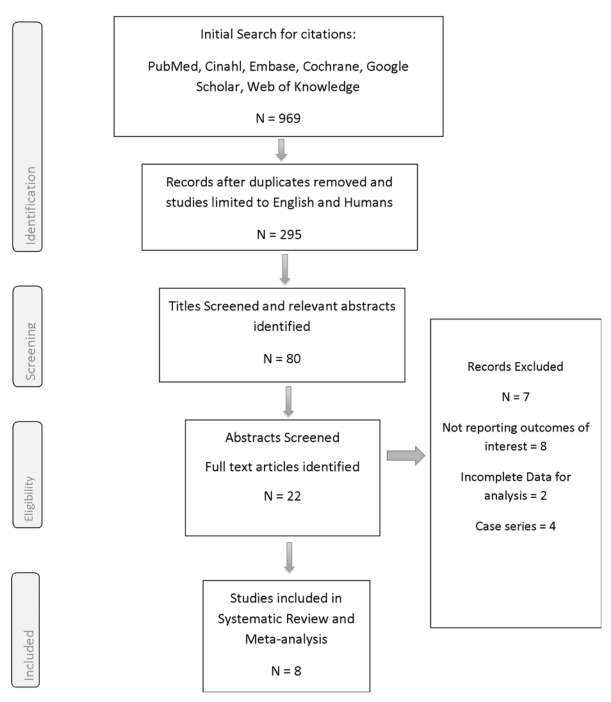


Fig 1. Prisma Flow Diagram. Eligibility for inclusion was determined by two researchers separately (KB, DH) by going through the abstracts of the relevant citations. Differences were settled by examining the full article by both authors, and then any remaining uncertainties regarding eligibility of studies were settled following a discussion with a third author (SRW).

doi:10.1371/journal.pone.0120154.g001



authors (KB, DH), and if remained unsettled, a third author was consulted to resolve the issue (SRW). The following characteristics regarding participants were recorded: age, sex, presence of co-morbidities, primary patency rate, secondary patency rate, maturation rate and postoperative complications (Haematoma, wound infection and steal syndrome). Usability of fistula for Haemodialysis, time to first use for HD and interventions needed to maintain patency were recorded when possible. We also extracted data on compliance with the Society for Vascular Surgery (SVS) recommended standards for reports dealing with arteriovenous haemodialysis accesses [29]. To this end, we assessed whether studies provided SVS standard-based grading of factors that affect outcomes and whether studies provided SVS standard-based grading of severity of arteriovenous access complications. The studies' inclusion and exclusion criteria were also recorded [Table 1].

# Quality assessment for risk of bias

The Downs and Black Tool was used for quality assessment [36]. This tool consists of a total of 27 questions assessing the quality of reporting, external validity and internal validity generating scores between 0 to 32 which includes a score of 0–5 for sample size justification, however, this has been modified by awarding one point for studies that reported on sample size calculations, and zero for those that did not report a methods of sample size calculation. Hence, the modified score ranged from 0 to 27, with higher scores reflecting higher quality. Details of the quality assessment can be found in a separate supplemental table [S1 Table].

# Data analysis

Statistical analyses were performed using Review Manager version 5.3 [37]. We used the random effects model of DerSimonian and Laird [38] to calculate pooled risk ratios for categorical outcomes measures. The Cochran's Q test was used to determine statistical heterogeneity among studies. 95% confidence interval and P-values < 5% were used to determine statistical significance. We compared between the fixed and random effects modelling to produce a sensitivity analysis aimed at detection of the influence of publication bias of small-study effects[39]. Regarding the meta-analysis we additionally, performed a sensitivity analysis limited to published articles only.

# Results

#### Study selection

The results of the study selection process are summarised in the PRISMA flow diagram [Fig. 1]. We started with a total of 969 citations. Following the removal of duplicates and limiting the search criteria to studies conducted on humans and in English language, we were left with 295 citations. We then screened the titles of those papers, and found 80 potentially relevant citations. The abstracts of those titles were examined for relevant outcomes, and 22 papers were evaluated for eligibility criteria, of those 8 citations met our criteria and were included in the systematic review [21,22,30–35]. Of those 8 studies, 1 was a randomised controlled studies (RCT). Five were retrospective cohort studies [22,30,31,33,34] and 2 studies were cohort studies but it was unclear whether they were retrospective or prospective [21,35]. This last citation was a conference presentation which we included in the review, however we also ran a group of sensitivity tests excluding the data from this citation and including data extracted from published papers only [35].

Six of the included studies compared outcomes between 1-stage versus 2-stage BB-AVF formation techniques, while Hossny et al compared 3 different groups, first group of patients had



_	
į	
š	
Studie	
ğ	
₫	
≥	
2	
2	
3	
7	
acteris	
ร	
<u>0</u>	
2	
_	
-	
Þ	
2	

φ		$c\widehat{\mathfrak{S}} \geqslant \overline{\mathfrak{o}} \qquad c\widehat{\mathfrak{c}} \qquad \mathfrak{o} \qquad \mathfrak{c} \qquad \mathfrak{c}$
Characteristics 2 stage	Mean age was 58 years (SD15) 44/84 were female. 33/84 had DM. 67/84 had S.Z. (SD7). 39/84 were black.  Mean vein size was 4.0mm (difference was in vein size (p = 0.041). Factors that affect outcome were not described in accordance with the SVS guidelines.	Mean age was 44.9 years (SD14) for men and 44.1 (SD13) for females. 36 59 were male. Mean duration of ESKD was (SD20) for men and 63.3 (SD21) for men and 63.2 (SD20) for men and 63.2 (SD20) for men and 63.2 (SD20) for men and since are smokers. Mean basilic vein diameter was 2.79 mm (SD0.1). The only significant difference between groups was in vein size (p<0.001). Factors that affect outcome were not accordance with the SVS guidelines.
Characteristics Number 2 stage 1 stage	84 two stage procedures were performed. Number of patients was unclear.	59 two stage procedures on 59 patients were included.
Characteristics 1 stage	Mean age was 58 years (SD15). 32/65 were female. 25/65 had had DM. 53/65 had had DM. 53/65 had mean BMI was 29 (SD6). 29/65 were black.  Mean vein size was 4.0mm (1.15D). The only significant difference was in vein size (p = 0.041). Factors that affect outcome were not described in accordance with the SVS guidelines.	Mean age was 43.1 years (SD16) for men and 42.5 years (SD13) for females. 28/47 were male. Mean duration of ESKD was 63.1 months (SD17) for men and 64.5 (SD18) for women. 15/47 had had heart disease. 2/47 had bwa Dasilic vein diameter was 3.46mm (SD0.2). The only significant difference between groups was in vein size (p.o.001). Factors that affect outcome were not accordance with the SVS guidelines.
Number 1 stage	Data were provided using using number of BBAVFs as the denominator rather than the number of patients. 65 one stage procedures were performed. Number of patients was unclear.	Data were provided using number of BVTs as the denominator rather than the number of patients. 47 one stage procedures on 47 patients were included and Total number of patients was 96 therefore some patients were patients were patients were was 96 therefore patients were patients were patients.
Main findings	Two stage I procedure patients had better functional primary, primary assisted and secondary patency rates at 1 and 2 years. Complication rates were similar.	Two stage procedure patients had a higher rate of maturation but 1 stage BVTs maturad faster. Thrombosis, bleeding, haematoma incidence were incidence were grage group. The two stage group required fewer intervention for patency within the first 10 days but after that there was no difference. Primary and secondary patency rates were better in the two stage group but no statistical analysis was performed for this outcome.
Outcomes assessed	Primary, primary assisted and secondary functional patency rates. Complications 3 such as thrombosis, haematoma, steal syndrome, infection, venous stenosis, mortality.	Primary and secondary patency rates, postoperative complications such as such as thrombosis, haemormage, haemormage, venous ameurysm development, mortality. Rate of fistula maturation and time to fistula maturation. Auxiliary patency.
Nature of the two stage procedure	First, BB-AVF created at the cubital fossa, then 4-6 weeks later a second procedure carried (following US assessment to determine if a second stage is necessary) for mobilisation and superficialisation of the fistula	First, BB-AVF created at the cubital fossa, then 4 weeks then 4 weeks procedure a second procedure carried for and superficialisation of the fistula
Nature of the one stage procedure	Basilic vein dissected and mobilised with preservation of the medial cutaneous nerve of the forearm. End to side arteriovenous anastomosis in the antecubital fossa.	Basilic vein dissected and mobilised with preservation of the medial curant. End to side arteriovenous anastomosis in the anatecubital fossa. HD was allowed after one month.
Exclusion	Specified	Specified specified
Inclusion	Consecutive patients who underwent BBAVF between January 1st 2009 and December 31st 2011.	Patients who underwent authors' austruction(s) between January 2007 and January 2012.
Date Key aspects of published design	Retrospective cohort study at King's College Hospital London. Mean follow up of BBAVF patients was 559 days (SD33). Median interval between first and second operations in the two stage group was 90 days. Allocation to groups was based upon preferences of the two surgeons. Patients with small veins mostly would mostly would have had 2 stage procedure.	Retrospective cohort study at the authors' institutions. Allocation to groups was based upon surgeon patients with basilic vein diameter <3mm had the 2 stage procedure. The second stage of the two stage procedure took place at 30 days. Mean follow up was for 36 months.
Date publishe	2013	2013
Study	[22]	Occan

_
G
rinnec
3
2
≔
-
ਨ
~~
$\mathbf{v}$
$\overline{}$
÷
e -
<u>le</u> 1.
ble 1.
able 1.

