Tuberculosis of the genitourinary system—Urinary tract tuberculosis: Renal tuberculosis—Part I

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Abstract

Tuberculosis (TB) remains a worldwide scourge and its incidence appears to be increasing due to various factors, such as the spread of human immunodeficiency virus (HIV) and acquired immunodeficiency syndrome (AIDS). The insidious onset and non-specific constitutional symptoms of genitourinary tuberculosis (GUTB) often lead to delayed diagnosis and rapid progression to a non-functioning kidney. Due to hematogenous dissemination of TB, there is a potential risk of involvement of the contralateral kidney too. Imaging plays an important role in the making of a timely diagnosis and in the planning of treatment, and thus helps to avoid complications such as renal failure. Imaging of GUTB still remains a challenge, mainly on account of the dearth of literature, especially related to the use of the newer modalities such as magnetic resonance imaging (MRI). This two-part article is a comprehensive review of the epidemiology, pathophysiology, and imaging findings in renal TB. Various imaging features of GUTB are outlined, from the pathognomonic lobar calcification on plain film, to finer early changes such as loss of calyceal sharpness and papillary necrosis on intravenous urography (IVU); to uneven caliectasis and urothelial thickening, in the absence of renal pelvic dilatation, as well as the hitherto unreported 'lobar caseation' on ultrasonography (USG). Well-known complications of GUTB such as sinus tracts, fistulae and amyloidosis are described, along with the relatively less well-known complications such as tuberculous interstitial nephritis (TIN), which may remain hidden because of its 'culture negative' nature and thus lead to renal failure. The second part of the article reviews the computed tomography (CT) and MRI features of GUTB and touches upon future imaging techniques along with imaging of TB in transplant recipients and in immunocompromised patients.

Key words: Renal tuberculosis; tuberculous interstitial nephritis; intravenous urography; ultrasonography; lobar caseation; uneven caliectasis

Introduction

Tuberculosis (TB) is the commonest worldwide cause of mortality from infectious diseases with nine million new cases and two million fatalities per year. Approximately 95% of cases occur in developing countries. In India, more than 1000 lives are lost every day due to TB despite the availability of modern diagnostic aids and treatment. The resurgence of TB has been noted in both endemic and non-endemic regions, mainly due to increased migration, the human immunodeficiency virus (HIV) pandemic, and the emergence of drug-resistant strains of Mycobacterium tuberculosis (MTB). A relative increase in extra-pulmonary TB has been reported due to a significant decline in pulmonary tuberculosis (PTB) and an only modest decline in extra-pulmonary TB.
The genitourinary tract is a primary target of hematogenous infections[8] and is the most common site of extra-pulmonary TB,[9] comprising 14-41% of the same.[10,11] Genitourinary tuberculosis (GUTB), a term coined by Wildbolz in 1937,[12] is a worldwide disease, but shows a more destructive behavior in developing countries. The kidney is the most common site of GUTB. An increased incidence of extra-pulmonary TB has been noted in acquired immunodeficiency syndrome (AIDS).[13]

Worldwide, 15% of TB patients are co-infected with HIV, and in HIV-endemic areas, as many as 75% of patients with GUTB are co-infected with HIV.[14,15] Hence, HIV-positive patients should be tested for TB, and patients with newly diagnosed TB should be assessed for HIV infection.[16] In fact, with the improving survival rates of AIDS patients, we can expect an increase in the incidence of urinary tract TB.[6]

Patient Population

GUTB usually affects adults between the second and fourth decades of life and is reported as being rare in children[17] and in the fifth and sixth decades. A mean age of 40.7 years (range: 5-90 years) has been noted.[8] There is often a long latent period (5-40 years) between the original pulmonary infection and the appearance of clinical renal disease,[6] which is probably why renal involvement is rare before the age of 20 years.

The youngest reported case of urinary tract tuberculosis (UTB) was 2 years old.[18] In India, it is not uncommon to see children with UTB. We have seen TB autonephrectomy in a—six year old girl.

Symptoms

UTB has an insidious onset, no specific symptoms and atypical presentations,[16] which lead to difficulty and delay in diagnosis.[1,9,19-21] Most patients present with local symptoms such as frequent voiding; dysuria,[21] pyuria,[17] back, flank, or abdominal pain,[22-24] and microscopic or macroscopic hematuria.[17] Systemic symptoms of fever, weight loss, and anorexia are less common.[17,22-24] Hematuria and culture-negative pyuria may be seen at urine analysis.[25] Urine analysis of sediment from a 24-hour specimen for acid-fast bacilli (AFB) is positive in 80-90% of cases of TB. Urine culture requires 6-8 weeks for diagnosis and there is a 10-20% false-negative rate.[19] Laboratory findings do not reveal the site or extent of disease, knowledge of which is imperative for further management. Imaging thus plays a major role, both in the initial workup as well as during follow-up.

