Utility of Blood Flow/Resistance Index Ratio ($Q_x$) as a Marker of Stenosis and Future Thrombotic Events in Native Arteriovenous Fistulas

Alessandro Colombo¹, Michele Provenzano², Laura Rivoli³, Cinzia Donato⁴, Marinella Capria², Giuseppe Leonardi², Salvatore Chiarella⁴, Michele Andreucci², Giorgio Fuiano², Davide Bolignano², and Giuseppe Coppolino²*

¹ Dialysis Unit, Hospital of Crotone, Crotone, Italy, ² Renal Unit, “Magna Graecia” University, Catanzaro, Italy, ³ Unit of Nephrology, Department of Internal Medicine, Chivasso Hospital, Turin, Italy, ⁴ Renal Unit, “Pugliese-Ciaccio” Hospital of Catanzaro, Catanzaro, Italy

Objective: The resistance index (RI) and the blood flow volume ($Q_a$) are the most used Doppler ultrasound (DUS) parameters to identify the presence of stenosis in arteriovenous fistula (AVF). However, the reliability of these indexes is now matter of concern, particularly in predicting subsequent thrombosis. In this study, we aimed at testing the diagnostic capacity of the $Q_a$/RI ratio ($Q_x$) for the early identification of AVF stenosis and for thrombosis risk stratification.

Methods: From a multicentre source population of 336 HD patients, we identified 119 patients presenting at least one “alarm sign” for clinical suspicious of stenosis. Patients were therefore categorized by DUS as stenotic ($n=60$) or not-stenotic ($n=59$) and prospectively followed. $Q_a$, RI, and $Q_x$, together with various clinical and laboratory parameters, were recorded.

Results: $Q_a$ and $Q_x$ were significantly higher while RI was significantly lower in non-stenotic vs. stenotic patients ($p<0.001$ for each comparison). At ROC analyses, $Q_x$ had the best discriminatory power in identifying the presence of stenosis as compared to $Q_a$ and RI (AUCs 0.976 vs. 0.953 and 0.804; $p=0.037$ and $p<0.0001$, respectively). During follow-up, we registered 30 thrombotic events with an incidence rate of 12.65 (95% CI 8.54–18.06) per 100 patients/year. In Cox-regression proportional hazard models, $Q_x$ showed a better capacity to predict thrombosis occurrence as compared to $Q_a$ (difference between c-indexes: 0.012; 95% CI 0.004–0.01).

Conclusions: In chronic haemodialysis patients, $Q_x$ might represent a more reliable and valid indicator for the early identification of stenotic AVFs and for predicting the risk of following thrombosis.

Keywords: blood flow, resistance index, stenosis, thrombosis, haemodialysis, Doppler ultrasound, arteriovenous fistula
INTRODUCTION

Arteriovenous fistula (AVF) is generally acknowledged as the best vascular access option for chronic haemodialysis (1). However, the clinical management of complications, principally stenosis, remains a major challenge for clinicians with a substantial impact on health resources worldwide (2–4). Stenosis not infrequently preludes to thrombotic events (5). Fluctuating shear stress in the stenotic region leads to intimal injury with a subsequent cascade of pro-inflammatory cytokines and proliferation of smooth-muscle cells, myofibroblasts and extracellular matrix that result in neo-intimal hyperplasia and subsequent risk of thrombosis (6).

Doppler ultrasound (DUS) surveillance is a practical and non-invasive approach to identify critical stenosis at risk of forthcoming thrombosis (7). Among DUS parameters, AVF blood flow volume ($Q_x$) is nowadays considered the best parameter to detect stenosis while the resistance index (RI) usually provides less reliable information (8). Stenosis progression overtime generates resistance to flow, thereby increasing RI and decreasing blood flow $Q_x$ (9). However, despite the lumen reduction, a high flow rate or a normal resistance index could paradoxically be observed in early stenotic AVFs, hence masquerading a good access patency (10, 11). Better indicators are therefore needed for improving critical stenosis identification and for stratifying the risk of thrombosis.

