P35
Percutaneous Nephrolithotomy in the Prone-Flexed Position: Is There an Optimal Access*
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Background: Percutaneous nephrolithotomy (PCNL) is the gold standard therapy for large renal calculi. The optimal puncture site for collecting system access remains controversial, with many suggesting increased morbidity from supra-costal puncture. We report a retrospective review of 318 consecutive PCNL cases over a sixty-month time period performed by two surgeons in a teaching environment.

Methods: Perioperative data was collected on 318 consecutive PCNL cases performed for intra-renal nephrolithiasis from May 2004 to June 2009; patient anatomic anomalies, calyceal diverticuli, endopyelotomy, antegrade ureteroscopy, and patients failing to return for follow-up were excluded. Data was analyzed in 2 groups: supra-costal and infra-costal. Success rates in the infra-costal group (90.8%) were equivalent to the infra-costal group (94.1%). Complication rates across groups were low, with no significant difference in complications between the infra-costal and infra-costal puncture groups, respectively, across Clavien grade I (12 vs. 1), grade II (4 vs. 11), grade IIIa (4 vs. 0), grade IIIb (2 vs 0), and grade IVa (1 vs 2), \( p = 0.067 \). All four (2.6%) Clavien grade III complications in the supra-costal group were pleural complications requiring chest drain insertion. No patients required a blood transfusion or angiembolisation.

Conclusion: When indicated, supra-costal access can provide excellent outcomes, particularly for patients with complex stone disease while obviating the need for multiple tracts or retreatment. Although complications, particularly pleural, occur more frequently in those patients undergoing supra-costal puncture, the incidence is low and acceptable considering the significant advantages in this patient population.

P36
Effective Radiation Exposure in follow-up of patients with Nephrolithiasis
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Background: Patients known to have previous nephrolithiasis are regularly followed in Stone Clinic with frequent imaging to rule out recurrence. Recently, there has been increasing awareness and concerns among both patients and health care professionals regarding long term risks from radiation exposure in follow-up of patients with benign disease. The aim of the present study was to quantify the yearly effective radiation doses associated with the follow-up of patients with nephrolithiasis in Stone Clinic at a single institution.

Methods: Retrospective chart review of 56 patients attending the Stone Clinic at a single academic centre between January and September 2009 was performed. Number and modality of diagnostic imaging studies in the previous 2 years were recorded. Effective radiation exposure doses (reported in mSv) were calculated from the Dose Length Product values (measured in mGy-cm) converted to mSv using a skin surface area of 1.73 m². Radiation exposure from Kidney Ureter Bladder Plain Films (KUBs), Intravenous Pyelograms (IVPs) and fluoroscopy procedures were estimated from previous published data.

Results: A total of 38 males and 18 females with a mean age of 49 years (range: 21-78 years) were included in this study. 137 KUBs, 9 IVPs, 47 fluoroscopic examinations and 73 CTs were reviewed. Median yearly calculated effective radiation exposure dose was 36.8 mSv (Mean: 33.86, range: 1.7 - 54.59 mSv) in 2008 and 21.64 mSv (Mean 22.42, range: 1.4-77.27 mSv) in 2009. A total of 7 (12.5%) patients received a calculated effective radiation exposure dose >50 mSv per year. Mean effective radiation exposure dose was significantly higher in 2008 when compared to 2009, but did not correlate with stone location, age or sex of the patient.

Conclusions: Currently there are no guidelines on imaging modality or frequency in follow-up of patients with nephrolithiasis. A significant portion of patients are receiving effective radiation doses that exceed the current occupational radiation hazard limits. Therefore, urologists should be cognizant of the radiation exposure of patients when ordering imaging studies for follow-up of patients with benign disease.
and multivariate analyses were performed using Pearson Chi-square, two-tailed t, and Fisher’s Exact tests. All variables were considered statistically significant at \( p < 0.05 \)

**Results:** Mean age and BMI were 54.15 years and 29.59 kg/m² respectively. Univariate analysis revealed that stone diameter (\( p = 0.0001 \)), complete staghorn stones (\( p < 0.001 \)), and upper pole stones (\( p < 0.0001 \)) were associated with having more residual stones while prior extracorporeal shockwave lithotripsy (\( p < 0.037 \)) and holmium laser use (\( p < 0.01 \)) were associated with improved stone-free status. In a sub-analysis of patients who had residual stones after the initial procedure, second look nephroscopy was associated with improved overall stone-free status (\( p < 0.019 \)) in comparison to patients who did not have a second-look.