A negative chest radiograph and tuberculin test cannot exclude the diagnosis of extra-pulmonary TB.[9] Only 36.5% of patients with UTB have a previous diagnosis of TB, or abnormal imaging studies.[5] Evidence of active TB or an abnormal chest radiograph is present in less than 50% of cases.[6,7] Only 20-30% of UTB patients will have a previous history of PTB; an additional 25-50% will have radiographic evidence of prior subclinical PTB.[16,26]

This article reviews the imaging findings on the basis of which an accurate non-invasive diagnosis of renal TB can be made.

Pathogenesis

Causative organisms

MTB, an obligate pathogen, is the usual cause, occasionally, Mycobacterium bovis,[27] and Mycobacterium avium intracellulare (MAIC). MAIC is transmitted via natural water sources, indoor water systems, pools, and hot tubs,[28-30] and can cause disseminated disease, particularly in immunosuppressed individuals, including renal transplant patients.[31]

Spread of tuberculosis to the urinary tract

Hematogenous dissemination of MTB occurs from a primary TB focus within the lungs, bone, or other organs and can involve both kidneys.[32] Bacille Calmette-Guerin (BCG)—a live, vaccine strain—can cause renal lesions via reflux, in 0.1% of patients undergoing intravesical instillation of BCG for the treatment of bladder cancer.[33,34]

Spread of infection at the time of primary tuberculosis

During the primary infection, alveolar macrophages phagocytose one or more mycobacteria lodged within an alveolus. Because of their high resistance to destruction, these virulent mycobacteria multiply within the macrophages, which may eventually result in lymphatic and hematogenous dissemination, with seeding of tubercular bacilli throughout the body.[35] The bacilli have stringent growth requirements and generally tend to proliferate only at a few sites, the kidney being one.[27]

The kidneys, and possibly the prostate and seminal vesicles, are often the primary sites of GUTB. All other genital organs, including the epididymis and bladder, become involved by ascent or descent of MTB from a source elsewhere in the genitourinary tract.[9] In most patients, acquired cellular immunity develops and there is inhibition of bacterial multiplication and containment of the disease by the formation of microscopic granulomas.[35] Healing may also occur as a result of anti-TB chemotherapy administered to control the clinically active focus.

In immune-competent patients, these granulomas heal or remain stable for many years.[36] As they usually do not exceed 3 mm in diameter,[37] they are difficult to visualize and may be missed unless carefully looked for.
Re-activation of tuberculosis
If there is a breakdown in host immunity, re-activation or re-infection occurs. It has been reported that a reduction in serum 25-OH-vitamin D levels leads to fall in cell-mediated immune defenses, which can result in activation of latent tuberculosis.[37,38] Hence, it would be worthwhile checking and restoring 25-OH-vitamin D levels in malnourished individuals with TB.

One or more tubercles may enlarge after years of inactivity.[39] This latent period varies considerably and may extend from 5 to 40 years.[22,25]

In the kidneys, the bacilli lodge in the periglomerular capillaries where they form microscopic granulomas, which may later grow into macroscopic granulomas.[39] This occurs bilaterally.

The morphology of the lesions depends on the site of infection, the virulence of the organism, and the immune status of the patient.[40] In immune-competent patients, granulomas are well formed and caseous necrosis is frequently seen. Various types of tuberculous involvement can occur in different areas of the same kidney.[41] or even in both kidneys. However, severe affection is more commonly unilateral. This is one possible cause of delay in patient presentation, leading to irreparable unilateral loss of renal function.

Parenchymal changes
The medullary portion of the renal parenchyma is usually spared initially. For unknown reasons, the upper and lower poles of the kidney are more commonly affected than other areas.[17] Cortical granulomas enlarge and coalesce, with the bacilli spilling down the nephrons and getting trapped in the narrow segment of the loop of Henle, establishing new foci of infection within the renal pyramid. These papillary lesions caseate and cavitate, frequently forming ulcero-cavernous lesions as they erode into the pelvicalyceal system (PCS).[35] Extensive papillary necrosis may develop with the formation of frank cavities and destruction of the adjacent renal parenchyma. These may also extend into the collecting system[24] via rupture, or cause parts of the papillae to become necrotic and slough.[41] A mass lesion may result from massive destruction and coalescence of granulomas, if they do not rupture into the adjoining calyx.[42] Alternatively, these granulomas may coalesce and form cavities after liquefaction. Hypercalcemia may occur, usually secondary to abnormal cortisol production by granulomatous tissue.[43] Although calcification is unusual in the early stages of the disease, nearly every end-stage tuberculous kidney contains calcification. Rarely, UTB can present as a well-circumscribed multi-septated cystic renal mass.[44] In immunosuppressed individuals, the granulomas may be less well formed and caseous necrosis is seen less frequently.[27]

Pelvicalyceal system changes
When bacilli are shed into the urine, the disease spreads antegradely to involve the urothelium of the renal pelvis, ureter, bladder and, at times, the adjacent genital tract.[17] Infection in the walls of the calyces, pelvis, and ureter produces significant inflammatory mucosal thickening, a commonly overlooked imaging finding. Single or multiple calyces may be involved in one or both kidneys. Microscopic granulomas may form here too. Ulceration soon follows.

