



Radiologic evaluation of the kidney transplant donor and recipient

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Abstract

The kidney is the most common solid organ transplant globally and rates continue to climb, driven by the increasing prevalence of end stage renal disease (ESRD). Compounded by advancements in surgical techniques and immunosuppression leading to longer graft survival, radiologists evermore commonly evaluate kidney transplant patients and candidates, underscoring their role along the transplant process. Multiphase computed tomography (CT) with multiplanar and 3D reformatting is the primary method for evaluating renal donor candidates, detailing renal size, vascular/collecting system anatomy, and identifying significant pathologies such as renal vascular diseases and nephrolithiasis. Ultrasound is the preferred initial postoperative imaging modality for graft evaluation due to its low cost, accessibility, noninvasiveness, and lack of radiation. CT and magnetic resonance imaging (MRI) may be useful adjunctive imaging techniques in diagnosing transplant pathology when ultrasound alone is not diagnostic. Kidney transplant complications are categorized by an approximate timeline framework, aiding in differential diagnosis based on onset, duration, and severity and include perinephric fluid collections, graft compression, iatrogenic injuries, vascular compromise, graft rejection, and neoplastic processes. This review discusses imaging strategies and important findings along the transplant timeline, from donor assessment to long-term recipient complications.

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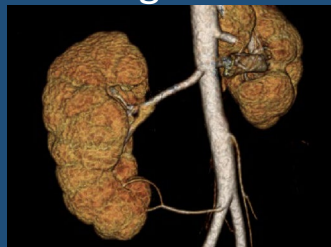
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Graphical Abstract

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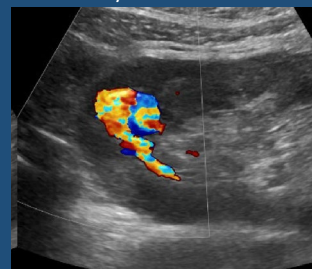
Accessory lower pole renal artery in a kidney donor candidate



Fibromuscular dysplasia (FMD) in a kidney donor candidate



Subcapsular hematoma of a transplanted kidney



Post-biopsy pseudoaneurysm in a transplanted kidney

- The kidney is the most common solid organ transplant globally
- Imaging strategies and findings vary along the transplant timeline
- Renal donor candidates are assessed for key anatomical variants and other surgically relevant pathologies
- Renal transplant recipients are at risk of various potentially graft threatening complications

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Keywords Renal donor · Kidney transplantation · Transplant complications

Abbreviations

CT	Computed tomography
CTA	Computed tomographic angiography
EMR	Electronic medical record
ESRD	End stage renal disease
FMD	Fibromuscular dysplasia
HU	Hounsfield unit
IVC	Inferior vena cava
MRI	Magnetic resonance imaging
MRA	Magnetic resonance angiography
MIPs	Maximum intensity projections
MPR	Multiplanar reconstruction
PSV	Peak systolic velocity
PACS	Picture archive and storage
PTLD	Post-transplant lymphoproliferative disorder
RACS	Renal allograft compartment syndrome
RI	Resistive index

Introduction

History and current state of kidney transplantation

The kidney was the first transplanted solid human organ and remains the most commonly transplanted today, driven by rising prevalence of end stage renal disease (ESRD) in the United States and globally (Fig. 1a) [1, 2]. Dialysis and kidney transplantation are the primary ESRD treatments. Although dialysis costs more than twice as much as transplantation with 5–6-fold increased mortality, dialysis is the primary therapy in the United States due to donor kidney scarcity (Fig. 1b) [2]. This shortage is somewhat alleviated by a steady rate of living kidney donation. Deceased donor kidney transplant rate has also increased over the past decade, partially due to young adult deaths from the opiate crisis and changes in donor and recipient eligibility criteria (Fig. 1c) [3]. As kidney transplant prevalence rises, so too does the need for radiologists' proficiency with kidney donor and recipient imaging.

Unlike deceased donors, who are explored surgically at organ harvest, living donor candidates are evaluated for key anatomy by radiologists, who also assess the post-transplant recipient for complications and help guide management by

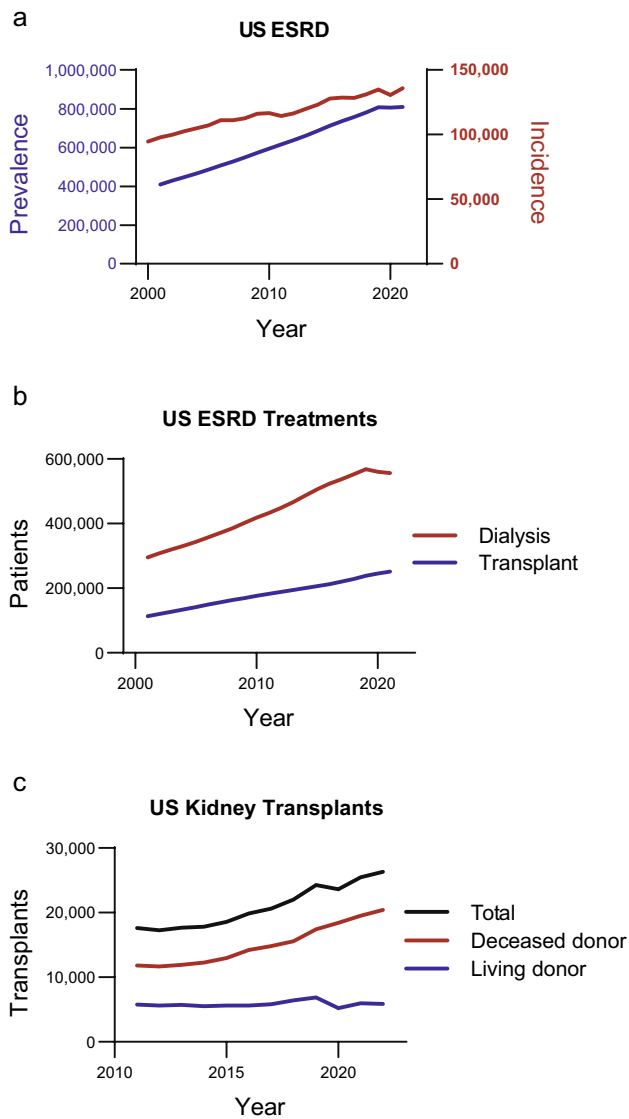


Fig. 1 Trends of ESRD in the United States. **a** Increasing incidence and prevalence of ESRD since 2001. **b** Prevalence of ESRD treatment modalities among living patients. **c** Rate of kidney transplants by donor type. Adapted from [2, 3]

the transplant team. This review covers imaging strategies and findings throughout the transplant timeline, from donor evaluation to chronic recipient complications.

Imaging techniques for the kidney donor

Computed tomography

Multiphase computed tomography (CT) is less invasive and more accurate than intravenous pyelography and angiography used historically to evaluate the kidneys, renal vasculature, and collecting systems. Donor evaluation

requires at least two contrast phases: arterial phase for assessing the renal cortical volume and arteries, and excretory phase for the ureters and parenchymal lesions. Arterial phase can be accomplished with injection timing or bolus tracking per institutional preference [4, 5].

Additional contrast phases and reconstructions vary by practice [4–8]. Stone detection can be performed on non-contrast acquisition, arterial and nephrogenic phases, or virtual non-contrast images from dual energy/spectral acquisitions which reduce radiation while maintaining at least 90% sensitivity [9]. Nephrogenic phase may help delineate the renal veins and increase sensitivity for renal and extrarenal pathology. Excretory phase images are either conventional CT stacks or a repeat topogram, which is sufficient for gross ureteral anatomy. Nephrogenic or delayed phase images may extend into the pelvis or be limited to the upper abdomen to reduce radiation. The protocol used at one of the authors' institutions is described in Table 1.

Multiplanar and 3D reconstruction

Modern helical, multidetector CT data can create volumetric, thin-slice images to allow for multiplanar reconstruction (MPR) by the interpreting radiologist on the picture archive and storage (PACS) system. While this obviates the need for separate coronal and sagittal images for the radiologist, 3 mm thick coronal and sagittal images are still usually sent to PACS for viewing in the electronic medical record (EMR) by the transplant team. These thicker coronal and sagittal images are not adequate to detect subtle but important findings such as vasculopathies, ostial calcifications, and small renal stones due to the Nyquist sampling theorem.