In the present study we tested the performance of a new surrogate Doppler parameter ($Q_x$), obtained as the $Q_x$/RI ratio, to detect the presence of AVF stenosis. We then evaluated the reliability of this new index to predict thrombosis occurrence in the brief to mid-term.

PATIENTS AND METHODS

Study Cohort

From January to December 2017, haemodialysis patients from four different Italian centers ("Magna Graecia" University and "Pugliese-Ciaccio" Hospital of Catanzaro, Crotone and Chivasso Hospitals) were screened for signs of AVF stenosis, such as difficult cannulation, thrombi aspiration during cannulation, increasing of pre-pump arterial and post-pump venous pressure, unfeasibility to reach adequate blood flow rate or elongation of haemostasis. Patients who were positive to at least one of these "alarm signs" underwent specific DUS surveillance (12). Patients were then categorized as stenotic or not-stenotic basing on recognized criteria: 2 major criteria [reduction of vessel size greater than 50% and a ratio between systolic peak velocity (SPV) in the stenotic region and SPV in pre-stenotic region major of 2 (>2)] and a supplementary finding like drop of the access arterial pressure and recirculation were recorded.

Exclusion/Inclusion Criteria

Patients included had a native vascular access created at least 4 months prior to our assessment to guarantee an adequate maturation period and were usually cannulated with classical rope-ladder technique. All subjects were on regular treatment haemodialysis with a rhythm of 4-h sessions three times a week with dry-weight stable for at least 2 months before entering the study and had achieved a normotensive oedema-free state. Exclusion criteria were the presence or a recent history of bleeding, malignancy, liver, thyroid or infectious diseases, alterations in leucocyte count or formula and/or treatment with steroids or immunosuppressors. The local ethics committee approved the study, and fully informed consent was obtained from all participants.

Procedure

Two different blinded experienced examiners performed an ultra-sonographic study of AVFs 1 h before dialysis. The patient was in sitting position with elbow pad to stabilize the upper limb. After physical examination, B-mode and color Doppler images were obtained from the feeding artery, arterial anastomosis and the outflow vein to characterize vascular access and identify the presence of stenosis. Patency and continuity of the artery and vein were assessed from the distal forearm to the upper arm. The following parameters were recorded for each included subject with a standardized protocol (1–4, 15):

- Blood Flow ($Q_x$) in the brachial artery by measuring the vessel diameter ($D$) and the time average velocity (TAV) through the formula: $Q_x$ (ml/minute) = TAV (cm/second) * $D$ (cm) * 60.
- Resistive index (RI) calculated by the following formula $RI = (A - B)/A$, ($A$ = Peak Systolic Velocity, $B$ = End Diastolic Velocity).

Finally, the $Q_x$ was computed as the $Q_x$/RI ratio. For each parameter, the mean of three different measures was considered for the statistical analysis.

After the baseline assessment, patients were then prospectively followed until the occurrence of a thrombosis event or up to the established end of the follow-up period (30th January 2020). Thrombosis was diagnosed by physical exam as no thrill, bruit or pulse and confirmed by DUS with an absence of flow on pulse-wave and color Doppler.

Statistical Analysis

The statistical analysis was performed using STATA version 14 (StatCorp, College Station, TX, USA), the GraphPad Prism and Med Calc package for figure depiction. Continuous variables were presented as mean ± SD or median (IQ range) as appropriate. Categorical variables were presented as percentage (%). Differences between groups were tested using the unpaired $t$-test for normally distributed values and Kruskal–Wallis analysis followed by a post-hoc test the Dunn’s test for nonparametric values. Receiver operating characteristics (ROC) analysis was employed to evaluate variable’s ($Q_x$, RI, and $Q_x$) ability for classifying disease status, which is, presence or absence of AVF critical stenosis. Comparison between ROC curves was assessed by a non-parametric approach (16). To find the best cut-off value for identifying the presence of AVF stenosis, the Youden index ($J$) was computed (17).