In advanced disease, in addition to loss of parenchyma by caseation, intra-renal scars and strictures lead to obstruction and dilatation of segments of the PCS. Strictures are more common at sites of normal narrowing, such as the calyceal neck, the pelvi-ureteric junction, and the ureterovesical junction. Early clarring is apparently reversible by appropriate steroid treatment, but end-stage fibrotic strictures are irreversible. Urinary obstruction from strictures along with renal parenchymal caseation destroys all or part of the kidney. The pattern of destruction depends on the relative rates of progression of parenchymal disease and urinary-flow obstruction. Parenchymal caseation, necrosis, and calcification may predominate, which causes the kidney to be destroyed [Figures 1 and 2]. Alternatively, obstruction may predominate, in which case massive hydrourephrosis or hydroureteritis may be the final stage. TB of the kidney thus reflects competing processes: (a) The destructive effects of the bacilli, leading to ulceration, cavitation, and fistulization and (b) the host’s secondary defense and healing mechanism leading to the formation of granulomas along with fibrosis, calcium deposition, and strictures, which may worsen the obstruction causing progressive renal dysfunction.[28] The final outcome is thus extremely variable.[45]

Usually, however, both processes occur concurrently and may lead to a non-functioning, calcified kidney of any size; this process is called autonephrectomy. Nephrectomy has been advised to remove the trapped dormant bacilli in such autonephrectomized kidneys.[46]

Tuberculous interstitial nephritis (TIN)
Occasionally, TB can affect the kidney more insidiously, causing TIN[27,47-49] which, if untreated, progresses to renal failure. Rupture of the bacilli into the interstitium can lead to isolated interstitial disease, without persistent pyuria, hematuria, or identifiable AFB in the urine, leading to diagnostic dilemmas. Histology reveals chronic tubulointerstitial nephritis, usually with granuloma formation, which may or may not be associated with caseation. With appropriate staining, AFB are identifiable on histology. Evidence of coexisting TB elsewhere may be the only clue to TB being the cause of the falling glomerular filtration rate (GFR). If the diagnosis has been made while useful renal function still remains, it may be possible to arrest the fall in GFR or even produce improvement, using a combination
of anti-tubercular treatment and corticosteroids.\textsuperscript{[27]} TIN has also been reported as a complication of intravesical BCG instillation for bladder cancers.\textsuperscript{[50,51]}

Renal failure

The overall incidence of renal failure reported in the literature is 24\%.\textsuperscript{[52]} There are three mechanisms by which TB can cause renal failure:

a. Renal parenchymal infection causing obliterative endarteritis with extensive dystrophic calcification or secondary renal amyloidosis, both leading to renal impairment.\textsuperscript{[53]}

b. Post-obstructive atrophy secondary to multiple strictures.\textsuperscript{[54,55]}

c. Insidious TIN destroying the renal parenchyma. This is a form of culture-negative renal TB. The only clue could be echogenic kidneys on US along with signs of TB elsewhere.

Renal tuberculosis with other renal diseases/lesions

There have been a number of case reports of TB associated with various forms of glomerulonephritis,\textsuperscript{[27]} including a case report of miliary TB complicated by focal proliferative glomerulonephritis, in which immune deposits were present, but no granulomas.\textsuperscript{[56]}

\textbf{Figure 1A:} Diagrammatic representation of the varied effects of tuberculosis on the urinary tract

\textbf{Figure 1B:} Diagrammatic representation demonstrating the pathological changes of renal tuberculosis

\textbf{Figure 1C:} Diagrammatic representation demonstrating the pathological changes of renal tuberculosis

\textbf{Figure 2:} Pathology specimen of end-stage renal tuberculosis: The basis for the ‘lobar caseation pattern’ is evident.

Confirmation is usually by AFB staining of histopathology/ Fine needle Aspiration Cytology (FNAC) samples.
Imaging of renal tuberculosis may occasionally be complicated by the concurrent presence of tumors, usually adenocarcinomas, although transitional cell carcinoma has also been reported. This association is purely coincidental, although it has been hypothesized that renal tumors may be responsible for reactivation of dormant TB foci. TIN occurring together with leukemic infiltration has also been reported.

Renal TB has also been noted in native adult polycystic kidneys, in immunocompromised transplant recipients in association with renal replacement lipomatosis, and in horse-shoe kidneys.

Complications

Extra renal spread

The TB disease process may spread to the perinephric and retroperitoneal areas. Fistulas may extend even beyond these confines, including into the gastrointestinal tract, skin, lymphatic vessels, and thoracic cavity (pleura, bronchus). Renal TB causing a liver abscess has been reported.