**Conclusions:** Preoperative and perioperative parameters can predict outcome of PCNL. Second look nephroscopy is associated with improved stone clearance. These results indicate that further studies with larger sample size are needed to construct preoperative prediction models.

**P38 Single Institution Review of the Incidence of Perinephric Hematoma after Shock Wave Lithotripsy**

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**Background:** Perinephric hematoma formation is a potentially serious complication of extracorporeal shock wave lithotripsy (SWL), with a reported incidence of between 0.2% and 30%. The aim of this study was to determine the incidence of and evaluate the risk factors for the development of clinically apparent post SWL renal hematoma with the latest generation shock wave lithotripter.

**Methods:** From April 2006 to September 2008, 3351 SWL treatments were performed using the Storz Modulith SLX-F2. Data was collected prospectively for patient age, body mass index, gender, stone size, stone location, number of shock waves, energy level, shock frequency, medications and the existence of hypertension and/or diabetes mellitus. A case-control analysis was then conducted to compare risk factors.

**Results:** Following SWL treatment, 12 patients developed clinically apparent renal hematomas for an overall incidence of 0.3%. All patients were male and 8 (66%) of the affected patients had known hypertension. Preoperative use of drugs with antiplatelet effect was associated with a greater risk of perinephric hematoma (\( p = 0.047 \)).

**Conclusions:** The incidence of clinical apparent hematomas following SWL with the SLX-F2 was 0.3%. Statistical significant risk factors for perinephric hematoma included male gender and use of drugs with antiplatelet effect.

**P39 Intermediate-Sized Urolithiasis: The Equivalency of Treatment Modalities**

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**Background:** Shock wave lithotripsy (SWL) is considered a standard treatment for upper tract stones less than 10 mm in diameter, whereas stones larger than 20 mm are best treated by percutaneous nephrolithotomy (PCNL). The treatment of stones between these sizes remains controversial. We review our modern series of SWL, RIRS and PCNL outcomes for intermediate-sized upper tract calculi.

**Methods:** Data from patients treated with SWL, RIRS and PCNL from June 2005 to June 2009 were reviewed. Analysis was restricted to those patients with a pre-treatment non-contrast CT scan conducted at our centre demonstrating an upper tract calculus measuring an area between 100 and 300 mm². Demographic, stone, patient, treatment and follow-up data were collected from a prospective database and review of CT and fluoroscopy imaging performed by two independent urologists and one radiologist. Data was analyzed with Chi square analysis and ANOVA where appropriate.

**Results:** 137 patients were referred with non-staghorn calculi with an area between 100-300 mm². Across groups, there were 89 males (65.0%), 61 right-sided stones (44.5%), and an overall mean age of 53.1 years (SD 14.2) and BMI of 29.0 kg/m² (SD 6.6). 53 (38.7%) patients were referred before SWL, while 41 (29.9%) and 43 (31.4%) underwent RIRS and PCNL, respectively. Mean stone area was higher in the PCNL group at 211.1 mm² (SD 56.8), compared with 172.6 mm² (SD 58.2) for the SWL group and 162.9 mm² (SD 54.9) for the RIRS group (\( p < 0.001 \)). Stone density, measured by Hounsfield units (HU) on CT, were significantly higher for SWL patients (1008 HU, SD 244) versus 786 HU (SD 289) for RIRS and 837 HU (SD 326), \( p = 0.002 \). Single treatment success rates were significantly better for PCNL at 95.3%, versus 87.8% for RIRS and 60.4% for SWL, \( p < 0.001 \). However, when up to two SWL treatments were administered, the success rate improved to 79.2%, thus removing any significant difference between the success of the three treatment modalities (\( p = 0.66 \)). Auxiliary treatments were more common after SWL (42.3%) vs. 9.8% and 7.0% in the RIRS and PCNL groups, respectively.

**Conclusions:** Although success rates are significantly higher with single treatment PCNL and RIRS when compared to SWL, when allowing for up to two SWL treatments there was no significant difference between treatment modalities. Thus, SWL is a reasonably successful treatment alternative for patients not fit for or not wishing a general anesthetic, provided they accept a higher number of treatments.