Date publishe	Date Key aspects of published design	Inclusion	Exclusion	Nature of the one stage procedure	Nature of the two stage procedure	Outcomes assessed	Main findings	Number 1 C	Characteristics 1 stage	Characteristics Number 2 stage 1 stage	Characteristics 2 stage
2010	Retrospective cohort study at Henry Ford Henry Ford USA on 173 consecutive patients who were scheduled for BYT. Allocation to groups was based on surgeon's preference. The length of follow up was not described explicitly although the report suggests that follow ended follow ended when fistulas were used in dialysis.	Patients who underwent BVT at the BVT at the authors' institution during a 5 year period between xx and xx.	specified specified	Basilic vein dissected and mobilised with preservation of the medial cutaneous nerve of the foream.  Arteriovenous anastomosis in the anastomosis in the brachilal rossa via the brachila for unnar artery.  HD was allowed only after least 6 weeks.	First, BB-AVF created at the cubital fossa, then 4-6 weeks then 4-6 weeks procedure carried for mobilisation and superficialisation of the fistula	Maturation rates and complications such as haematomas, dehiscence, infection, steal syndrome, venous hypertension. 30 day mortality.	One stage procedures had significantly higher complication rates. Haematomas and venous hypertension occurred significantly more often in one stage procedures. Maturation rates were similar although time to although time to although time to although time to stage group.	Data were provided using humber of a the number of denominator is at the number of denominator is a the number of destinct BVT general which this patient is not destinct BVT additional and was and the number of distinct BVT additional and was and was and was and the number of distinct BVT additional and was and was and the number of distinct BVT additional and was and was and the procedures is this patient and which and a this patient and a this	Mean age was 59 years (SD15). Adv/76 were male. 61/76 were black. 45/76 had DM. 51/76 had DM. 51/76 were black. 45/76 were previous didalysis accosss. 61/76 were performed under local anaesthesia. The only significant baseline differences were patients in one stage group had anaesthesia. The only significant baseline patients in one stage group had anaesthesia. An history of previous access and they also were more likely to have general anaesthesia. Notably there were no data on baseline vein diameters. Factors that affect outcome were not date on baseline vein described in accordance with the SVS guidelines.	98 patients underwent two stage procedures. 98 had the first stage and 72 subsequently underwent the study had two distinct BVT procedures and was thus included twice but it was not clear which procedures this partient underwent.	Mean age was 62 years (SD16). 41/98 were male. 73/98 male. 73/98 had DM. 30/98 had previous dialysis access. 14/98 were previous patients. 4/98 were performed under local anaesthesia and general anaesthesia. Notably there were no data on baseline vein diameters. Factors that affect outcome were not data on baseline vein diameters. Factors that affect outcome were not data on baseline vein diameters. Factors that affect outcome were not data on baseline vein diameters. Secondance with the SVS guidelines.

(Continued)

=
ě
ŧμ
0
9
÷
<u>e</u>
<u>a</u>
_

Characteristics Number 2 stage Characteristics 1 stage	20 patients who Mean age was underwent 20 two 35.8 years stage procedures (SD7.3). 11/20 were included. Wean period of occluded in the follow up was interval between 14.8 (SD5). stages and thus factors that was excluded. Were not described in accordance with the SVS guidelines.
s Number 2 sta	
Characteristics 1 stage	
Number 1 stage	
Main findings	Early patency was achieved in 112/20 in the one stage group versus 18/20 in the two stage group. Patency at end of follow was 10/20 versus 16/20. The authors did not use an intention to treat analysis. When an intention to treat analysis was used, the difference was not significant. There was no significant infection or aneurysm rates.
Outcomes assessed	Patency at 4 weeks and patency at end of follow up period. Aneurysm formation and infection.
Nature of the two stage procedure	First, BB-AVF created at the cubital fossa by anastomosing a mobilised segment of basilica vein to the brachial artery. Then 2-4 weeks later a second procedure carried for mobilisation and superficialisation of the fistula.
Nature of the one stage procedure	BB-AVFs were made First, BB-AVF using the traditional one created at the stage technique.  cubital fossa b anastomosing mobilised segment of basilica vein to the brachial artery. Then 2 weeks later a second procedure carried for mobilisation are superficialisation of the fistula.
Exclusion	Specified specified
Inclusion	It involved 40 None patients who specif were admitted for secondary vascular access procedures between June 1993 and December 1995.
Key aspects of Inclusion design	Prospective randomised controlled trial at El Menoufia University Hospital Egypt. Allocation to groups was performed randomly and groups were matched for age and gender. No details on the randomisation process were provided. Follow up was for 6–24 months.
Date Key asp published design	1998
Study	Mallah 32 32

(Continued)



outcome were not described in accordance with the SVS guidelines. Mean age was 54 years (SD14. 29/77 were male. 67/77 had current catheter usage at the time of surgery. 16/77 had prior Characteristics fistula. Average BMI was 28.1, neart failure. 27, ad prior failure coronary artery nad congestive 77 had GA and 42/77 had DM, hypertension, 21/77 had access. 39/77 block. Factors arteriovenous disease, 7/77 regional arm others had 71/77 had hat affect psilateral 2 stage of an Characteristics Number 2 stage underwent the 77 patients two stage procedure. was in regards to history of catheter use and prior ipsilateral 14/29 were male. 16/29 had 54 years (SD21). current catheter usage at the time of the congestive heart access. Average hypertension, 5/ 29 had coronary regional arm block. The only accordance with between groups ad GA and the surgery. 16/29 had prior Mean age was 16/29 had DM, artery disease, affect outcome BMI was 28.1, characteristics differences in failure. 13/29 described in Factors that others had procedure. 28/29 had significant quidelines. psilateral the SVS 2/29 had baseline 1 stage were not Number 1 29 patients one stage BVT underwent stage primary patency rates were better in the one stage group. Other outcomes were not significantly Primary patency Main findings and assisted primary assisted three years, reinterventions, mortality, major complications, infections, steal and secondary maturation and patency up to complications Outcomes assessed syndrome, such as fistula The anastomosis and, subsequently, the vein wastransposed was createdin the first stage Nature of the in the second procedure two stage anastomosis along with the superficialtransposition, sall in the same tprocedure Nature of the one stage procedure Brachial artery tobrachial vein Exclusion None specified BVT between June 2006 and June 2010. It is computerised cohort study on 106 patients 106 patients who who underwent BVT at underwent the Methodist BVT betweer DeBakey Heart & June 2006 Vascular Centre and June 106 patients consecutive. whether the Data came cases were Inclusion It involved database. from a unclear Key aspects of design of one stage BVT or two stage BVT Follow up was for in Texas. Choice was based upon oreference. surgeon published 2012 Date Study Syed

(Continued)

Table 1. (Continued)



