We are IntechOpen, the world's leading publisher of Open Access books Built by scientists, for scientists



186,000

200M



Our authors are among the

TOP 1% most cited scientists





WEB OF SCIENCE

Selection of our books indexed in the Book Citation Index in Web of Science™ Core Collection (BKCI)

# Interested in publishing with us? Contact book.department@intechopen.com

Numbers displayed above are based on latest data collected. For more information visit www.intechopen.com



# Current Status of Synthetic and Biological Grafts for Hemodialysis

 Purav P. Patel<sup>1</sup>, Maria Altieri<sup>2</sup>, Tarun R. Jindal<sup>3</sup>, Steven R. Guy<sup>4</sup>, Edward M. Falta<sup>5</sup>, Eric A. Elster<sup>5</sup>, Frank P. Hurst<sup>5</sup>, Anton N. Sidawy<sup>2</sup> and Rahul M. Jindal<sup>2,5</sup> <sup>1</sup>Medical University of Lublin <sup>2</sup>George Washington School of Medicine <sup>3</sup>Indiana University School of Medicine <sup>4</sup>Department of Surgery, Drexel University, Philadelphia <sup>5</sup>Walter Reed Army Medical Center, Washington DC <sup>1</sup>Poland <sup>2,3,45</sup>USA

# 1. Introduction

Patients with chronic kidney disease (CKD) stage 4 or 5 have the option of kidney transplantation, hemodialysis (HD), peritoneal dialysis (PD), or conservative management.<sup>1</sup> National Kidney and Urologic Disease Information Clearinghouse reported that in 2007, there were 368, 544 U.S. residents with ESRD who were receiving dialysis, of whom 341, 264 were undergoing HD.<sup>2</sup> Quality of life and long-term survival of patients with CKD who are on HD depends on the successful placement of vascular access, as autogenous arteriovenous access, prosthetic arteriovenous access, or tunneled central venous catheter. DOQI guidelines are emphatic that autogenous arteriovenous access placement should be considered first, as it provides the optimal vascular access, followed by prosthetic grafts if autogenous arteriovenous access placement is not possible. There is a great deal of controversy regarding the choice of synthetic or biological material, as the guidelines suggest that it should be based on surgeon's experience and preference. The evidence to support the superiority of tapered versus uniform tubes, thick- versus thin- walled characteristics, elastic versus non-elastic arterial, stretch vs. standard PTFE, externally supported vs. unsupported grafts, is still evolving.

An ideal vascular graft would have the following characteristics: 1) appropriate size to match host vessels, 2) mechanical strength, 3) low thrombogenicity/ complete endothelialization, 4) rapid/ complete healing, 5) ease of handling, 6) resistance to infection, 7) structural durability in face of repeated needle puncture, 8) low incidence of hyperplastic intimal changes and 9) low cost.<sup>3</sup>

In this chapter, we will review the development of vascular grafts over the years and discuss the advantages of one over the other.

# 2. Prosthetic grafts for hemodialysis

Although the first synthetic graft used for HD access in the United States was made of Dacron dating back to the early 1970s, unfavorable results and the availability of better prosthetic materials like expanded PTFE (ePTFE) forced its abandonment. In 1976, Dr. Baker presented the first results of ePTFE grafts in 72 HD patients. Since then ePTFE remains the graft of choice for vascular access.<sup>4</sup> ePTFE is considered the material of choice due to the fact that it is readily accessible, ease of implantation, good medium term patency, and relatively low complication rates compared to other synthetic and biological materials.<sup>5</sup> The medical community has made strenuous efforts to increase the use of autogenous arteriovenous access, prevalence has increased from 22% in 1995 to 57.7% in 2011. However, the use of synthetic grafts still remains significant (18.4% prevalence in 2011).<sup>6,7</sup>

# 2.1 Indications for use of prosthetic grafts

Autogenous arteriovenous access are clearly superior in terms of long term patency to grafts, but not feasible in many patients undergoing HD<sup>8</sup>. The indications for prosthetic grafts include: lack of suitable vessels particularly in elderly and diabetic patients, need for immediate cannulation and in children who cannot tolerate multiple painful venipunctures.<sup>5</sup>

# 2.2 Complications associated with prosthetic grafts

Graft failures are typically caused by stenosis (leading cause of graft failure) due to thrombosis and neointimal hyperplasia at site of anastamosis, as well as graft infection (contributes to 10-15% of graft failure).<sup>9,10</sup> Other less common complications of prosthetic accesses include; steal syndrome, seromas, aneurysm formation, central vein stenosis and bleeding.<sup>11</sup> Thrombosis seems to occur soon after implantation due to technical problem, with a clot typically forming at the surface of the graft when it is first exposed to blood. The clot is formed initially of platelet aggregates, and then fibrin and thrombin is laid down via activation of the coagulation cascade. Platelet activity is generally most intense during the first 24 hours and subsides to a very low level after 1 week.<sup>9</sup> Neointimal hyperplasia in prosthetic conduits can be attributed to upstream and downstream events. The upstream factors include; hemodynamic stress at the graft-vein anastamosis, compliance mismatch between the graft material and vein (more studies required to establish this factor), arterial injury at the time of graft placement, intrinsic properties of the synthetic graft itself (shown to attract macrophages which then secrete specific cytokines bFGF, PDGF, and VEGF), graft injury from dialysis needles, as well as the presence of uremia (causing endothelial dysfunction even prior to synthetic graft implant).<sup>12-14</sup> The downhill events are essentially a consequence of the upstream events. Pro-inflammatory cells release cytokines promoting the migration of smooth muscle cells and myofibroblasts from the adventitia media into the intimal layer, where they proliferate and cause lesions of neointimal hyperplasia.<sup>13</sup> These stenotic lesions are usually treated by percutaneous angioplasty (PTA) or open patch angioplasty, which unfortunately, predisposes the patient to restenotic lesions due to endothelial and smooth muscle cell injury.<sup>15</sup>

Infection is the second most common complication of synthetic grafts and can lead to further complications such subacute bacterial endocarditis, epidural or brain abscess.<sup>11</sup> These complications can lead to graft failure in up to 35% of patients.<sup>16</sup> Graft infections have an incidence rate as high as 2%, and are 4 times as prevalent in synthetic grafts when compared to autogenous veins.<sup>9</sup> Common causative organisms are Staphylococcus aureus (26.32% of

Cumulative patency of  $67\% \pm 6\%$  (12 mo),  $50\% \pm 7\%$  (24 mo), In PTFE-silicone group, mean Bleeding after needle removal after early and late punctures time to first use was 1.3 days index (GPAI) of CL graft was 7- yr cumulative patency 55% was significantly decreased Graft platelet accumulation of PTFE-silicone grafts vs. conventional PTFE grafts. conventional PTFE grafts Comments significantly reduced. vs. 2 to 4 weeks for 43% ± 9% (48 mo). (p < 0.001).7.12%; Perigraft hematoma 2.01%; Graft thrombosis (21%), infection pseudoaneurysm formation (8%), (25%), or stenosis by neointimal (>6 weeks) 17.52%; Infection Early thrombosis (<6 weeks) Pseudoaneurysm 0% vs. 17.1 Thrombosis 36.6% vs. 28.5%; Complications Infection 13.3% vs. 11.4% 9.58%; Late thrombosis N/Asteal syndrome (3%) Steal syndrome 15 hyperplasia (34%), Seroma 3% vs. 0% Steal 3% vs. 0% 75% vs. 67% Secondary patency at 1 yr N/AN/AN/APATENCY ASSISTED PRIMARY N/AN/AN/AN/A**Primary Patency** 63% vs. 66% at 1 yr N/AN/AN/AAUTHOR COUNTRY YEAR NO. AND GRAFT TYPE vs. 20 control PTFE vs. 12 control PTFE high porous grafts 548 Impra® grafts 65 (30 Vasutek vs. 35 control PTFE 67 (55 Gore-Tex®, 12 Impra® grafts) carbon lined (CL) Experiment 2: 12 Experiment 1: 20 grafts) grafts grafts 19891992 1983 1983 Japan USA USA USA et al.<sup>49,50</sup> Tsuchida Schanzer Johnson et al.<sup>47</sup> Munda et al.<sup>29</sup> et al.<sup>48</sup>

