An effective and ultrarapid technique for kidney transplantation in the rabbit is introduced. Vascular anastomosis was completed using a novel cuff technique in which mating cuffs were used to join the delicate renal vein. The ureter was reconstructed by spatulated end-to-end anastomosis, with special attention to the rabbit's unique ureteral vascular anatomy. The total vascular anastomosis time was 3.4 ± 1.3 min, and there were no episodes of bleeding or thrombosis. The ureter complication rate was 7.3%. Kidneys transplanted after 5 h of cold storage using the new technique yielded better postoperative creatinine results than similar preserved kidneys transplanted using previously described methods. We suggest this technique for studies of long- and short-term kidney preservation and transplantation in the rabbit, as well as for veterinary transplantation in which donor kidneys must be stored for only a short time before use.

The rabbit is an attractive model for kidney transplantation research, in view of its docility, convenient size, easy maintenance, low cost, and renal anatomical and functional similarity to human kidneys. We were further attracted to the rabbit kidney transplant model due to the overwhelming use of this model in studies involving renal cryopreservation. Most of the publications describing rabbit kidney transplantation techniques utilized microsurgical techniques developed 25–30 years ago, especially by Green and Dunn. In those studies, the warm ischemic time was reported to be between 20–30 min. Since then, few publications have commented on the warm ischemic time. Khirabadi and Fahy adopted the traditional cuff technique in their rabbit kidney transplantation model, cutting the time for vascular reconstruction to 10–15 min, but this still represents substantial warm ischemia.

The difficulty of rabbit renal vascular anastomosis is mainly due to the delicate nature of the renal vein. We describe a technique that permits ultrarapid (~3 min) vascular anastomosis, the fastest method so far published, with excellent postoperative patency. The new anastomosis method extends earlier cuff approaches by adding two mating venous cuffs that can be “plugged” together and secured in a process lasting less than 1 min, without using a loupe or suturing. Another common technical problem with rabbit kidney transplantation is ureter obstruction, which ranges from 10–30% postoperatively. We therefore also report on a modified technique for ureter reconstruction that better preserves ureteral blood flow and yields improved patency rates.

These improvements collectively provide a new, controllable, and simplified rabbit renal transplantation model.

MATERIALS AND METHODS

All procedures involving animals were consistent with the guidelines established by the Animal Care and Use Committee of 21st Century Medicine, Inc., and the principles set forth in the Guide for the Care and Use of Laboratory Animals (Institute of Laboratory Animal Resources, National Research Council, DHHS, publication no. (NIH) 86-23 (1985)) and USDA guidelines.

Male New Zealand White rabbits (Harlan Sprague Dawley, Indianapolis, IN) weighing 2.5–3.1 kg were housed in separate cages under standard vivarium conditions, and were allowed free access to water and chow. On the day of surgery, rabbits were premedicated with intramuscular ketamine (30 mg/kg) and xylazine (5 mg/kg) (Henry Schein, Inc., Melville, NY). General anesthesia was maintained by inhalation of 1.5–5% isoflurane in oxygen at a flow rate of 1.5 L/min via a face mask. Rectal temperature was maintained between 36–38.5°C, using a heating pad and warm saline for irrigating abdominal tissues. Heart rate and saturated oxygen status were
monitored by a pulse oximeter (Vet/OX® 4404, Heska Corp., Fort Collins, CO), with the probe placed on the tail. Arterial pressure was measured using a 20-gauge i.v. canula inserted into the central ear artery.

Two separate surgeries were performed for each transplant: a morning surgery for organ procurement, and an afternoon surgery for transplantation. During the morning surgery, the rabbit was hydrated with about 50 ml of lactated Ringer’s solution by slow i.v. drip. During the afternoon surgery, each rabbit received about 150 ml of lactated Ringer’s solution in a similar fashion.