<b>PLOS</b>	ONE
· • · • • • • • • • • • • • • • • • • •	ı

Study	Date Key ask published design	Key aspects of I design	Inclusion	Exclusion	Nature of the one stage procedure	Nature of the two stage procedure	Outcomes assessed	Main findings	Number 1 stage	Characteristics Number 2 stage 1 stage	Number 2 stage	Characteristics 2 stage
Agarwal	2014	Retrospective cohort study involving 144 consecutive patients who underwent BVT at a US hospital. Patients with basalic vein diameter of <4mm were chosen for the two stage procedure. Mean follow up duration was unclear. Some patients were followed for greater than 4 years.	It involved consecutive patients who underwent BVT creation between January 2005 and who received all access-fatted care (surgical and radiological) up to a 4 year follow up point in December 2013.	Patients were excluded if interventions or follow up had taken place at an outside institution.	Not specified	Not specified	Maturation rates, mean time to inlitation of fistula use, intensity of percutaneous interventions per patient year on dialysis, primary assisted patency, primary assisted patency and secondary patency annually.	Modest reduction in primary and secondary patency rates in the two stage group compared to the one stage group	61 patients underwent 61 one stage BVTs	Mean age was 59.1 years. No other were provided on baseline characteristics. Factors that affect outcome were not described in accordance with the SVS guidelines.	83 patients underwent 83 two stage BVTs.	Mean age was 61.5 years. No other data were provided on baseline characteristics. Factors that affect outcome were not described in accordance with the SVS guidelines.
Hossny [21]	5003	Cohort study involving 70 patients at Menofia University Egypt. University Egypt. It is unclear whether it was prospective or retrospective or retrospective. The study compared basilic vein transposition vein transposition vein transposition procedure versus a one stage elevation procedure. It is unclear on what grounds patients were selected for different procedures. Mean follow up time was 25.8 Mean follow up time was 25.8	t involved 70 brachiobasilic sixulas that were performed in 70 patients over an unspecified 2 year period at the author's institution.	spedified	30 fistulas created using a traditional one stage BVT to create a BB-AVF20 fistulas created in a one stage elevation technique, the basilic vein was brought superficial to the deep fascia and subcutaneous tissue rather than through a subcutaneous tunnel	created in a two fistula for stage elevation dialysis, technique, the basilic vein was brought application deep fascia and loedema, subcutaneous harmatoma tissue rather than thrombosis, though a venous subcutaneous hypertensio tunnel hypholeakk perioperativ mortality,	Ability to access fiftstula for dialysis, cumulative secondary patency, complications (coeping and an analysis), who thrombosis, which is a secondary thrombosis, who thrombosis, which is a secondary thrombosis, who thrombosis, who thrombosis, who thrombosis, who thrombosis, who thrombosis, who thrombosis, which is a secondary thrombosis,	The one stage BVT had a lower complication rate and was favoured by the dalysis staff compared to basalic vein superficialisation techniques	20 patients underwent 20 one stage BVTs, while 20 patients underwent one stage one stage passilic vein elevation procedure	For the one stage BVT: Wean Age = 45.7 (1-98), 12/40 were created in male patients, 17/20 had had diabetes and 10/20 had hypertension/For the one stage basilic vein elevation: Mean age = 49.3 (26- 77.1, 12 had diabetes and 7 had hypertension. Factors that affect outcome were not described in accordance with the SVS guidelines.	20 patients underwent two stage basilic vein procedure	Mean Age = 54 created in male created in male patients. Diabetics = 4/20 and 5/20 had Hypertension. Factors that affect outcome were not adescribed in accordance with the SVS guidelines.



Study	Date published	Date Key aspects of published design	Inclusion	Exclusion	Nature of the one stage procedure	Nature of the two stage procedure	Outcomes assessed	Main findings	Number 1 stage	Characteristics 1 stage	Characteristics Number 2 stage 1 stage	Characteristics 2 stage
Effat [35] 2013	2013	Cohort study involving 104 patients who underwent 106 Brachiobasilic fistulas at Zagazig University University Hospital from October 2010 to December 2011. It is unclear whether it was prospective or retrospective or retrospective or retrospective. Comparison between one stage BVT, two stag	Scheduled for brachiobasilic fistula with a basilic vein >2.5mm diameter and a brachial artery >3mm.	Scheduled for Patients were brachlobasilic excluded if istula with a vein diameter basilic vein - {2.5mm, failure of diameter and mature in artery >3mm. staged or massive venous hypertension after creation of the brachlobasilic shunt and failed to be corrected, patients who refused the second stages or who were lost to follow up between stages.	All fistulas created using a traditional one stage BVT to create a BB-AVF	38 fistulas were created using a two stage BVT technique, stage nor involved forming a BB-AVF, the second stage involved mobilisation and superficialisation of the fistula. In 40 fistulas, they carried a two stage superficialisation procedures without transposing the basilic vein	functional patency (ability to access the fistula for haemodialysis), mean time to use the fistula, complications such as haematomas requiring exploration, wound dehiscence or infection, thrombosis, steal syndrome, venous hypertension requiring intervention, failure to mature.	Lower patency rates for the one stage technique and increased chance of developing postoperative complications compared to the two stage technique	28 one stage BVTs performed. Wumber of patients unclear.	28 one stage Mean age = 43.6 38 two stage BVTs  BVTs  ± 11.9, 13/28  BVTs and 40  performed.  were male, 13, stage  Number of 28 had diabetes superficializa  patients and 16/28 had  noclear.  Factors that  affect outcome  were not  described in  accordance with  the SVS  guidelines.	38 two stage BVTs and 40 two stage superficialization. Number of patients unclear	For the two stage BVT: Mean age = 48.4 ± 10.2, 20/38 were created in male patients. 19/38 had diabetes and 22/38 had had have the two stage elevation: Mean age 475 ± 18.4, 16/40 were created in male patients, 23/40 had diabetes and 26/40 had hypertension. Factors that affect outcome were not described in accordance with the SVS guidelines.

doi:10.1371/journal.pone.0120154.t001



traditional 1-stage basilic vein transposition (BVT), while the second group had 1-stage basilic vein elevation, and the third group underwent a 2-stage BB-AVF. For the sake of this meta-analysis, we pooled the first 2 groups from this particular study together [21]. Similarly, Effat had 3 groups of patients in his conference paper, the first group had standard 1-stage BVT, whereas the second group had a 2-stage BVT and the last group consisted of patients who had 2-stage superficialization of the basilic vein to create BB-AVF. We pooled the data from the 1-stage procedures in this last study together in the meta-analysis [35].

# **Participants**

The studies included a total of 849 patients who had 859 fistulas, of those 366 fistulas were formed using a 1-stage technique, while the remaining 493 fistulas were created in a 2-stage technique. Overall, 432 were male patients versus 417 female patients. Kakkos et al [31] did not specify the male to female ratio in the 72 patients who underwent a 2-stage procedure in their study, however, in the remaining studies, 181 men had 1-stage fistula procedure compared to 164 in the 2-stage group. Similarly, 226 in the 1-stage group were female patients compared to 150 in the 2-stage group. Of the 6 [21,22,30,31,33,35] studies that reported past history of diabetes, 143/295 patients were in the 1-stage group while 202/390 were in the 2-stage group. History of hypertension was reported in 5 studies [21,22,30,33,35], with 123/219 patients in the 1-stage group and 202/390 in the 2-stage group having the diagnosis. All studies reported on findings in adult patients with end stage renal disease (ESRD), El-Mallah [32] had the youngest patients (23.5  $\pm$  5.8 years for the 1-stage group, and 35.8  $\pm$  7.3 years for patients in the 2-stage group), while the remaining studies included patients in their fifties and sixties [Table 1]. Inclusion and exclusion criteria of studies among other characteristics are outlined in [Table 1]. Main outcomes reported in studies are summarised in [Table 2].

#### Successful maturation rate

Successful maturation rates were reported in 6 of included studies [21,22,31–34]; the criteria used for reporting maturation are found in [Table 3]. Those studies had a combined total of 683 fistulas, 301 of those were created in the one stage group, whereas 382 were created in the two stage group. The difference between the two groups was not significant in pooled analysis (Pooled risk ratio = 0.95 [0.82, 1.11], 95% CI, P = 0.53) [Fig. 2]. Heterogeneity was detected statistically (Cochran's Q = 14.48; degree of freedom (DF) = 5; P = 0.001;  $I_2 = 65\%$ ). The significance of the results was not altered when using the fixed effects analysis model as a sensitivity test to detect publication bias (Pooled risk ratio = 0.92 [0.84, 1.01], 95% CI, P = 0.07).

#### Postoperative complications

**Haematoma.** The incidence of postoperative wound haematoma was reported in 6 of the included studies [21,22,30,31,33,35] with a total of 711 fistulas, of those, 295 fistulas were created in the 1-stage group and 416 fistulas in the 2-stage group. Analysis of pooled data showed the difference was not significant (Pooled risk ratio = 0.73 [0.34, 1.58], 95% CI, P = 0.43) [Fig. 3]. Heterogeneity was not detected statistically (Cochran's Q = 9.76; degree of freedom (DF) = 5; P = 0.08;  $I_2 = 49\%$ ). The results were not changed significantly when using the fixed effects analysis model as a sensitivity test to detect publication bias (Pooled risk ratio = 0.67 [0.41, 1.11], 95% CI, P = 0.12). A sensitivity test by excluding the data from the conference paper by Effat [35] was carried out, and no significant difference was found in the incidence of postoperative haematoma between the two groups (Pooled risk ratio = 0.67 [0.27, 1.64], 95% CI, P = 0.38).