With respect to the risk of thrombosis, two multivariable Cox proportional hazards models were built including $Q_x$ and $Q_x$. 
respectively. We used Cox analyses to estimate hazard ratio (HR) and 95% CI of AVF thrombosis over time. For the model building process, univariate analysis testing the association between the main clinical variables and the onset of thrombosis was assessed by means of logistic regression analysis. The variables with $p < 0.150$ at univariate analysis were selected and included in the first multivariable Cox proportional hazard model. Next, backward variable selection method with an elimination criterion of $p < 0.10$ was performed to fit the second Cox model (Model with $Q_a$ or $Q_x$). Such as stringent cut-off for variables inclusion was used in order to avoid model overfitting, due to the limited sample of the cohort. The final model was adjusted by: systolic blood pressure, $Q_a$ or $Q_x$ and AVF dysfunction.

To assess the performance of the two models ($Q_a$ vs. $Q_x$), we compared the discrimination ability of each model by calculating the c-index for Cox proportional hazard model. CI of difference between c-indexes was calculated by the bootstrap method; bootstrap CI was calculated with 1,000 replicates and using the percentile method (18).

**RESULTS**

**Patients’ Flow**

The source population consisted of 336 chronic haemodialysis patients. During the screening phase, 143 patients displayed at least one clinical sign of possible AVF stenosis. Twenty-four patients were excluded because lost at follow-up or because underwent a percutaneous angioplasty/surgical re-intervention before a thrombotic event. The final study cohort therefore consisted of 119 individuals who resulted almost equally distributed among the stenotic ($n = 60$) and non-stenotic...
(n = 59) group according to DUS examination. Figure 1 depicts the patients’ flow.

### Baseline Assessment

The whole population included a moderate-high percentage of patients with type-II diabetes and CVD (29.4 and 58.0%, respectively). The principal types of AVF were distal (Radiocephalic fistulas) (63.0%) or proximal (Brachial artery–to–transposed basilic vein fistulas) (31.1%) with a marginal part represented by mid-arm (Brachiocephalic fistula) (5.9%) access. Overall Qx, RI, and Qx were 1,437 (820–2,260) mL/min, 0.55 ± 0.09 and 2,727 (1,371–4,385) mL/min/RI, respectively.

The stenotic and non-stenotic groups were homogeneous for sex, CVD, smoking habit and laboratory parameters. Conversely, stenotic group showed a significant higher prevalence of diabetes, lower BMI, and higher frequency of previous access dysfunction as compared to non-stenotic group. Mean levels of Qx and Qx were significantly higher [2,255 (1,755–3,128) vs. 820 (663–1,304) mL/min and 4,377 (3,486–6,342) vs. 1,488 mL/min/RI (1,000–2,320)] while RI was significantly lower (0.49 ± 0.07 vs. 0.59 ± 0.08) (p < 0.001 for all comparisons) in not stenotic as compared to stenotic AVFs. Main baseline data and comparisons are summarized in Table 1.

ROC curves (Figure 2) showed that Qx had a better discriminatory power in identifying the presence of AVF stenosis as compared to Qx and RI. Values of Area Under the Curves (AUC) were 0.976, 0.953, and 0.804 for Qx, Qx, and RI, respectively, and the difference between curves was statistically significant (AUC Qx vs. Qx = 0.037; AUC Qx vs. RI = 0.001). When the cut-points J were derived from the ROC curves, the threshold of 3,333 mL/min/RI for Qx (Sensitivity = 99.9%, Specificity = 83.3%) displayed the best accuracy in detecting the presence of stenosis as compared to that of Qx (1,615 mL/min; Sensitivity = 94.9%, Specificity = 81.7%) and RI (0.53; Sensitivity = 81.3%, Specificity = 68.3%).