Comments	Stretch ePTFE has better primary patency rates and less stenosis due to intimal hyperplasia vs. standard ePTFE grafts.	Graft survival in late camulation group (beyond 14 days) was 93, 92, and 91% vs. 80, 78 and 73% for the early camulation group (earlier than 14 days) at 1, 2, and 3 yr. respectively (p=0.0008).	Lower extremity AV dialysis accesses are associated with multiple complications and should be placed only if significant patient morbidity can be accepted and justified.	Cumulative primary patency estimates for early camulation were 89%, 82%, and 70% vs. 86%, 78% and 74% for the late camulation group at 3, 6, and 12 mo. respectively. Early camulation of prosthetic dialysis grafts did not increase perioperative morbidity rates or decrease 12-month cumulative primary patency rates.	
Complications	Thrombotic events: 40% (standard) vs. 12% (stretch)	Thrombosis: 14% of early cannulation group vs. 7% in late cannulation group. Infection: 7% early cannulation group vs. 1% late cannulation group.	Thrombosis (33%) graft infection (18%), distal limb ischemia (16%), aneurysmal dilatation of the graft requiring revision (7%), and fistula-induced congestive heart failure symptoms (4%)	Thrombosis occurred before cannulation in 2.0% of early cannulation and 3.2% of late cannulation group.	
Secondary patency	N/A	N/A	N/A	N/A	
ASSISTED PRIMARY PATENCY	N/A	N/A	N/A	N/A	
Primary Patency PATENCY	59% vs. 29% (1 yr)	N/A	59% (12 mo) 47% (24 mo)	N/A	
COUNTRY YEAR NO. AND GRAFT TYPE	37 (17 stretch ePTFE vs. 20 standard ePTFE grafts)	270 Gore-Tex® stretch graft (96 cannulated within 14 days, 174 cannulated after 14 days)	45 lower extremity access grafts (39 ePTFE, 6 modified bovine heterographs)	79 stretch ePTFE grafts (48 underwent early cannulation and 31 late cannulation)	
YEAR	1995	1996	1996	1997	
COUNTRY	Netherlands 1995	NSA	NSA	USA	
AUTHOR	Tordoir et al. <sup>20</sup>	Dawidson et al <sup>22</sup> .	Taylor et al. <sup>51</sup>	Hakaim et al. <sup>21</sup>	

prostheses reduces downstream Infection rate (16%) vs. forearm hrombectomy or in the number performance of 6-mm standard number of days before the first of thrombectomies or revisions nanufacturer, whether Gore-No difference between Gore-Mean survival time for thigh **TDMAC-heparin** coating of ePTFE grafts on the basis of PTFE grafts was 84.6 weeks Tex® and Impra® in the performed better than Comments Non-reinforced PTFE PTFE microvascular No difference in the infection rates (20%). per graft (P > 0.50). Tex® or Impra®. reinforced PTFE nicroemboli. (SD 76.1). mean of 2.45 thrombosis/graft with 1.09 revisions/graft required 1.20 revisions/graft, while non-reinforced Impra® only had a Reinforced Gore-Tex® had a mean Steal Syndrome: 6% in each group Mean number of emboli during a control group, 84 for the heparin-20-minute period was 91 for the Infection: 11% Impra® and 14% irrigated group, and 22 for the of 2.75 thrombosis/graft and Ccomplications was similar Complications Infection: 16% [DMAC-heparin group. between the two grafts. Thrombosis Gore-Tex® Gore-Tex® 41%) 49%, Gore-Tex® mpra® grafts at 69%) and at 2 yr between Gore-Impra® 33%, Secondary nonreinforced 2 yr (P > 0.13)No difference yr (Impra® patency N/AN/AFex® and reinforced (p = 0.15)at 1 year vs. 77% 80% ASSISTED Primary Patency PRIMARY PATENCY N/AN/AN/AN/AN/AImpra® grafts at 2 yr (P > 0.53) 43%, Gore-Tex® 47%) and at 2 yr mpra® grafts at Gore-Tex® 26%) Greater primary nonreinforced between Gore-Impra® 30%, No difference yr (Impra® yr. (p=0.02) N/AN/Apatency for [ex® and (p = 0.78)NO. AND GRAFT and 64 Gore-Tex® i male rats per test Heparin irrigated neparin irrigated) reinforced Gore-29 (65 Impra® group (control, 632 grafts (438 mpra® grafts) nonreinforced 90 (100 Gore-74 thigh PTFE mpra® grafts and TDMAC TYPE Tex® vs. 194 lex® vs. 90 grafts grafts AUTHOR COUNTRY YEAR 1997 1997 1997 1997 1998 Australia USA USA USA USA Schuman et al<sup>23</sup> Kaufman Hurlbert et al.<sup>18</sup> Khadra et al.<sup>19</sup> et al.<sup>53</sup> et al.<sup>52</sup> Ritter

Current Status of Synthetic and Biological Grafts for Hemodialysis

Comments	Luminal surface of PU grafts took 4 weeks to completely endothelialize, whereas ePTFE grafts took 24 weeks (P <.05).	Geometric design of the new graft-end was based on the reduction of three time- and area-averaged hemodynamic parameters (including the wall shear stress gradient, wall shear stress angle gradient, and radial pressure gradient).		
Complications	Neointimal cell proliferation lower in PU grafts compared with ePTFE at 56 days (1.4 +/- $0.1\mu$ versus 8.6 +/- $1.5\mu$ , P <.001) and at 6 mo (0.15 +/- $0.002\mu$ versus $3.4$ +/- $0.5\mu$ , p <.001). Neointimal thickness at 6 mo after implantation was $3.2$ +/- 0.8 $\mu$ for PU compared with $10.3$ +/- $3.1\mu$ for ePTFE (P <.05).	₽/N	Thrombotic occlusion lower in cuff group (0.68 ppy) than in control (0.88 ppy) (p=0.0007) Stenosis- CG: 0.22 ppy vs. control: 0.14 ppy; Infection- CG: 0.06 ppy vs. control: 0.01 ppy; Hemorrhage- CG: 0.10 ppy vs. control: 0.01 ppy; Pseudoaneurysm- CG: 0.04 ppy vs. control: 0.02 ppy; Ischemia- CG: 0.04 ppy vs. control: 0.00 ppy; Venous hypertension- CG: 0.00 ppy vs. control: 0.01 ppy; Seroma- CG: 0.04 ppy vs. control: 0.04 ppy vs. control: 0.01 ppy; Seroma- CG: 0.04 ppy vs. control: 0.04 ppy	
Secondary natencv	N/A	N/A	Control: 76%, 62%, 52%, 43%, Control: 86%, vs. cuff group: group: 71%, 89%, 81%, 76%, 60%, 53%, and 65% and 41% at 6 mo, 18 mo, 2 at 6 mo, 18 mo, 2 mo, 2 yr respectively (p=0.53)	
ASSISTED PRIMARY	N/A	N/A	Control: 76%, 62%, 52%, 43% ws. cuff group: 71%, 60%, 53%, and 41% at 6 mo, 18 mo, 2 yr respectively (p=0.53)	
Primary Patency	N/A	N/A	Control: 69%, 56% , 42%, 34% vs. cuff group: 62%, 43%, 30%, and 19% at 6 mo, 18 mo, 2 yr respectively (P = .097)	
AUTHOR COUNTRY YEAR NO. AND GRAFT	69 [37 polycarbonate polyurethane (PU) vs. 32 ePTFE] grafts into abdominal aortas of male Sprague- Dawley rats	Computer Hemodynamic Analysis of Venaflo® II	120 (59 cuffed PTFE vs. 61 standard PTFE grafts)	
YEAR	1999	2000	2000	
COUNTRY	USA	USA	Netherlands 2000	
AUTHOR	Jeschke et al <sup>54</sup>	Longest et al. <sup>25</sup>	Lemson et al. <sup>24</sup>	

Current Status of Synthetic and Biological Grafts for Hemodialysis

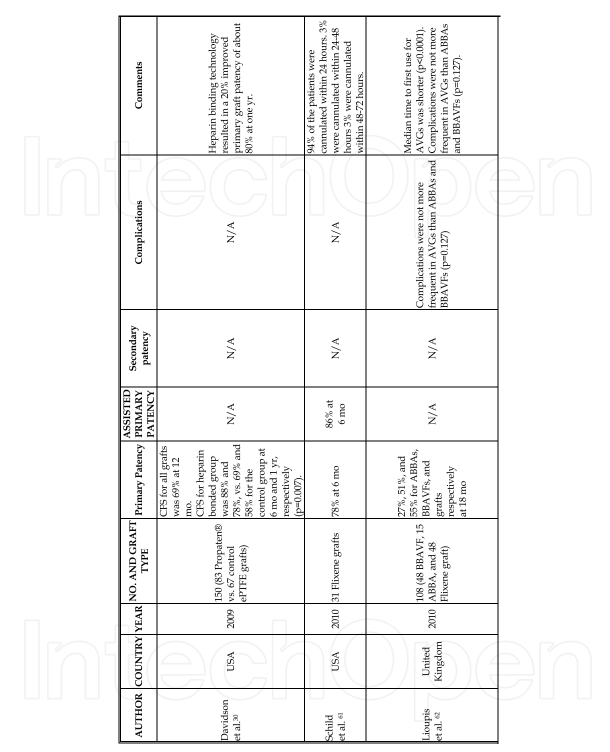
Comments	At ≥5 minutes, more PUU cannulation sites achieved hemostasis compared with ePTFE sites (P <.0001). 53.9% of all PUU grafts were cannulated before 9 days versus none with the ePTFE grafts (P <.001).	Cuffed ePTFE graft provided stable blood flow and satisfactory graft patency during 2 yr of follow-up, even in high risk patients with a prior history of vascular access thrombosis.	Graft patency at 12 mo. was 64% vs. 32% (P = 037) and 58% vs. 21 vs. 21 mo. (P = 0213) for Venaflo® II and Gore-Tex® respectively. Incidence of graft failure was lower in the cuffed ePTFE graft group (P = 039).	
Complications	Higher incidence of graft kinking in PUU group (p<0.001) Infection, anastomotic obstruction or thrombosis, pseudoaneurysm, kinking, stenosis, wound healing	No early postoperative complications. One graft was lost to thrombosis in the first yr; two grafts were lost to thrombosis in the second yr.	Graft-vein anastamosis stenosis: 15% Venaflo® II vs. 41% Gore-Tex®	
Secondary patency	87% vs. 90% (6 mo) and 78% vs. 80% (12 mo). (P >.05).	Graft patency rates were 90.9% at 1 yr and 68.2% at 2 yr	Significant improvement observed in the Venaflo® II group compared with the Gore- Tex® group at both 12 months (+32%, P = .037) and 24 months +37%, P = .021).	
ASSISTED PRIMARY PATENCY	N/A	N/A	N/A	
ASSISTED ASSISTED Primary Patency PATENCY	55% vs. 47% (6 mo) and 44% vs. 36% (12 mo) (P >.05).	N/A	A significant advantage in primary graft patency was observed with the Venaflo® II at 12 months after placement (P < 01).	
COUNTRY YEAR NO. AND GRAFT TYPE	142 (71 Vectra® vs. 71 Gore-Tex® or Impra® grafts)	12 Venaflo® II grafts	48 (24 Venaflo® II vs. 24 Gore-Tex® grafts)	
YEAR	2001	2001	2002	
	USA	USA	NSA	
AUTHOR	Glickman et al. <sup>35</sup>	Nyberg et al. <sup>27</sup>	Sorom et al. <sup>26</sup>	

