

## How Does Your PICCOMPARE? A Pilot Randomized Controlled Trial Comparing Various PICC Materials in Pediatrics

Tricia Kleidon, RN, MNursSci<sup>1,2\*</sup>, Amanda J. Ullman, RN, MAppSci, PhD<sup>2,3</sup>, Li Zhang, MBBS, PhD<sup>2</sup>, Gabor Mihala, MEng<sup>2,4,5</sup>, Brett Chaseling, MBBS (Hons), FANZCA<sup>1,6</sup>, Jason Schoutrop, BSc (Hons), MBBS, FANZCA<sup>1,6</sup>, Claire M. Rickard, RN, PhD<sup>2,7</sup>

<sup>1</sup>Children's Health Queensland, Lady Cilento Children's Hospital, South Brisbane, Queensland, Australia; <sup>2</sup>Alliance for Vascular Access Teaching and Research Group, Menzies Health Institute Queensland, Nathan Campus, Queensland, Australia; <sup>3</sup>School of Nursing and Midwifery, Griffith University, Nathan Campus, Queensland, Australia; <sup>4</sup>Centre for Applied Health Economics, Menzies Health Institute Queensland, Nathan Campus, Queensland, Australia; <sup>5</sup>School of Medicine, Griffith University, Queensland, Australia; <sup>6</sup>University of Queensland, St. Lucia, Queensland, Australia; <sup>7</sup>School of Nursing and Midwifery, Griffith University, Nathan Campus, Queensland, Australia.

**BACKGROUND:** Despite the popularity of peripherally inserted central catheters (PICCs), recent literature highlights their potential injurious complications. Innovative PICC materials have been developed to prevent thrombosis and infection formation (Endexo<sup>®</sup>) and antireflux valves to prevent occlusion (pressure-activated safety valve<sup>®</sup>). No large randomized controlled trial has assessed these technologies. Our primary aim was to evaluate the feasibility of a large randomized controlled efficacy trial of PICC materials and design to reduce PICC complication in pediatrics.

**METHODS:** A randomized controlled feasibility trial was undertaken at the Lady Cilento Children's Hospital in South Brisbane, Australia, between March 2016 and November 2016. Consecutive recruitment of 150 pediatric participants were randomly assigned to receive either (1) polyurethane PICC with a clamp or (2) BioFlo<sup>®</sup> PICC (AngioDynamics Inc, Queensbury, New York). Primary outcomes were trial feasibility, including PICC failure (thrombosis, occlusion, infection, breakage,

or dislodgement). Secondary outcomes were PICC complications during use.

**RESULTS:** Protocol feasibility was established, including staff and patient acceptability, timely recruitment, no missing primary outcome data, and 0% attrition. PICC failure was 22% (16 of 74, standard care) and 11% (8 of 72, BioFlo<sup>®</sup>) corresponding to 12.6 and 7.3 failures per 1000 hours (risk ratio 0.58; 95% confidence interval, 0.21-1.43;  $P = .172$ ). PICC failures were primarily due to thrombosis (standard care 7% versus BioFlo<sup>®</sup> 3%) and complete occlusion (standard care 7% versus BioFlo<sup>®</sup> 1%). No blood stream infections occurred. Significantly fewer patients with BioFlo<sup>®</sup> had PICC complications during use (15% vs 34%;  $P = .009$ ).

**CONCLUSION:** BioFlo<sup>®</sup> PICCs appear potentially safer for pediatrics than traditional standard care PICCs with a clamp. Further research is required to definitively identify clinical, cost-effective methods to prevent PICC failure and improve reliability. *Journal of Hospital Medicine* 2018;13:517-525. Published online first February 8, 2018.

Peripherally inserted central catheters (PICCs) have evolved since their inception in the early 1970s and are used with increasing frequency for pediatric inpatients and outpatients.<sup>1-3</sup> Emerging literature, including a meta-analysis of international observational studies,<sup>4</sup> reports PICC failure (complications necessitating premature removal)

occurs in up to 30% of PICCs, most commonly due to infection, thrombosis, occlusion, and fracture.<sup>4-7</sup> Raffini et al.<sup>7</sup> report the increasing incidence of pediatric PICC-related thrombosis increases morbidity and mortality<sup>8</sup> and negatively impacts future vessel health and preservation.<sup>9</sup>