**Donor Kidney Procurement**

A midventral laparotomy was performed under sterile conditions. After the intestine was covered and protected with warm saline-soaked gauze, the right kidney was exposed. Chlorpromazine (2.5 mg dissolved in 2 ml saline) was injected around the renal vessels to avoid vasospasm during dissection, and then the right kidney and its vasculature were gently dissected free from surrounding fat and connective tissue. The ureter was transected approximately 5 cm from the hilum. A short cannula (18-gauge Vialon i.v. catheter, 2.0 cm long, Becton Dickinson, Sandy, UT) was inserted into the ureter and ligated in place to permit ready visualization of urine output. Mannitol (2.4 ml of a 25% w/v solution) and furosemide (0.2 ml of 25 mg/ml) were administered intravenously to establish diuresis before ligating the renal artery and then the vein. Immediately after ligation, the renal vein was severed, and the artery was incised and cannulated in situ with a 16-gauge i.v. catheter (inner diameter (ID), 1.16 mm; outer diameter (OD), 1.70 mm; Becton Dickinson). The kidney was flushed at a pressure of 60 mmHg with 50 ml of 0°C storage solution (Eurocollins, UW, or others) containing heparin (10 U/ml) and chlorpromazine (50 μg/ml). During the initial cold flush, the artery was ligated onto the cannula and cut between the body and the cannula, at which point the kidney was removed and transferred into the selected cold solution at 0°C. A second flush was then carried out, using another 50 ml of cold solution containing no added drugs. The laparotomy was closed in two layers with 2-0 Vicryl for the abdominal wall and 3-0 Vicryl for the skin. The kidneys were stored in the storage solution (900 ml) at 0°C for 5 h. The time for kidney retrieval surgery (from “skin to skin”) was 50—60 min. The warm ischemic period during renal procurement was about 1 min.

**Kidney Transplantation**

Kidney transplantation was performed approximately 5 h after nephrectomy. The donor rabbit was reanesthetized and reopened under sterile conditions. A peritoneal flap covering the left kidney and ureter was prepared by cutting along the left renal vessels and the aorta to expose the left ureter and kidney. After injection of chlorpromazine around the renal artery and vein, both vessels were isolated. The kidney was stripped of surrounding connective tissue. The left renal vessels were clampedatraumatically at their origins close to the aorta and vena cava using microvascular clamps, and transected as close as possible to the hilum. The ureter was ligated close to the kidney to avoid damaging the ureteral vein connecting the middle ureter to the vena cava, and to ensure sufficient ureteral length to preclude traction during the subsequent ureteroureterostomy. The left kidney was then removed and discarded. The lumens of the left renal artery and vein remaining in the animal were flushed with warm saline. Both vessels were then passed through and everted over cuffs and secured with circular 6-0 sutures (Fig. 1A). The arterial cuff was made of a 14-gauge Surflo i.v. catheter segment (ID, 1.73 mm; OD, 2.17 mm; length, ~5 mm; Terumo Medical Corp., Elkton, MD). The venous cuff was about 5 mm in length and was made of 10-gauge Angiocath™ I.V. catheter (ID, 3.0 mm; OD, 3.2 mm; Becton Dickinson), with the ID enlarged by manual distension to ~3.2 mm along most of its length, and to ~3.3 mm at the end designated for receiving the kidney.
The vein of the preserved kidney was cuffed while the kidney remained ex vivo in the 0°C storage solution (Fig. 1B). The ex vivo venous cuff was made from 10-gauge Angiocath™ material without expanding its diameter, so that its outer diameter approximated the inner diameter of the in vivo venous cuff. The ex vivo venous cuff was secured with two 6-0 sutures. The two venous cuffs were designed so as to allow the ex vivo (male) cuff and the in vivo (female) cuff to mate.

Shortly before grafting the kidney, 1.6 ml of 25% mannitol solution were administered intravenously. The preserved kidney was then placed in the peritoneal cavity at the site of the original left kidney. Arterial anastomosis was performed by passing the donor kidney’s artery over the recipient arterial cuff and securing it with a prepositioned circular 6-0 prolene suture. Venous anastomosis was then completed by simple insertion of the donor cuffed vein into the recipient cuffed vein, with special attention to vascular orientation on both sides of the anastomosis. The initial cuff-positioning step (Fig. 1C) was facilitated by the expanded (3.3 mm ID) segment of the female cuff, allowing easy insertion and secure orientation of the male cuff. To create a permanent anastomosis, the male cuff was then driven into the female cuff by exerting greater force, applied for example by a spatula pressing on the trailing edge of the advancing male cuff, until the male cuff was fully seated in the female cuff (Fig. 1D). Generally, no further securing sutures were required to maintain the venous anastomosis, but a single suture was sufficient to accomplish stabilization when desired. The vascular clamps were released immediately after fully seating the male cuff in the female cuff.