Comments	Patency rate for upper extremity AVF in adults is superior to that for PTFE counterparts, although the overall quality of the studies in the meta-analysis was less than ideal.	PVG is an acceptable alternative to the stretch ePTFE.	PVG were first cannulated at a median time of 19 days after implantation, with 12% used within 3 days.	8% of preimplantation heparin activity remained on heparin grafts after 2 hours and only 2% after 7 days.	Patients implanted with a PU graft were dialyzed within hours after surgery.	
Complications	Y/N	78.7% vs. 79.9% Thrombosis: 26.7% vs. 35.7%;   78.7% vs. 79.9% Stenosis: 23.3% vs. 28.6%;   (1 yr); 78.7% vs. Infection: 6.7% vs. 14.3%; Seroma:   69.3% (2 yr) 0% vs. 3.6%; False aneurysm: 0%   PVG and Gore- vs. 3.6%; Kinking: 3.3% vs. 3.6%;   Tex® stretch Arterial Steal: 0% vs. 3.6%;   graft respectively PVG and Gore-Tex® stretch graft	32.7% graft infection rate; 30% graft median time of 19 days after thrombosis rate. Infection caused implantation, with 12% used 61.5% of graft loss within 3 days.	₽/N	N/A	
Secondary patency	86% vs. 76% (6 mo) 77% vs. 55% (18 mo)	78.7% vs. 79.9% (1 yr); 78.7% vs. 69.3% (2 yr) PVG and Gore- Tex® stretch graft respectively	86% at 1 yr; 64% at 3 yr	N/A	N/A	
ASSISTED PRIMARY PATENCY	N/A	N/A	N/A	N/A	N/A	
Primary Patency PATENCY	72% vs. 58% (6 mo) 51% vs. 33% (18 mo)	60.7% vs. 56.5% (1 yr); 54.7% vs. 51.8% (2 yr) PVG and Gore- Tex® stretch graft respectively	73% at 1 yr	87.5% heparin coated vs. 50% control (7 days) P=0.28	N/A	
AUTHOR COUNTRY YEAR NO. AND GRAFT TYPE	34 study meta- analysis (autogenous vs. various PTFE access)	58 (30 Thoratec® PVG vs. 28 Gore- Tex® stretch grafts)	163 PVG	10 bilateral aortoiliac grafting in dogs (5 heparin adsorbed Carboflo® vs. 5 control Carboflo® grafts)	5 PU grafts	
YEAR	2003	2003	2003	2003	2004	
COUNTRY	USA	Japan	Singapore	USA	Turkey	
AUTHOR	Huber et al. <sup>55</sup>	Kiyama et al. <sup>56</sup>	Peng et al. <sup>57</sup>	Laredo et al. <sup>28</sup>	Hazinedaro glu et al. <sup>58</sup>	

292

l			s	(%		of	
	Comments	Within the first 4 days after graft placement, 108 of 133 grafts (81%) were cannulated.	Degree of venous stenosis was significantly reduced.	Infection was less $(10\% vs. 45\%)$	Cuffed ePTFE graft provided better long-term outcome.	Use of Vectra® grafts and TBB fistulas started after a median of 14 (7-30) and 70 (52-102) days, respectively (P < .001)	
	Complications	24% patients died (unrelated to78% and 61% at 6 graft placement), 49% (n=50) hadmo and 1 yrgraft thrombosis (94% or 47/50respectivelyunderwent successfulpercutaneous thrombectomy)	Mean degree of stenosis in non- cuffed graft group (44.95%) was greater than that of the cuffed graft group (22.76%)	10% (n=3) developed infection	Thrombosis as a cause of complete graft failure was higher $(34\%)$ in the standard group than in the cutfied group $(9\%)$ (p = 0.0125). Infection of the graft was observed in 12% of the cutfied group and in 6% of the standard group (p = 0.55)	6.6% graft infection rate Postoperative complications more frequent in TBB fistulas and late complications in Vectra® grafts.	
	Secondary patency	78% and 61% at 6 mo and 1 yr respectively	N/A	N/A	81.8% vs. 56.1% at 1 yr, 61.8% vs. 46.3% at 2 yr, 51.5% vs. 33.1% at 3 yr (p=0.047)	97% vs. 100% (1 mo); 81% vs. 88% (12 mo); 87% vs. 84% (18 mo) P=0.91	
	ASSISTED PRIMARY PATENCY	N/A	N/A	N/A	N/A	93% vs. 100% (1 mo); 70% vs. 82% (12 mo); 58% vs. 78% (18 mo) P=0.033	
	Primary Patency	51% and 33% at 6 mo and 1 yr respectively.	N/A	42% at 12 mo	37.7% vs. 25.7% at 1 yr, 35% vs. 10.3% at 2 yr, 28% vs. 5.1% at 3 yr (p = 0.086)	92% vs. 100% (1 mo); 50% vs. 46% (12 mo); 26% vs. 31% (18 mo) P=0.62	
	AUTHOR COUNTRY YEAR NO. AND GRAFT TYPE	133 PUU grafts	39 (17 Venoflo II vs. 19 Gore-Tex® grafts)	30 Vectra® grafts in HIV + patients	67 (41 Venaflo® II vs. 26 control ePTFE grafts)	117 (76 Vectra® grafts vs. 41 TBB)	
	YEAR	2005	2006	2007	2008	2008	
	COUNTRY '	USA	Taiwan	USA	USA	USA	
	AUTHOR	Jefic et al. <sup>34</sup>	Liu et al. <sup>59</sup>	Schild et al. <sup>36</sup>	Tsoulfas et al. <sup>60</sup>	Kakkos et al. <sup>33</sup>	

Current Status of Synthetic and Biological Grafts for Hemodialysis



**Abbreviations**: polytetrafluoroethylene (PTFE), expanded polytetrafluoroethylene (ePTFE), carbon lined (CL), graft platelet accumulation index (GPAI), high porous (HP), tridodecylmethylammonium chloride (TDMAC), polycarbonate polyurethane (PU), per patient year (PPY), polyurethaneurea (PUU), polyurethane vascular graft (PVG), transposed brachio-basilic fistula (TBB), clot free survival (CFS), autogenous brachial vein-brachial artery access (ABBA), brachio-basilic arteriovenous fistula (BBAVF), Not available (N/ A)

Table 1. Clinical trials of prosthetic grafts for hemodialysis.

infections cultured), methacillin resistant Staphylococcus aureus (21.05%), followed by Pseudomonas aeruginosa (5.26%).<sup>11</sup> The largest number of infections occur when patients are going under routine dialysis (more than 50% of patients) and as a complication of chronic cannulation, rather than postoperative complications.<sup>11</sup> Reducing *Staphylococcus aureus* carrier state in patients undergoing HD and improving antiseptic technique may reduce the rate of infections in grafts.