# Table 2. Main outcomes from included studies.

PLOS ONE

		1-sta	1-stage procedure				2-stage	2-stage procedure		
Study	Number of 1-stage Patency fistulas	· Patency	Haematoma	Wound Infection	Steal	Number of 2-stage fistulas	Patency	Haematoma	Wound infection	Steal
Vrakas [22]	65 one stage procedures were performed. Number of patients was unclear.	Primary functional patency at 1 and 2 years was 71% and 53%. Assisted Primary functional patency at 1 and 2 years was 77% and 57%. Secondary functional patency at 1 and 2 years was 79% and 57%. The definitions for patency outcomes were based upon the SVS guidelines.	3/65No SVS grading was provided	3 / 65	2 / 65No SVS grading was provided	84 two stage procedures were performed. Number of patients was unclear.	Primary functional patency at 1 and 2 years 87% and 75%. Assisted Primary functional patency at 1 and 2 years was 95% and 77%. Secondary functional patency at 1 and 2 years was 95% and 77% at 1 and 2 years	3/84No SVS grading was provided.	2 / 84	3/84
Ozcan [30]	47 one stage procedures on 47 patients were included and Total number of patients was 96 therefore some patients were included twice.	Primary patency at 1, 2 and 3 years was 33/47 (70%), 30/47 (64%), and 27/47 (54%) Secondary patency at 1, 2 and 3 years was 36/47 (76%), 43/47 (72%), and 31/47 (66%). The definitions for patency outcomes were unclear.	8 / 47No SVS grading was provided	6 / 47No SVS grading was provided	4 / 47No SVS grading was provided but all required surgical management	59 two stage procedures on 59 patients were included.	Primary patency at 1, 2 and 3 years was 41/59 (84%), 36/59 (73%), and 34/59 (69%). Secondary patency at 1, 2 and 3 years was 44/59 (90%), 40/59 (82%), and 38/59 (77%)	3/59No SVS grading was provided	5 / 59No SVS grading was provided	3 / 59No SVS grading was provided but all require surgical treatment
Kakkos [31]	76 one stage procedures were performed. One patient in the study had two distinct BVT procedures and was thus included twice but it was not clear which procedures this procedures this patient underwent.	Not reported	grading was SVS grading provided. Across was provided the whole study, most were grade 1 or 2 and 3 were grade 3	5 / 76No SVS grading was provided	3 / 76No SVS grading was provided but all were managed conservatively	98 patients underwent two stage procedures.98 had the first stage and 72 subsequently underwent the second stage. One patient in the study had two distinct BVT procedures and was thus included twice but it was not clear which underwent.	Not reported	3 / 72No SVS 0 / 72No grading was SVS grading provided. Across was provided the whole study, most were grade 1 or 2 and 3 were grade 3	0 / 72No SVS grading was provided	2 / 72No SVS grading was provided but all were managed conservatively
EI Mallah	20 patients who underwent 20 one stage procedures were included.	Early patency (4 weeks) = 12/20 (60%), Overall patency (at the end of follow-up) = 10/20 (50%), The definitions for patency outcomes were unclear.		3 / 20No SVS grading was provided but they were described as mild infections	0 / 20	20 patients who underwent 20 two stage procedures were included. One fistula occluded in the interval between stages and thus was excluded.	Early patency (4 weeks) 2-stage = 18/20 (90%). Overall patency at the end of the study = 16/20 (80%)		1 / 20No SVS grading was provided but they were described as mild infections	0 / 20



		1-sta	1-stage procedure				2-stage	2-stage procedure		
Study	Number of 1-stage Patency fistulas		Haematoma	Wound	Steal	Number of 2-stage fistulas	Patency	Haematoma	Wound infection	Steal
Syed [33]	29 patients underwent 20 one stage BVT	Primary patency at 1, 2 and 3 years was 82%, 81%, and 51%. Assisted primary patency at 1, 2 and 3 years was 91%, 77%, and 48%. Secondary patency at 1, 2 and 3 years was 91%, and 58%. The definitions are similar to those in the SVS guidelines.	2 / 29No SVS grading was provided.	0 / 29	0 / 29	77 patients underwent the two stage procedure.	Primary patency at 1, 2 and 3 years was 67%, 27%, and 18%. Assisted primary patency at 1, 2 and 3 years was 77%, 41%, and 24%. Secondary patency at 1, 2 and 3 years was 81%, 61%, and 45%	6 / 77No SVS grading was provided.	3 / 77No SVS grading was provided.	3 / 77No SVS grading was provided.
Agarwal	61 patients underwent 61 one stage BVTs	Primary unassisted patency at 1 and 2 years was 26% and 7%. Primary assisted patency at 1, 2, 3 and 4 years was 67%, 38%, 21% and 8%. Secondary patency at 1, 2, 3 and 4 years was 86%, 56%, 69% and 57%. The definitions are similar to those in the SVS guidelines.				83 patients underwent 83 two stage BVTs.	Primary unassisted patency at 1 and 2 years was 13% and 0%. Primary assisted patency at 1, 2, 3 and 4 years was 66%, 5econdary patency at 1, 2, 3 and 4 years was 76%, 71%, 49% and 25%.			
Hossny [21]	J 20 patients underwent 20 one stage BVTs, while 20 patients underwent one stage basilic vein elevation procedure	The study reported 87% cumulative secondary patency rate at 1 year across all groups, with 86.7% for the BVT group, 90% for the 1-stage elevation group and 84.2% for the 2-stage elevation group. 1 death was stage elevation final analysis.  Cumulative secondary patency rate at 2 years for rate at 2 years for all groups was 75%, with 82.8k for the BVT group, 70% for the 1-stage elevation group and 68.4% for the 2-stage elevation group. 2 deaths were excluded from final analysis.	6 / 50No SVS grading was provided.		0 / 20	20 patients underwent two stage basilic vein procedure	The study reported 87% cumulative secondary patency rate at 1 year across all groups, with 86.7% for the BVT group, 90% for the 1-stage elevation group and 84.2% for the 2-stage elevation group. 1 death was succondary patency rate at 2 years for all groups was 75%, with 82.8% for the BVT group, 70% for the 1-stage elevation group and 68.4% for the 2-stage elevation.	5 / 20No SVS grading was provided.		0 / 20
										(F. C.)

Table 2. (Continued)



Table 2. (Continued)

	±1	-stage procedure				2-staç	2-stage procedure		
Study	Number of 1-stage Patency fistulas	Haematoma	Wound Infection	Steal	Number of 2-stage fistulas	Patency	Haematoma	Wound infection	Steal
Effat conference [35]	28 one stage BVTs Not reported performed. Number of patients unclear.	2 /28No SVS grading was provided.	2 / 28No SVS grading was provided	2 / 28No 2 / 28No SVS SVS grading grading was was provided provided	38 two stage BVTs and Not reported 40 two stage superficialization. Number of patients unclear	Not reported	7 / 78No SVS 12 / 78No 0 / 78No SVS grading was SVS grading grading was provided provided	12 / 78No SVS grading was provided	0 / 78No SVS grading was provided

doi:10.1371/journal.pone.0120154.t002



Table 3. Maturation.