PICCs have progressed from relatively simple, silicone-based catheters with an external clamp to chemically engineered polyurethane with pressure-activated valves placed at the proximal or distal catheter hub with the intent to reduce occlusion.<sup>10</sup> Further modernization of PICC material occurred with the incorporation of antithrombogenic (AT) material (Endexo<sup>®</sup>). These PICCs are designed to contain a nonstick polymer, which is designed to reduce the adherence of blood components (platelets and clotting factors) and inhibit thrombus formation (and hence prevent deep vein thrombosis and occlusion, as well as inhibit microbial biofilm attachment [and subsequent infection]).<sup>11</sup>

In addition to new materials, other aspects of this PICC design have been the addition of a pressure-activated safe-

**\*Address for correspondence:** Tricia Kleidon, Lady Cilento Children's Hospital, level 7, Department of Anaesthetics, 501 Stanley Street, South Brisbane, 4101, Qld, Australia; Telephone: +61(0)407175301; Fax: +61(0)730684419; E-mail: tricia.kleidon@health.qld.gov.au

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ty valve (PASV<sup>®</sup>) built into the proximal hub. Pressure-activated valve technology promises to prevent catheter occlusion by reducing blood reflux into the PICC; the valve opens with pressure during infusion and aspiration and remains closed with normal venous pressure, circumventing the need for clinicians to manually clamp the PICC and reducing human error and the potential for thrombosis, occlusion, and fracture development.<sup>12</sup> Hoffer et al.<sup>13</sup> reported half as many occlusions of valved PICCs (3.3%) compared with nonvalved or clamped PICCs (7.1%); although not statistically significant ( $P = .10$ ), perhaps due to the small sample, overall complications, including occlusion and infection, were significantly lessened with the valved PICC (35% vs 79%;  $P = .02$ ). Comparatively, Pittiruti et al.<sup>14</sup> conducted a trial of 2 types of valved PICCs with an open-ended, nonvalved PICC and found no reduction in PICC occlusion or catheter malfunction.

Today, PICC use is common for patients who require short-to-medium intravenous therapy. PICCs are increasingly recognized for their significant complications, including thrombosis and infection.<sup>15</sup> Novel PICC technology, including the incorporation of AT material such as Endexo<sup>®</sup> and PASV<sup>®</sup>, may reduce complications; however, the clinical efficacy, cost-effectiveness, and acceptability of these innovations have not been tested through randomized trials in pediatric patients. In accordance with Medical Research Council guidelines<sup>16</sup> for developing interventions, we pilot tested the feasibility of the BioFlo<sup>®</sup> PICC, including intervention acceptability, compliance, recruitment, and initial estimates of effect, in anticipation of a subsequent full-scale efficacy randomized controlled trial. Our secondary aim was to compare the effectiveness of the BioFlo<sup>®</sup> PICC with Endexo<sup>®</sup> and PASV<sup>®</sup> technology in reducing PICC complications and failure.

## METHODS

### Design

We undertook a pilot randomized controlled trial comparing the standard polyurethane PICC (with external clamp) with the BioFlo<sup>®</sup> PICC (with internal valve) in preventing catheter failure in pediatric patients. The study was prospectively registered with the Australian Clinical Trials Registry (ACTRN12615001290583), and the research protocol was published.<sup>17</sup>

### Study Setting

The study commenced in March 2016 at the Lady Cilento Children's Hospital in South Brisbane, Australia, a tertiary-level, specialist, pediatric teaching hospital in Queensland, Australia, providing full-spectrum health services to children and young people from birth to 18 years of age. Recruitment, including data collection, was completed in November 2016.

### Sample

The target sample size was 110 participants, 50 participants per group plus 10% for potential attrition, as determined by standard pilot trial sample size recommendations.<sup>18</sup> With ethics approval, the sample size was later increased to 150 participants in order to adequately pilot a microbiological substudy

method (published separately).<sup>17</sup> Participants were consecutively recruited if they met the inclusion criteria: PICC insertion, age <18 years, predicted hospital stay >24 hours, single-lumen PICC, and written informed consent by an English-speaking, legal parent or guardian. Patients were excluded if they had a current (<48 hours) blood stream infection (BSI), vessel size <2 mm, could not speak English without an interpreter, required a multilumen PICC, or were previously enrolled in the study.

### Interventions

Participants were randomized to receive either of the following PICCs: (1) standard care: Cook<sup>™</sup> polyurethane, turbo-ject, power-injectable PICC (Cook Medical, Bloomington, IN) or (2) comparison: BioFlo<sup>®</sup> polyurethane with Endexo<sup>®</sup> technology (AngioDynamics Inc, Queensbury, New York).