# 2.3 Characteristics of PTFE grafts

While PTFE is available in various configurations and is produced by various manufacturers, very few have proven to be more beneficial in improving patency in randomized clinical trials for long-term.<sup>1</sup>

# 2.3.1 Effect of wall thickness

In order to examine the effect of wall thickness on patency, Lenz et al.<sup>17</sup> investigated both standard wall and thin wall configuration of PTFE. Although the incidence of complications and mortality did not statistically differ amongst the 2 groups, standard ePTFE had better patency rates. Studies comparing 2 manufacturers of ePTFE grafts: Gore-Tex® (W.L. Gore and Associates, Flagstaff, AZ) and Impra® (C. R. Bard Inc., Murray Hill, NJ) did not find any difference in the performance of 6-mm standard ePTFE grafts.<sup>18 19</sup>

# 2.3.2 Effect of stretch characteristics

In an attempt to reduce kinking of the graft in areas of angulation and to improve intraoperative handling, the graft was modified to stretch (Gore- Tex® Stretch). Tordoir et al. reported a cumulative primary patency rate of 59% in the stretch ePTFE group compared to 29% in standard ePTFE group at 1 year (p < 0.01). In addition, there were significantly fewer thrombotic events for the stretch ePTFE grafts as opposed to the standard ePTFE grafts (40%vs. 12%, p<0.001).<sup>20</sup> Early cannulation of stretch ePTFE grafts was not found to increase peri-operative morbidity rates or decrease 12-month cumulative primary patency rates.<sup>21</sup> In contrast, another study comparing the patency of early cannulation with late cannulation in Gore-Tex® stretch grafts showed that graft patency after thrombosis formation was significantly higher in the late cannulation group (p=0.0002).<sup>22</sup>

# 2.3.3 Effect of ringed reinforcement

Ring reinforced grafts were created to reduce kinking at the apex of loop grafts and decrease incidence of thrombosis associated with external compression. In a retrospective study in which 632 reinforced and non-reinforced PTFE grafts were compared for patency and complications, it was found that non-reinforced grafts had higher primary and secondary patency rates.<sup>23</sup>

# 2.3.4 Effect of cuff or hood on venous ourflow

One of the few modifications that improved patency rates in PTFE vascular grafts was placing a cuff or hood on the venous outflow. The main objective of placing a cuffed PTFE graft is to enlarge the outflow, and reduce mechanical sheer stress in order to reduce thrombotic occlusion caused by neointimal hyperplasia.<sup>1</sup> In a computer simulated model, Venaflo® II (C. R. Bard Inc., Murray Hill, NJ), a flared-end ePTFE graft to simulate a vein

Comments	Overall patency rate at 1 yr was 60% (no obvious superiority of umbilical vein over expanded Teflon grafts)	Survival rate: 3 mo. (93% vs. 84%); 1 yr. (84% vs. 65%); 2 yr (72% vs. 63%); 3 yr (54% vs. 56%) for BCAH and PTFE respectively.	AVF remains the procedure of choice. PTFE appears to be a reasonable second choice.	22 patients with a 20 mo follow-up confirms better survival of BVMG than PTFE (p < 0.04).	
Complications	Thrombosis: 26.1%; Infection: 13.0%	No difference between BCAH and PTFE	Overall Complication rate: 20.71% AVF, 66.6% PTFE, 55.93% sheep collagen, 37.33 bovine xenograft Diffetion: 0% AVF, 10% PTFE, 2% sheep collagen, 1% bovine xenograft Thrombosis: 12% AVF, 45% PTFE, 44% Sheep collagen, 27% bovine collagen, 27% bovine xenograft.	N/A	
Secondary patency	N/A	N/A	AVF: 74%. 67%. 64%, 56%, 36% at 1, 2, 3, 5, and 10 yr. respective-ly. Grafts: Bovine Xenografts: 56%, 29%, 24%; PTFE: 58%, 47%, 40%; 58%, 47%, 40%; Sheep Collagen: 71%, 58%, 45% at 1, 2, 3 yr. respectively	50% at 12 mo and 6.6% at 50 mo for PTFE; 81.9% at 19 mo for BMVG; AVF: 52.4% at 50 mo and 46.5% at 90 mo	
ASSISTED PRIMARY PATENCY	N/A	N/A	N/A	N/A	
Primary Patency	N/A	N/A	720 (429 AVF 59%, 51%, 32% at 1, and 291 grafts   and 291 grafts 2, 3, 5, and 10 yr.   [150 bovine 2, 3, 5, and 10 yr.   respectively. respectively.   xenografts, 69 Grafts: Bovine   PTFE grafts, 59 Xenografts: 51%, autologous and   autologous and Sheep Collagen: 11, 23%, 39% at 1, vein grafts])   2, and 10 yr. 27, 44%, 39% at 1, vein grafts])	17.4 % (PTFE) vs. 23.9% (BMVG) at 12 mo AVF: 43% at 12 mo	
NO. AND GRAFT TYPE	23 Dardik Biograft <sup>TM</sup>	140 (62 Artegraft® vs. 78 PTFE [Gore-Tex® or Impra®] grafts)	720 (429 AVF and 291 grafts [150 bovine xenografts, 69 PTFE grafts, 59 sheep collagen grafts, 10 autologous and autologous and thomologous and vein grafts])	53 PTFE, 10 reinforced PTFE, and 22 ProCol® grafts 404 native AVF	
YEAR	1981	1983	1996	2001	
COUNTRY YEAR	Canada	USA	Switzerland	Italy	
AUTHOR	Wellington 63	Hurt el al. <sup>64</sup>	Enzler et al. <sup>65</sup> Switzerland	Bacchini et al. <sup>42</sup>	

296

Current Status of Synthetic and Biological Grafts for Hemodialysi	sis
---	-----

Comments	80% of the ePTFE grafts were abandoned vs. 34% of the BMVG group.	At 10 weeks, SG showed fibroblast cell migration and proliferation None of the SG grafts became infected, but ePTFE graft group became infected within 54 days of implantation. At 10 weeks, SG showed fibroblast with incorporation into the surrounding subcutaneous tissue, and elongated cells expressing the contractile protein smooth muscle actin were also observed. After 24 weeks, procollagen synthesis was demonstrated in the fully colonized graft matrix.	Intervention rate was lower in the ProCol® group (0.97 versus 1.37) vs. synthetic grafts (p = 0.003).	
Complications	Infection and thrombosis in the BMVG group were lower than the ePTFE group.	None of the SG grafts became infected, but ePTFE graft group became infected within 54 days of implantation.	Complication rates, including dilation, seroma, infection, and thrombosis, were lower for the ProCol® vs. synthetic grafts (p < 0.001).	
Secondary patency	45% BMVG vs. 18% ePTFE at 1 yr (p=0.011) (p=0.006)	N/A	65.6% ProCol® vs. 55.5% synthetic at 12 mo. 42.9% synthetic at 24 mo (p =0.036).	
ASSISTED PRIMARY PATENCY	45% BMVG vs. 18% ePTFE at 1 yr (p=0.011)	N/A	N/A	
Primary Patency	33% in BMVG vs. 18% in ePTFE group at 1 yr (p=0.120)	72.6% and 58.6% for SG vs. 57.4% and 54.7% for the ePTFE grafts at 6 mo and 12 mo respectively.	35.6% ProCol® vs. 28.4% synthetic grafts at 12 mo (p=0.524)	
NO. AND GRAFT TYPE	74 (59 ProCol® vs. 15 ePTFE grafts [Impra®, Gore-Tex® or Boston Scientific])	19 SynerGraft® in 12 canines (11 between carotid artery and jugular vein and 8 between femoral artery and vein)	276 (183 ProCol® and 93 synthetic grafts [90 ePTFE, 1 silicone, 2 PUUJ])	
YEAR	2003	2004	2005	
COUNTRY YEAR	USA	USA	USA	
AUTHOR	Glickman et al. <sup>43</sup>	Matsuura et al. <sup>39</sup>	Katzman et al. <sup>44</sup>	

Complications	No infections were seen in either group, but 2 aneurysms occurred in the SG group. SG group. No significant difference in the number of the number of proup to maintain patency, patients with SG had a higher total cost thrombectomies between the SG and PTFE groups than those with PTFE grafts. (1.2 vs. 2.0, $p = 0.3$ ) (1.2 vs. 2.0, $p = 0.3$ ) (1	Patient 4: True and N/A pseudoaneurysm	
Secondary patency	No significant differences	N/A	
ASSISTED PRIMARY PATENCY	No significant differences	N/A	
Primary Patency	No significant differences	Patient 1 and 2: Radial artery- brachial vein implantation, and had a patency of 5 mo and 6 mo respectively. Patient 3 and 4: Brachial arteryaxillary vein implantation and had patency of 8 mo and 7 mo respectively.	
NO. AND GRAFT TYPE	54 (27 54 (27 SynerGraft® vs. 27 PTFE grafts)	4 total SynerGraft®	
YEAR	2005	2006	
COUNTRY YEAR	USA	Turkey	
AUTHOR	Madden et al.66	Emrecan et al.67	

298

	Comments	SG is an alternative when autologous vein is not available.	The BMVG is a viable alternative for HD access in patients where autologous construction is not possible, and should be given priority in patients with a failed ePTFE graft or high risk for infection.	Both grafts were adequate conduits for HD and amenable to repair. Anticipated advantages for SG were not seen in either patency or stability.	
	Complications	5% infection rate at 1 yr	Patency rates did not Less severe complications differ in the BMVG group	4% vs. 9% infection rate for SG and ePTFE at 1 yr respectively (p=0.410)	
	Secondary patency	81% at 1 yr	Patency rates did not differ	57% vs. 68% at 1 yr P = 0.370	
	ASSISTED PRIMARY PATENCY	45% at 1 yr	N/A	52% vs. 64% at 1 yr P = 0.430	
	Primary Patency	29% at 1 yr	Patency rates did not differ	28% for SG vs. 48 % for ePTFE at 1 yr P = 0.290	
	NO. AND GRAFT TYPE	25 SynerGraft®	46 (23 ProCol® vs. 23 ePTFE grafts)	56 (29 SynerGraft® vs. 27 ePTFE grafts)	
Int	COUNTRY YEAR	United Kingdom	Switzerland 2007	United 2009 Kingdom	en
	AUTHOR CC	Darby l et al. <sup>40</sup> K	Tahami et al. <sup>68</sup>	Chemla I et al. <sup>41</sup> K	

**Abbreviations**: bovine carotid artery heterograph (BCAH), polytetrafluoroethylene (PTFE), arteriovenous fistula (AVG), bovine mesenteric vein graft (BMVG), preserved saphenous vein (PSV), Endoscopic Vessel Harvesting System (EVHS), hemodialysis (HD), SynerGraft® (SG), polyurethaneurea (PUU), Not available (N/ A).