The ureter was reconstructed over an 8-mm length of 18-gauge Vialon i.v. catheter (Becton Dickinson) placed into the adjoining ureter segments. Special attention was paid to preserving the unique ureteral vein in rabbits that connects the middle ureter to the vena cava. Eight interrupted sutures of 9-0 nylon were used to complete a tension-free spatulated anastomosis.

At the end of vascular and ureteral anastomosis, the reconstructed ureter and the kidney were covered by the peritoneal flap to avoid postoperative adhesions and to secure their positions, and the flap was sutured in place. The abdominal muscle was closed, as described earlier. The skin was sutured in running fashion with a 3-0 Ethilon suture. Renal transplantation required 80–100 min from “skin to skin.”

Postoperative Maintenance and Follow-Up

After recovering from anesthesia, rabbits were allowed free access to water and chow. Blood samples (0.5 ml) were drawn daily via the ear artery for determination of serum levels of blood urea nitrogen (BUN), creatinine, and other chemistries. After approximately 2 weeks of postoperative monitoring, rabbits were again anesthetized and opened, and the grafts were quickly removed and flushed free of blood, using 50 ml of saline containing 5,000 units of heparin at 60–80 mmHg. Kidneys were then fixed by perfusion with 50 ml of modified Karnovsky’s fixative (~730 mOsm) at the same pressure. Fixed kidneys were sent to Paragon Bioservices (Baltimore, MD) for routine histological processing and periodic acid-Schiff (PAS) staining.

RESULTS

The results of 113 consecutive renal autografts are summarized. Of these, 109 (96.5%) rabbits survived the anesthesia and transplantation procedure, and 84.4% of the surviving rabbits were followed for 14–30 days without incident. The combined arterial and venous vascular anastomosis time was 3.4 ± 1.3 min (mean ± SD). There were no episodes of venous or arterial thrombosis or leakage in this series. Incisional hernia, sepsis, wound infection, and hind limb paralysis were not observed in the present series. The only complication noted was ureter blockage, which was evidenced by rising serum creatinine levels in 8 cases out of 109 (7.3%) on postoperative days 5–10. Most of these incidents recovered spontaneously.

The functional performance of kidneys preserved at 0°C in Eurocollins solution for 5 h and transplanted as described here is compared in Table 1 to historical data obtained using our previous surgical model and to results obtained with sham-operated controls. The postoperative peak creatinine level resulting from short-term preservation with the present surgical model was significantly lower than previously reported results (P < 0.01), and was in fact even lower than the peak observed in previous sham-operated controls. In addition, serum creatinine returned to normal in 2 days in this model, as compared to 9 days in the previous study.

DISCUSSION

In this model, rabbits underwent two major surgical interventions in 1 day. The overall survival rate was 96.5%, which exceeds previously reported survival rates. The only surgical complication was ureter blockage, which occurred in 7.3% of the procedures, compared to an incidence of 10–30% in the literature. The mean peak creatinine level for kidneys stored in Eurocollins solution for 5 h was significantly lower than previous results, and even better than results
Renal Vascular Anastomosis

Warm ischemia during renal transplantation probably contributes to the overall damage observed after renal cold storage and should therefore be minimized. In this study, the warm ischemic time attributable to nephrectomy was about 1 min, and vascular anastomosis contributed approximately another 3 min on average to the total warm ischemic time. To our knowledge, this is the fastest vascular reconstruction so far reported in renal transplantation, with normal anastomosis times ranging from about 9–25 min in rabbit models and 5–30 min in the rat. The method of vascular anastomosis described here is a further development of our previously described cuff technique, but using two mated cuffs for renal vein anastomosis. The nonsuture external cuff method was first described in 1900 by Payr. Modifications of the technique were extensively and successfully used in experimental organ transplantation research involving rabbits and rats. Theoretically, the cuff technique is faster than end-to-end suturing and avoids problems such as intimal damage, imperfect intima-to-intima contact, complications from the presence of intraluminal suture, and incomplete sealing of the anastomosis. One past problem with realizing these theoretical advantages in the rabbit has been the delicate nature of the rabbit renal vein, which has required at least 10 min to complete cuffing and anastomosis, with a patency rate of 83–99%. We overcame this time limitation by using two mating venous cuffs and by preparing all cuffs while the kidney was still in the cold preservation solution. During the actual transplantation procedure, the two mating cuffs were simply “plugged” together in a process which could be completed in less than 1 min. The renal artery, on the other hand, is easy to pass over the arterial cuff and secure, and thus does not need the mating cuff technique for fast anastomosis.