3D images created by a technologist are useful to referring clinicians without PACS access to help conceptualize the spatial relationships between the anatomic structures. These can include curvilinear planes following key vessels, maximum intensity projections (MIPs) showing single-image vascular relationships, and 3D rendered rotating anatomic models. The choice of images sent to the EMR must be made in context of referring transplant team's expectations as well as 3D technologist availability or, without this resource, the time and technology constraints of the radiologist.

Magnetic resonance imaging

Magnetic resonance imaging (MRI) is an alternative for donor evaluation, offering advantages like the absence of ionizing radiation to typically young and healthy donors, no risk of contrast-induced nephropathy, lower risk of allergic-like contrast reactions, and better characterization of incidental lesions. Studies suggest lower test performance metrics with contrast-enhanced magnetic resonance angiography (MRA)

Table 1 Example of a kidney donor evaluation CT protocol

Contrast	Oral: water Intravenous: 370 mg/mL iodine, mixed and injected by patient weight 50 mL, 75% contrast + 25% saline, 3.5 mL/s (under 250 lb) 80 mL, 100% contrast, 4 mL/s (over 250 lb) 40 mL saline, 4 mL/s after contrast (all weights)
Acquisitions	1. Topogram 2. (If single energy CT) precontrast, left atrium to iliac arteries 3. Postcontrast bolus tracking series, upper abdominal aorta, 150 HU trigger with 10 s diagnostic delay 4. Arterial phase, left atrium to iliac arteries 5. Topogram (6 min) 6. Delay phase (6 min), left atrium to iliac arteries
Reconstructions	1. (If single energy CT) precontrast 2 mm thick and spacing in transaxial, coronal, and sagittal planes 2. (If dual energy CT) arterial phase: (a) 50 kEv, 1.25 mm thick, 0.625 mm spacing (b) virtual noncontrast, 2.5 mm thickness and spacing, (c) iodine maps 2.5 mm thickness and spacing 3. (If single energy CT) atrial phase: 1.25 mm thick, 0.625 mm spacing 4. 6 min delay: 5 mm thickness and spacing 5. Coronal and sagittal: arterial phase, 3 mm thickness and spacing (archival) 6. 3D lab cortical volumes 7. 3D lab batch oblique coronals through right and left renal hila (archival) 8. 3D lab MIP single images (minimum 4) of renal arteries, renal veins (archival)
Other	Radiologist saves archival images of each measurement and curvilinear vessel reconstructions, if performed

compared to CT angiography (CTA), especially in detecting arterial variants [10–12]. Image quality depends on field strength: 3 Tesla magnets can produce higher-quality images while underestimating arterial diameter, overestimating renal artery stenosis, and potentially suffering from medical hardware-induced artifacts in some patients [13].

Nuclear renal scintigraphy

Renal scintigraphy was the first widely used modality for quantifying split renal function [14]. More recently, cortical volume measurement by CT has demonstrated high correlation with scintigraphy [15, 16], thus CT has supplanted routine scintigraphy in many transplant centers. Nuclear studies may still be used to delineate split function if there is > 10% volume discrepancy on CT. At one author's institution, this is typically performed by injecting 370 MBq (10 mCi) ^{99m}Tc -mercapto acetyl triglycine (MAG3) and measuring peak renal uptake over 30 min. If split function discrepancy > 10% is confirmed, the smaller kidney is still considered based on intended recipient size. If it is too small, the donor can still donate to a smaller-bodied recipient as part of a paired- or chained-donor arrangement, thus allowing the originally intended recipient to receive a different, larger kidney.

Assessing the renal donor

Living donor surgical approaches

Understanding the surgical approach for donor nephrectomy helps the radiologist distill pertinent anatomic details on

CT. Nearly all nephrectomies (> 98%) are performed using minimally invasive techniques including pure laparoscopic, hand-assisted laparoscopic, and robot-assisted techniques through either a transperitoneal or retroperitoneal approach [3] (Fig. 2a). Hand-assisted laparoscopic nephrectomy is the most common approach and may have lower complication rates but longer hospital stays compared to robot-assisted nephrectomy [17, 18]. All three techniques may utilize an anterior approach through a low midline, transverse suprapubic, or ipsilateral incisions for gel, camera, and instrument ports [17]. Left nephrectomy is preferred as the left renal vein is usually longer, making for easier implantation. In an anterior approach left nephrectomy, the left colon, spleen, and pancreatic tail are medialized to expose the renal fossa, while on the right, the liver is retracted and the right colon and duodenum are medialized. The retroperitoneal approach provides a more direct route to the kidney but can be less accessible due to trans-muscular route and a tighter working space, possibly limiting harvestable renal vessel length [19].

Renal and vascular measurements

Renal length should be measured using MPR images aligned to the principal and secondary orthogonal long axes, given that kidneys often lie obliquely, making coronal measurements unreliable [20] (Fig. 2b). Kidney length is a poor predictor of renal function [21, 22], so parenchymal or cortical volumes should also be reported—both are valid as proxies for renal function [23]—as assessed by the radiologist, a 3D technologist, or automated segmentation software. These volumes

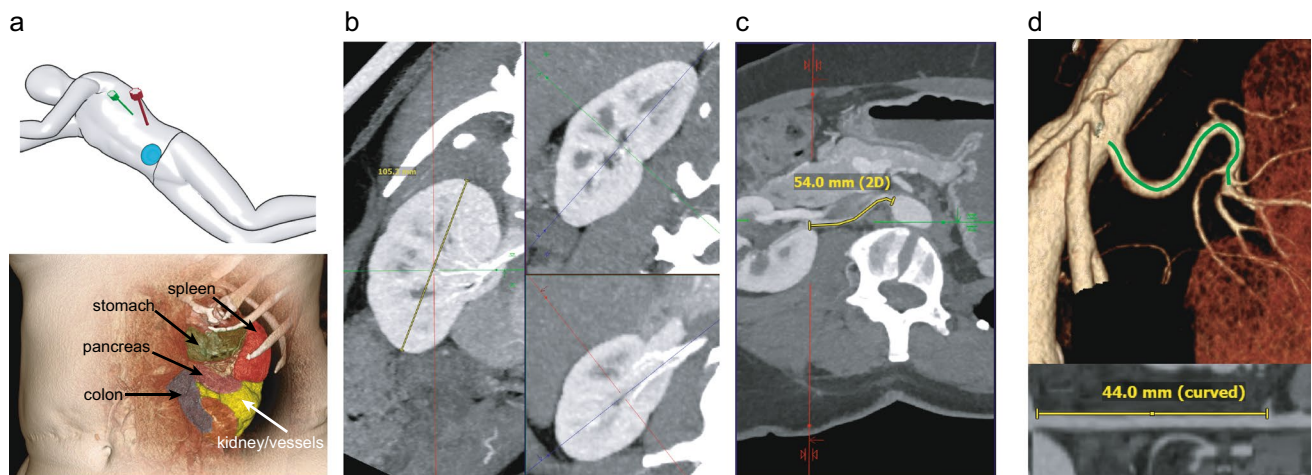


Fig. 2 Renal donor surgical and measurement techniques. **a** (above) Surgical approach of hand-assisted laparoscopic donor nephrectomy with suprapubic gel port (light blue) and camera/manipulation ports (green and red). (below) 3D CT reconstruction showing anterolateral view from a camera port angle with the kidney/renal vessels (yellow,

white arrow) partially obscured by viscera, requiring retraction. **b** Double-oblique MPR technique for measuring true renal length. **c** single plane curved measurement of a renal artery. **d** centerline curvilinear measurement of a tortuous renal artery

are used to estimate split function for triaging which kidney to donate, as current published data are too preliminary to support an absolute renal volume cutoff for transplantation [24–26].