Study	One stage	Two stage	Source of data	comments
Vrakas [22]	36 / 65	49 / 84	Primary failure rates were reported. This was defined as an AVF that was never used for dialysis. Primary failure may have resulted from inadequate maturation, early thrombosis, failure of first cannulation, and other complications which made AVF unusable. Successful maturation rates were derived from these data.	The number of AVFs that required intervention to assist maturation is unclear.
Kakkos [31]	67 / 76	69 / 98	Maturation rates were reported. Maturation was based upon clinical judgement (development of basilic vein dilatation and thrill for a sufficient length).	Includes fistulas that required intervention to assist maturation for dialysis. 7 one stage fistulas required such intervention and 3 two stage fistulas required such intervention.
El-Mallah [32]	12 / 20	18 / 20	Patency at 4 weeks was reported and we used this figure to determine successful maturation. The authors did not provide a definition for patency.	The number of AVFs that required intervention to assist maturation is unclear.
Syed [33]	6 / 29	14 / 77	Maturation rates were reported. Fistula maturation was defined as dilation of the vein to allow cannulation and support dialysis at a minimum flow rate of 350ml/min for at least 3 sessions.	The number of AVFs that required intervention to assist maturation is unclear.
Agarwal	55 / 61	62 / 83	Maturation rates were reported.  Maturation was defined as the use of the fistula for haemodialysis for any amount of time or, if it was not used, documentation in surgical or renal records that the fistula was mature and ready for use based upon successful cannulation and/or physical examination by vascular surgery.	Includes an unspecified number of fistulae that needed percutaneous intervention to assist maturation.
Hossny [21]	47 / 50	19 / 20	Numbers of fistulas that were successfully used for dialysis at 6 weeks were reported.	No patients needed reintervention to assist achievement of successful dialysis at 6 weeks.

doi:10.1371/journal.pone.0120154.t003

**Wound infection.** Six of the included studies [30-33,35] reported on the incidence of postoperative wound infection with a total number of 681 fistulas, 265 of those belonged to the 1-stage group, while 416 consisted of 2-stage fistulas. Meta-analysis of the pooled data showed the difference between groups not to be significant (Pooled risk ratio = 0.77 [0.35, 1.68], 95% CI, P = 0.51) [Fig. 4]. There was no evidence of statistical heterogeneity (Cochran's Q = 5.76; degree of freedom (DF) = 5; P = 0.51;  $I_2 = 13\%$ ). The results were not changed significantly when using the fixed effects analysis model (Pooled risk ratio = 0.73 [0.39, 1.37], 95% CI, P = 0.32). A sensitivity test by excluding the data from the conference paper by Effat [35] was carried out, and no significant difference was found in the incidence of postoperative wound infection between the two groups (Pooled risk ratio = 0.57 [0.25, 1.27], 95% CI, P = 0.17).

**Steal syndrome.** Six of the studies [22,30–33,35] reported on the risk of developing significant postoperative ischaemia (steal syndrome). Those studies had a combined total of 681



Fig 2. Successful maturation rate.

doi:10.1371/journal.pone.0120154.g002

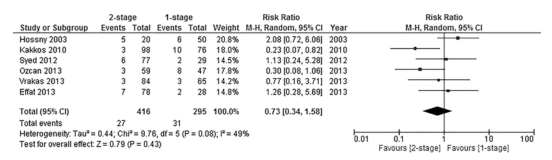


Fig 3. Postoperative Haematoma.

doi:10.1371/journal.pone.0120154.g003

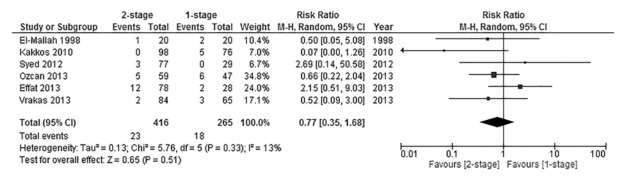


Fig 4. Postoperative wound infection.

doi:10.1371/journal.pone.0120154.g004

patients, of those, 265 belonged in the 1-stage group, while 416 belonged in the 2-stage group. Analysis of pooled data showed the difference was not significant (Pooled risk ratio = 0.65 [0.27, 1.53], 95% CI, P = 0.32) [Fig. 5]. Heterogeneity was not detected statistically (Cochran's Q = 3.42; degree of freedom (DF) = 4; P = 0.49;  $I_2$  = 0%). The results were not changed significantly when using the fixed effects analysis model (Pooled risk ratio = 0.64 [0.29, 1.40], 95% CI, P = 0.26). A sensitivity test by excluding the data from the conference paper by Effat [35] was carried out, and no significant difference was found in the incidence of postoperative haematoma between the two groups (Pooled risk ratio = 0.79 [0.32, 1.94], 95% CI, P = 0.60).



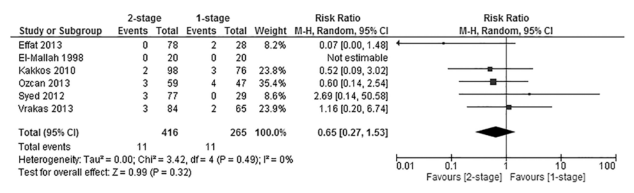


Fig 5. Steal syndrome.

doi:10.1371/journal.pone.0120154.g005

# **Systematic Review**

El-Mallah in his paper titled "Staged basilic vein transposition for dialysis angioaccess" published in 1998 [32] compared outcomes in two groups randomly allocated to receive either 1-stage BVT or 2-stage BVT. The difference in early patency rates was significant and favoured the 2-stage approach (60% of 1-stage vs 90% of 2-stage, P < 0.05), as well as overall patency rates at the end of follow-up (50% of 1-stage vs 80% of 2-stage, P < 0.05) [32]. Postoperative wound infection rate also favoured the 2-stage approach with one case compared to 3 in the 1-stage group. There was no difference in postoperative aneurysmal dilatation, and there was no significant ischaemia (steal syndrome) reported in either of the two groups [32].

Hossny looked at the different surgical techniques used in creation of a BB-AVF in 2003 [21]. He compared patency rates and dialysis related complications in 70 patients divided in 3 groups, 30 of those patients had traditional BVT whereas 20 patients had 1-stage BB-AVF with elevation and the remaining 20 patients had 2-stage fistula with elevation of the vein [21]. Cumulative secondary patency rates were comparable among the 3 groups at 1 year and 2 years; at 1 year (86.7% for the BVT group, 90% for the 1-stage elevation group and 84.2% for the last group), while at 2 years (82.8%, 70%, and 68.4%, respectively) [21]. Similarly, no significant difference was found in his study in postoperative early thrombosis across the groups. Postoperative arm oedema occurred in a total of 14 patients, all of whom had temporary subclavian access catheters sited on the same arm on which the new fistula was created. All 14 cases were managed conservatively with success. The difference in developing postoperative haematoma significantly favoured the traditional BVT approach compared to the two elevation methods with fewer haematomas reported in the first group. Interestingly, the dialysis staff were more satisfied with the 1-stage BVT technique, whereas only 53.3% reported satisfaction with the elevation technique (1-stage and 2-stage techniques) (P < .001) [21].

Kakkos et al also tried to answer the question of "What is the Optimal Technique" for performing a BVT in a retrospective study of 173 patients published in 2010 [31]. They found that the incidence of venous hypertension (17% vs 4%, P = 0.004), wound infection (13% vs 3%, P = 0.012) and all complications (43% vs 11%, P < 0.001) were significantly higher in patients who had 1-stage BVT when compared to those who had a 2-stage BVT [31]. Time to fistula use in HD was—expectedly—shorter in the 1-stage group (Median = 68 (49–103) days) compared to the 2-stage group (Median = 132 (102–166) days). This difference was significant (P < 0.001) [31]. However maturation rates were similar (85% for the 1-stage groups versus 82% for the 2-stage group, P = 0.49). Median time to use the fistulas for HD in the 2-stage group was 66 in patients who developed postoperative complications, compared to 50 days in those who did not



(P = 0.019). They also found that wound infection occurred more in patients who were operated under general anaesthetic compared to those who had their procedures done under local anaesthetic (OR 38, P < 0.001). Also, venous hypertension was found to occur more frequently in patients who developed postoperative wound haematoma, but the difference was not statistically significant (18% vs 6%, P 0.12) [31]. A trend was noted towards steal being more common in patients with previous vascular access than in those who did not have such access (4.9% vs 1.1%, P = 0.19) [31].

Syed et al carried out a similar study comparing the outcomes of 1-stage (29 patients) and 2-stage (77 patients) BVT and published their findings in 2012 [33]. 79% of patients in the 1-stage group had a history of a previously failed access for HD compared to 51% of the 2-stage group. They found that the rate of primary failure was comparable between both groups (21% vs 18%) [33]. In their study, patients who had 1-stage BVT had better patency rates when compared to those who had 2-stage procedures at 1 year (82% vs 67%), 2 years (81% vs 27%) and 3 years (51% vs 18%) respectively. The same finding was reported for secondary patency (91%, 80% and 58% for 1-stage BVT compared to 81%, 61% and 45% for the 2-stage group at 1, 2 and 3 years respectively) [33]. Reintervention rate in this study was 62% for the 1-stage vs 66% for the 2-stage group. It is worth noting that 87% of the patients in the 2-stage group were using catheters for dialysis, whereas 55% of the 1-stage group were dialysing through a catheter at the time of access formation [33].