### Prospective Phase and Analyses

During a median follow-up of 24.5 (21.7–26.9) months, we registered 30 thrombotic events with an incidence rate of 12.65

---

**Table 1** | Demographic and clinical characteristics of patients at basal visit.

|                         | Overall | Stenosis | p          |
|-------------------------|---------|----------|------------|
| Number (%)              | 119 (100) | 60 (50.4) | 59 (49.6) | –         |
| Age (years)             | 62.7 ± 11.8 | 62.9 ± 12.3 | 62.6 ± 11.3 | 0.897     |
| Males (%)               | 58.0 | 52.5 | 63.3 | 0.233     |
| Diabetes (%)            | 29.4 | 37.3 | 21.7 | 0.061     |
| History of CVD (%)      | 58.0 | 64.4 | 51.7 | 0.169     |
| Smoking (%)             | 7.6 | 6.8 | 8.3 | 0.749     |
| BMI (kg/m²)             | 26.3 ± 4.1 | 25.6 ± 3.0 | 27.0 ± 4.8 | 0.070     |
| Systolic blood pressure (mmHg) | 136 ± 19 | 137 ± 18 | 135 ± 20 | 0.538     |
| Diastolic blood pressure (mmHg) | 82 ± 8 | 82 ± 7 | 82 ± 9 | 0.770     |
| Hemoglobin (g/dL)       | 10.8 ± 0.8 | 10.8 ± 0.8 | 10.8 ± 0.9 | 0.655     |
| Haematocrit (%)         | 33 ± 4 | 33 ± 3 | 33 ± 3 | 0.734     |
| Platelet (×10^3)        | 204 ± 52 | 206 ± 56 | 203 ± 48 | 0.799     |
| Calcium (mg/dL)         | 9.4 ± 0.6 | 9.4 ± 0.5 | 9.4 ± 0.7 | 0.660     |
| Phosphorus (mg/dL)      | 5.8 ± 1.2 | 5.90 ± 0.90 | 5.80 ± 1.42 | 0.639     |
| Total cholesterol (mg/dL) | 190 ± 40 | 192 ± 43 | 186 ± 38 | 0.412     |
| Antihypertensive drugs (n) | 1.5 ± 1 | 1.5 ± 1 | 1.5 ± 1 | 0.755     |
| HD vintage (months)     | 30.0 ± 13.9 | 31.4 ± 14.4 | 28.7 ± 13.4 | 0.292     |
| Vintage of AV access (years)** | 26.2 ± 12.7 | 28.7 ± 13.5 | 23.8 ± 12.0 | 0.037     |
| Types of AVF            | – | – | – | 0.003     |
| Distal (Radiocephalic fistulas) (%) | 63.0 | 78.0 | 48.3 |           |
| Mid-arm (Brachiocephalic fistulas) (%) | 5.9 | 5.1 | 6.7 |           |
| Proximal (Brachial artery–to–transposed basilic vein fistulas) (%) | 31.1 | 16.9 | 45.0 |           |

| Qx (mL/min)             | 1,437 (820–2,260) | 820 (663–1,304) | 2,255 (1,755–3,128) | <0.001     |
| RI                      | 0.55 ± 0.09 | 0.59 ± 0.08 | 0.49 ± 0.07 | <0.001     |
| Qx (mL/min/RI)          | 2,727 (1,371–4,385) | 1,488 (1,000–2,320) | 4,377 (3,486–6,342) | <0.001     |
| History of access dysfunction (%) | 26.9 | 39.0 | 15.0 | 0.003     |

* CVD, cardiovascular diseases: myocardial infarction, stroke, peripheral artery disease, chronic heart failure. **Dialysis vintage, length of time on dialysis.
FIGURE 2 | Receiver operating characteristics curves of AVF blood flow volume (Qa), resistance index (RI) and Qa/RI Resistance Index ratio (Qx) considering the presence of AVF stenosis as status variable. The area under the curve (AUC) for Qa, Qx, and RI were, respectively, 0.953, 0.976, and 0.804 (AUC Qa vs. Qx p = 0.037; AUC Qx vs. RI p = 0.0001). The best cut-off values to identify the presence of stenosis were: 1,615 mL/min for Qa, with 94.9% (95% CI, 91.7–97.8) sensitivity and 81.7% (95% CI, 69.8–93.4) specificity; 3,333 mL/min/RI for Qx, with 99.9% (95% CI, 98.2–100) sensitivity and 83.3% (95% CI: 72.2–95.1) specificity; 0.53 for RI with 81.3% (95% CI, 69.1–90.3) sensitivity and 68.3% (95% CI 55–79.7) specificity.