The simplest method for measuring straight or short vessels involves aligning an MPR axis along the vessel length and taking a linear measurement. Curved vessel segments contained within a single plane can be measured using multiple segments or curved measurement tools found in many PACS toolsets. Slight deviations from a double oblique plane (within a vessel diameter) can be "flattened" using MIPs with minimal loss of accuracy (Fig. 2c). More tortuous vessel segments may require curved plane reconstruction or vessel centerline segmentation, either native in PACS or with dedicated 3D software (Fig. 2d).

Vascular measurements are tailored to the surgical approach and structures that may limit access to the renal vessels. The hilar arteries are measured from the aorta to the first segmental branch or until they enter the renal sinus, noting any earlier branching polar or capsular arteries (Fig. 3a).

The right renal vein length is determined from the union of its last segmental tributaries to the inferior vena cava (IVC). The left renal vein is measured four times: from each of two upstream points (the last segmental junction and the insertion of the left gonadal vein) to each of two points downstream (the IVC and the left aorta margin) (Fig. 3a). These measurements aid in planning both anterior and retroperitoneal surgical approaches.

Vascular anatomy and variants

Renal arteries are classified as hilar, entering the renal sinus to perfuse the kidney centrifugally, or polar, traversing the capsular cortex to perfuse centripetally (Fig. 3b). One to three renal arteries are typical, the largest defined as the main renal artery and smaller arteries as accessory regardless of their perfusion distribution. Early segmental branching is a variant with the first left renal artery segmental branch within 1 cm of the aorta or the right branch within 1 cm of the right IVC margin, which may require special surgical consideration. Retrocaval branching of the right renal artery should also be reported, as this limits renal artery manipulation from anterior approach (Fig. 3c).

Small upper polar arteries, especially smaller than 3 mm, may be sacrificed during transplant, but lower polar and hilar arteries are preserved to prevent ischemic ureteral necrosis, necessitating detailed description of lower pole accessory arteries including diameter (Fig. 3d). Lower polar arteries can arise from the distal aorta or, rarely, the iliac arteries (Fig. 3e). Also rarely, a precaval accessory artery can congenitally enlarge to become a main pre-caval right renal artery, decreasing retroperitoneal accessibility (Fig. 3f). Smaller capsular arteries, branching from renal arteries or the aorta and perfusing the capsule and adjacent tissues, are generally not preserved during transplant but must be accurately identified to avoid confusion with polar arteries.

The renal veins are more variable than the renal arteries due to their embryologic origin as bridges between the ventral (subcardinal) and dorsal (supracardinal) venous

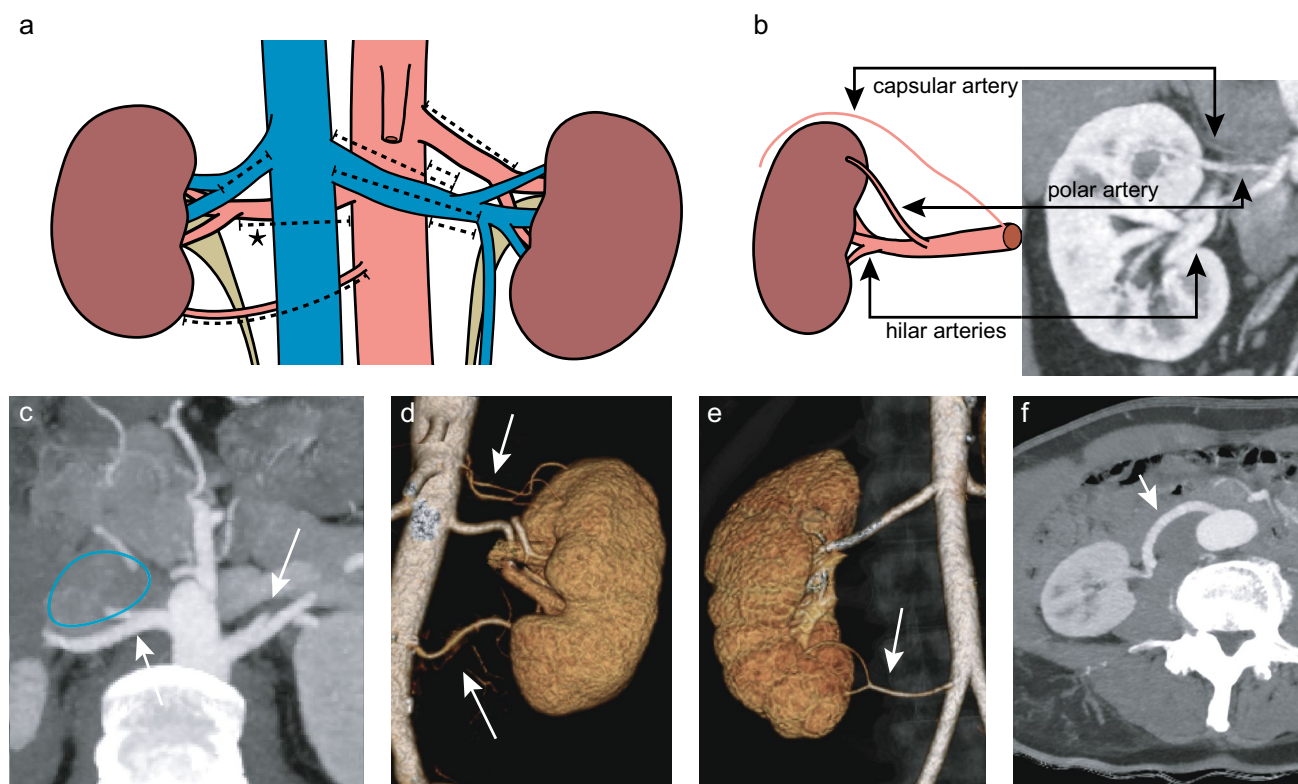


Fig. 3 Normal and variant vascular anatomy. **a** Arterial measurements from the aorta to first segmental branches, including a measurement from the right IVC margin (*) if there is < 1 cm early segmental branching. Venous measurements from the IVC to last segmental tributaries, IVC to left gonadal vein, and the same left landmarks to the left aortic wall. **b** Hilar, polar, and capsular artery diagram mirroring an example from a renal donor CT. Any of these can arise from the main renal artery or directly from the aorta. **c**

Olique transaxial CT of bilateral early arterial segmental branching, with a retrocaval variant on the right, the inferior vena cava contour shown in blue. **d** 3D volume rendering of left upper and lower polar arteries arising from the aorta, the lower pole artery important for ureteral perfusion. **e** 3D volume rendering of a right lower pole artery arising from the common iliac artery. **f** Precaval main right renal artery, important if retroperitoneal approach is a surgical consideration

systems [27]. The right renal vein typically forms from this ventral-dorsal anastomosis, leading to duplicated veins in about 1 in 6 patients [28]. The left renal vein originates from a collar of three venous bridges around the aorta: the intersubcardinal, intersupracardinal, and sub-supracardinal anastomoses. During normal development, the supracardinal anastomosis regresses, leaving the pre-aortic left renal vein composed of the remaining two anastomoses. A circum-aortic vein results if both intercardinal anastomoses persist, while a retroaortic vein forms if only the intersubcardinal anastomosis regresses. Any retroaortic component often communicates extensively with the lumbar/azygos system, also of supracardinal origin (Fig. 4a). The retroaortic vein can also insert low on the IVC or left iliac vein (Fig. 4b). These variants are well-reported, each occurring in about 1 in 30 patients. If circum-aortic, the diameters and lengths of each component should be reported, as should any lumbar plexus bridging veins > 5 mm in diameter that increase risk of hemorrhage. Other venous variations such as duplicated, left, or absent IVC should also be described in detail with

renal vein length measured to the first downstream central vein (Fig. 4c).

Renal vascular diseases

Atherosclerosis, the most common vascular disease in living renal donors, is increasingly common as donors over 55 years old have risen from 1 in 6 donors to 1 in 4 since 2011 [3]. Mild atherosclerosis does not preclude donation, but the more affected vessel is typically transplanted to reduce donor risk. An exception is made for coarse ostial calcification, which can complicate vessel closure during harvest, potentially causing hemorrhage or intimal laceration leading to thromboembolism (Fig. 5a). While there are no definitive criteria for donor eligibility based on vessel condition, radiologists should document the area percentage of cross-sectional narrowing at significant stenoses and measure the proximity of major calcifications to the arterial ostium.