Table 2. Clinical trials of biological conduits for hemodialysis

cuff, showed measurable improvements in reducing wall shear stress gradient, wall shear stress angle gradient, and radial pressure gradient.<sup>25</sup> Sorom et al.<sup>26</sup>found that Venaflo® II was associated with increased blood flow rates during HD and improved graft patency compared with ePTFE graft. Similarly, in a smaller study it was found that a flared-end ePTFE graft provided stable blood flow and satisfactory graft patency during 2 years of follow-up, even in high risk patients with a prior history of vascular access thrombosis.<sup>27</sup> However, a European study did not show improvement in patency rate despite a reduction in thrombotic occlusion and stenosis.<sup>24</sup>

## 2.3.5 Effect of coating PTFE

Another technique meant to improve PTFE graft has been coating the PTFE vascular grafts with carbon or heparin to prevent early graft failure and improve overall patency rates.<sup>28</sup> In a canine model, Tsuchida et al. showed that the graft platelet accumulation index (GPAI) was significantly (p<0.05) lower in the carbon lined PTFE group when compared to the control PTFE group.<sup>29</sup> They concluded that carbon lining decreases platelet accumulation on PTFE grafts. Propaten®, ePTFE with bioactive heparin covalently bound to it (W.L. Gore and Associates) has also been shown to be effective in improving graft patency. It is the only vascular graft of its kind approved for HD access on the market. Davidson et al. <sup>30</sup> found 20% improved primary graft patency of about 80% at one year when comparing Propaten® to standard ePTFE. In order to diminish the risk of neointimal hyperplasia. Cagiannos et al.<sup>31</sup> studied the effects of coating an ePTFE graft with rapamycin in a porcine model. They showed that the rapamycin coated grafts significantly (P < 0.0001) lowered cross sectional narrowing in the outflow graft when compared to non-coated grafts; as well as no evidence of medial necrosis or aneurysmal degeneration. After a four week observation period, coated grafts showed features of diminished neointimal hyperplasia compared to untreated ePTFE grafts. Researchers have also analyzed the effect of a bioabsorbable vascular wrap mesh containing paclitaxel on neointimal hyperplasia in a sheep model. Paclitaxel coated mesh significantly reduced neointimal hyperplasia and neointimal capillary density without toxicity to adjacent vessels.32

### 2.3.6 Self-sealing grafts

K/ DOQI recommends that PTFE grafts should not be routinely used until at least 2 weeks after placement and not until swelling has subsided so that palpation of the course of the graft can be performed. This time is needed for tissue-to-graft incorporation, reducing peri graft hematoma.<sup>1</sup> Due to this complication, newer "self-sealing" grafts have been designed that can be cannulated sooner.<sup>11</sup> Vectra® vascular grafts (C. R. Bard Inc., Murray Hill, NJ) made of a proprietary blend of segmented polyetherurethaneurea and a siloxane containing a surface-modifying additive (SMA), were designed to provide early cannulation, reducing the need for temporary central venous catheters to provide access until the graft matures. In a study of 76 patients, in which Vectra® grafts were compared to transposed brachio-basilic vein (TBB) autogenous access, Kakkos et al.<sup>33</sup> found that aggressive graft surveillance and endovascular treatment methods resulted in equivalent long-term secondary patency rates. The advantage of earlier use of Vectra® graft must be balanced against the need for more frequent secondary interventions and the risk of graft infection. In a single center study, Jefic et al. obtained 81% (108 of 133 grafts) cannulation rate within 4 days of Polyurethane (PU) [Thoratec® Vascular Access Graft] graft placement in which none of the recipients required a temporary catheter. A shorter mean bleeding time after withdrawal of dialysis needle was also acquired in the PU graft group (4.0 minutes vs. 9.2 minutes in ePTFE group).<sup>34</sup> Similarly, Glickman et al. showed that PU grafts had achieved better hemostasis at cannulation sites compared to ePTFE sites when the two were compared at 5 minutes or less after dialysis (P<0.0001). Also, they showed that 53.9% of all PU grafts were cannulated before 9 days vs. none of the ePTFE grafts (P<0.001).<sup>35</sup> In the HIV- positive ESRD patient population, reduction of temporary catheter use and prevention of infection is critical. A study of 30 consecutive HIV positive patients receiving Vectra® graft implantation showed a lower infection rate (10% vs. 45%) than published reports of infection in PTFE grafts. It was concluded that the unique self-sealing property of the Vectra® grafts reduced the development of perigraft hematoma and may have accounted for decreased infection.<sup>36</sup>

# 3. Biological conduits for hemodialysis

Biological graft materials tend to have less intimal hyperplasia at the venous anastamosis, reduced tendency to thrombose, and a lower risk of infection when compared to PTFE.<sup>11</sup> Butler et al. compared bovine heterographs to PTFE and found that the synthetic graft had significantly fewer late thromboses, increased resistance to infection, easier to repair and had comparable longevity.<sup>37</sup> Anderson et al. found that bovine heterographs required twice as many revisions per dialysis month to maintain patency.<sup>38</sup>

SynerGraft® 100 (SG100 [CryoLife Inc., Atlanta, GA]) is a modified bovine ureter graft with some similarities to synthetic graft (similar internal diameter and strong tissue matrix). This graft has been processed to remove xenograft cells while maintaining a collagen matrix that is not chemically cross-linked by aldehydes allowing re-population by autologous cells. Matsuura et al. reported a primary patency rate of 72.6% and 58.6% for SG 100 vs. 57.4% and 54.7% for the ePTFE grafts at 6 months and 12 months, respectively. SG 100 graft showed fibroblast cell migration and proliferation with incorporation into the surrounding subcutaneous tissue after 10 weeks, and procollagen synthesis demonstrated at 24 weeks; while the ePTFE graft had no evidence of cellular ingrowth.<sup>39</sup> In a study of 23 patients receiving SG 100 grafts, Darby et al. found that the bovine ureter graft was a stable vascular access conduit, providing a suitable graft alternative when autologous vein was not available. Their study showed 29% primary, 45% primary assisted, and 81% secondary patency rates at 1 year, with only a 5% infection rate.<sup>40</sup> On the other hand, Chemla et al. found that both grafts were adequate conduits for HD, the anticipated advantages for SG 100 were not seen in either patency or stability.<sup>41</sup>

# 4. Future developments in prosthetic and biological conduits for hemodialysis

The use of pharmacological agents may hold the promise of long-term graft patency. Treatment with 200 mg of dipyridamole and 25 mg of aspirin twice daily resulted in significant improvement of patency rates while adverse effects in both the treatment and placebo groups remained the same.<sup>45</sup> Other agents, such as fish oil and anticoagulants have also been tried with limited success (table 4).

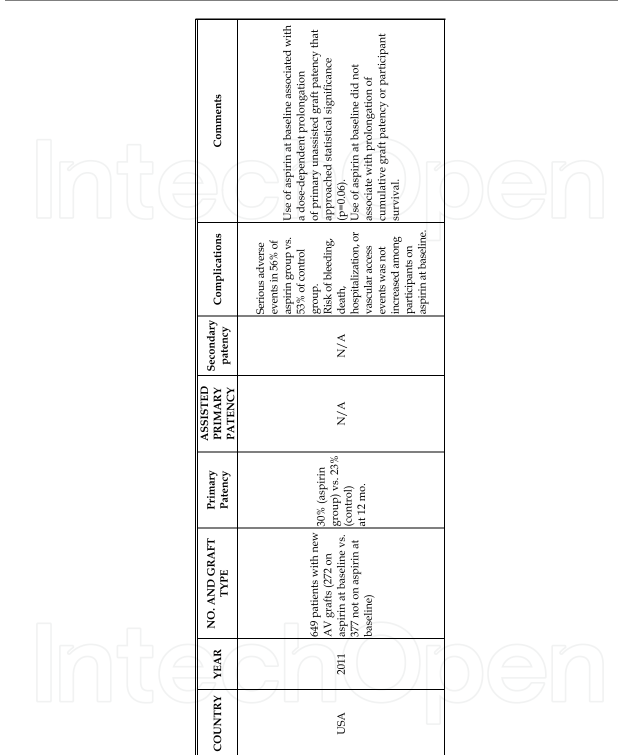
Comments	Rapamycin-eluting ePTFE grafts decrease neointimal hyperplasia in a porcine model.	Paclitaxel-eluting mesh significantly reduced neointimal hyperplasia and neointimal capillary density without toxicity to the adjacent vein.	New graft design addresses two major problems responsible for the development of venous stenosis of prosthetic grafts.	
Complications	N/A	N/A	N/A	
Secondary patency	N/A	N/A	N/A	
ASSISTED PRIMARY PATENCY	N/A	N/A	N/A	
Primary Patency	N/A	N/A	N/A	
NO. AND GRAFT TYPE	22 Mongrel pigs into 3 groups: untreated ePTFE (n=6), adhesive- only coated ePTFE (n = 6), or adhesive- and Rapamycin-coated ePTFE (n = 10)	40 male sheep into 5 groups: no mesh; or a 3-cm x 6-cm mesh with $0.0, 0.3, 0.7$ , or $1.2 \mu$ g/mm <sup>2</sup> of paclitaxel for a total dose of $0.0, 0.6, 1.3,$ or $2.2 $ mg, respectively.	In-vitro study of double outflow (bi-flow) grafts	
YEAR	2005	2007	2010	
COUNTRY	NSA	NSA	Germany	
AUTHOR	Cagiannos et al <sup>31</sup>	Kohler et al <sup>32</sup>	Heise et al <sup>46</sup>	