Ureter Anastomosis

A very common technical problem of renal transplantation in the rabbit is the maintenance of adequate ureteral patency. A wide variety of techniques has been reported. Most techniques involve either end-to-end uretostomy or uretero-cystostomy. No matter which technique is used, the ureter complication rate varies from 10–30% to 5–30 min in the rat. Although edema may contribute to ureteral stenosis, ischemia of the suture line and subsequent stricture formation are the main causes of ureter obstruction. On this basis, we adopted a spatulated end-to-end reconstruction technique, which generates a more distributed anastomosis line that is less susceptible to occlusive stricture. An internal stent facilitated the approximation of the two sides of the ureter and precluded accidental suturing together of the dorsal and ventral ureteral walls. This small stent can be passed into the bladder and then voided without incident most of the time.

We also felt that attention to the anatomy of the rabbit ureter’s vasculature could be important for minimizing ureteral ischemia. The rabbit ureter is supplied predominantly by a branch of the renal artery proximally and by a branch of the vesicular artery distally. The middle segment of the rabbit ureter has no reported extrinsic blood supply, and is drained by a

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Table 1. Postoperative Serum Creatinine Levels

<table>
<thead>
<tr>
<th></th>
<th>EC (5 h) (n = 6) (this model)</th>
<th>EC (5 h) (n = 7) (historical data)</th>
<th>Sham operation (n = 6) (historical data)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peak serum creatinine (mg/dl)</td>
<td>2.02 ± 0.72c</td>
<td>6.81 ± 3.13</td>
<td>3.76 ± 3.43</td>
</tr>
<tr>
<td>Days to peak</td>
<td>1.0 ± 0.0</td>
<td>2.7 ± 1.1</td>
<td>1.5 ± 0.6</td>
</tr>
<tr>
<td>Days to new baseline</td>
<td>2.2 ± 0.4c</td>
<td>9.3 ± 1.1</td>
<td>5.8 ± 1.8</td>
</tr>
<tr>
<td>Preoperative creatinine (mg/dl)</td>
<td>0.68 ± 0.12</td>
<td>1.41 ± 0.10</td>
<td>1.15 ± 0.07</td>
</tr>
<tr>
<td>Final creatinine (mg/dl)</td>
<td>1.20 ± 0.22</td>
<td>1.84 ± 0.15</td>
<td>1.58 ± 0.13</td>
</tr>
<tr>
<td>Final minus initial creatinine</td>
<td>0.52</td>
<td>0.43</td>
<td>0.43</td>
</tr>
</tbody>
</table>

*Mean ± SD (mg/dl).

Data from Khirabadi and Fahy.11

Significantly different from historical Eurocollins (EC) data, P < 0.01 by Student’s t-test.

Days to return creatinine to values equal to final values achieved for each specific rabbit.
single vein emptying into the vena cava. However, we observed that this “emptying vein” is accompanied by a tiny artery originating from either the aorta or the vesiculac artery (unpublished observations). A single group of longitudinal arteries and veins runs the full length of the ureter within the adventitia. Branches of these longitudinal vessels pass tangentially through the muscularis to supply a vascular complex within the lamina propria. Therefore, in addition to minimizing any manipulation of the tissue at the hilum, the blood vessels supplying the middle segment of the recipient ureter have to be preserved. In the present series of rabbit renal autografts, in which attention to these considerations was given, only 7.3% of the grafts showed some degree of ureteral obstruction on postoperative days 5–10, and most of these complications resolved without active intervention.

CONCLUSIONS

We suggest this ultrarapid nonsuture mated cuff technique for studies of long- and short-term kidney preservation and transplantation in the rabbit, as well as for veterinary transplantation in which donor kidneys must be stored for only a short time before use.

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REFERENCES