(95% CI 8.54–18.06) per 100 patients/year. From the univariate linear regression analysis (Table 2), covariates with $p < 0.150$ (presence of stenosis, Qa or Qx, RI, AVF dysfunction and systolic blood pressure) were included in the multivariate linear regression model. After backward selection of variables, with an elimination criterion of $p < 0.100$, systolic blood pressure, the presence of AVF dysfunction, low Qa and Qx were independently and significantly associated to the development of thrombosis over time (Table 3).

C index of model with Qx was higher than model with Qa, with the difference being statistically significant after applying the bootstrap method (0.012; 95%CI 0.004–0.019). This indicates a better discrimination ability of the model with Qx as compared to that with Qa on thrombosis occurrence.

DISCUSSION

In the present study, we have demonstrated the potential usefulness of Qx as a new surrogate ultrasound marker to detect stenosis and predict thrombosis episodes in haemodialysis AVF. Qx combines information from Qa and RI, two other surrogate markers that have already been validated in literature and extensively used in daily practice. Yet, in our study, the
Colombo et al.

**TABLE 2 | Univariate logistic regression analysis for the onset of thrombosis.**

| Characteristics | β Coefficient (95% CI) | p   |
|-----------------|------------------------|-----|
| Age, years      | 0.007 (–0.029–0.042)   | 0.708|
| Gender          |                        |     |
| male (vs. female) | 0.112 (–0.731–0.954)   | 0.796|
| Stenosis        |                        |     |
| (yes vs. no)    | 2.401 (1.263–3.538)    | <0.001|
| Q_s (mL/min)    | –0.002 (–0.002 to –0.001) | <0.001|
| Q_r (mL/min/RI) | –0.001 (–0.002 to –0.001) | <0.001|
| RI              | 7.76 (2.77–12.76)      | 0.002|
| Time from AVF creation, months | 0.016 (–0.016–0.048) | 0.332|
| AVF dysfunction (yes vs. no) | 1.44 (0.55–2.33) | 0.001|
| CVD, %          | 0.690 (–0.195–0.1575)  | 0.127|
| Systolic BP, mmHg | –0.024 (–0.047 to –0.001) | 0.040|
| Diastolic BP, mmHg | –0.023 (–0.074–0.027) | 0.356|
| Calcium, mg/dL  | 0.009 (–0.686–0.704)   | 0.980|
| Phosphorus, mg/dL | 0.043 (–0.303–0.388)  | 0.809|
| Cholesterol, mg/dL | –0.001 (–0.012–0.009) | 0.788|
| Hemoglobin, g/dL | –0.353 (–0.885–0.179) | 0.193|

**Bold values are statistical significant.**

**TABLE 3 | Multivariable Cox models on the risk of thrombosis.**

|                  | Model with Q_s | Model with Q_r |
|------------------|----------------|----------------|
|                  | HR 95% CI      | HR 95% CI      |
| Systolic blood pressure, mmHg | 0.97 0.95–0.99 | 0.98 0.95–0.99 |
| AVF dysfunction, yes vs. no | 2.39 1.15–4.96 | 2.35 1.13–4.86 |
| Q_s or Q_r       | 0.99 <0.001    | 0.99 <0.001    |
| c-index          | 0.766          | 0.778          |

**Bold values are statistical significant.**

Q_s/RI ratio showed an improved diagnostic performance with respect to the single parameters considered alone. This finding may be of high clinical relevance as it may help overcoming the accuracy limitations that characterizes Q_s and RI, often leading to a misleading interpretation of AVF patency. In fact, lower RI values are not infrequently found in spite of set lumen narrowing in outflow vein, while a high flow rate is often maintained in early stenosis with lumen reduction (10, 19).

Different guidelines suggest variable criteria to approach AVF stenosis. In particular, it is still highly debated whether an immediate revascularization should be preferred over a “wait and see” strategy in order to prevent thrombosis occurrence (8, 20, 21).