**P40 Ambulatory percutaneous nephrolithotomy: Initial series**

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**Introduction:** Percutaneous nephrolithotomy (PCNL) is the gold standard for management of large renal stones. Traditionally, patients are admitted post-operatively with a large-bore nephrostomy tube, which is removed after a normal nephrostomy on post-operative day two. Although tubeless PCNL has been described previously, there have been no reports of ambulatory tubeless PCNL. The aim of the present study was to assess the safety and feasibility of ambulatory tubeless PCNL. Here the initial 10 patients are presented.

**Methods:** The initial series of 10 patients undergoing ambulatory tubeless PCNL was included in the present study. Patient information including age, sex, fluoroscopy time, operating room time, stone size (using largest diameter), and Hounsfield Units (HU) were collected prospectively and analyzed retrospectively. Furthermore, number of needle punctures, number of tracts, and stone free status were ascertained. Amount of narcotic administered in the recovery room (mg morphine equivalents), amount of time spent in recovery room (minutes), amount of narcotics used at home, and complications were recorded and documented. Criteria for same day discharges were: single tract, stone free status, absence of post-operative pain, and satisfactory post-operative chest X ray and CBC. All patients had antegrade double J stents placed intra-operatively. Male patients were discharged home with Foley catheter. Follow-up office visit was done on post operatively 2 days for trial of void and removal of the flank dressing. Double J stents were removed a week later cystoscopically.

**Results:** Out of the 10 patients undergoing ambulatory PCNL, 2 had established nephrostomy tracts. The rest of the 8 patients had nephrostomy tract established intra-operatively by the urologist. The median operating and fluoroscopy times were 83.5 and 4.45 minutes, respectively. The median stone diameter was 20 mm with a median of 800 HU. Patients spent a median of 240 minutes in the recovery room and received a median of 19.25 mg of morphine equivalents. There were no intra-operative complications and none of the patients required transfusions. There were two post-operative complications. The first was a Deep Vein Thrombosis requiring anticoagulation. The second was re-admission for multi-resistant E. Coli UTI requiring intravenous antibiotics.

**Conclusion:** In highly-selected patients, ambulatory tubeless PCNL is safe and feasible. More patients are needed to verify criteria for patients undergoing ambulatory approach.
P41
Radiation Exposure During Percutaneous Nephrolithotomy in a Contemporary Series
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Introduction: Minimizing radiation exposure during urologic procedures has gained importance recently. Traditionally, percutaneous nephrolithotomy (PCNL) has been associated with the highest radiation exposure in endourologic procedures. Therefore, the aim of the present study was to document radiation exposure during a contemporary series of PCNL and to determine factors influencing fluoroscopy time.

Methods: All patients with large renal stones presenting for PCNL between July 31, 2009 and November 20th, 2009 were included in the study. Patient information including age, sex, fluoroscopy time (seconds), operating room time (minutes), stone size (using largest diameter in mm), and stone density measured in Hounsfield Units (HU) were collected prospectively and analyzed retrospectively. Linear regression was used to determine whether fluoroscopy time varied with length of surgery, stone size and stone density.

Results: There were a total of 19 patients with a median age of 51 yrs old (14 males and 5 females). Median OR time was 85 minutes and the median fluoroscopy time was 317 ± (5.28) minutes (range 127-720 s). Median stone size was 25 mm with a median of 800 HU. Length of fluoroscopy was independent of the length of surgery (p > 0.05) and stone size (p > 0.05). There was a trend towards less fluoroscopy time with denser stones (-19%/100 HU). However, this did not reach statistical significance (p = 0.06). This could be due to the small sample size.

Conclusions: Urologists should be cognizant of radiation exposure during PCNL. More patients are needed to verify whether fluoroscopy times decrease with higher stone densities.

P42
Tibial Nerve Neuromodulation of Bladder Activity in Cats
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Background: Posterior tibial nerve stimulation (PTNS) is a relatively new approach used to treat medically refractory overactive bladder. However, its efficacy and poststimulation effects are not yet fully recognized. The goals of this study were to therefore test the efficacy of different stimulation parameters and to determine the post-stimulation effects of PTNS.