Amyloidosis

Chronic TB is occasionally complicated by amyloidosis which, in India, is an important cause of renal disease. Prompt treatment of the underlying TB focus can prevent progression to end-stage renal disease.

Squamous metaplasia

Keratinizing squamous metaplasia, which is a potential risk factor for the development of squamous carcinoma, may develop as a late complication of chronic inflammation and infection of the renal pelvis.

Imaging studies

Imaging findings in renal TB depend upon the extent of the disease process. Familiarity with various imaging features permits early diagnosis and prompt management, thereby reducing patient morbidity. There is a correlation between the timing of the diagnosis and the severity of UTB. Delay in diagnosis may lead to end-stage renal disease, necessitating long-term dialysis.

Although it is usually stated that imaging studies are only suggestive of the disease and should not be used for the confirmation or exclusion of UTB, with sufficient experience, one can confidently diagnose UTB on the basis of the imaging findings alone.

The intravenous urogram (IVU) remains the gold standard in imaging early renal TB. Ultrasonography (USG), computed tomography (CT), and magnetic resonance imaging (MRI) lack the spatial resolution necessary to demonstrate the fine erosive changes that affect the urothelium. However, each imaging modality has its own strengths and even the simple plain film has much to offer.

The plain radiograph

Although UTB most commonly results from hematogenous spread of a pulmonary focus, the chest radiograph will be negative in half. In the other half, there is evidence of healed/active pulmonary disease. However, only 10% of radiographs show signs of active PTB. Extra-pulmonary manifestations such as calcification of lymph nodes, adrenals, prostate, seminal vesicles, or vas deferens; psoas abscesses; calcified granulomas in the liver or spleen; as well as spinal abnormalities, may noted. These additional findings lend support to the diagnosis of renal TB.

Radiographic identification of calcification associated with renal TB is becoming less common. It is noted on conventional radiography in 24-44%. It may be the first sign that TB is present. Fine calcifications that were previously unidentifiable are now much better seen with CT. Although calcification is unusual in the early stages of the disease, nearly every end-stage tuberculous kidney contains calcification. Renal calcification can take a variety of patterns, varying from few minute areas of calcification to a complete cast of the kidney. Initially, the calcifications are faint and punctate, but eventually coalesce. Early calcification may be amorphous, granular, or curvilinear, occurring typically within the renal parenchyma. Focal globular calcification involving a renal lobe is frequently associated with a granulomatous mass. Triangular ring-like calcifications that are characteristic of papillary necrosis may be noted within the collecting system.

Calified caseous tissue characteristically appears to be very homogeneous and only moderately dense, looking like ground glass; this is often referred to as ‘putty kidney’. Labeled calcification ‘putty-like’ if any area of faint calcification of uniform density was greater than 1 cm in diameter. A lobar pattern of calcification, with calcific rims outlining the periphery of distorted renal lobes, is pathognomonic of TB. [Figure 3A and B]. This occurs in far-advanced renal TB, accompanied by autonephrectomy. As TB often independently involves each lobe of the kidney, renal destruction takes place lobe by lobe. Hydrocalicosis combined with caseation pushes the residual normal renal parenchyma peripherally. As calcification occurs at the boundary between necrotic and viable tissue, the appearance is that of lobar rims.

Gow believed that calcification in renal TB has an unfavorable prognosis and, if left alone, would result in an increase in the size of the calcification and deterioration of renal function. Renal or ureteric calculi have been noted in up to 19% of cases. Apperson et al. emphasized the difficulty of differentiating calcifications from calculi in renal TB. In their cases of renal TB, 9.3% had discrete calculi and 8.7% had parenchymal calcification. They also
Renal imaging studies – Intravenous urography