### Outcomes

The primary outcome was feasibility of a full-efficacy trial established by composite analysis of the elements of eligibility (>70% of patients will be eligible), recruitment (>70% of patients will agree to enroll), retention and attrition (<15% of participants are lost to follow-up or withdraw from the study), protocol adherence (>80% of participants receive their allocated, randomly assigned study product), missing data (<10% of data are missed during data collection), parent and healthcare staff satisfaction, and PICC failure effect size estimates to allow sample size calculations.<sup>18,19</sup> PICC failure was defined as the following complications associated with PICC removal: (1) catheter-associated BSI,<sup>8,20-22</sup> (2) local site infection,<sup>22</sup> (3) venous thrombosis,<sup>23</sup> (4) occlusion,<sup>24,25</sup> (5) PICC fracture, or (6) PICC dislodgement.<sup>25,26</sup> Parents (or caregivers) and healthcare staff were asked to rate their level of confidence with the study product and ease of PICC removal by using a 0 to 100 numeric rating scale (NRS) of increasing confidence and/or ease. These data were collected at the time of PICC removal. Operators were also asked to rate their levels of satisfaction with the insertion equipment and ease of PICC insertion immediately upon completion of the insertion procedure (both 0-100 NRS of increasing satisfaction and/or ease). Secondary outcomes included individual PICC complications (eg, occlusion) occurring at any time point during the PICC dwell (including at removal), adverse events, pain, redness at the insertion site, and overall PICC dwell.

### Study Procedures

The research nurse (ReN) screened operating theater lists for patients, obtained written informed consent, and initiated the randomization. Randomization was computer generated, and web based via Griffith University (<https://www151.griffith.edu.au/random>) to ensure allocation concealment until study entry. Patients were randomly assigned in a 1:1 ratio with computer-generated and randomly varied block sizes of 2 and 4. Data were collected by the ReN on the day of insertion, at day 1 postinsertion, then every 2 to 3 days thereafter so that PICCs were checked at least twice per week until study completion. Participants were included in the trial until 12 weeks post-PICC

insertion, study withdrawal or PICC removal (whichever came first), with an additional 48 hours follow-up for infection outcomes. Patient review was face to face during the inpatient stay, with discharged patients' follow-up occurring via outpatient clinics, hospital-in-the-home service, or telephone.

Data collection was via Research Electronic Data Capture (<http://project-redcap.org/>). The ReN collected data on primary and secondary outcomes by using the predefined criteria. Demographic and clinical data were collected to assess the success of randomization, describe the participant group, and display characteristics known to increase the risk of PICC complication and thrombosis. A blinded radiologist and infectious disease specialist reviewed and diagnosed thrombosis of deep veins and catheter-associated BSI outcomes, respectively.

### PICC Procedures

Extensive prestudy education for 2 months prior to trial commencement was provided to all clinicians involved with the insertion and care of PICCs, including the study products. PICCs were inserted in an operating theater environment by a qualified consultant pediatric anesthetist, a senior anesthetic registrar or fellow in an approved anesthetic training program, or pediatric vascular access nurse practitioner. Ultrasound guidance was used to assess a patient's vasculature and puncture the vessel. The operator chose the PICC size on the basis of clinical judgment of vessel size and patient needs and then inserted the allocated PICC.<sup>27</sup> Preferred PICC tip location was the cavoatrial junction. All PICC tip positions were confirmed with a chest x-ray before use.

Postinsertion, PICCs were managed by local interdisciplinary clinicians in accordance with local practice guidelines.<sup>27-31</sup> PICC care and management includes the use of 2% chlorhexidine gluconate in 70% alcohol for site antisepsis and neutral displacement needleless connectors (TUTA Pulse; Medical Australia Limited, Lidcombe, New South Wales, Australia); normal saline was used to flush after medication administration, and if the device was not in use for 6 hours or longer, heparin is instilled with securement via bordered polyurethane dressing (Tegaderm 1616; 3M, St Paul, Minnesota) and a sutureless securement device (Statlock VPPCSP; Bard, Georgia).

### Statistical Analyses

Data were exported to Stata 15<sup>32</sup> for cleaning and analysis. Data cleaning of outlying figures and missing and implausible data was undertaken prior to analysis. Missing data were not imputed. The PICC was the unit of measurement, and all randomly assigned patients were analyzed on an intention-to-treat basis.<sup>33</sup> Descriptive statistics (frequencies and percentages) were used to ascertain the primary outcome of feasibility for the larger trial. Incidence rates (per 1000 catheter days) and rate ratios, including 95% confidence intervals (CIs), were calculated. The comparability of groups at baseline was described across demographic, clinical, and device characteristics. Kaplan-Meier survival curves (with log-rank tests) were used to compare PICC failure between study groups over time. Associations between baseline characteristics and failure were described by

calculating hazard ratios (HRs). Univariable Cox regression was performed only due to the relatively low number of outcomes. *P* values of <.05 were considered statistically significant.