Methods: Ten female cats anesthetized with alpha-chloralose underwent PTNS via a nerve cuff electrode. Initially, the bladder was infused with saline via a urethral catheter to a volume about 100-120% of bladder capacity in order to induce rhythmic bladder contractions. CMG was then performed 5 times in the ensuing 1-1.5 hours without PTNS in order to determine the post-stimulation effect on the micturnition reflex. Finally, PTNS was applied during CMG to determine its maximal inhibitory effect on bladder activity.

Results: PTNS at both 5 Hz and 30 Hz significantly (p < 0.05) inhibited isovolumetric bladder contractions at a stimulation intensity 2-3 times that of the threshold (T) voltage—the minimum voltage required for inducing low movement at 5 Hz. After continuous 30 minute PTNS, bladder capacity was significantly (p < 0.05) increased to 134±2.1% (30 Hz) and 137.8 ± 1.2% (5 Hz) of the control capacity. There was no significant difference between 5 Hz and 30 Hz stimulation. In the post-stimulation period, PTNS at 5 Hz further increased bladder capacity to 177 ± 10% of control capacity.

Conclusions: This study demonstrated that PTNS could significantly inhibit bladder activity at both low (5 Hz) and high (30 Hz) stimulation frequencies. Furthermore, continuous 30 minute PTNS could induce a post-stimulation inhibitory effect on the bladder lasting 1-1.5 hours, supporting the clinical observation of PTNS having a long-lasting inhibitory effect on bladder overactivity.

P43
Withdrawn

P44
Standard of Care Outcomes using Mesh Products (“Perigee” and “Apogee”) for Treating Female Pelvic Prolapse in 170 patients - Single Surgeon Experience
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Background: The POWER Registry was established as an observational registry measuring standard of care practices assessing safety and efficacy of type 1 mesh products to treat female pelvic prolapse, across different specialties with sites from USA and New Zealand. This report is from one local center with one surgeon that participated in this registry.

Methods: Eligible patients are adult females with at least one AMS product implanted for prolapse repair and no other restrictions on product type. Prolapse was quantified via Baden-Walker measurement in this center. The serial follow-up period was for two years.

Results: Patients were enrolled from May 2005 and the study was completed in August 2009. During this period 170 patients were enrolled in this center. The median procedure time was 78 min (53-123 IQK). Areas of prolapse repair included cystocele (89%), rectocele (49%), vault (45%), and enterocele (6%). The rates of concomitant hysterectomy and incontinence repair were 52% and 90%, respectively. More than 95% percent of patients were Caucasian. Mean patient age was 60 years and mean gravity was 2.8 with 1.33 standard deviation (range 0-8). The mean intra-operative complication rate was 1.2%. Most patients (83%) were free of prolapse device-related events. Conversely 17% of patients experienced at least one prolapse device-related event and one with bladder perforation during insertion requiring closure as a serious event, but with no subsequent issues. In another, the insertion needle was noted in the bladder, and after repositioning, did not require bladder repair. The most frequent event, mesh extrusion through the vaginal mucosa, occurred in 6.5% of all patients. All of them were satisfactorily managed with estrogen cream and/or trimming and secondary closure. Mesh erosion involving the urethra/bladder did not occur. Incontinence issues were noted in 3.5% of patients. Other events included healing issues, pain, granuloma formation (4%), and dehiscence (2.4%).

In 63 patients who completed 19-24 months follow-up, the anterior prolapse significantly improved at least one grade (ABW Cystocele -1.94, p < 0.001). Similar improvement was seen in the posterior area (ABW Rectocele -1.12, p < 0.001, ABW Vault -1.69, p < 0.001, and ABW Enterocele -0.13 p < 0.001).

Conclusions: Standard of care outcomes using minimally invasive, type 1 mesh for treating prolapse exhibited a good safety profile with low complication rates and good efficacy. The most common prolapse device-related complication was mesh extrusion through the vaginal mucosa that was satisfactorily managed with no subsequent or long-standing issues.