Ozcan et al published a paper in 2013 with preliminary results from their study comparing 1-stage and 2-stage BVT to create AVF access in HD patients [30]. They retrospectively divided their patients to those with a basilic vein > 3 mm and who had a 1-stage BVT procedure, and those with a basilic vein < 3 mm who had a 2-stage procedure. Although the diameter of the basilic vein was statistically higher in the first group  $(3.46 \pm 0.2 \text{ mm})$  compared to the second group (2.79  $\pm$  0.1 mm) (P < 0.005), the rate of fistula maturation was significantly lower in the first group (66% vs 77%, P < 0.005) [30]. Also, postoperative complications were significantly higher among the first group of patients who had 1-stage BVT. Thrombosis occurred in 34% compared to 23% of patients who had a 2-stage procedure, haemorrhage in (36% vs 14%) and haematoma in (17% vs 6%) respectively. Time required for the fistula to mature was significantly shorter in the first group (Mean 41 ± 14 days) compared to the second group (Mean 64  $\pm$  28 days) (P < 0.05) [30]. Early interventions ( $\leq$  10 days) for fistula thrombosis occurred more frequently in the first group (21% vs 12%, P < 0.05), although there was no significant difference in terms of late interventions ( $\geq 10$  days) required to deal with access thrombosis (20% in the first group vs 22% in the second) [30]. Also they reported superior primary patency rates at 6, 12, 18, 24, 30 and 36 months for those who had 2-stage BVT fistulas compared to the first group of patients (1-stage 83%, 70%, 68%64%, 60% and 57% versus 88%, 84%, 80, 73%, 71% and 69% for the 2-stage respectively). Similarly, the 1-stage had lower secondary patency rates at 6, 12, 18, 24, 30 and 36 months when compared to the 2-stage group (85%, 76%, 74%, 72%, 70% and 66% versus 94%, 90%, 84%, 82%, 80% and 77% respectively) [30].

Similarly, Vrakas et al evaluated the difference in outcomes between 1-stage (65 fistulas) and 2-stage (84 fistulas) BB-AVFs performed in 141 patients [22]. They performed ultrasound scans 4–6 weeks after the first stage procedure to determine if a second stage was required. Patients who had their fistulas created in a 1-stage approach had a bigger preoperative basilic vein diameter  $(4.0 \pm 1.1 \text{ mm vs } 3.6 \pm 1.3 \text{ mm}, P = 0.041)$  [22]. There was no difference in primary failure between the groups (45% vs 42%, P = 0.718), however the 1-stage BB-AVF had significantly lower primary (71% vs 87%; P = 0.034), assisted primary (77% vs 95%; P = 0.017), and secondary (79% vs 95%; P = 0.026) functional patency rates compared to the 2-stage BB-AVF [22]. Multivariate Cox regression analysis showed that the 1-stage procedure was 3.2 times more likely to fail (P = 0.028), and male gender was associated with loss of access (P = 0.054).



66% of the firstulas in the first group were used successfully for HD compared to 60% in the 2-stage group (P = 0.407), and intervention before first successful HD session was equivalent between both groups (21% vs 11%, P = 0.201) [22]. Overall, 93 (62%) fistulas were successfully used for HD (66% 1-stage vs 60% 2-stage; P = 0.407), of the remaining 56 (38%), 19 fistulas (34%) failed before needling, 2 (4%) received a renal transplant, 7 (13%) died, and 28 (50%) BBAVFs remain patient in patients awaiting to start HD [22].

Agarwal et al examined the outcomes of 1-stage vs 2-stage BVT AVF. They included patients who underwent percutaneous angioplasty (assisted maturation) in calculating the overall maturation rate which was 90% for the 1-stage group (55/61 patients) compared to 75% of the 2-stage group (62/83 patients) (P = 0.02). Subgroup analysis showed that both men (54/66 patients) and women (64/78) in this study had a maturation rate of 82% (P = 0.97)[34]. Primary unassisted patency rates were comparable between the groups (69%, 52%, 26%, and 7% for the 1-stage BVT at 3 months, 6 months, 1 and 2 years; compared to 58%, 35%, 13%, and 0% of the 2-stage group, respectively (P = 0.12) [34]. Similarly, no significant difference was found in secondary patency on an intent to-treat basis (86%, 75%, 69%, and 57% at 1, 2, 3, and 4 years for 1-stage group; compared to 76%, 71%, 49%, and 25% of the 2-stage group, respectively); (P = 0.12) [34]. The intensity of percutaneous interventions in their study was 1.84/patient-year of dialysis (PYD) for the 1-stage group versus 2.15/PYD for the 2-stage group (P = 0.57) [34]. They suggested that although the 2-stage BB-AVF technique resulted in modest reduction in maturation and patency rates, it should still be favoured to the use of synthetic grafts in patients who would not be suitable for a 1-stage BB-AVF procedure [34].

The number of AVFs that failed to progress from the first stage to the second stage in the two-staged BVT approach were unclear in four studies [22,30,34,35]. In the remaining four, El-Mallah [32] reported 1/20 patient which had an occluded shunt, while Hossny [21] also had 1/20 patient failing to progress due to spontaneous thrombosis within the first 4 weeks postoperatively. Syed [33] had 2/77 patients that never progressed to the second stage of the procedure. Kakkos [31] reported that 26/98 of his patients never had a second stage procedure (thrombosed (n = 4), failed to mature and was abandoned during the re-exploration (n = 12), patient refused the procedure (n = 3), lost to follow-up (n = 1), died (n = 2), venous hypertension (n = 2), venous monomelic neuropathy (n = 1) requiring ligation, moved out of state (n = 1)).

Number of interventions required to maintain patency or to improve the fistula maturation rates were not reported clearly in all studies. Hossny reported that in the one-stage group one patient underwent ligation and another had a surgical revision, same numbers occurred in the two-stage group. All ligations were done to treat venous hypertension, whereas all revisions were performed to improve poor flow [21]. Kakkos reported that in the two-stage group 6 patients had endovenous angioplasty interventions, 3 had surgical revisions compared to 3 patients and 1 patient in the one-stage group respectively [31]. Syed et al performed 37 angioplasty interventions, 9 surgical revisions and 5 thrombectomy procedures in their two-stage group, compared to 14, 2 and 2 patients respectively [33]. The remaining studies either did not report data related to fistula salvage procedures or it was reported in poor details making it difficult to quantify those interventions.

With the exception of the studies by El-Mallah et al [32] and the one by Vrakas et al [22] which both reported significantly superior patency rates in the two-stage groups, and the paper by Syed [33] which conversely reported a significantly better patency rates in the one-stage BVT group, the remaining studies all reported comparable patency rates [21,30,31,34,35] [Table 2]. However it is important to point out that patency rate data were reported as percentages with the lack of clearly identifiable denominators in the majority of those studies, thus making pooling those data in a meta-analysis not feasible. Also, definitions used in individual studies included in this review for patency rates (primary, assisted primary and secondary) differed significantly.



#### **Discussion**

The number of patients with end stage renal disease (ESRD) requiring haemodialysis (HD) is steadily rising, a trend that is expected to continue [2]. A well-functioning AVF is superior to grafts and central catheters in providing access for haemodialysis efficiently and at the same time with the least rate of access related complications. This has lead vascular surgeons to resort to the basilic vein which by virtue of its anatomical position is less likely to be damaged by repeated cannulation as with the more superficial veins of the arm and forearm. However, a consensus on how to form a brachiobasilic AVF does not exist as some surgeons choose to do this in a one-stage operation, while others prefer a two-stage procedure with the first procedure usually involving making the anastomosis between the basilic vein and the brachial artery, while in the second stage the arterialised vein is mobilised and brought closer to the skin surface to facilitate cannulation for HD sessions.

This review identified eight studies [21,22,30-35], including data from a conference paper [35] in order to increase the rigour of the review. The pooled data referred to 849 patients with a total of 859 fistulas, 366 of those fistulas belonged to patients who underwent a 1-stage BB-AVF, while 493 fistulas were performed using a 2-stage technique to create the access. The data from 6 of the included studies [21,22,30,31,33,35] were used to compare the difference between the two groups in developing postoperative haematoma which was not significant (Pooled risk ratio = 0.69 [0.30, 1.56], 95% CI, P = 0.37). Excluding the data from the conference paper by Effat [35] in a sensitivity test did not alter the result (Pooled risk ratio = 0.61 [0.23, 1.60], 95% CI, P = 0.31).