### Ethics

The Children's Health Service District, Queensland (Human Research Ethics Committee/15/QRCH/164), and Griffith University (2016/077) Human Research Ethics Committees provided ethics and governance approval. Informed consent was obtained from parents or legal guardians, with children providing youth assent if they were 7 years or older, dependent upon cognitive ability.

## RESULTS

### Participant and PICC Characteristics

Participant and PICC characteristics are described in Table 1. The majority of participant and PICC characteristics were balanced between intervention groups. The mean patient age was 7.3 years (standard deviation 5.0; range 0-18). PICC insertion was most commonly for a respiratory diagnosis (*n* = 98; 65%). Most PICCs were placed in the basilica vein (*n* = 115; 79%), with insertion being successful on the first attempt (*n* = 125; 86%). There was some imbalance (>10% absolute difference between groups) in nurse practitioner and registrar insertions (standard care 35% and 23% vs BioFlo® 51% and 8%, respectively) and patients with leucocytes <1000  $\mu$ l (standard care 10% vs BioFlo® 22%). Optimal PICC tip location at the cavoatrial junction was higher with BioFlo® than standard care, although this difference was <10%.

### Feasibility Outcomes

As shown in Figure 1, the majority of feasibility criteria were met, with 94% of 188 screened patients being eligible to participate and 97% of eligible patients consenting to enroll. Of 150 patients randomly assigned, 4 (1 in standard care and 3 in BioFlo®) were unable to have a PICC inserted or the procedure was cancelled. Demographic data only were collected for these 4 patients. No participants were lost to follow-up, and no primary outcome data were missing. Staff satisfaction with insertion kit and ease of insertion, ease of removal of the PICC, and parental confidence in the PICC product were similar across both groups (Table 2).

### PICC Failure and Complications

In total, 24 of 146 participants (16%) experienced PICC failure. There were 16 (22%) failures of standard care PICCs and 8 (11%) failures of BioFlo® PICCs. This corresponded to incident rates of 12.6 and 7.3 per 1000 catheter days (incident rate ratio 0.58; 95% CI, 0.21-1.43; *P* = .172; Table 2). Failure was most commonly from thrombosis (*n* = 5; 7%) or occlusion (*n* = 5; 7%) in the standard care group, with lower incidences in the BioFlo® group (*n* = 2 [3%] and *n* = 1 [1%], respectively). Figure 2 displays survival from PICC failure.

Considering the entire PICC dwell, of the 74 standard care patients, 49 (66%) had no complications, 9 (12%) had complications during the dwell but none at removal, 2 (3%) had no complications during the dwell but had a complication (ie, failure)

TABLE 1. Participant (n = 150) and PICC Characteristics (n = 144)

Participant characteristics	N	Standard Care	BioFlo®
		n = 75	n = 75
Age (years) <sup>a</sup>	150	7.5 (4.9) (0.0-18.0)	7.1 (5.1) (0.0-17.0)
Weight (kg) <sup>a</sup>	133	27.6 (16.2) (5.2-78.0)	28.4 (17.8) (4.9-70.0)
Sex (male)	149	41 (55)	45 (61)
Insertion on dominant side	101	17 (34)	16 (31)
Comorbidities:	150		
none		19 (25)	12 (16)
1		47 (63)	53 (71)
2 or more		9 (12)	10 (13)
Diagnosis:	150		
respiratory		51 (68)	47 (63)
medical		6 (8)	5 (7)
oncology		4 (5)	5 (7)
surgical		5 (7)	3 (4)
gastroenterology		1 (1)	5 (7)
haematology		0 (0)	1 (1)
other		8 (11)	9 (12)
Previous deep vein thrombosis	136	4 (6)	6 (9)
Leucocytes <1000/μl	112	6 (10)	12 (22)
Assistance to mobilise	150	7 (9)	4 (5)
Confused, agitated, or drowsy	150	3 (4)	4 (5)
Placement:	146	59 (80)	56 (78)
basilica		7 (9)	11 (15)
brachial			
cephalic		6 (8)	3 (4)
axilla		2 (3)	2 (3)
Subsequent insertion	138	40 (57)	36 (53)

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at removal, and 14 (19%) had complications during the dwell and at removal. For the 72 BioFlo® patients, 61 (85%) had no PICC complications, 3 (4%) had complications during the dwell but none at removal, 4 (5.5%) had no complications during the dwell but had a complication (ie, failure) at removal, and 4 (5.5%) had complications during the dwell and at removal.