The IVU has been considered as one of the most useful tests for obtaining anatomical and functional details of the kidneys. It can show a broad range of findings, depending on the severity of infection. In a series of 45 patients, the IVU pointed to the diagnosis of urinary TB in 88%. However, approximately 10-15% of patients who present with active renal TB may have normal urographic findings. Isolated parenchymal miliary tubercles usually produce urographic findings only when a calyx is involved. The earliest urographic change occurs in the minor calyces, with subtle initial signs such as minimal calyceal dilatation, and mild loss of calyceal sharpness due to mucosal edema. As the disease progresses, the calyceal outline becomes more irregular, fuzzy, and ragged and, later, feathery and moth-eaten in appearance. Although calyceal erosion has been described as the first IVU sign in renal TB, in the author’s opinion, in practice, early papillary necrosis may be the first detectable sign. It has been stated that erosion of the papillae from TB is more ragged than that from other causes, but this need not always be true. Caseating parenchymal tuberculomas may rupture into an adjacent calyx, resulting in an irregular cavity that fills with contrast, resembling renal papillary necrosis on IVU or retrograde pyelography. TB papillary necrosis results not only from ischemia, which is the basis of change in most renal papillary necrosis, but also as a result of direct tissue destruction. We have seen classic early forniceal and even central papillary necrosis in numerous proven cases of renal TB that cannot be differentiated from papillary necrosis due to other causes. In TB, the central type is probably due to ischemia, and the forniceal, usually due to direct erosion. Ischemic papillary necrosis in renal TB could be caused by a small granuloma eroding or impinging upon adjacent vessels, or be the result of TB endarteritis. Medullary cavitation with communication to the collecting system has been described as a frequent finding by Kollins et al. It may be the sole radiographic abnormality and, at times, may not be differentiable from papillary necrosis due to other causes. It may involve one or more papillae, unilaterally or bilaterally, and can vary in appearance from small and smooth to large and irregular [Figure 6A-C]. Irregular pools of contrast material may thus be seen adjacent to dilated calyces. Focal or global compromise of renal function may be noted [Figure 9A and B].
The abnormalities outlined so far, i.e., calyceal dilatation, loss of sharpness, fuzziness, papillary necrosis, and initial cavitation, are the early changes of renal TB. The late or advanced manifestations include extensive cavitation, fibrotic strictures, cortical scars, mass lesions, calcification, autonephrectomy, perinephric abscess, and fistula formation.

**Cavitation**
Cicatricial deformity of the calyces may lead to pinching of the tips of the minor calyces. Continued destruction may transform the minor calyx into a large pocket of necrotic caseous material, with or without fibrous obstruction of the infundibulum.[73] Cavities may be differentiated as obstructive or non-obstructive.[95] In the former, contrast medium does not enter the cavity on retrograde pyelography, but vague visualization may occur on the IVU; in the latter, opacification occurs only on the retrograde study [Figure 10], making it difficult to differentiate a dilated, diseased calyx from a cavity that has ruptured into the calyx. The obstructive type needs close follow-up with US. Gentle balloon dilatation may be attempted if the narrowing gets worrisome. Successful antegrade balloon dilatation of tuberculous strictures of the urinary tract has been performed by Kim et al.[86,87] The lipping type of cavity, projecting medially, is considered diagnostic of TB.[88]

Renal parenchymal cavitations result from caseation of enlarging tuberculomas. Kidneys with extensive cavitations are referred to as ulcero-cavernous kidneys, and usually do not excrete contrast medium.[36] If the cavity ruptures into the collecting system, pronounced TB bacilluria results, increasing the likelihood of the disease process spreading to other parts of the urothelium, with the possibility of a schirrous reaction that may later cause stenosis and obstruction of the collecting system.[69]

**Strictures/scars**
Renal functional damage due to strictures is greater

**Figure 5:** Intravenous urogram revealing an upward pointing (arrow) renal pelvic calculus, suggesting the presence of a hiked up renal pelvis. Multiple discrete calcifications are noted in an upper polar tuberculosis cavity (circled area)

**Figure 6 (A-C):** (A) Intravenous urogram revealing lower infundibular (arrow) and renal pelvic scarring (curved arrow). Note areas of papillary necrosis in the circled area, (B) Intravenous urogram revealing papillary necrosis in the upper group of calyces, with irregularity of the calyceal margins and the lateral margin of the upper infundibulum (dotted circle), indicating spread of infection from the calyx to the infundibulum. (Healing fornicial papillary necrosis of non-TB origin noted in a lower calyx (arrow), (C) Intravenous urogram revealing multiple parenchymal cavities (black arrows) with areas of papillary necrosis (white arrow) in the upper group calyces, bilaterally. The (L) upper group (lateral division) calyceal outline is destroyed by adjacent granulomatous tissue (arrowheads)
The three danger points of fibrosis are: The lower ureter, the pelvi-ureteric junction, and the neck of a calyx. Fibrosis is the result of healing and may thus develop during treatment. Strictures may affect the calyceal neck, infundibulum, or the renal pelvis and result in hydrocalyx, regional hydrocalicosis, or generalized dilatation of the calyces and infundibula, respectively. Commonly, a number of strictures are present, and the renal pelvis is small and contracted. Obstruction of areas not directly affected by tuberculous ulcerations and kinking of the renal pelvis can occur due to traction from a strictured infundibulum and parenchymal fibrosis. These are known as Kerr’s kinks. Scarring causes various kinds of calyceal deformities, some of which are probably unique to TB. A stricture of the inferior margin of the renal pelvis...
and its cephalic retraction, the so-called ‘hiked-up pelvis,’ may be seen [Figure 8A].

Obstruction from strictures leads to dilatation of the PCS and pressure atrophy of the renal tissue. Such hydronephrosis tends to have irregular margins and reveal filling defects, which are due to caseous debris [Figure 12]. TB infection in the dilated calyces results in a closed pyelocalyx and leads to caseation of the surrounding renal tissue. A completely stenosed infundibulum or calyx can cause complete failure of contrast excretion by the involved renal parenchyma (‘phantom calyx’) [Figure 13]. If such an area is small and represents the only abnormal focus within the kidney, the urogram may erroneously be interpreted as normal. A tiny infundibular stump (amputated calyx) or spike may be a good clue in such instances.