Fig. 4 Renal vein variants. **a** Circumaortic left renal vein with equal pre- and retroaortic veins, a preserved sub-supracardinal anastomosis near the renal hilum (arrow), and a prominent retroaortic-lumbar communication (arrowhead). **b** A low retroaortic left renal vein inserting on the left common iliac vein (arrow) and via a large lumbar-IVC plexus vein (arrowheads). **c** A left renal vein inserting on a left IVC with empty retroperitoneal fat to the right of the infrarenal aorta

Fibromuscular dysplasia (FMD) is a non-inflammatory vasculopathy affecting medium to large vessels, predominantly diagnosed in middle-aged females, and most commonly involving the renal arteries [29]. This condition can lead to dissections, aneurysms, and stenosis and is likely underdiagnosed due to its imaging similarities with atherosclerosis and its tendency to be asymptomatic [30]. On CT, FMD can present as beading from alternating strictures and aneurysms, long smooth stenoses from thrombosed dissections, or fenestrated dissection flaps [31] (Fig. 5b). Mild FMD may be challenging to detect on CT [32] and can progress post-transplant to disease

threatening renal function, making it a conundrum when approving renal donors [33, 34].

Despite FMD's association with progressive renal insufficiency, data from FMD-positive donors show no significant long-term risks post-donation, likely in part due to selection bias for only mild cases approved for donation [35]. As incidental FMD in potential donors typically lacks symptoms and prior imaging, further assessment by a vascular specialist is crucial to manage the elevated risks of renal hypertension, stroke, and peripheral vascular disease associated with FMD and similar vasculopathies regardless of if the donor is committee-approved for surgery. Imaging appearances of FMD and inflammatory vasculitides such as polyarteritis nodosa can overlap but differ greatly by treatment, further emphasizing the need for specialist referral to further evaluate the vascular abnormality.

Collecting system variants

Small cadaveric studies have reported partial ureteral duplication in 2–5% of surveyed kidneys [36] and a larger study of candidate kidney donors showed single bilateral ureters in >98% of patients [37]. While ureteral duplication does not disqualify a donor, surgeons must be aware of the duplication to avoid damaging the common circulation and causing ischemic necrosis. With modern techniques, duplicated ureters are not linked to a higher risk of ureteral complications [38], although the surgeon may instead take the other kidney if there are no contraindications (Fig. 5c).

Nephrolithiasis

Historically, renal stones were often a contraindication for donation. Today, after recognizing that about 8% of asymptomatic adults have stones, most stones being small and few in number, and data show comparable graft survival rates in stone-bearing allografts, transplant teams are now less apprehensive of donor stones [39–41]. With a history of stones or stones on pre-donation CT, a potential donor may undergo 24 h urine collection analysis to assess the stone etiology and risk of future stones. In cases of unilateral cortical stones <5 mm, the affected kidney is preferentially donated and a ureteral stent is left in place post-transplant. Larger stones may be treated post-nephrectomy through ex vivo methods like back table pyelotomy, ureteroscopy, and lithotripsy before graft implantation [42]. Stones >5 mm may increase risk of recipient complications (Fig. 5d, e).

Calcified renal stones can be detected on non-enhanced, virtual non-enhanced dual-energy, or contrast-enhanced images depending on institutional preference. The location, number, and size of stones should be reported based on measurements using bone window/levels to prevent blooming artifact from inflating stone size. If cortical stones

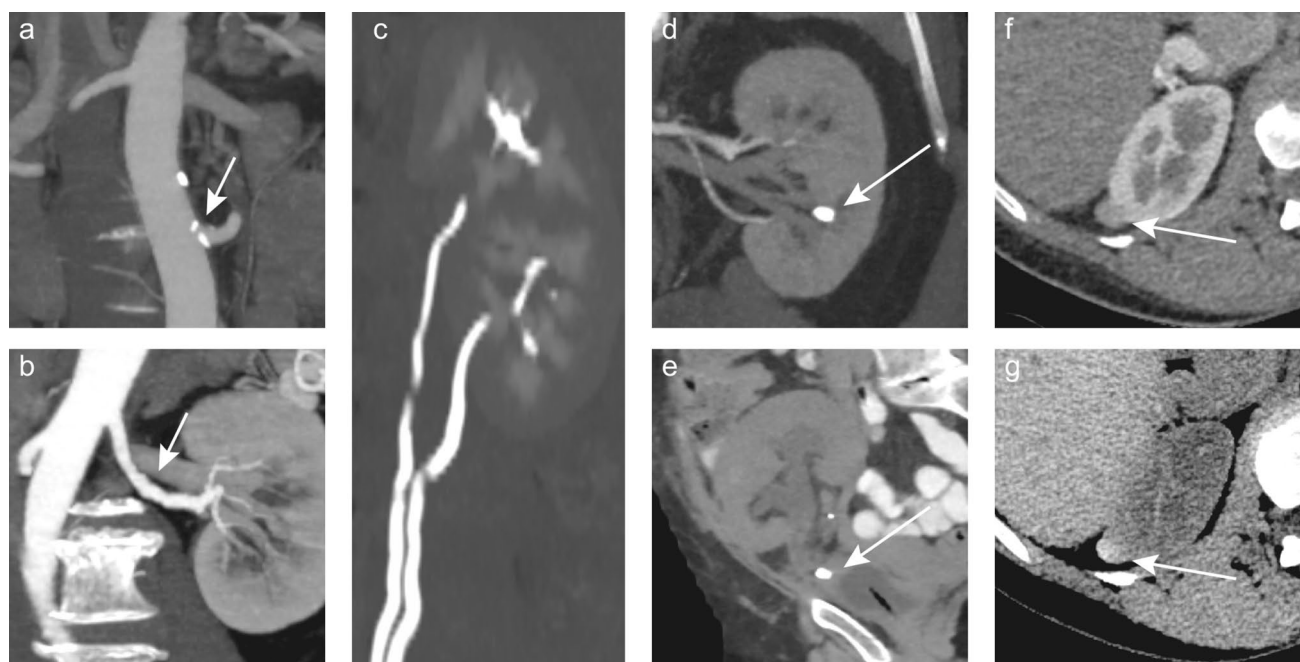


Fig. 5 Significant donor abnormalities. **a** Ostial atherosclerosis in a low main left renal artery. The patient was declined as a donor due to renal artery atherosclerosis. **b** Fibromuscular dysplasia with mid renal artery beading. The patient was declined as a donor due to severity and multifocality of FMD lesions. **c** Duplicated left ureters. The right kidney with normal vessels and a single ureter was instead donated.

d A 7 mm stone in the candidate donor kidney. The right kidney was not donated due to vascular anatomy. **e** The same stone 5 months later in the recipient, now migrated into and obstructing the ureter. **f** An indeterminate lesion, possibly a mass on arterial phase. **g** Virtual nonenhanced image from the CT showing intrinsic hyper attenuation in keeping with a proteinaceous cyst, later confirmed on MRI

are present, the ureters and bladder (if imaged) warrant extra scrutiny to exclude stone migration into the collecting system.

Renal lesions

Transplanting kidneys with small suspected renal tumors, including renal cell carcinoma, is a viable strategy to expand the donor pool to help meet the growing need for kidney transplants [43]. Studies show that the risk conveyed by possible cancer recurrence in recipients of kidneys with renal cell carcinoma is very low, particularly when compared to the risks of remaining on hemodialysis [44]. Renal masses, large cysts, and benign mixed solid-cystic lesions can be removed via partial nephrectomy on the back table prior to transplantation depending on lesion size and location as well as other donor and recipient characteristics.

Multiphasic CT for donor evaluation provides a non-enhanced, arterially enhanced, and variably a nephrogenic and/or excretory phase set of images for enhancement assessment, increasing specificity compared to routine monophasic CT (Fig. 5f, g). While MRI can further characterize indeterminate renal lesion contents, it is not commonly employed due to the low risks associated with transplanting kidneys after back-table mass excision. Further

diagnostic workup should be pursued if the patient is not approved for donation to minimize their own future risk.