Incidence of postoperative wound infection was reported in five studies [30-33,35], and the difference between the 1-stage group and the 2-stage group was not found to be significant (Pooled risk ratio = 0.82 [0.31, 2.18], 95% CI, P = 0.69). This remained unchanged when excluding the data by Effat [35] in a sensitivity test (Pooled risk ratio = 0.57 [0.21, 1.51], 95% CI, P = 0.27).

Similarly, the difference between the two groups was not found to be significant when it came to postoperative ischaemia (steal syndrome) in the 6 studies which reported this complication [ $\underline{22,30-33,35}$ ] (Pooled risk ratio = 0.51 [0.20, 1.30], 95% CI, P = 0.16). We performed a sensitivity test by excluding the data by Effat [ $\underline{35}$ ] from the pooled data, and the result was not altered (Pooled risk ratio = 0.63 [0.23, 1.69], 95% CI, P = 0.35).

Ozcan et al [30] allocated patients to groups based on vein diameter, with those with basilic vein > 3 mm receiving a 1-stage BVT, while patients with basilic vein < 3 mm received a 2-stage BVT. Even with this seemingly advantageous difference in favour of the 1-stage approach, they reported superior patency rates and maturation rates in patients who had a 2-stage procedure with primary patency at 1, 2 and 3 years for the 1-stage group of (70%), (64%), and (54%) versus (84%), (73%), and (69%) in the 2-stage group. Secondary patency rates at 1, 2 and 3 years for the 1-stage group were (76%), (72%), and (66%) versus (90%), (82%), and (77%) in the 2-stage group.

Similarly, Vrakas et al reported a smaller mean vein diameter of  $(3.6 \pm 1.3 \text{ mm})$  for the 2-stage, versus  $(4.0 \pm 1.1 \text{ mm})$  fistulas created in the 1-stage group, yet their results favoured the 2-stage approach with primary functional patency at 1 and 2 years for the 1-stage group of 71% and 53% versus 87% and 75% in the 2-stage group. Assisted Primary functional patency at 1 and 2 years for the 1-stage group was 77% and 57% versus 95% and 77% in the 2-stage group, while secondary functional patency at 1 and 2 years for the 1-stage group was 79% and 57% versus 95% and 77% in the 2-stage group.

Conversely, in study by Agarwal et al  $[\underline{34}]$ , their patients in the 1-stage group achieved better maturation rate than those who had a 2-stage BVT fistulas (90% vs 75%, P = 0.02). They did



not include any analysis between the two groups based on vein diameter. Vein diameter has been shown to negatively influence maturation and patency rates in AVFs, and is one of the main predictors of those outcomes in fistulas [40,41], and indeed has been shown to be the only independent predictor of maturation in some studies[42,43]. Syed et al reported similar findings with better primary and cumulative patencies in the 1-stage group with primary patency at 1, 2 and 3 years (82% vs 67%), (81% vs 27%) and (51% vs 18%), while (secondary patency at 1, 2 and 3 years (91%, 80% and 58% for 1-stage BVT and 81%, 61% and 45% for the 2-stage group. Variations in vein diameter between the two groups were not reported in this study.

Kakkos et al [31] did not find a significant difference in maturation between the two groups, as 15% of fistulas in the 1-stage group did not mature, compared to 18% in the second group (P=0.49). They did however find significant difference in developing postoperative haematoma (13% vs 3%, P=0.012), venous hypertension (17% vs 4%, P=0.004) and overall complications (43% vs 11%, P<0.001), all in favour of the 2-stage BVT technique.

Kim et al compared the 2-stage approach to all other AVF procedures including 1-stage BVT, radiocephalic and brachiocephalic fistulas. All of the 2-stage BB-AVFs in their study successfully matured compared to a pool consisting of all different types that showed a combined maturation rate of 52% (P = 0.001). Fistula failure occurred in 7% of the 2-stage BVT compared to 59% of other fistulas (P = 0.001), and more 2-stage BVT fistulas were used successfully for HD compared to all other fistula types (87% vs 48%, P = 0.024). Also, the patency rate at 1 year was superior in the 2-stage group compared to other AVFs (91% vs 47%, P = 0.003).

One of the limitations of this review is the low number of randomised trials included—1 study was randomised—while the remaining 7 were cohort studies. Most of these studies were retrospective. Another limitation is the variation in surgical approaches, those variations include technical differences in performing the procedure, as well as differences in equipment used and expertise among participating surgeons. Those limitations can be addressed by conducting a large randomised multi-centre trial that would adhere to a rigid protocol in patients' selection process and performing the procedures. Another limiting factor is the lack of sufficient sub-group analysis among included studies, particularly analysis taking into account factors that are known to be associated with fistula maturation such as vein diameter. Finally, we highlight that included studies were not compliant with SVS reporting recommendations regarding baseline factors that affect outcomes or severity of complications.

#### Conclusion

Although more studies seem to favour the 2-stage BVT approach, evidence in the literature is not sufficient to draw a final conclusion as the difference between the 1-stage and the 2-stage approaches for creation of a BB-AVF is not statistically significant in terms of the overall maturation rate and postoperative complications. Patency rates (primary, assisted primary and secondary) were comparable in the majority of studies. Large randomised properly conducted trials with adequate sub-group analysis are needed before making a final recommendation. Future studies should aim for compliance with established reporting standards.

## **Supporting Information**

**S1 PRISMA Checklist. PRISMA 2009 Checklist.** (DOCX)

S1 Table. Quality assessment score of individual studies. (DOCX)



### **Author Contributions**

Conceived and designed the experiments: KB DH LB MW MCM EK PB SRW. Performed the experiments: KB DH SE LB MW MCM EK PB SRW. Analyzed the data: KB DH SRW. Wrote the paper: KB DH SE SRW.

#### References

- Bashar K, Healy D, Browne LD, Kheirelseid EA, Walsh MT, Moloney MC, et al. Role of far infra-red therapy in dialysis arterio-venous fistula maturation and survival: systematic review and meta-analysis. PLoS One. (2014); 9: e104931. doi: 10.1371/journal.pone.0104931 PMID: 25115802
- Frankel A. Temporary access and central venous catheters. Eur J Vasc Endovasc Surg. (2006); 31: 417–422. PMID: 16360326
- Spergel LM, Ravani P, Roy-Chaudhury P, Asif A, Besarab A. Surgical salvage of the autogenous arteriovenous fistula (AVF). J Nephrol. (2007); 20: 388–398. PMID: <u>17879203</u>
- NKF-DOQI clinical practice guidelines for vascular access. National Kidney Foundation-Dialysis Outcomes Quality Initiative. Am J Kidney Dis. (1997); 30: S150–191. PMID: 9339150
- Lynch JR, Mohan S, McClellan WM. Achieving the goal: results from the Fistula First Breakthrough Initiative. Curr Opin Nephrol Hypertens. (2011); 20: 583–592. doi: <a href="https://doi.org/10.1097/MNH.0b013e32834b33c4">10.1097/MNH.0b013e32834b33c4</a>
   PMID: 21897231
- Ethier JH, Lindsay RM, Barre PE, Kappel JE, Carlisle EJ, Common A. Clinical practice guidelines for vascular access. Canadian Society pf Nephrology. J Am Soc Nephrol. (1999); 10 Suppl 13: S297–305. PMID: 10425613
- Fluck R, Kumwenda M. Renal Association Clinical Practice Guideline on vascular access for haemodialysis. Nephron Clin Pract. (2011); 118 Suppl 1: c225–240. doi: 10.1159/000328071 PMID: 21555898
- McCann M, Einarsdottir H, Van Waeleghem JP, Murphy F, Sedgewick J. Vascular access management 1: an overview. J Ren Care. (2008); 34: 77–84. doi: <a href="https://doi.org/10.1111/j.1755-6686.2008.00022.x">10.1111/j.1755-6686.2008.00022.x</a> PMID: 18498572
- Centofanti G, Fujii EY, Cavalcante RN, Bortolini E, de Abreu LC, Valenti VE, et al. An experience of vascular access for hemodialysis in Brazil. Int Arch Med. (2011); 4: 16. doi: <a href="https://doi.org/10.1186/1755-7682-4-16">10.1186/1755-7682-4-16</a>
   PMID: 21569616
- Leblanc M, Saint-Sauveur E, Pichette V. Native arterio-venous fistula for hemodialysis: What to expect early after creation? J Vasc Access. (2003); 4: 39–44. PMID: <u>17642058</u>
- NKF-KDOQI. 2006 Updates Clinical Practice Guidelines and Recommendations. (2006); pp. PMID: 17044433
- Sultan S, Hynes N, Hamada N, Tawfick W. Patients on hemodialysis are better served by a proximal arteriovenous fistula for long-term venous access. Vasc Endovascular Surg. (2012); 46: 624–634. doi: 10.1177/1538574412462635 PMID: 23064823
- Ilhan G, Esi E, Bozok S, Yurekli I, Ozpak B, Ozelci A, et al. The clinical utility of vascular mapping with Doppler ultrasound prior to arteriovenous fistula construction for hemodialysis access. J Vasc Access. (2013); 14: 83–88. doi: 10.5301/jva.5000097 PMID: 23032950
- 14. Heye S, Fourneau I, Maleux G, Claes K, Kuypers D, Oyen R. Preoperative mapping for haemodialysis access surgery with CO(2) venography of the upper limb. Eur J Vasc Endovasc Surg. (2010); 39: 340–345. doi: 10.1016/j.ejvs.2009.11.036 PMID: 20080420
- 15. Allon M. Current management of vascular access. Clin J Am Soc Nephrol. (2007); 2: 786–800. PMID: 17699495
- Rayner HC, Pisoni RL, Gillespie BW, Goodkin DA, Akiba T, Akizawa T, et al. Creation, cannulation and survival of arteriovenous fistulae: data from the Dialysis Outcomes and Practice Patterns Study. Kidney Int. (2003); 63: 323–330. PMID: 12472799
- Pisoni RL, Young EW, Dykstra DM, Greenwood RN, Hecking E, Gillespie B, et al. Vascular access use in Europe and the United States: results from the DOPPS. Kidney Int. (2002); 61: 305–316. PMID: 11786113
- Ethier J, Mendelssohn DC, Elder SJ, Hasegawa T, Akizawa T, Akiba T, et al. Vascular access use and outcomes: an international perspective from the Dialysis Outcomes and Practice Patterns Study.
   Nephrol Dial Transplant. (2008); 23: 3219–3226. doi: 10.1093/ndt/gfn261 PMID: 18511606
- Hirth RA, Turenne MN, Woods JD, Young EW, Port FK, Pauly MV, et al. Predictors of type of vascular access in hemodialysis patients. JAMA. (1996); 276: 1303–1308. PMID: 8861988