More than twice as many standard care patients as BioFlo® patients had a complication during the PICC dwell, and this difference was statistically significant (25 of 74, 34% vs 11 of 72, 15%;  $P = .009$ ; Table 2). These results are consistent with the Kaplan-Meier curve, which shows longer complication-free survival with BioFlo® (Figure 2A and 2B). The median BioFlo® dwell was 1 day longer (13.8 vs 12.9 days), and the median time to first complication was 1 day later (4.0 BioFlo® vs 3.0 standard care; Table 2).

As per supplementary Table 1, univariate Cox regression identified PICC failure as significantly associated with tip placement in the proximal superior vena cava (SVC) compared to

the SVC–right atrium junction (HR 2.61; 95% CI, 1.17-5.82;  $P = .024$ ). Reduced risk of PICC failure was significantly associated with any infusion during the dwell (continuous fluid infusion,  $P = .007$ ; continuous antibiotic,  $P = .042$ ; or intermittent infusion,  $P = .046$ ) compared to no infusion. Other variables potentially influencing the risk of failure included PICC insertion by nurse specialist compared to consultant anesthetist (HR 2.61; 95% CI, 0.85-5.44) or registrar (HR 1.97; 95% CI, 0.57-6.77). These differences were not statistically significant; however, baseline imbalance between study groups for this variable and the feasibility design preclude absolute conclusions.

## DISCUSSION

This is the first pilot feasibility trial of new PICC materials and valve design incorporated in the BioFlo® PICC in the pediatric population. The trial incorporated best practice for randomized trials, including using a concurrent control group, centralized and concealed randomization, predetermined feasibility

TABLE 1. Participant (n = 150) and PICC Characteristics (n = 144) (continued)

Participant characteristics	N	Standard Care	BioFlo®
		n = 75	n = 75
Multiple insertion attempts:	146		
1 (success at first insertion)		60 (81)	65 (90)
2		11 (15)	5 (7)
3 or more		3 (4)	2 (3)
Ultrasound guidance used	146	74 (100)	72 (100)
Catheter tip location:	146	52 (70)	57 (79)
SVC–RA junction		22 (30)	15 (21)
SVC			
Vein size (mm) <sup>b</sup>	103	3.7 (0.9)	3.1 (0.9)
Catheter size:	146		
3 Fr		57 (77)	58 (81)
4 Fr		17 (23)	14 (19)
Catheter-to-vein ratio (%; n = 103) <sup>c</sup>	103	32.3 (27.6-35.0)	36.2 (30.9-43.7)
Treatment <sup>d</sup> :			
intermittent bolus medication	146	63 (84)	61 (85)
cont. antibiotic infusion	146	27 (36)	31 (43)
intermittent infusion	146	23 (31)	27 (38)
cont. nonantibiotic infusion	146	11 (15)	15 (21)
Thrombolytic treatment	146	4 (5)	5 (7)
Number of blood samples <sup>b</sup>	146	0.57 (0.50)	0.71 (0.70)
Infection at recruitment:			
respiratory	150	31 (41)	30 (40)
wound	150	2 (3)	7 (9)
osteomyelitis	150	3 (4)	2 (3)
positive BC (48 hours prior)	150	2 (3)	2 (3)
shunt, cerebrospinal fluid	150	1 (1)	1 (1)
urinary	150	1 (1)	0 (0)
other	150	4 (5)	9 (12)

<sup>a</sup> Mean, standard deviation, and range shown.

<sup>b</sup> Mean and standard deviation.

<sup>c</sup> Median, interquartile range (shown as 25th and 75th percentiles) and maximum shown.

<sup>d</sup> Ever received, multiple treatment types possible per patient.

NOTE: Frequencies (column percentages) are shown unless otherwise noted. Percentages are calculated with the number of nonmissing observations in the denominator. Abbreviations: BC, blood culture; cont, continuous; Fr, French gauge size; N, number of nonmissing observations; PICC, peripherally inserted central catheter; RA, right atrium; SVC, superior vena cava; tx, therapy;  $\mu$ l, microliter.

criteria, and a registered and published trial protocol.<sup>17</sup> As in other studies,<sup>15,24,34</sup> PICC failure and complication prevalence was unacceptably high for this essential device. Standard care PICCs failed twice as often as the new BioFlo® PICCs (22% vs 11%), which is a clinically important difference. As researchers in a pilot study, we did not expect to detect statistically significant differences; however, we found that overall complications during the dwell occurred significantly more with the standard care than BioFlo® PICCs ( $P = .009$ ).