Imaging techniques for the transplant recipient

Ultrasound imaging protocol

Ultrasound is the preferred initial imaging modality for evaluating renal grafts due to low cost, accessibility, noninvasiveness, and lack of radiation, which is crucial given the frequent and long-term monitoring needed for transplant recipients. It is used both routinely to establish a postoperative baseline and as needed for diagnosing complications. Examinations should be performed with a linear transducer, frequency range 2–18 MHz, and ideally with the patient well-hydrated. Standard protocol includes grayscale and color Doppler imaging of the graft and urinary bladder.

Grayscale assessment of the graft evaluates for size, parenchymal echotexture, hydronephrosis, and perinephric collections. Color Doppler assesses global graft perfusion and patency of the graft vasculature, including renal vein, renal artery at the hilum, mid-artery, and anastomosis, and

interlobar arteries at the polar and interpolar regions. The renal vein should show continuous flow with respiratory variability. The renal artery and interlobar arteries should exhibit a waveform with rapid systolic upstroke and continuous, antegrade diastolic flow. The renal artery resistive index (RI) quantifies distal resistance, calculated as (peak systolic velocity – end diastolic velocity)/peak systolic velocity, with normal values of 0.50–0.70. Finally, urinary bladder evaluation should document bladder distention, ureteral stent location (common postoperatively), a ureteral jet, and any filling defects.

Magnetic resonance and computed tomography considerations

CT and MRI are less commonly used in kidney transplant evaluation in part due to radiation and contrast induced nephropathy with CT and length of exam time and poor spatial resolution with MRI. However, both can be adjunctive techniques, providing extra diagnostic information after initial ultrasound evaluation if needed.

For example, if ultrasound cannot clearly assess graft vasculature due to bowel gas shadowing or poor sound penetration, CTA or MRA can measure vessel patency. CT protocols can be custom tailored to minimize risk of contrast-induced nephropathy. CT and MRI offer 3D reformatting capabilities and enable more comprehensive graft and surrounding anatomic evaluation, for example when assessing deep perinephric fluid collections. CT and MRI urography can also help localize urinary leaks, which is difficult with ultrasound.

Post-transplant complications

Complication timeline

Kidney transplantation can involve various complications such as iatrogenic injuries, vascular compromise, urinary issues, perinephric fluid collections, graft rejection, and infectious or neoplastic processes, many of which will be discussed later. Kidney transplant complications can be categorized by postoperative timing into early (including hyperacute and acute), intermediate, and late stages [45] (Fig. 6). Although complications may vary in onset timing, duration, and severity, this timeline framework helps narrow the differential diagnosis in evaluating kidney transplant pathology.

Perinephric collections

Perinephric collections are common posttransplant complications with typical postoperative timeline occurrence windows, from earliest to latest: hematoma, urinoma, abscess, and lymphocele. Although imaging features may overlap, correlating clinical factors with color Doppler ultrasound and adjunctive CT or MRI can confirm the diagnosis.

Perinephric hematomas typically occur from minutes to 5 days post-transplantation [45] but can occur anytime due to the graft’s vulnerable anterior pelvic position. Perinephric hematomas appear on grayscale ultrasound as complex collections with heterogeneous echotexture and possible internal septations or retractile clot (Fig. 7a) without internal vascularity on color Doppler. Noncontrast CT shows hyperattenuating blood products > 30 HU (Fig. 7b), and

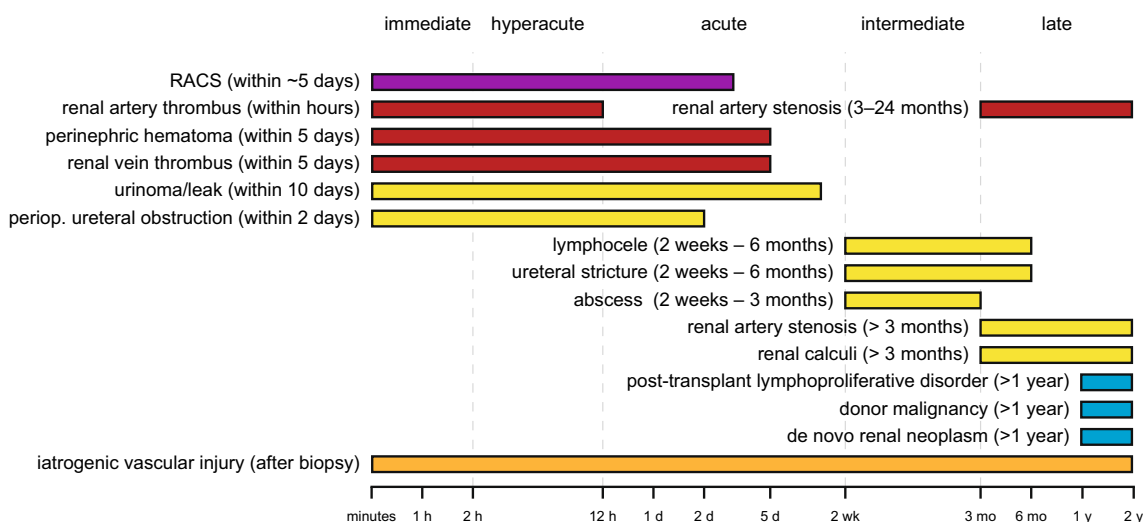


Fig. 6 Approximate timeline of post-transplant complications assessed by imaging. Adapted from [45]



Fig. 7 Imaging of perinephric collections. **a** Ultrasound of a collection (arrow) along the renal hilum 2 days after transplant and **b** CT the following day showing a hematocrit level (arrow) indicating a hematoma. **c** Delayed phase CT 3 days after transplant showing extravasation at the ureteral anastomosis. **d** Renal scintigraphy and **e**

cystogram performed on a different patient on post-transplant day 6 confirming an active urine leak (arrows). **f** Ultrasound 1 month after transplant showing a heterogeneous collection, followed by **g** CT at 4 months showing organization of the collection into an abscess

multiphase CTA may be used to find active bleeding sources in expanding hematomas. Most perinephric hematomas resolve spontaneously, though some require treatment if they significantly compress the graft, as discussed below.

Perinephric urinomas are collections of urine leaking from the renal collecting system, typically from the renal pelvis, ureter, or ureteral anastomosis. This complication

often occurs within 10 days of transplantation, due to inadequate blood supply to the ureter, urinary tract pressures from obstruction, or caliceal/forniceal rupture [45, 46]. Urinomas typically are anechoic collections between the graft and bladder on grayscale ultrasound. Excretory phase CT/MRI can confirm urine leaks by showing extraluminal contrast (Fig. 7c). Retrograde urography and nuclear renal

scintigraphy can also detect urine extravasation but are less commonly used (Fig. 7d, e). Conservative measures like nephrostomy and stenting often resolve leaks, but severe cases may necessitate surgical interventions such as ureteral reimplantation, psoas hitch, bladder flap, or ileal conduit urinary diversion.

Perinephric abscesses often arise from graft infections or extend from abdominal wall/peritoneal infections weeks to months post-transplantation. Fever and leukocytosis with a perinephric fluid collection suggest abscess, confirmed with ultrasound showing a thick-walled, complex collection with septations, debris, and wall hypervascularity on color Doppler. Contrast-enhanced CT or MRI can assess the full extent of the collection and its impact on surrounding pelvic structures (Fig. 7f, g), possibly beyond the depth visualized by ultrasound. Like abscesses elsewhere, treatment includes systemic antibiotics and percutaneous drainage.

Lymphoceles are the most common perinephric collection, typically appearing along lymphatic pathways 2 weeks to 6 months post-transplant. They appear as thin-walled anechoic collections with internal septations on ultrasound and are usually asymptomatic but may grow enough to cause mass effect. CT shows a well-circumscribed and homogeneously hypoattenuating collection, while MRI shows T1 hypointense and T2 hyperintense fluid and often thin septations within the collection.