The main aim of our investigation was to explore the validity of a new DUS parameter for critical stenosis to overcome hurdles that usually do not allow early intervention.

Such a tool could help in the decision process for a timely corrective procedure, like angioplasty, in order to maintain AVF patency for a longer time (22). Implications for health-care systems would be of utmost importance as vascular access failure remains a major cause of mortality and hospitalizations in the uremic population (23) with a consequent high economic burden (24).

In addition, stratifying stenotic lesions according to the risk of thrombosis remains a demanding priority. Several studies quantified the increased risk of infection and cardiovascular events after AVF failure (25) while other studies assessed the impact of conversion from native AVF to a catheter on mortality (26–29). Recently the AURORA trial (30) displayed a strong association between AVF thrombosis and worsened patients’ survival, emphasizing the impact of cardiovascular risk in the uremic population. The high mortality rate after AVF thrombosis was attributed to ionic imbalance related to the inability to perform a regular dialysis session, difficulties associated with central catheter placement as well as the revascularization procedure itself. In this respect, there is extensive agreement in current literature about the need of developing easy and validated tools for a better stratification of the thrombotic risk of AVF. Shintaku et al. found flow rate and resistance index immediately after AVF creation to be better predictors of early access failure and AVF maturation as compared to lumen diameter measurement. In fact, these two parameters may reveal functional changes preluding to morphological alterations leading to thrombosis (31).

According to our findings, Q_s could represent a possible way to improve even the detection accuracy of a single Q_a measurement, simply by performing a RI-based "normalization." This was confirmed by the robust results obtained at Cox multivariate analyses in which a model including Q_s predicted the thrombotic event independently from other confounders such as age, sex and type of AVF and in a stronger manner than Q_a.

The main strength of our study was the ample cohort and the long follow-up after the baseline assessment that allowed us to catch an adequate number of thrombotic events and, hence, to perform reliable multivariate adjustments and elaborated comparison of models. In addition, despite multicentre, the study population resulted quite homogeneous and the DUS procedures were highly reproducible across the different centers involved. The main limitation is probably represented by the lack of an angiographic exam as reference to confirm steno-thrombotic events. Digital subtraction angiography is nowadays recognized...
as the gold standard exam to characterize stenotic vascular lesions. However, the procedure is invasive, time- and cost-consuming, and the nephrotoxic radiopaque contrast medium injected to map vessels is not free from possible harmful effects.

Nevertheless, patients have been carefully checked and examined for a long list of possible clinical and DUS signs of stenosis/thrombosis and findings have been confirmed by a double check. Despite this, however, due to the intrinsic aim of the study and the overall characteristics of the cohort, the possible influence of selection bias on results cannot be fully ruled out.

In conclusion, Qs has been found to be a more precise way than a simple Q or RI measurement for the rapid identification of a stenotic AVF and for stratifying the risk of a following thrombosis. Future, larger studies are needed to validate this parameter and to confirm whether such a simple measurement may really improve patterns of clinical practice by driving early re-vascularization or angioplasty before an overt thrombosis occurs.