Parenchymal scars are common, being seen in over 50% of patients. They can be seen either overlying blunted calyces or in between the calyces.

**Mass lesion**
As the disease progresses it may become difficult to differentiate between hydronephrosis and TB granulomas, as both are masses that do not opacify with contrast at urography/pyelography. US, CT, and MRI are helpful in such situations.

**Calcification**
Calcification has been dealt with in detail in the plain film section. The characteristic lobar pattern of calcification should be remembered [Figure 3A and B]. There is an increased incidence of renal calculi and intrarenalcalyceal dense calculi-like calcifications-‘pseudo-calculi’.

Wang et al. noted three imaging patterns in 94% of IVUs with multiple findings: (a) multiple stricture sites; (b) a single stricture with one other imaging finding; and (c) autonephrectomy with any other imaging finding, barring stricture [Figures 9, 11 and 13].

**Autonephrectomy**
The late phase of progression of granulomatous destruction of the kidney, with subsequent obstructive uropathy, can lead to an autonephrectomy. This is considered typical of end-stage renal TB. There are two types: (1) the caseo-cavernous autonephrectomized kidney, i.e., an enlarged kidney converted into a caseous filled sac, with or without calcification; and (2) the shrunken, fibrotic, and often calcified kidney. In both instances, there is usually obstruction of the ureter at some point, but this is not essential in type (1). However, both types will be non-functional on the IVU [Figure 14].

Non-functioning kidneys [Figure 9A-C] in TB can be seen in: (A) autonephrectomy, (B) obstruction (ureteric obstruction, including post-treatment fibrosis), and (C) renovascular hypertension due to renal artery disease in a fibrotic kidney filled with cavities. In (C), autopsy usually reveals conical stenosis or complete obstruction of the renal artery, usually due to medial hypertrophy with intimal or subintimal sclerosis. Surgery (nephrectomy) has been extremely effective in relieving such hypertension.

US or CT plays an important role not only in evaluating non-functioning kidneys but also in patients with complications of renal TB.
The other important complications of renal TB are: (1) perinephritis, (2) perinephric abscess, (3) fistulae, (4) psoas abscesses and, rarely, (5) renal failure.

**Perinephric and psoas abscesses**
Tuberculous pyocalicosis, parenchymal abscesses, or pyonephrosis may perforate into the perinephric region to cause perinephric abscess. Renal TB may spread to the psoas muscles via the perinephric space, although involvement from the spine or via a hematogenous route is more common [Figure 15A and B].

IVU features of perinephric abscess include restricted renal movement on comparative deep inspiration and expiration films, or on intentional double exposure films. Those of psoas abscess include lateral renal and ureteric displacement. These are better evaluated on USG, CT, and MRI.

**Fistulae and sinus tracts**
Fistulae involving the kidney may communicate with adjoining structures such as bowel, skin, blood vessels, and lymphatic vessels, as well as the thoracic cavity (pleura, bronchus). Renal fistulas can be classified into those that communicate with the renal collecting system via the parenchyma (reno or nephro) and those that communicate with the renal pelvis (pyelo). Extension of perinephric abscesses to adjacent viscera or tissues results in sinus tracts or fistulae such as nephrogastric, pyeloduodenal, nephrocolonic, and nephrococutaneous sinus/fistula.

A fistula leading to the stomach or descending colon originates from the left kidney and that to the duodenum and ascending colon from the right kidney. Such fistulae are uncommon, the nephrogastric type being amongst the least common. Unusual complications also include reno-gluteal fistula.

Quite often, urography shows no excretion of contrast from the affected kidney and the diagnosis is established by retrograde pyelography. Rarely, TB of the intestinal tract involves the urinary tract, producing entero-renal and entero-vesical fistulae.

Fistulae between the renal collecting system at the fornix and the relatively abundant lymphatic drainage around it have been described. These eventually communicate with the retroperitoneal lymphatic system via the peripelvic system and lead to chyluria. Renal tuberculosis and filariasis are the leading causes of chyluria worldwide. Pyelosinus and/or pyelo-interstitial backflow from a fragile calyx, if accompanied by pyelo-lymphatic backflow, is a useful pointer toward renal TB [Figure 10]. A calyx may be labeled as ‘fragile’ if it reveals fornical rupture despite adequate precautions having been taken to avoid excessive abdominal compression during urography [Figure 11].
On USG, a mass may be missed if its (a) the more commonly seen infiltrative pattern, with calcifications. Two patterns of TB have been described: parenchymal masses, cavities, urothelial thickening, and 59% of cases. However, carefully performed USG can show most of and inability to define the extent of perirenal spread.