Graft compression

Renal allograft compartment syndrome

Renal allograft compartment syndrome (RACS) is a rare but serious cause of early graft dysfunction or loss from intracompartmental hypertension in the confined iliac fossa which induces ischemia, often requiring immediate reoperation hours to days after transplant. Risk factors include extrinsic vascular compression or mismatch between donor renal size and recipient body habitus. Imaging shows edematous graft parenchyma with reduced or absent cortical flow and high-resistance waveforms, often with a perinephric hematoma (Fig. 8a, b).

Subcapsular hematoma

Subcapsular hematomas, different from perirenal hematomas, are collections of blood between the renal capsule and parenchyma, which can compress the parenchyma causing graft dysfunction and hypertension from renin release (eponymously the “Page phenomenon”), even if small. Often iatrogenic, this complication usually follows a percutaneous biopsy and presents with pain, new hypertension, graft dysfunction, and a variable hemoglobin

drop. Imaging typically shows a complex, crescentic, heterogeneous collection on ultrasound and hyperdense on noncontrast CT (Fig. 8c). Spectral Doppler may show elevated resistive indices and decreased or reversed diastolic flow in otherwise normal hilar vasculature (Fig. 8d). Subcapsular hematoma with renal dysfunction or hypertension necessitates prompt surgical evacuation to save the graft.

Rejection

Graft rejection is identified and classified using the evolving Banff criteria [47], with a simpler, less specific classification—hyperacute, acute, and chronic rejection—used in clinical discussions. Hyperacute rejection, a rare event caused by pre-formed antibodies, leads to immediate microvascular occlusion and graft loss. Its imaging is non-specific and may show signs on ultrasound such as high resistance waveforms and parenchymal edema, much like with primary vascular complications. Acute rejection, typically occurring around one-week post-transplant and mediated by T-cells, also presents non-specific imaging findings such as graft edema and vascular compromise. Chronic rejection can occur over many months or years with ultrasound findings typical of chronic renal disease, such as a thinned and hyperechoic cortex.

Ultimately, the radiologist provides information for the transplant team to integrate with clinical and laboratory data to determine if abnormalities are due to rejection or a surgically intervenable cause. Histology is a valuable tool in making this determination, but the value of graft biopsy should be judiciously weighed against the risk of iatrogenic complications.

Iatrogenic complications

Intraoperative or percutaneous renal graft biopsies are common in posttransplant diagnostic workups. Standard technique uses a 16-gauge or 18-gauge biopsy device to target the renal cortex, usually the lower pole, via a cortical tangential approach to minimize injury [48]. Optimal samples contain renal cortex without medullary tissue. Major iatrogenic complications, like bleeding requiring blood transfusion or vascular embolization, are rare (0.24–4%) [45, 49–52] when performed by a subspecialized proceduralist. Less severe complications like arteriovenous fistulae and pseudoaneurysms are more common. These complications are best evaluated by ultrasound, though CTA or MRA are valuable for detailing vascular anatomy before any endovascular therapy.

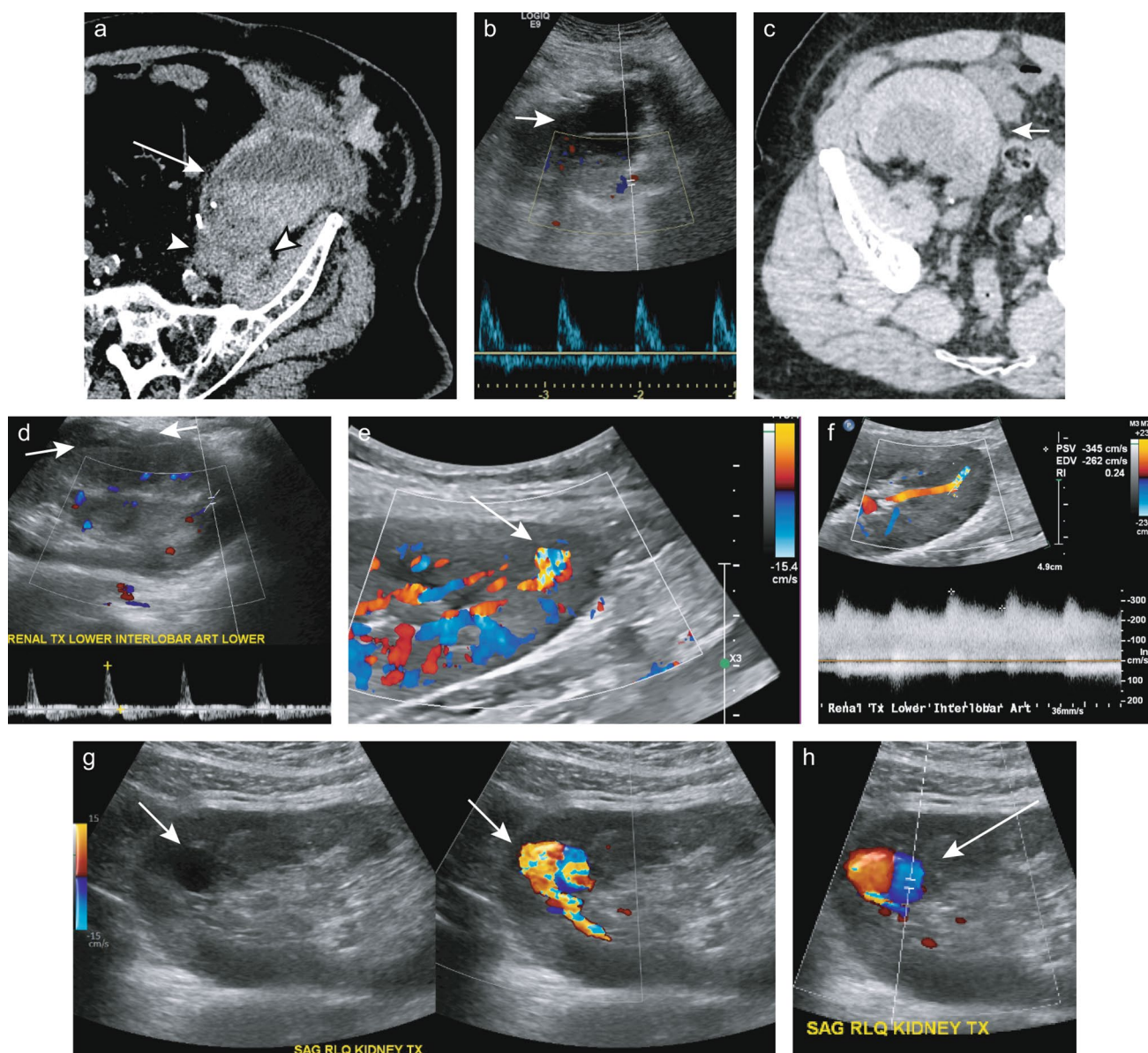


Fig. 8 Vascular injuries and consequences. **a** CT performed day-of-surgery with a large, layering hematoma (arrow) displacing and compressing the graft kidney posteriorly (arrowheads). **b** Immediately subsequent ultrasound showing the hematoma (arrow) and high resistance waveform with diastolic reversal in the renal arcuate arteries, indicating RACS. **c** Non-enhanced CT after ultrasound-guided cortical biopsy showing a crescentic, hyperattenuating collection compressing the renal cortex and effacing

the sinus fat. **d** Ultrasound of the same patient showing the crescentic hematoma (arrows), sinus fat effacement, and high resistance arterial waveform with diastolic flow reversal. **e** Doppler ultrasound of a nidus of turbulent, high-velocity flow at a cortical biopsy site. **f** Doppler of an efferent vein showing high-velocity, arterIALIZED flow of an AVF. **g** Grayscale and color Doppler ultrasound with a hypoechoic structure at a biopsy site with turbulent cortical flow and **h** a yin-yang, or Pepsi sign of a pseudoaneurysm

Iatrogenic arteriovenous fistula

Arteriovenous fistulae (AVFs) are abnormal connections between an artery and a vein, bypassing the capillary bed. This iatrogenic complication occurs in up to 15% of patients [53], is commonly asymptomatic, and may be detected incidentally on routine post-biopsy ultrasounds. Larger, clinically significant AVFs may cause hematuria,

hypertension, graft dysfunction, and in severe cases, high-output heart failure. Small AVFs are commonly occult on grayscale imaging but are revealed by disorganized blood flow extending beyond normal vessel margins on color Doppler. Ultrasound may also show a feeding artery with a high-velocity, low-resistance waveform (Fig. 8e) and a vein with aliasing and arterIALIZED waveforms (Fig. 8f). Since most iatrogenic AVFs are small, management is largely

observation and follow-up imaging, as 77% naturally regress [54]. Transcatheter embolization may be necessary for larger or symptomatic AVFs.