**Abbreviations:** expanded polytetrafluoroethylene (ePTFE), Not available (N/ A) Table 3. Clinical trials of experimental conduits for hemodialysis

Current Status of Synthetic and Biological Grafts for Hemodialysis

Comments	Type I patients: At the end of the 18 mo follow-up, cumulative rates of thrombosis were: 21± 9% on dipyridamole, alone, 25±11% on dipyridamole and aspirin combination, 42±13% on placebo, and 80±12% on aspirin alone. RR of thrombosis with dipyridamole was 0.35 (P = 0.02) with 95% CI of 0.15 and 0.36. The RR with aspirin was 1.99 with 95% CI of 0.88 and 4.48 (not significant, P=0.18). Type II patients: high thrombosis rates regardless of treatment group. Overall, 78% thrombosis in Type II patients and no statistical differences between study groups.	Use of preoperative single-dose IV vancomycin prophylaxis for hemodialysis vascular graft procedures reduces the risk of postoperative access infection	
Complications	Angina pectoris, GI bleeding, headache, nausea, and vomiting reported	Access infection developed in 1% of vancomycin group and in 6% of control group (P = 0.006). All 14 infections occurred in upper extremity PTFE grafts.	
Secondary patency	N/A	N/A	
ASSISTED PRIMARY PATENCY	N/A	N/A	
Primary Patency	N/A	N/A	
NO. AND GRAFT TYPE	Dipyridamole, Aspirin, Dipyridamole + Aspirin, or placebo in: 84 Type I (new ePTFE graft) patients 23 Type II (previously revised ePTFE graft) patients	408 various vascular access procedures (206 pre-operation treatment with vancomycin vs. 202 non-medicated control group)	
YEAR	1994	1997	
COUNTRY	USA	NSA	
AUTHOR	Sreedhara et al. <sup>69</sup>	Zibari et al. <sup>70</sup>	

	lents	aled a significant a oil-treated and (0.03), with a power decreased venous systemic BP, values.	ot yet published	idamole plus nt but modest risk of stenosis ration of primary newly created	th a 27% (P < 0.001) (opidogrel with a se in mortality, and 0.02) increase in red with patients medications. a drugs was P < 0.0001) increase cally significant umong warfarin, 1 on survival.	
	Comments	Survival analysis revealed a significant difference between fish oil-treated and Gas, bloating most untreated patients (P < 0.03), with a power of 90%.commonof 90%.Fish oil treatment also decreased venous outflow resistance and systemic BP, compared with control values.	Results of study not yet published	Graft failure, death, and seriousTreatment with dipyridamole plusadverse events (including bleeding) did not between studyasignificant but modest effect in reducing the risk of stenosis and improving the duration of primary differ significantly unassisted patency of newly created grafts	Warfarin associated with a 27% ( $P < 0.001$ ) increase in mortality, clopidogrel with a 24% ( $P = 0.002$ ) increase in mortality, and aspirin with a 6% ( $P = 0.02$ ) increase in mortality when compared with patients receiving none of these medications. Prescription of multiple drugs was associated with a 22% ( $P < 0.0001$ ) increase in mortality. No statistically significant interaction was found among warfarin, clopidogrel, and aspirin on survival.	
	Complications	Gas, bloating most common complaints	N/A	Graft failure, death, and serious adverse events (including bleeding) did not differ significantly between study groups	N/A	
	Secondary patency	N/A	N/A	N/A	N/A	
	ASSISTED PRIMARY PATENCY	N/A	N/A	N/A	N/A	
	Primary Patency	75.6% fish oil- treated group vs 14.9% control group at 1 yr	₽/N	23% placebo vs. 28% dipyridamole/ aspirin group at 1 yr	N/A	
	NO. AND GRAFT TYPE	24 total new Gore- Tex® graft patients (12 fish oil vs. 12 control oil group)	232 new grafts for access (116 fish oil treatment vs. 116 control group)	nts: 321 ble and vs. 328 up.	Anticoagulant and antiplatelet medications on 41,425 dialysis patients (8.3% on warfarin 8.3%, 10.0% on clopidogrel, 30.4% on aspirin, 8.1% patients on at least two of these drugs, and 59.7% not medicated)	
	YEAR	2002	2007	2009	2009	
COUNTRY		USA	NSA	USA	USA	
	AUTHOR	Schmitz et al. <sup>71</sup>	Lok et al. <sup>72</sup>	Dixon et al. <sup>45</sup>	Chan et al. <sup>73</sup>	



 $\label{eq:Abbreviations: expanded polytetrafluoroethylene (ePTFE), polytetrafluoroethylene (PTFE), Not available (N/ A).$ 

Dixon et al.<sup>74</sup>

Table 4. Effects of various medications on vascular access grafts

AUTHOR

In an approach of reducing neointimal hyperplasia by decreasing mechanical sheer stress, a new double channel (Bi-Flow) graft was designed. These grafts showed laminar flow and lower levels of turbulence, leading to lower risk of stenosis.<sup>46</sup>

# 5. Conclusions

Prosthetic grafts should be reserved for situations where autogenous vein is not available to perform an access. The most commonly used prosthetic graft is e-PTFE based. Newer advances in medication bonding to decrease thrombosis and formation of intimal hyperplasia may be promising. In addition, various graft characteristics such as flared-end and stretch may provide better patency. Biologic grafts are being tested; however, at this point data are lacking to show superiority over prosthetic grafts. This area is a fertile ground for randomized clinical trials in the search for a man made or biologic graft that would equal autogenous vein in patency and complication rates.

# 6. Abbreviations

Chronic kidney disease (CKD), hemodialysis (HD), peritoneal dialysis (PD), arteriovenous fistula (AVF), polytetrafluoroethylene (PTFE), expanded PTFE (ePTFE), transposed brachialbasilic vein (TBB), percutaneous angioplasty (PTA)

# 7. Definitions

Primary Patency: Interval of time from access placement to any intervention necessary tomaintain patency of access. Assisted Primary Patency: Interval of time from access placement to time of intervention necessary to maintain the functionality of the access. Secondary Patency: Interval of time from access placement to access abandonment including intervening surgical or endovascular manipulations

Graft	Graft Material	Manufacturer
Artegraft®	Bovine Carotid Artery Heterograft	Artegraft Inc., New Brunswick, NJ
Atrium Adventa™ VXT	Reinforced ePTFE	Atrium Medical Corp., Hudson, NH
Boston Scientific	ePTFE	Boston Scientific Corp., Natick, MA
Carboflo®	Carbon impregnated ePTFE	C.R. Bard Inc., Murry Hill, NJ
CryoVein®	Cryopreserved femoral vein	CryoLife Inc., Atlanta, GA
Flixene™	Trilaminate membrane	Atrium Medical Corp., Hudson, NH

# 8. Manufacturer and graft

306

Graft	Graft Material	Manufacturer	
Gore-Tex®	ePTFE	W.L. Gore and Associates, Flagstaff, AZ	
Gore-Tex® Intering® graft	reinforced ePTFE with radial support	W.L. Gore and Associates, Flagstaff, AZ	
Gore-Tex® stretch graft	Stretch ePTFE	W.L. Gore and Associates, Flagstaff, AZ	
Gore-Tex® stretch graft with removable rings	Stretch ePTFE with removable rings	W.L. Gore and Associates, Flagstaff, AZ	
Impra®	ePTFE	C.R. Bard Inc., Murry Hill, NJ	
Dardik Biograft™	Modified human umbilical vein	Meadox Medicals Inc., Oakland, NJ	
ProCol®	Bovine mesenteric vein heterograph	Hancock Jaffe Laboratories Inc., Irvine, CA	
Propaten®	Bioactive heparin convalently bound to ePTFE	W.L. Gore and Associates, Flagstaff, AZ.	
SynerGraft® 100	Bovine Ureter Graft	CryoLife Inc., Atlanta, GA	
Thoratec® Vascular Access Graft	Polyurethane	Thoratec Corp., Pleasanton, CA	
VascuLink™	Self-sealing polycarbonate urethane graft	Lemaitre Vascular Inc., Burlington, MA	
Vascutek®	Self-sealing ePTFE	Tarumo Interventional Systems, Somerset, NJ	
Vectra®	Proprietary blend of segmented polyetherurethaneurea and a siloxane	C.R. Bard Inc., Murry Hill, NJ	
Venaflo® II	Cuffed ePTFE	C.R. Bard Inc., Murry Hill, NJ	

# 9. Disclaimer

The views expressed in this article are those of the authors and do not necessarily reflect the official policy of the Department of Army, the Department of Defense, or the US government.