to calcifications, inability to evaluate renal function, isoechoic parenchymal masses, relatively less sensitivity difficulties in detecting subtle urothelial changes and or CT in the evaluation of renal TB

It has been stated that USG is less sensitive than IVU or CT in the evaluation of renal TB because of the difficulties in detecting subtle urethelial changes and isoechoic parenchymal masses, relatively less sensitivity to calcifications, inability to evaluate renal function, and inability to define the extent of perirenal spread. However, carefully performed USG can show most of the spectrum of morphologic abnormalities in urinary TB. In a large series, USG established the diagnosis in 59% of cases. Sonographic features in renal TB include parenchymal masses, cavities, urothelial thickening, and calcifications. Two patterns of TB have been described: (a) the more commonly seen infiltrative pattern, with increased echogenicity due to calcifications, infected debris, and/or abscesses; or (b) hydronephrosis or pyonephrosis, with dilated calyces and a small renal pelvis. Unfortunately, the demarcation between the two is often not very clear, and the picture is often a combination of both processes.

Each individual finding can also be caused by other disease processes, and the key to correct diagnosis is overall pattern recognition. The distinguishing feature is the visualization of multiple abnormalities involving various sites and the presence of different stages of disease in the same patient. Rui et al. have attempted to classify the USG features of renal tuberculosis into six types.

The normal renal cortex is homogenous and hypoechoic to the liver and spleen. The medulla can easily be differentiated from the cortex by the presence of the small rounded hypoechoic medullary pyramids adjacent to the bright central echo complex of the renal sinus. Das et al. reported bilateral disease in 30% of their cases on USG and attributed their improved detection rate to the use of high-frequency transducers that improved tissue characterization allowing visualization of early focal lesions. This emphasizes the need for careful scanning of the contralateral kidney.

The most frequently encountered sonographic parenchymal abnormality is a focal renal lesion—the parenchymal granuloma. Small (5-15 mm) focal lesions are either echogenic or have an echogenic border with a central area of low echogenicity, whereas larger focal lesions (>15 mm) have mixed echogenicity and poorly defined borders. Granulomas can be better appreciated on color flow imaging, as the ‘cut off’ of the vasculature highlights the presence of granulomas. Masses with cavitation/calcification may also be noted. Vijayraghavan et al. noted masses of mixed echogenicity with multiple punctuate calcifications. Caseation occurring within the masses leads to parenchymal cavities (that can be detected on USG) which could rupture into the PCS. Occasionally, an appearance like that of ‘lobar nephronia’ may be noted. Lobar nephronia is usually well delineated and is accompanied by prolonged pain and fever—clinical findings that is usually only minimal or conspicuous by its absence in tuberculous lesions. Whenever required, an USG-guided FNAC (with culture of the aspirate) may be performed to confirm the diagnosis, especially in patients with negative urine cultures. USG-guided FNAC also aids in defining the nature of sonographically visible lesions in patients with positive urine cultures, as TB, acute focal bacterial nephritis (AFBN), xanthogranulomatous pyelonephritis (XPN), and small benign and malignant tumors may have a similar appearance on USG. Modern laboratory tests such as the polymerase chain reaction (PCR) aid quick diagnosis. The
high sensitivity and specificity, in addition to the potential for rapid detection of mycobacteria, make PCR (both IS6110:MTB species-specific DNA insertion sequence and 16S rRNA: *Mycobacterium* genus-specific sequence encoding ribosomal ribonucleic acid) a useful tool in the clinical management of mycobacterial infection in the urinary tract.\[^{[108]}\] New PCR based technologies such as cartridge based nucleic acid amplification techniques [CBNAAT] (GeneXpert – Cepheid USA) and Line Probe Assay (LPA) systems, diagnose TB much earlier (in 2 hours and 2 days respectively).

Papillary involvement may be seen as an echogenic non-shadowing medullary mass in close proximity to the calyces, into which it commonly ruptures, to produce a cavitative lesion that communicates with a calyx via a thin or wide anechoic tract [Figure 20A and B]. Its appearance is dependent upon the extent of destruction.\[^{[32,43]}\] Sloughing is accompanied by an echogenic flap, which is separated from the calyceal wall. Intracalyceal filling defects may be caused by sloughed papillae as well as blood clots, fungal balls, or other debris; however, other features of renal TB help distinguish the cause. In doubtful cases, FNAC may be performed.\[^{[32]}\]

Focal caliectasis resulting from infundibular stenosis is
degrees of fibrosis and obstruction affecting different sites [Figures 22 and 23A-C]. The accompanying urothelial thickening is quite evident if looked for. When the renal pelvis and ureter are involved by TB, the hydronephrosis becomes severe.