Iatrogenic pseudoaneurysm

Unlike AVFs, pseudoaneurysms are a contained bleed or vascular injury affecting all three layers of the artery. Iatrogenic pseudoaneurysms may be asymptomatic if small, but may cause hematuria and graft dysfunction when larger. Sonographically, small pseudoaneurysms may resemble AVFs, but larger ones appear as simple or complex cystic structures on grayscale (Fig. 8g) with to-and-fro flow and the yin-yang sign/Pepsi sign swirls on color Doppler (Fig. 8h). Pseudoaneurysm treatment varies by size; those under 2 cm are usually managed conservatively, while larger

pseudoaneurysms often require transcatheter embolization [45].

Renal artery complications

Renal artery stenosis

Transplant renal artery stenosis is the most common vascular complication after kidney transplantation (1–3%) [55], typically occurring 3 months to 2 years post-transplant and presenting with refractory arterial hypertension, sometimes with declining graft function. Classic ultrasound features of transplant renal artery stenosis include focally elevated peak systolic velocity (PSV, > 340–400 cm/s) with turbulent flow aliasing at the stenosis (Fig. 9a) and main renal artery to external iliac artery PSV ratio elevation (> 1.8). Downstream

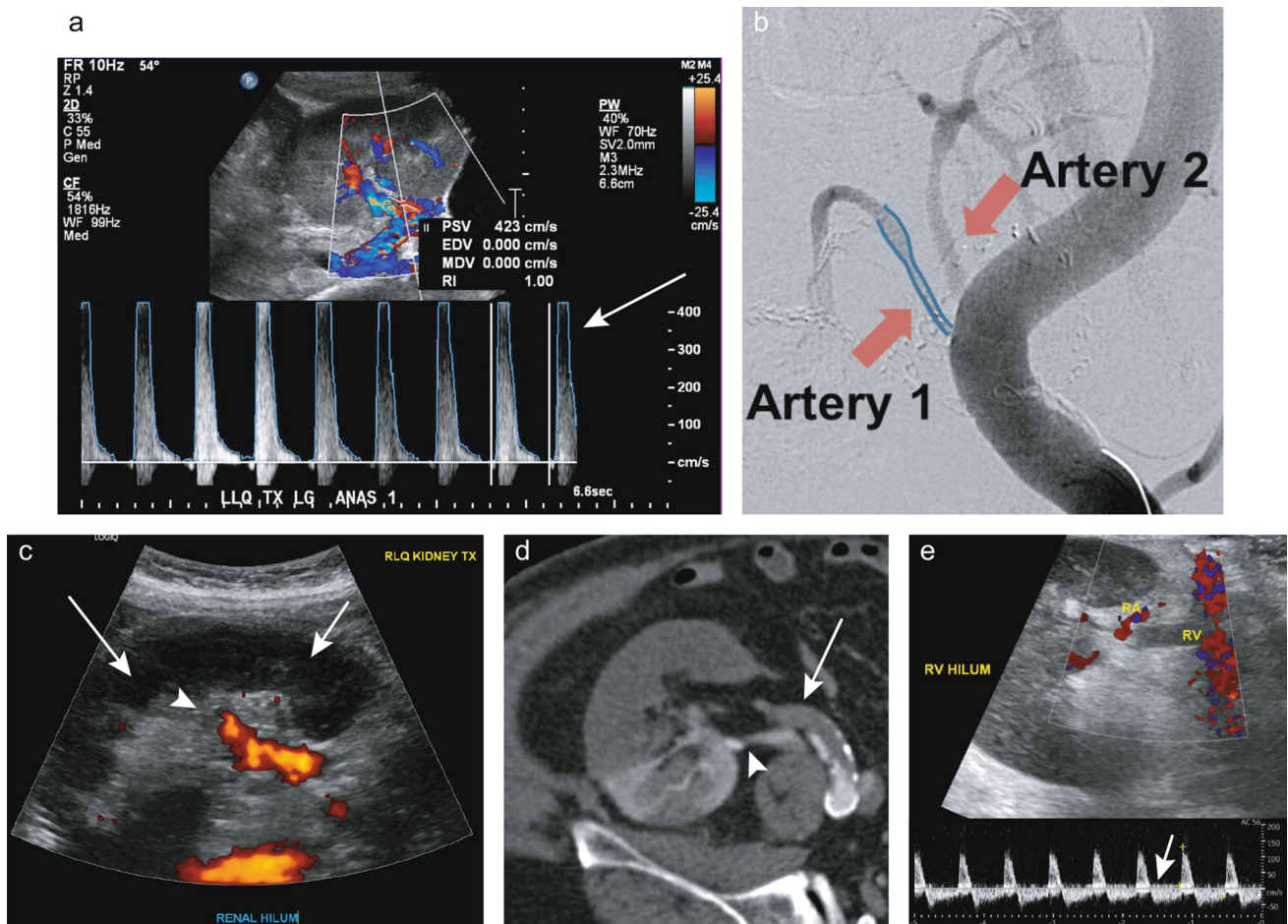


Fig. 9 Vascular compromise. **a** Elevated peak systolic velocity > 400 cm/s (arrow) with resistive index of 1.00, followed by **b** digital subtraction angiography showing stenosis of one of the paired renal arteries while artery 2 is widely patent. **c** Renal artery thrombosis with power Doppler with absent flow in the renal parenchyma (arrow) and abrupt cutoff of the renal artery and minimal

arterial flow in the renal sinus (arrowhead). **d** CTA of renal artery thrombosis (arrow) with absent enhancement from the lower pole to the interpolar kidney, sparing the upper pole via a small accessory polar branch (arrowhead). **e** Renal vein thrombosis with absent flow in the vein and reversal of diastolic flow in the adjacent artery (arrow)

renal and segmental arteries often show *tardus parvus* waveforms with low RIs (<0.50) on spectral analysis.

A common pitfall is postoperative edema at the arterial anastomosis that transiently elevates PSV, mimicking arterial stenosis. Therefore, early postoperative elevated PSV requires careful interpretation of downstream spectral waveforms and short-term follow-up ultrasound to confirm resolution. When confirmation of renal artery stenosis suspected by ultrasound is needed, options include CTA/MRA or digital subtraction angiography using iodinated contrast (Fig. 9b) or carbon dioxide. Angiography is useful diagnostically and therapeutically allowing interventions like transluminal angioplasty and stent placement. In complex or refractory cases, surgical intervention may be required to repair the stenosis.

Renal artery thrombosis

Transplant renal artery thrombosis, a rare but serious complication post-kidney transplantation (<0.5% incidence) [56], typically occurs within hours of transplant with sudden cessation of urine output, increased hypertension, and possibly lower quadrant pain from an inflammatory reaction to the ischemic graft. Potential causes of transplant arterial thrombosis include anastomotic occlusion, hyperacute graft rejection, arterial compression or kinking, vasculitis, or an intraluminal arterial flap.

Sonographic signs of transplant renal artery thrombosis are segmental or global loss of parenchymal perfusion and absent flow in transplant renal vasculature on Doppler analysis (Fig. 9c). Rarely, ultrasound is equivocal and CTA/MRA may be used, showing nonopacification of the transplant vessels and diminished or absent contrast enhancement of the parenchyma (Fig. 9d). CTA/MRA also allows greater assessment of transplant vascular anatomy, including accessory vessels. Timely diagnosis is vital to prevent graft loss which typically requires immediate surgical reoperation.

Renal vein complications

Renal vein thrombosis

Transplant renal vein thrombosis, a major cause of early graft failure, occurs in less than 5% of adults and up to 8.2% of pediatric patients [45]. It typically develops within the first 5 days post-transplant, peaking at 48 h, presenting with sudden pain, oliguria/anuria, and possible ipsilateral lower extremity edema from thrombus propagation to the iliac veins.