BioFlo® PICC material offers a major advancement in PICC material through the incorporation of AT technologies into

catheter materials, such as PICCs. Endexo® is a low molecular-weight, fluoro-oligomeric additive that self-locates to the top few nanometers of the material surface. When added to power-injectable polyurethane, the additive results in a strong but passive, nonstick, fluorinated surface in the base PICC material. This inhibits platelet adhesion, suppresses protein procoagulant conformation, and thereby reduces thrombus formation in medical devices. Additionally, Endexo® is not a catheter coating; rather, it is incorporated within the polyurethane of the PICC, thereby ensuring these AT properties are present on the internal, external, and cut surfaces of the PICC. If this tech-

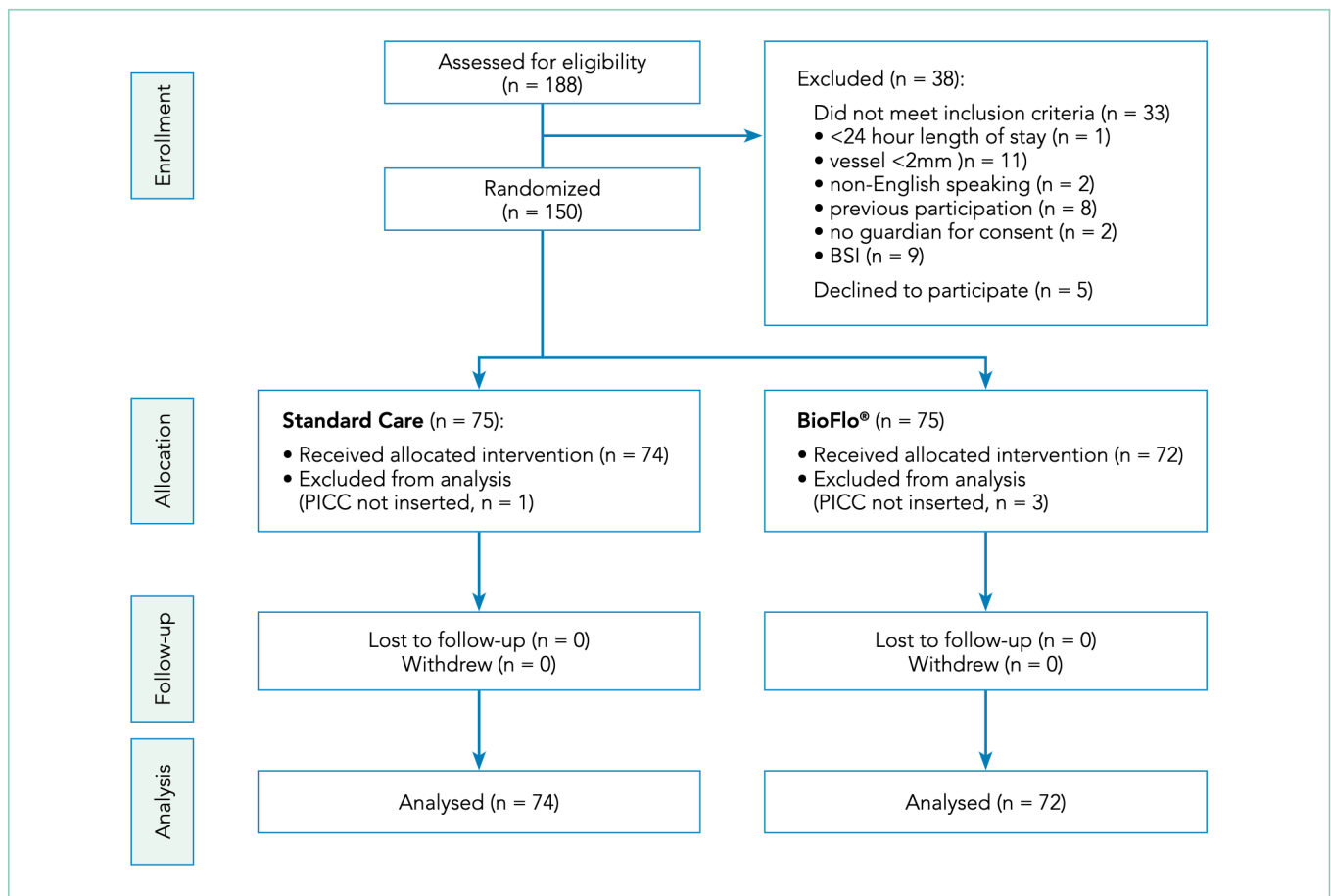


FIG 1. CONSORT Flowchart of study participants

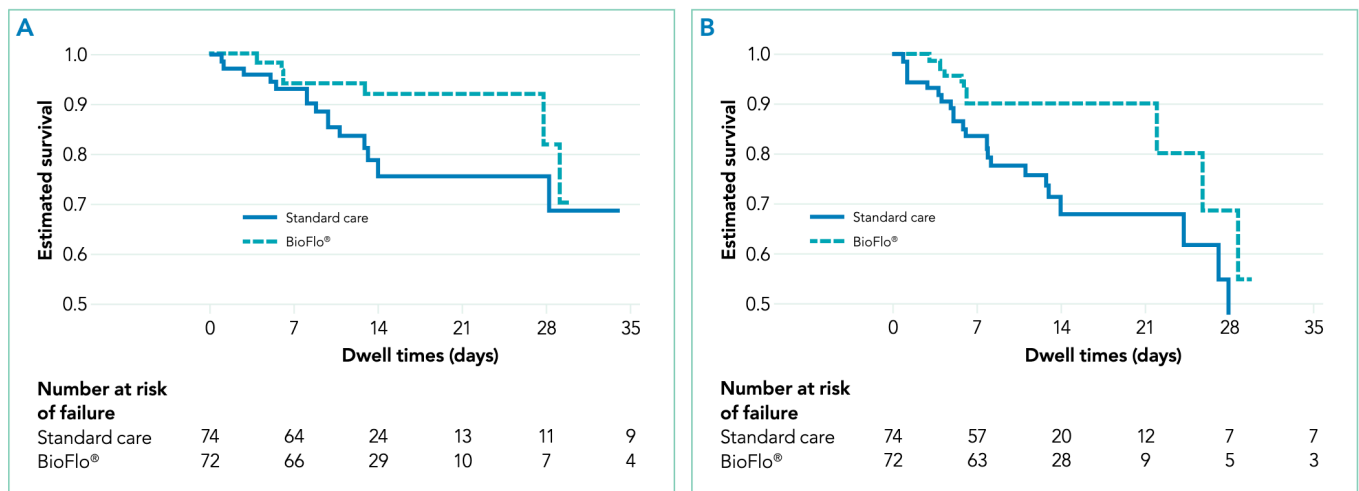


FIG 2. (A) Kaplan-Meier curves of PICC failure. (B) Kaplan-Meier curves of PICC complication

nology can reduce complication during treatment and reduce failure from infection, thrombosis, occlusion, fracture, and dislodgement, it will improve patient outcomes considerably and lower health system costs. Previous studies investigating valve technology in PICC design to reduce occlusion have been inconclusive.<sup>12-14,35,36</sup> Occlusion (both partial and complete) was less frequent in our study with the BioFlo® group (n = 3; 4%)

compared to the standard care group (n = 6; 8%). The results of this pilot study suggest that either the Endexo® material or PASV® technology has a positive association with occlusion reduction during PICC treatment.