# 10. References

- [1] Clinical Practice Guidelines and Clinical Practice Recommendations 2006 Update. National Kidney Foundation, 2006. (Accessed at
- http://www.kidney.org/professionals/kdoqi/guideline\_uphd\_pd\_va/index.htm.) [2] USRDS: Annual Data Report. 2010. (Accessed at http://www.usrds.org/adr.htm.)

- [3] Wilhelmi M., A. H. Materials Used for Hemodialysis Vascular Access: Current Strategies and a Call to Action. Graft 2003;6:6-15.
- [4] Konner K. History of vascular access for haemodialysis. Nephrol Dial Transplant 2005;20:2629-35.
- [5] Akoh JA. Prosthetic arteriovenous grafts for hemodialysis. JVasc Access 2009;10:137-47.
- [6] Finelli L, Miller JT, Tokars JI, Alter MJ, Arduino MJ. National surveillance of dialysisassociated diseases in the United States, 2002. Semin Dial 2005;18:52-61.
- [7] Fistula First Data. Arteriovenous Fistula First, 2011. (Accessed at http://www.fistulafirst.org/AboutFistulaFirst/FFBIData.aspx.)
- [8] Hakim R, Himmelfarb J Hemodialysis access failure: a call to action. Kidney Int 1998;54:1029-40.
- [9] Ku DN, Allen RC. Chapter 128: Vacular Grafts. 2nd ed: Boca Raton: CRC Press LLC; 2000.
- [10] Budu-Grajdeanu P, Schugart RC, Friedman A, Valentine C, Agarwal AK, Rovin BH. A mathematical model of venous neointimal hyperplasia formation. Theor Biol Med Model 2008;5:2.
- [11] Wilson SE. Vascular Access: Principles and Practice. 5th ed: Wolters Kluwer Lippincott Williams & Wilkins; 2010.
- [12] Roy-Chaudhury P, Kelly BS, Miller MA, et al. Venous neointimal hyperplasia in polytetrafluoroethylene dialysis grafts. Kidney Int 2001;59:2325-34.
- [13] Roy-Chaudhury P, Sukhatme VP, Cheung AK. Hemodialysis vascular access dysfunction: a cellular and molecular viewpoint. JAm Soc Nephrol 2006;17:1112-27.
- [14] Mezzano D, Pais EO, Aranda E, et al. Inflammation, not hyperhomocysteinemia, is related to oxidative stress and hemostatic and endothelial dysfunction in uremia. Kidney Int 2001;60:1844-50.
- [15] Chang CJ, Ko PJ, Hsu LA, et al. Highly increased cell proliferation activity in the restenotic hemodialysis vascular access after percutaneous transluminal angioplasty: implication in prevention of restenosis. Am JKidney Dis 2004;43:74-84.
- [16] Schutte WP, Helmer SD, Salazar L, Smith JL. Surgical treatment of infected prosthetic dialysis arteriovenous grafts: total versus partial graft excision. Am J Surg 2007;193:385-8; discussion 8.
- [17] Lenz BJ, Veldenz HC, Dennis JW, Khansarinia S, Atteberry LR. A three-year follow-up on standard versus thin wall ePTFE grafts for hemodialysis. J Vasc Surg 1998;28:464-70; discussion 70.
- [18] Kaufman JL, Garb JL, Berman JA, Rhee SW, Norris MA, Friedmann P. A prospective comparison of two expanded polytetrafluoroethylene grafts for linear forearm hemodialysis access: does the manufacturer matter? JAm Coll Surg 1997;185:74-9.
- [19] Hurlbert SN, Mattos MA, Henretta JP, et al. Long-term patency rates, complications and cost-effectiveness of polytetrafluoroethylene (PTFE) grafts for hemodialysis access: a prospective study that compares Impra versus Gore-tex grafts. Cardiovasc Surg 1998;6:652-6.
- [20] Tordoir JH, Hofstra L, Leunissen KM, Kitslaar PJ. Early experience with stretch polytetrafluoroethylene grafts for haemodialysis access surgery: results of a prospective randomised study. Eur JVasc Endovasc Surg 1995;9:305-9.

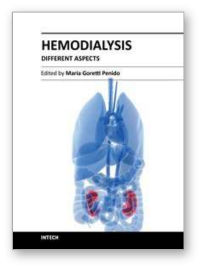
- [21] Hakaim AG, Scott TE. Durability of early prosthetic dialysis graft cannulation: results of a prospective, nonrandomized clinical trial. JVasc Surg 1997;25:1002-5; discussion 5-6.
- [22] Dawidson IJ, Ar'Rajab A, Melone LD, Poole T, Griffin D, Risser R. Early use of the Gore-Tex Stretch Graft. Blood Purif 1996;14:337-44.
- [23] Schuman ES, Standage BA, Ragsdale JW, Gross GF. Reinforced versus nonreinforced polytetrafluoroethylene grafts for hemodialysis access. Am JSurg 1997;173:407-10.
- [24] Lemson MS, Tordoir JH, van Det RJ, et al. Effects of a venous cuff at the venous anastomosis of polytetrafluoroethylene grafts for hemodialysis vascular access. J Vasc Surg 2000;32:1155-63.
- [25] Longest PW, Kleinstreuer C. Computational haemodynamics analysis and comparison study of arterio-venous grafts. JMed Eng Technol 2000;24:102-10.
- [26] Sorom AJ, Hughes CB, McCarthy JT, et al. Prospective, randomized evaluation of a cuffed expanded polytetrafluoroethylene graft for hemodialysis vascular access. Surgery 2002;132:135-40.
- [27] Nyberg SL, Hughes CB, Valenzuela YM, et al. Preliminary experience with a cuffed ePTFE graft for hemodialysis vascular access. ASAIO J2001;47:333-7.
- [28] Laredo J, Xue L, Husak VA, Ellinger J, Greisler HP. Silyl-heparin adsorption improves the in vivo thromboresistance of carbon-coated polytetrafluoroethylene vascular grafts. Am JSurg 2003;186:556-60.
- [29] Tsuchida H, Cameron BL, Marcus CS, Wilson SE. Modified polytetrafluoroethylene: indium 111-labeled platelet deposition on carbon-lined and high-porosity polytetrafluoroethylene grafts. JVasc Surg 1992;16:643-9; discussion 9-50.
- [30] Davidson I, Hackerman C, Kapadia A, Minhajuddib A. Heparin bonded hemodialysis e-PTFE grafts result in 20% clot free survival benefit. JVasc Access 2009;10:153-6.
- [31] Cagiannos C, Abul-Khoudoud OR, DeRijk W, et al. Rapamycin-coated expanded polytetrafluoroethylene bypass grafts exhibit decreased anastomotic neointimal hyperplasia in a porcine model. JVasc Surg 2005;42:980-8.
- [32] Kohler TR, Toleikis PM, Gravett DM, Avelar RL. Inhibition of neointimal hyperplasia in a sheep model of dialysis access failure with the bioabsorbable Vascular Wrap paclitaxel-eluting mesh. JVasc Surg 2007;45:1029-37; discussion 37-8.
- [33] Kakkos SK, Andrzejewski T, Haddad JA, et al. Equivalent secondary patency rates of upper extremity Vectra Vascular Access Grafts and transposed brachial-basilic fistulas with aggressive access surveillance and endovascular treatment. J Vasc Surg 2008;47:407-14.
- [34] Jefic D, Reddy PP, Flynn LM, Provenzano R. A single center experience in the use of polyurethaneurea arteriovenous grafts. Nephrol News Issues 2005;19:44-7.
- [35] Glickman MH, Stokes GK, Ross JR, et al. Multicenter evaluation of a polytetrafluoroethylene vascular access graft as compared with the expanded polytetrafluoroethylene vascular access graft in hemodialysis applications. J Vasc Surg 2001;34:465-72; discussion 72-3.
- [36] Schild AF, Perez EA, Gillaspie E, Patel AR, Noicely K, Baltodano N. Use of the Vectra polyetherurethaneurea graft for dialysis access in HIV-positive patients with endstage renal disease. Vasc Endovascular Surg 2007;41:506-8.