P45
Pelvic Floor Reconstruction with Customized Mesh
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Background: Pelvic floor reconstruction with mesh graft is a currently popular procedure to correct the vault prolapse and cystocele. Most of the commercial products are designed with only a “one-size-fits-all female patient” in mind. Since the pelvic size and shape of each individual is different, “one-size” product will not fit every female patient exactly. It is suggested that this is the possible cause of recurrent prolapse (incompletely fit a larger pelvis) and mesh erosion/extrusion (bunched up mesh to fit a smaller pelvis). Using 3D pelvic CT scan with cystogram, the pelvic size and shape is able to exactly measured and the mesh can be tailoring to fit each individual’s pelvis. The above complications can be avoided.
Results: perineal ligament to correct hypermobility of the urethra. Extrusion of mesh are observed. Pelvic muscular pain occurred in 4 posterior urethrovesical angle is also resolved by anchoring the mesh to the cystocele were studied past 2-3 years. A 3D pelvic CT scan is performed. The pelvic size is measured by Vitea 2,3D processing software (Fig. 1). A 15x15 cm polypropylene mesh (pore size 1121, w-66 c-90) is tailoring according to the above measurement (Fig. 2). The mesh is anchoring into the sacrospinous ligament and each side of Arch Tendinous Pelvic Fascia (ATPF) by Capio devic. The normal posterior urethrovesical angle is also resolved by anchoring the mesh to the insertion area of ATPF to the pelvic ramus. The mesh also sutured to peri-urethral ligament to correct hypermobility of the urethra.

Methods: Twenty female patients with vault prolapse and grade III or IV cystocele were studied past 2-3 years. A 3D pelvic CT scan with cystogram is performed. The pelvic size is measured by Vitea 2,3D processing software (Fig. 1). A 15x15 cm polypropylene mesh (pore size 1121, w-66 c-90) is tailoring according to the above measurement (Fig. 2). The mesh is anchoring into the sacrospinous ligament and each side of Arch Tendinous Pelvic Fascia (ATPF) by Capio devic. The normal posterior urethrovesical angle is also resolved by anchoring the mesh to the insertion area of ATPF to the pelvic ramus. The mesh also sutured to peri-urethral ligament to correct hypermobility of the urethra.

Results: All twenty patients have not recurrent prolapse. No erosion or extrusion of mesh are observed. Pelvic muscular pain occurred in 4 patients and usually resolved in three months. Two patients developed ureteral obstruction from the procedure and had lysis procedure. Ureteral obstruction can be prevented with ureteral catheter placement prior to the procedure.

Conclusions: The pelvic floor reconstruction with individual customized mesh graft can provided a better result than the one-size-fit mesh graft. More study is needed.

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**P46**

A Trial of Bilateral Percutaneous Nephrostomy Tube Placement in Refractory Non-ulcerative Interstitial Cystitis Patients Prior to Simple Urinary Diversion

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**Background:** Urinary diversion (with or without cystectomy) as a treatment modality for end-stage classic ulcerative interstitial cystitis (IC) has been reported to be beneficial. Diversion with cystectomy for non-ulcerative IC fails to benefit the majority of patients and can add significant morbidity. Recent reports suggest that simple urinary diversion without cystectomy could benefit non-ulcerative IC patients. Simple diversion could improve IC symptoms by preventing bladder stretching and removing chemical irritation from urine. We offered a trial of bilateral percutaneous nephrostomy tubes with Fogarty ureteral occlusion balloons to four patients with refractory non-ulcerative IC to determine if their symptoms would significantly decline enough to be offered a simple diversion.

**Methods:** Four female patients, who fulfilled the National Institute of Arthritis, Diabetes, Digestive, and Kidney Diseases criteria for non-ulcerative IC and who had failed previous medical and intravesical treatments, underwent placement of bilateral 8 French percutaneous nephrostomies with 4 French Fogarty ureteral occlusion balloons. Pre- and post-procedure pain scores were reviewed for each patient. Three patients had significant reduction of pain scores and subsequently underwent ileal conduit urinary diversion (2 open, 1 robotic assisted laparoscopic approach).

**Results:** Mean duration of disease at time of nephrostomy placement was 7 years (range 5-11). Mean duration of nephrostomy placement was 27 days (range 9-50). Mean decrease in pain scores at time of nephrostomy was 5.3 (range 3-8). Mean time between nephrostomy placement and ileal conduit urinary diversion was 154 days (range 103-227). Mean hospital length of stay after diversion was 18 days (range 6-29). At six month postoperative visits, patients were re-evaluated. One patient (age = 63) had complete resolution of pelvic symptoms with pain score = 0. One patient (age = 32) went back to her pre-nephrostomy pain score of 9; her pain is exacerbated by bowel movements and she continued on extensive narcotic regimen and intravesical therapy. One patient (age = 27) intermittently has pre-nephrostomy pain scores but at six month visit was using significantly less narcotics for symptom control and was overall pleased with surgical outcome.