The pelvis is usually scarred. This pattern of diffuse uneven caliectasis (without renal pelvic dilatation) accompanied by urothelial thickening, is a good pointer of renal TB [Figure 24A and B], especially in the absence of a renal pelvic calculus. The renal size is generally maintained. Frank renal swelling is more commonly seen with non-TB infections. However, it is not entirely unknown in renal TB [Figure 19]. A TB renal abscess may be noted as an irregular sonolucent cavity, with a semi-solid echogenicity and a thick ill-defined wall. Necrotic debris and scattered echogenic foci may be noted [Figure 18C]. The renal abscess can extend outward and rupture, leading to a perinephric abscess [Figure 25A and B] and later to a cutaneous fistula. A thorough analysis of the retroperitoneal compartments and, in particular, the

frequently encountered [Figure 21], often accompanied by varying degrees of urothelial thickening. This may create a fairly characteristic sonographic pattern of a focally dilated collecting system containing debris. Characteristic uneven caliectasis is seen, which is caused by varying
Merchant, et al.: Renal tuberculosis-Part I (IVU and USG)

Figure 23 (A-C): (A) Ultrasonography (USG) image revealing hyperechogenic areas of caseation interspersed with the echogenic sinus echoes. (coronal scan), (B) Oblique ultrasonography (USG) scan reveals uneven caliectasis (white arrows) with a hazy interface and urothelial thickening in the upper calyces. The lower calyceal region is replaced by hyperechogenic caseous tissue, (C) Comparative ultrasonography (USG) image of regular (evenly dilated) caliectasis with hyperechoic fungal balls (white arrows) in a HIV-positive patient (note the hyperechogenic material is lying within clearly dilated calyces and are not replacing them as happens in tuberculous caseation)

Figure 24: (A) Moderate-to-severe urothelial thickening noted throughout the visualized urothelium. This is well visualized on account of the dilatation due to a tuberculous ureteric stricture, (B) ultrasonography (USG) image revealing uneven caliectasis with ragged urothelial thickening (arrowheads). Note significant debris in the lower calyces

Figure 25: (A) Ultrasonography (USG) image revealing left tuberculous perinephric collection due to a ruptured upper polar tuberculous abscess. (A) Grey scale image, (B) Ultrasonography (USG) image revealing left tuberculosis perinephric collection due to a ruptured upper polar tuberculous abscess. Color flow image revealing lateral extent of the renal parenchyma

psosas muscle sheath, is required as these are possible sites of a migrating abscess. Restriction of renal movement is a good USG pointer toward perinephric spread. We have seen a case where frank renal swelling was noted along with diffuse spread to the perirenal and posterior pararenal spaces, resulting in innumerable multiseptated collections therein. USG, being a rapid, dynamic, and safe modality, can also be used to detect the presence of TB elsewhere in the abdomen. Findings of omental caking, especially in conjunction with septated ascites and peritoneal, mesenteric, or bowel wall thickening and lymphadenopathy, are virtually diagnostic of abdominal TB.\[110\]

Evolutive multifocal TB also gives rise to diffuse heterogeneous changes, with both parenchymal and central architecture being altered [Figure 26]. Fluid-filled cavities are seen, corresponding either to dilated calyces, or to TB abscesses. When diffuse, the abnormalities are rather similar to those brought about by extensive diffuse urothelial tumors.\[99\] Caseation is fairly well recognized on USG once one is aware of its pattern [Figure 27A]. Color flow imaging
Focal calcification may be seen as highly echogenic areas with distal shadowing and, in most cases, will represent a healed granulomatous process. Parenchymal calcification may be accompanied by overlying cortical scars.

Calcification may be punctate or dense, within a visible renal mass, or in the infected parenchyma. In the PCS, speckled or curvilinear calcifications may also be noted in the walls of the calyces, pelvis, or ureter. Faint/early ureteral calcification, a very good pointer toward UTB, may be missed on USG. A careful look at the plain film will help avoid this pitfall. USG is also unable to distinguish between the various types of calcifications that occur. However, if looked carefully, the characteristic ‘lobar pattern’ of calcification is easily demonstrable on USG [Figure 28A-C]. Involvement of even a single lobe can be diagnostic of renal TB. Small calculi may not be identifiable at times, as they do not shadow in the midst of surrounding fibrotic tissue. USG elastography/tissue strain analytics [including exciting new variants such as acoustic radiation force impulse (ARFI) technology, for evaluation of deep tissues not accessible by superficial compression elastography] have the potential to evaluate fibrosis, and thus to contribute further to treatment planning.

In addition to dense areas of calcification, approximately 25% of patients with renal TB demonstrate tumefactive masses similar to those seen in XPN [Figure 19]. Air within
the collecting system may indicate the presence of an enteric fistula. Residual functional tissue left in a densely calcified kidney is poorly seen on USG and is best evaluated by CT.\(^{[76]}\) with nuclear scintigraphy and MRI being the other options.

CT urography (CTU) performed on the current high-end MDCT scanners have the potential to detect early TB changes that were usually noted on IVU. Hence, familiarity with the IVU changes of renal TB remains necessary. These, along with other CT and MRI findings in renal TB, will be covered in part II of this review article.

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