REFERENCES

1. Riella MC, Roy-Chaudhury P. Vascular access in haemodialysis: strengthening the Achilles’ heel. Nat Rev Nephrol. (2013) 9:348–57. doi: 10.1038/nrneph.2013.76
2. Astor BC, Eustace JA, Powe NR, Klag MJ, Fink NE, Coresh J, et al. Type of vascular access and survival among incident hemodialysis patients: the choices for healthy outcomes in caring for ESRD (CHOICE) study. J Am Soc Nephrol. (2005) 16:1449–55. doi: 10.1681/ASN.2004090748
3. Arora P, Kausz AT, Obredor GT, Ruthazer R, Khan S, Jenulesen CS, et al. Hospital utilization among chronic dialysis patients. J Am Soc Nephrol. (2000) 11:740–6. Available online at: https://jasn.asnjournals.org/content/11/4/740.long
4. Coppolino G, Lucisano G, Bolignano D, Buemi M. Acute cardiovascular complications of hemodialysis. Minerva Urol Nefrol. (2010) 62:67–80.
5. Schwarz C, Mitterbauer C, Boczula M, Maza T, Funovics M, Heinze G, et al. Flow monitoring: performance characteristics of ultrasound dilution vs. color Doppler ultrasound compared with fistulography. Am J Kidney Dis. (2003) 42:539–45. doi: 10.1053/s0272-6386(03)00786-8
6. Brahmbhatt A, Remuzzi A, Franzoni M, Misra S. The molecular mechanisms of hemodialysis vascular access failure. Kidney Int. (2016) 89:303–16. doi: 10.1016/j.kint.2015.12.019
7. Solleymanian T, Sheik H, Tareh F, Argani H, Ossareh S. Hemodialysis vascular access failure. Kidney Int. (2017) 18:35–42. doi: 10.5301/jva.5000610
8. Gallieni M, Hollenbeck M, Inston N, Kumwenda M, Powell S, Tordoir J, et al. Clinical practice guideline on peri- and postoperative care of arteriovenous fistulas and grafts for hemodialysis in adults. Nephrol Dial Transplant. (2019) 34:i1–i2. doi: 10.1093/ndt/giz072
9. Hoeben H, Abu-Alfa AK, Reilly RF, Arunny JE, Bouman K, Perazella MA. Vascular access surveillance: evaluation of combining dynamic venous pressure and vascular access blood flow measurements. Am J Nephrol. (2003) 23:403–8. doi: 10.1159/000074297
10. Paulson WD, Mois L, Lok CE. Vascular access surveillance: an ongoing controversy. Kidney Int. (2012) 81:132–42. doi: 10.1038/ki.2011.337
11. White JJ, Ram SJ, Jones SA, Schwab SJ, Paulson WD. Influence of luminal diameters on flow surveillance of hemodialysis grafts: insights from a mathematical model. Clin J Am Soc Nephrol. (2006) 1:972–8. doi: 10.2215/CJN.00582006
12. Campos RP, Chula GC, Perreto S, Riella MC, do Nascimento MM. Accuracy of physical examination and intra-access pressure in the detection of stenosis in hemodialysis arteriovenous fistula. Semin Dial. (2008) 21:269–73. doi: 10.1111/j.1525-139X.2007.00419.x
13. Roca-Tey R, Ibeas I, Moreno T, Guuss E, Merino JL, Vallespin J, et al. Dialysis arteriovenous access monitoring and surveillance according to the 2017 Spanish Guidelines. J Vasc Access. (2018) 19:422–9. doi: 10.1177/1129729818761307
14. Brandt AH, Jensen J, Hansen KL, Hansen P, Lange T, Rix M, et al. Surveillancce for hemodialysis access stenosis: usefulness of ultrasound vector volume flow. J Vasc Access. (2016) 17:483–8. doi: 10.5301/jva.5000589
15. Malik J, Kudlicka J, Novakova L, Adamecz J, Malikova H, Kavan J. Surveillance of arteriovenous accesses with the use of duplex Doppler ultrasonography. J Vasc Access. (2014) 15 (Suppl. 7):S28–32. doi: 10.5301/jva.5000240
16. DeLong ER, DeLong DM, Clarke-Pearson DL. Comparing the areas under two or more correlated receiver operating characteristic curves: a nonparametric approach. Biometrics. (1988) 44:837–45. doi: 10.2307/253195
17. Ruopp MD, Perkins NJ, Whitcomb BW, Schisterman EF. Youden index and optimal cut-point estimated from observations affected by a lower limit of detection. Biom J. (2008) 50:419–30. doi: 10.1002/bimj.200710415
18. Harrell FE, Jr., Lee KL, Mark DB. Multivariable prognostic models: issues in developing models, evaluating assumptions and adequacy, and measuring and reducing errors. Stat Med. (1996) 15:361–87. doi: 10.1002/(SICI)1097-0258(19960229)15:4<361::AID-SIM168>3.0.CO;2-4
19. Rajabi-Jaghargh E, Banerjee RK. Combined functional and anatomical diagnostic endpoints for assessing arteriovenous fistula dysfunction. World J Nephrol. (2015) 4:6–18. doi: 10.5527/wjn.v4.i1.6
20. Brown RS, Patibandla BK, Goldfarb-Rumyantzev AS. The survival benefit of ”Fistula First, Catheter Last” in hemodialysis is primarily due to patient factors. J Am Soc Nephrol. (2017) 28:645–52. doi: 10.1681/ASN.201610019
21. Tessitore N, Lipari G, Poli A, Bedogna V, Baggio E, Loschiavo C, et al. Can blood flow surveillance and pre-emptive repair of subclinical stenosis prolong the useful life of arteriovenous fistula? A randomized controlled study. Nephrol Dial Transplant. (2004) 19:2325–33. doi: 10.1093/ndt/gfh316
22. Vascular Access Work G. Clinical practice guidelines for vascular access. Am J Kidney Dis. (2006) 48 (Suppl. 1):S248–73. doi: 10.1053/j.ajkd.2006.04.040
23. Gil Giraldó Y, Munoz Ramos P, Ruano P, Quiroga B. Vascular access-related mortality in hemodialysis patients during and after hospitalization. Ther Apher Dial. (2020) 24:688–94. doi: 10.1111/1744-9987.13479
24. Al-Balas A, Lee T, Young CJ, Kepes JA, Barker-Finkel J, Allon M. The clinical and economic effect of vascular access selection in patients initiating