Thrombosis was the primary failure type for the standard care PICCs, comprising one-third of failures. All but one patient with radiologically confirmed thrombosis required the removal of the

TABLE 2. Study Outcomes

Outcomes	Standard Care n = 74	BioFlo® n = 72	P Value
Failure necessitating removal	16 (22)	8 (11)	.087 <sup>a</sup>
Catheter-days	1268	1097	
IRR (per 1000 days, 95% CI)	Referent	0.58 (0.21-1.43)	.172 <sup>b</sup>
Dwell time (days) <sup>c</sup>	12.9 (9.0-14.1; 104)	13.8 (10.0-17.3; 44)	
Complications resulting in failure <sup>d</sup> :			
CVAD-associated thrombosis	5 (7)	2 (3)	
occlusion, complete	5 (7)	1 (1)	
occlusion, partial	3 (4)	2 (3)	
CVAD breakage	3 (4)	0 (0)	
dislodgement, complete	0 (0)	2 (3)	
dislodgement, partial	1 (1)	0 (0)	
local infection, confirmed	0 (0)	1 (1)	
CVAD-associated BSI	0 (0)	0 (0)	
Complications during treatment <sup>d</sup> :			
any, per patient	25 (34)	11 (15)	.009 <sup>a</sup>
occlusion, partial	11 (15)	5 (7)	
occlusion, complete	10 (14)	2 (3)	
thrombosis	6 (8)	2 (3)	
CVAD breakage	3 (4)	0 (0)	
dislodgement, complete	0 (0)	2 (3)	
dislodgement, partial	2 (3)	0 (0)	
local infection, confirmed	0 (0)	1 (1)	
CVAD-associated BSI	0 (0)	0 (0)	
Pain (at any time)	9 (12)	4 (6)	
Redness (at any time)	8 (11)	1 (1)	
Time to first complication (days, n = 36) <sup>e</sup>	7.8 (4.0-13; 92)	6.1 (4.2-26; 38)	
Ease of insertion (0 = worst, 100 = best) <sup>f</sup>	85.2 (16.0)	88.3 (14.9)	
Satisfaction with ins. kit (0 = lowest, 100 = highest) <sup>f</sup>	86.2 (12.2)	93.6 (7.7)	
Difficulty of removal (0 = worst, 100 = best) <sup>f</sup>	91.3 (5.0)	90.6 (7.1)	
Confidence in product (0 = lowest, 100 = highest) <sup>f</sup>	84.4 (14.2)	89.3 (7.2)	