- [37] Butler HG, 3rd, Baker LD, Jr., Johnson JM. Vascular access for chronic hemodialysis: polytetrafluoroethylene (PTFE) versus bovine heterograft. Am J Surg 1977;134:791-3.
- [38] Anderson CB, Sicard GA, Etheredge EE. Bovine carotid artery and expanded polytetrafluroethylene grafts for hemodialysis vascular access. J Surg Res 1980;29:184-8.
- [39] Matsuura JH, Black KS, Levitt AB, et al. Cellular remodeling of depopulated bovine ureter used as an arteriovenous graft in the canine model. J Am Coll Surg 2004;198:778-83.
- [40] Darby CR, Roy D, Deardon D, Cornall A. Depopulated bovine ureteric xenograft for complex haemodialysis vascular access. Eur JVasc Endovasc Surg 2006;31:181-6.
- [41] Chemla ES, Morsy M. Randomized clinical trial comparing decellularized bovine ureter with expanded polytetrafluoroethylene for vascular access. Br J Surg 2009;96:34-9.
- [42] Bacchini G, Del Vecchio L, Andrulli S, Pontoriero G, Locatelli F. Survival of prosthetic grafts of different materials after impairment of a native arteriovenous fistula in hemodialysis patients. ASAIO J2001;47:30-3.
- [43] Glickman MH, Lawson JH, Katzman HE, Schild AF, Fujitani RM. Challenges of hemodialysis access for high risk patients: Impact of mesenteric vein bioprosthetic graft. JVasc Access 2003;4:73-80.
- [44] Katzman HE, Glickman MH, Schild AF, Fujitani RM, Lawson JH. Multicenter evaluation of the bovine mesenteric vein bioprostheses for hemodialysis access in patients with an earlier failed prosthetic graft. JAm Coll Surg 2005;201:223-30.
- [45] Dixon BS, Beck GJ, Vazquez MA, et al. Effect of dipyridamole plus aspirin on hemodialysis graft patency. N Engl JMed 2009;360:2191-201.
- [46] Heise M, Kirschner P, Rabsch A, Zanow J, Settmacher U, Heidenhain C. In vitro testing of a newly developed arteriovenous double-outflow graft. J Vasc Surg 2010;52:421-8.
- [47] Johnson JM, Anderson JM. Reasonable Expectations for PTFE Grafts in Hemodialysis Access. Dialysis & Transplantation 1983;12:238-40.
- [48] Munda R, First MR, Alexander JW, Linnemann CC, Jr., Fidler JP, Kittur D. Polytetrafluoroethylene graft survival in hemodialysis. JAMA 1983;249:219-22.
- [49] Schanzer H, Martinelli G, Chiang K, Burrows L, Peirce EC, 2nd. Clinical trials of a new polytetrafluoroethylene-silicone graft. Am JSurg 1989;158:117-20.
- [50] Schanzer H, Martinelli G, Burrows L, Chiang K, Peirce EC, 2nd. Clinical trial of a selfsealing PTFE-silicone dialysis graft. ASAIO Trans 1989;35:211-3.
- [51] Taylor SM, Eaves GL, Weatherford DA, McAlhany JC, Jr., Russell HE, Langan EM, 3rd. Results and complications of arteriovenous access dialysis grafts in the lower extremity: a five year review. Am Surg 1996;62:188-91.
- [52] Ritter EF, Kim YB, Reischl HP, Serafin D, Rudner AM, Klitzman B. Heparin coating of vascular prostheses reduces thromboemboli. Surgery 1997;122:888-92.
- [53] Khadra MH, Dwyer AJ, Thompson JF. Advantages of polytetrafluoroethylene arteriovenous loops in the thigh for hemodialysis access. Am JSurg 1997;173:280-3.

- [54] Jeschke MG, Hermanutz V, Wolf SE, Koveker GB. Polyurethane vascular prostheses decreases neointimal formation compared with expanded polytetrafluoroethylene. JVasc Surg 1999;29:168-76.
- [55] Huber TS, Carter JW, Carter RL, Seeger JM. Patency of autogenous and polytetrafluoroethylene upper extremity arteriovenous hemodialysis accesses: a systematic review. JVasc Surg 2003;38:1005-11.
- [56] Kiyama H, Imazeki T, Kurihara S, Yoneshima H. Long-term follow-up of polyurethane vascular grafts for hemoaccess bridge fistulas. Ann Vasc Surg 2003;17:516-21.
- [57] Peng CW, Tan SG. Polyurethane grafts: a viable alternative for dialysis arteriovenous access? Asian Cardiovasc Thorac Ann 2003;11:314-8.
- [58] Hazinedaroglu SM, Kayaoglu HA, Ayli D, Duman N, Yerdel MA. Immediate postimplant hemodialysis through a new "self-sealing" heparin-bonded polycarbonate/ urethane graft. Transplant Proc 2004;36:2599-602.
- [59] Liu YH, Hung YN, Hsieh HC, Ko PJ Impact of cuffed, expanded polytetrafluoroethylene dialysis grafts on graft outlet stenosis. World JSurg 2006;30:2290-4.
- [60] Tsoulfas G, Hertl M, Ko DS, et al. Long-term outcome of a cuffed expanded PTFE graft for hemodialysis vascular access. World JSurg 2008;32:1827-31.
- [61] Schild F. APHECS II Trail. Atriums Propective Hemodialysis Early Cannulation Trial with the Flixene Vascular Graft. 2010.
- [62] Lioupis C, Mistry H, Rix T, Chandak P, Tyrrell M, Valenti D. Comparison among transposed brachiobasilic, brachiobrachial arteriovenous fistulas and FlixeneTM vascular graft. JVasc Access 2010.
- [63] Wellington JL. Umbilical vein grafts for vascular access in patients on long-term dialysis. Can JSurg 1981;24:608-9.
- [64] Hurt AV, Batello-Cruz M, Skipper BJ, Teaf SR, Sterling WA, Jr. Bovine carotid artery heterografts versus polytetrafluoroethylene grafts. A prospective, randomized study. Am JSurg 1983;146:844-7.
- [65] Enzler MA, Rajmon T, Lachat M, Largiader F. Long-term function of vascular access for hemodialysis. Clin Transplant 1996;10:511-5.
- [66] Madden RL, Lipkowitz GS, Browne BJ, Kurbanov A. A comparison of cryopreserved vein allografts and prosthetic grafts for hemodialysis access. Ann Vasc Surg 2005;19:686-91.
- [67] Emrecan B, Yilik L, Ozbek C, Gurbuz A. Bovine ureter graft for haemodialysis access surgery. Nephrol Dial Transplant 2006;21:2290-1.
- [68] Tahami VB, Hakki H, Reber PU, Widmer MK, Kniemeyer HW. Polytetrafluoroethylene and bovine mesenterial vein grafts for hemodialysis access: a comparative study. J Vasc Access 2007;8:17-20.
- [69] Sreedhara R, Himmelfarb J, Lazarus JM, Hakim RM. Anti-platelet therapy in graft thrombosis: results of a prospective, randomized, double-blind study. Kidney Int 1994;45:1477-83.
- [70] Zibari GB, Gadallah MF, Landreneau M, et al. Preoperative vancomycin prophylaxis decreases incidence of postoperative hemodialysis vascular access infections. Am J Kidney Dis 1997;30:343-8.

- [71] Schmitz PG, McCloud LK, Reikes ST, Leonard CL, Gellens ME. Prophylaxis of hemodialysis graft thrombosis with fish oil: double-blind, randomized, prospective trial. JAm Soc Nephrol 2002;13:184-90.
- [72] Lok CE, Allon M, Donnelly S, et al. Design of the fish oil inhibition of stenosis in hemodialysis grafts (FISH) study. Clin Trials 2007;4:357-67.
- [73] Chan KE, Lazarus JM, Thadhani R, Hakim RM. Anticoagulant and antiplatelet usage associates with mortality among hemodialysis patients. J Am Soc Nephrol 2009;20:872-81.
- [74] Dixon BS, Beck GJ, Dember LM, et al. Use of Aspirin Associates with Longer Primary Patency of Hemodialysis Grafts. JAm Soc Nephrol 2011;22:773-81.





Hemodialysis - Different Aspects Edited by Prof. Maria Goretti Penido

ISBN 978-953-307-315-6 Hard cover, 321 pages **Publisher** InTech **Published online** 14, November, 2011 **Published in print edition** November, 2011

The book provides practical and accessible information on all aspects of hemodialysis, with emphasis on dayto-day management of patients. It is quite comprehensive as it covers almost all the aspects of hemodialysis. In short it is a valuable book and an essential aid in the dialysis room.

# How to reference

In order to correctly reference this scholarly work, feel free to copy and paste the following:

Purav P. Patel, Maria Altieri, Tarun R. Jindal, Steven R. Guy, Edward M. Falta, Eric A. Elster, Frank P. Hurst, Anton N. Sidawy and Rahul M. Jindal (2011). Current Status of Synthetic and Biological Grafts for Hemodialysis, Hemodialysis - Different Aspects, Prof. Maria Goretti Penido (Ed.), ISBN: 978-953-307-315-6, InTech, Available from: http://www.intechopen.com/books/hemodialysis-different-aspects/current-status-ofsynthetic-and-biological-grafts-for-hemodialysis



# InTech Europe

University Campus STeP Ri Slavka Krautzeka 83/A 51000 Rijeka, Croatia Phone: +385 (51) 770 447 Fax: +385 (51) 686 166 www.intechopen.com

# InTech China

Unit 405, Office Block, Hotel Equatorial Shanghai No.65, Yan An Road (West), Shanghai, 200040, China 中国上海市延安西路65号上海国际贵都大饭店办公楼405单元 Phone: +86-21-62489820 Fax: +86-21-62489821 © 2011 The Author(s). Licensee IntechOpen. This is an open access article distributed under the terms of the <u>Creative Commons Attribution 3.0</u> <u>License</u>, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

# IntechOpen

# IntechOpen