**Conclusions:** Patients who suffer from refractory non-ulcerative IC present a treatment challenge for urologists. Therapy with irreversible urinary diversion can be beneficial but long term success is not easily predicted by a preoperative trial of bilateral nephrostomy tubes with ureteral occlusion balloon catheters. Our limited series suggest that further criteria may need to be fulfilled prior to a patient being eligible for simple diversion.

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**P47**

Modified Scrotal (Bianchi) Mid-Raphe Single-Incision Orchidopexy for Palpable Undescended Testis

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**Background:** To compare the results of a low trans-scrotal mid-raphe orchidopexy in patients with palpable undescended testes (UDT), to a high-scrotal incision (Bianchi) and to the conventional inguinal approach.

**Methods:** We used a retrospective cohort study design. All orchidopexies performed between 2003 and 2009 with a minimum of 3-month follow-up were included. All palpable UDT that could be brought down manually into the upper third of the scrotum under general anaesthesia, were then reviewed (group 1: high-scrotal incision, group 2: low-scrotal incision) and compared to the inguinal two-incision technique (group 3). We excluded cases who had undergone previousinguinal surgery or with concomitant surgeries. We comprehensively reviewed the charts and focused on the following outcomes: operative time, success as defined by mid or lower scrotal position of the testicle, and complications at 6-12-weeks and 1-year after surgery.
Results: A total of 286 orchidopexies were performed in 214 patients with palpable UDT. In group 1, a high-scrotal incision was performed in 44 patients for 60 UDT (success 59/60, 98%) with one recurrence. A modification to the technique was adopted and since 2005, patients in group 2 had a trans-scrotal orchidopexy through a single low-scrotal incision on the median raphe. It was performed in 81 patients for 125 UDT. All tests except 1 (99%) were located in a good position within the scrotum. In group 3, a standard inguinal two-incision orchidopexy was performed in 89 patients for 101 UDT (success 100%). The mean operative time for unilateral UDT was significantly shorter for the low trans-scrotal orchidopexy (mean 28 min vs. mean 37 min; \( p < 0.001 \)) than for the inguinal orchidopexy but equivalent to a high scrotal incision (27 min; \( p = 0.59 \)). One patient approached by high-scrotal incision required conversion to a traditional inguinal approach. All patent processes vaginalis were ligated, regardless of their size. In all 160 children followed at 1 year, no long term atrophy or secondary reascent were observed. Postoperative complications included transient postoperative scrotal hematomata in a single patient from group 1 and 2 wound infections in group 3.

Conclusion: Low trans-scrotal mid-raphe orchidopexy appears to be an excellent alternative to the high-scrotal incision or the standard inguinal orchidopexy for low palpable UDT especially for bilateral cases. Scrotal orchidopexy is simple, safe, and effective in selected cases.

P48
Renal Perfusion Pump vs. Cold Storage for Donation after Cardiac Death Kidneys: A Systematic Review

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Introduction: The use of a renal perfusion pump vs. cold storage for the preservation of kidneys obtained from donors after cardiac death (DCD) is controversial. Herein, we examine the impact of using renal perfusion pumps on delayed graft function and graft survival using a systematic review.

Methods: The PUBMED database was searched and reference lists from 1960 to 2009 were consulted. Randomized controlled trials as well as prospective and retrospective cohort studies that compared delayed graft function and graft survival were included. Studies excluded from review were unable to discriminate between DCD and neurologically deceased donor (NDD) deaths. Eleven studies that followed a total of 3377 participants qualified for review. Statistical analysis was carried out using the random effects criteria and odds ratios calculated using the Cochrane database software.

Results: We found that perfusion pumped kidneys from DCD donors had reduced DGF rates vs. kidneys that were placed in cold storage (Fig. 1) \( (p = 0.008; \text{ odds ratio} 0.4, \text{ CI} 0.20-0.82) \). In addition, graft function at 1 year post-transplantation favored perfusion pumped kidneys \( (p = 0.04, \text{ odds ratio} 1.96, \text{ CI} 1.02-3.77) \). No differences in primary non-function rates between groups were noted.

Conclusion: This review supports the use of perfusion pumps for DCD kidney transplantation with respect to delayed graft function rates and graft survival.