- Dagher F, Gelber R, Ramos E, Sadler J. The use of basilic vein and brachial artery as an A-V fistula for long term hemodialysis. J Surg Res. (1976); 20: 373–376. PMID: 933493
- Hossny A. Brachiobasilic arteriovenous fistula: different surgical techniques and their effects on fistula patency and dialysis-related complications. J Vasc Surg. (2003); 37: 821–826. PMID: 12663983
- Vrakas G, Defigueiredo F, Turner S, Jones C, Taylor J, Calder F. A comparison of the outcomes of one-stage and two-stage brachiobasilic arteriovenous fistulas. J Vasc Surg. (2013); 58: 1300–1304. doi: 10.1016/j.jvs.2013.05.030 PMID: 23810301
- Koksoy C, Demirci RK, Balci D, Solak T, Kose SK. Brachiobasilic versus brachiocephalic arteriovenous fistula: a prospective randomized study. J Vasc Surg. (2009); 49: 171–177 e175. doi: 10.1016/j.jvs. 2008.08.002 PMID: 18945577
- 24. Harper SJ, Goncalves I, Doughman T, Nicholson ML. Arteriovenous fistula formation using transposed basilic vein: extensive single centre experience. Eur J Vasc Endovasc Surg. (2008); 36: 237–241. doi: 10.1016/j.ejvs.2008.02.012 PMID: 18396069
- Taghizadeh A, Dasgupta P, Khan MS, Taylor J, Koffman G. Long-term outcomes of brachiobasilic transposition fistula for haemodialysis. Eur J Vasc Endovasc Surg. (2003); 26: 670–672. PMID: 14603429
- Anaya-Ayala JE, Younes HK, Kaiser CL, Syed O, Ismail N, Naoum JJ, et al. Prevalence of variant brachial-basilic vein anatomy and implications for vascular access planning. J Vasc Surg. (2011); 53: 720–724. doi: 10.1016/j.jvs.2010.09.072 PMID: 21144691
- 27. Glickman M. Basilic vein transposition: review of different techniques. J Vasc Access. (2014); 15 Suppl 7: S81–84. doi: 10.5301/jva.5000260 PMID: 24817461
- Liberati A, Altman DG, Tetzlaff J, Mulrow C, Gotzsche PC, Ioannidis JP, et al. The PRISMA statement for reporting systematic reviews and meta-analyses of studies that evaluate health care interventions: explanation and elaboration. PLoS Med. (2009); 6: e1000100. doi: 10.1371/journal.pmed.1000100 PMID: 19621070
- Sidawy AN, Gray R, Besarab A, Henry M, Ascher E, Silva M Jr., et al. Recommended standards for reports dealing with arteriovenous hemodialysis accesses. J Vasc Surg. (2002); 35: 603–610. PMID: 11877717
- Ozcan S, Gur AK, Yener AU, Odabasi D. Comparison of one- and two-stage basilic vein transposition for arterio-venous fistula formation in haemodialysis patients: preliminary results. Cardiovasc J Afr. (2013); 24: 364–368. doi: 10.5830/CVJA-2013-077 PMID: 24337214
- Kakkos SK, Haddad GK, Weaver MR, Haddad RK, Scully MM. Basilic vein transposition: what is the optimal technique? Eur J Vasc Endovasc Surg. (2010); 39: 612–619. doi: 10.1016/j.ejvs.2010.01.006 PMID: 20172751
- El Mallah S. Staged basilic vein transposition for dialysis angioaccess. Int Angiol. (1998); 17: 65–68.
   PMID: 9754891
- Syed FA, Smolock CJ, Duran C, Anaya-Ayala JE, Naoum JJ, Hyunh TT, et al. Comparison of outcomes of one-stage basilic vein transpositions and two-stage basilic vein transpositions. Ann Vasc Surg. (2012); 26: 852–857. doi: 10.1016/j.avsg.2011.12.013 PMID: 22794333
- Agarwal A, Mantell M, Cohen R, Yan Y, Trerotola S, Clark TW. Outcomes of single-stage compared to two-stage basilic vein transposition fistulae. Semin Dial. (2014); 27: 298–302. doi: <a href="https://doi.org/10.1111/sdi.12170">10.1111/sdi.12170</a>
   PMID: 24320222
- 35. Effat M. Staging of Brachiobasilic Arteriovenous Fistulae, Does it Worth Effort? VSE. (2013); 9.
- 36. Downs SH, Black N. The feasibility of creating a checklist for the assessment of the methodological quality both of randomised and non-randomised studies of health care interventions. J Epidemiol Community Health. (1998); 52: 377–384. PMID: 9764259
- The Nordic Cochrane Centre TCc. Review Manager (RevMan) [Computer Program]. Version 5.2. (2012).
- DerSimonian R, Laird NM. Evaluating the Effect of Coaching on SAT Scores: A Meta-Analysis. Harvard Educational Review. (1983); 53: 1–15.
- Sterne JAC, Egger M, Moher D. Chapter 10: Addressing reporting biases. Higgins JPT, Green S (editors) Cochrane Handbook for Systematic Reviews of Intervention (2011); Version 5.1.0
- Feldman HI, Joffe M, Rosas SE, Burns JE, Knauss J, Brayman K. Predictors of successful arteriovenous fistula maturation. Am J Kidney Dis. (2003); 42: 1000–1012. PMID: 14582044
- Mendes RR, Farber MA, Marston WA, Dinwiddie LC, Keagy BA, Burnham SJ. Prediction of wrist arteriovenous fistula maturation with preoperative vein mapping with ultrasonography. J Vasc Surg. (2002); 36: 460–463. PMID: 12218967



- 42. Lauvao LS, Ihnat DM, Goshima KR, Chavez L, Gruessner AC, Mills JL Sr. Vein diameter is the major predictor of fistula maturation. J Vasc Surg. (2009); 49: 1499–1504. doi: 10.1016/j.jvs.2009.02.018 PMID: 19497513
- 43. Maya ID, O'Neal JC, Young CJ, Barker-Finkel J, Allon M. Outcomes of brachiocephalic fistulas, transposed brachiobasilic fistulas, and upper arm grafts. Clin J Am Soc Nephrol. (2009); 4: 86–92. doi: 10. 2215/CJN.02910608 PMID: 18945990