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by Comitato Etico Calabria Centrale. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

AC and GC designed the study. DB and MP analyzed the data. GC had primary responsibility for the collection, analysis, and interpretation of the data and the final content. All authors contributed to drafting and writing the manuscript, read, and approved the final manuscript.
hemodialysis with a catheter. *J Am Soc Nephrol.* (2017) 28:3679–87. doi: 10.1681/ASN.2016060707

25. Ravani P, Palmer SC, Oliver MJ, Quinn RR, MacRae JM, Tai DJ, et al. Associations between hemodialysis access type and clinical outcomes: a systematic review. *J Am Soc Nephrol.* (2013) 24:465–73. doi: 10.1681/ASN.2012070643

26. Bradbury BD, Chen F, Furniss A, Pisoni RL, Keen M, Mapes D, et al. Conversion of vascular access type among incident hemodialysis patients: description and association with mortality. *Am J Kidney Dis.* (2009) 53:804–14. doi: 10.1053/j.ajkd.2008.11.031

27. Allon M, Daugirdas J, Depner TA, Greene T, Ornt D, Schwab SJ. Effect of change in vascular access on patient mortality in hemodialysis patients. *Am J Kidney Dis.* (2006) 47:469–77. doi: 10.1053/j.ajkd.2005.11.023

28. Leporini C, Pisano A, Russo E, G DA, de Sarro G, Coppolino G, et al. Effect of pentoxifylline on renal outcomes in chronic kidney disease patients: a systematic review and meta-analysis. *Pharmacol Res.* (2016) 107:315–32. doi: 10.1016/j.phrs.2016.03.001

29. Coppolino G, Leonardi G, Andreucci M, Bolignano D. Oxidative stress and kidney function: a brief update. *Curr Pharm Des.* (2018) 24:4794–9. doi: 10.2174/1381612825666190112165206

30. Girerd S, Girerd N, Frimat L, Holdaas H, Jardine AG, Schmieder RE, et al. Arteriovenous fistula thrombosis is associated with increased all-cause and cardiovascular mortality in haemodialysis patients from the AURORA trial. *Clin Kidney J.* (2020) 13:116–22. doi: 10.1093/ckj/sfz048

31. Shintaku S, Kawanishi H, Morishi M, Ago R, Banshodani M, Hashimoto S, et al. Postoperative day 1 access blood flow and resistive index can predict patency in distal forearm arteriovenous fistula. *J Vasc Access.* (2017) 18:371–8. doi: 10.5301/jva.5000777

**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

*Copyright © 2021 Colombo, Provenzano, Rivoli, Donato, Capria, Leonardi, Chiarella, Andreucci, Fusano, Bolignano and Coppolino. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.*