<sup>a</sup> Chi-square test.<sup>b</sup> Log-rank test.<sup>c</sup> Median and interquartile range (25<sup>th</sup> and 75<sup>th</sup> percentiles) shown.<sup>d</sup> More than 1 complication per patient possible.<sup>e</sup> Maximum value shown.<sup>f</sup> Mean (and standard deviation) shown.

NOTE: Frequencies (and column percentages) are shown unless otherwise noted. Percentages are calculated with the number of nonmissing observations in the denominator. Abbreviations: BSI, bloodstream infection; CI, confidence interval; CVAD, central venous access device; ins, insertion; IRR, incidence rate ratio; PICC, peripherally inserted central catheter.

PICC prior to completion of treatment. The decision to remove the PICC or retain and treat conservatively remained with the treating team. Raffini et al.<sup>7</sup> found thrombosis to increase in patients with one or more coexisting chronic medical condition. Slightly more standard care than BioFlo® patients were free of such comorbidities (25% vs 16%), yet standard care patients still had the higher number of thromboses (7% vs 3%). Morgenthaler

and Rodriguez<sup>37</sup> reported vascular access-associated thrombosis in pediatrics to be less common than in adults but higher in medically complex children. Worryingly, Menendez et al.<sup>38</sup> reported pediatric thrombosis to be largely asymptomatic, so the true incidence in our study is likely higher because only radiologically confirmed thromboses were recorded.

Occlusion (partial or complete) was the predominant com-

plication across the study, being associated with one-third of all failures. When occlusion complications during the dwell (some of which were resolved with treatment), in addition to those causing failure, were considered, this number was even greater. Occlusion complications are prevalent and costly. Smith et al.<sup>24</sup> reported that occlusion was the most common reason for PICC removal and the most likely complication to delay treatment. Both the BioFlo<sup>®</sup> and standard care PICCs are pressure rated with good tensile strength; however, fracture occurred in 4% (n = 3) of standard care PICCs compared to no fractures in BioFlo<sup>®</sup> PICCs. Although the numbers are small, it may suggest a superior tensile strength of the BioFlo<sup>®</sup> material.

This study reinforces previously published results<sup>24,38</sup> that PICC tip position is important and can influence complications, such as occlusion and thrombosis. In addition, we found a significant association with failure when PICCs did not have a continuous infusion. These findings reinforce the need for optimal tip location at insertion and ongoing flushing and maintenance of PICCs not used for infusions.

Limitations of this study include the small sample size, which was not designed to detect statistical differences in the primary outcome between groups. Despite randomization, there were slight imbalances at baseline for inserter type and leukocyte count, although these were not significantly associated with PICC failure in the Cox regression (data not shown), and thus were unlikely to influence findings. Additionally, a difference of <10% was associated with PICC tip position, favoring the BioFlo<sup>®</sup> group. PICC tip position outside the cavoatrial junction was positively associated with failure; therefore, the effect of tip positioning on outcomes is difficult to ascertain given the small sample size and feasibility nature of the study. Further study is warranted to further explore this effect. The population sampled was pediatric medical and surgical inpatients with a vessel size >2 mm attending the operating theater suite for PICC insertion, thereby limiting the study's generalizability to adults and other populations, including neonates and those with PICCs inserted in the pediatric intensive care unit. The

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study could not be blinded because study products had to be visible to the clinical and research staff. However, it is unlikely that staff would intentionally sabotage PICCs to bias the study. Blinding was possible for the assessment of blood culture and ultrasound reports to diagnose infection and thrombosis. Strengths of this study included 100% protocol adherence, and no patients were lost to follow-up.

## CONCLUSION

These results confirm that PICC failure is unacceptably high and suggest that the innovative BioFlo<sup>®</sup> PICC material and design holds promise to improve PICC outcomes by reducing complications and overall PICC failure. Trials of this technology are feasible, safe, and acceptable to healthcare staff and parents. Further trials are required, including in other patient populations, to definitively identify clinical, cost-effective methods to prevent PICC failure and improve reliability during treatment.

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