Renal vein thrombosis appears as edematous or engorged renal parenchyma with lost corticomedullary differentiation on grayscale ultrasound. Typical color Doppler findings

include absent flow in the vein and reversed arterial diastolic flow (Fig. 9e), however this flow reversal is nonspecific, also seen with complications including graft torsion, subcapsular hematoma compression, severe rejection, or acute tubular necrosis. Like arterial thrombosis, prompt diagnosis and treatment of renal vein thrombosis are critical for graft salvage. Acute renal vein thrombosis requires open surgical thrombectomy but still results in a high rate of graft loss. Delayed partial thrombosis may be treated with endovascular thrombolysis, albeit with a higher bleeding risk [57].

Graft tumors and mimics

Transplant recipients, due to ongoing immunosuppression, have higher risk of all malignancies, though de novo graft renal neoplasms are exceedingly rare with incidence <0.5% [58]. Careful interpretation of transplant renal ultrasound is crucial to detect small renal neoplasms, appearing as isoechoic or hypoechoic masses with vascularity on color Doppler. CT detection can be challenging due to frequent noncontrast exams to preserve renal function (Fig. 10a); MRI is often more effective for both detection and characterization. Focal rejection, an important graft tumor mimic, may appear as a pseudomass on any modality (Fig. 10b, c).

Kidney transplant recipients are over twice as likely to be diagnosed with primary malignancies than the general population due to several factors, including oncogenic virus infections, carcinogenic medications, suppressed immune anti-tumor response, and enhanced detection from frequent imaging. The most significantly increased risks are for squamous cell carcinoma and Kaposi sarcoma, though most organs and tissues outside the brain are susceptible [59]. Post-transplant lymphoproliferative disorder (PTLD), typically associated with Epstein–Barr virus, presents with a range of imaging findings from nonspecific lymphadenopathy and masses in various tissues, including the graft (Fig. 10d) [60]. Early detection by imaging can lead to prompt immunosuppression-reduction and increased odds of response without further need for chemotherapy [61].

Future considerations

Radiologists are encountering an increasing number of kidney transplant patients, a trend that will continue as kidney transplantation follows the growing demand. Even more, we may be entering a new phase in transplant medicine highlighted by recent advancements in porcine renal xenotransplantation, with the first and second xenotransplants into humans performed at Massachusetts General Hospital and New York University, respectively [62, 63]. Momentum is building in light of new genome

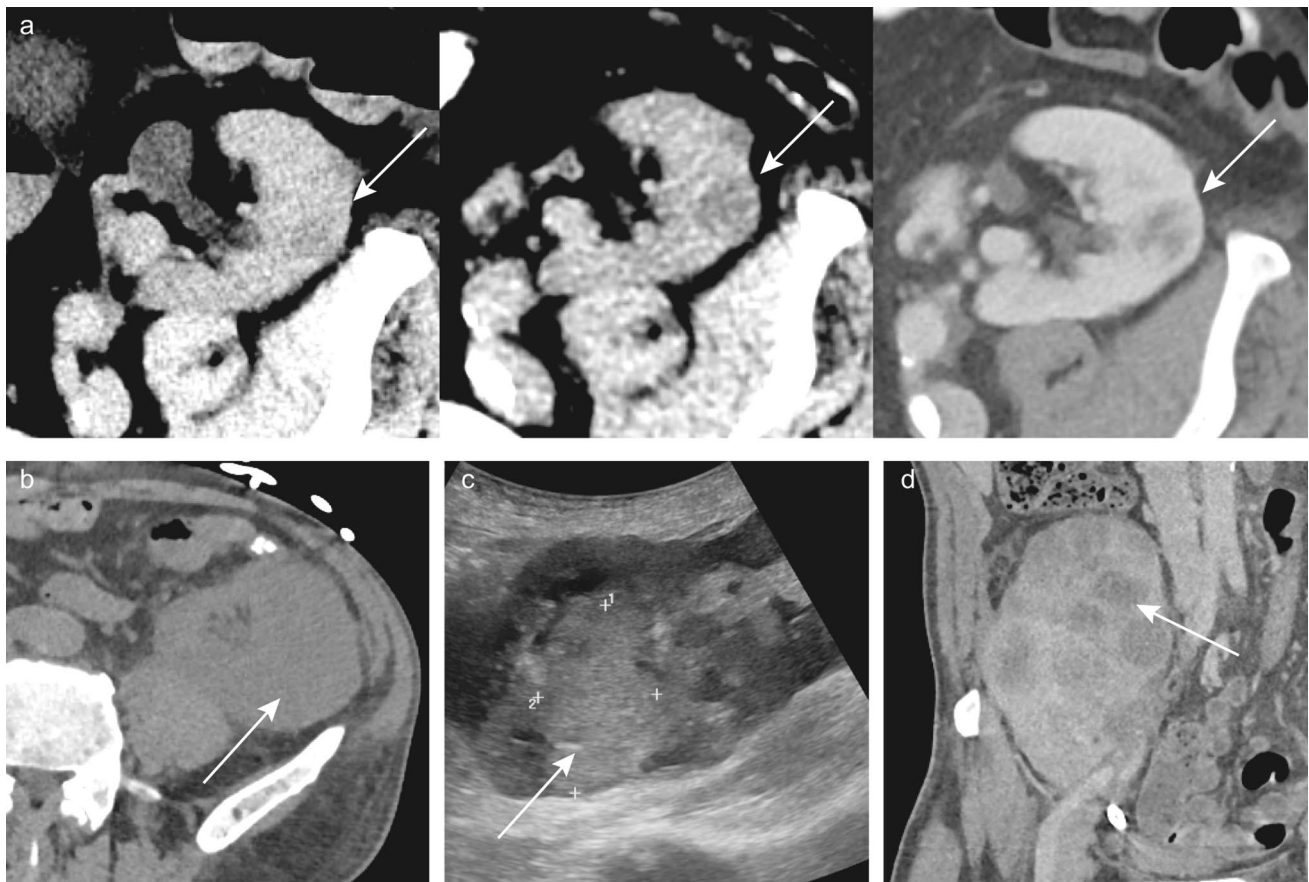


Fig. 10 Post-transplant mass lesions. **a** Growth of a solid mass in the allograft unnoticed among cysts on noncontrast CTs for two years before showing enhancement on contrast-enhanced CT. Biopsy showed oncocytoma. **b** Mass-like expansion of the graft on non-contrast CT suspicious for neoplasm, confirmed on **c** follow up

ultrasound as an irregular mass. Biopsy showed focal rejection and no malignancy. **d** Post-transplant lymphoproliferative disorder (PTLD) presenting as numerous masses infiltrating the right lower quadrant renal allograft (arrow)

editing technologies [64]; xenografts will possibly provide a large source of future donor kidneys. The imaging characteristics of normal and abnormal xenografts have yet to be reported and will be one of many future research opportunities in the field.

Radiomics, advanced MRI techniques, and machine learning together are expanding the radiologist's toolkit in assessing renal graft health. Early data suggest multiple diffusion-weighted imaging parameters predict graft dysfunction independently of urine and blood chemistries, causing a push for new research in the field [65, 66]. Early reports of machine learning being used to combine clinical data with the large volume of serial imaging data are promising, further indicating the need for more research in the area [67].

Conclusions

Kidney transplantation is increasing in the United States and abroad, emphasizing the need for radiologists to know best practices of donor and post-transplant recipient evaluation. Transplant patient assessment is multidisciplinary; radiologists gain from knowing the surgical approach of donor nephrectomy and implications of post-operative complications to better communicate with referring transplant surgeons. Knowledge of the clinical manifestations of imaging findings aids radiologists in adjusting pre-test probabilities and prompting additional imaging if initial studies are inadequate. With the recent introduction of genetically

modified renal xenografts as a promising solution to the organ shortage, familiarity with transplant imaging may become further vital to the radiologist.

Declarations

Conflict of interest RJG, JD, and LAD, have no financial or other disclosures. AK- Research Grant – GE Healthcare, PanCAN, Bayer; Consultant- Bayer.

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