Introduction
This article is based on a prospective 10 years observational study on loin pain haematuria syndrome (LPHS) complicating symptomatic nephroptosis (SN) and the results of renal sympathetic denervation and nephropexy (RSD&N) surgery [1,2]. The objective here is to demonstrate that renal pedicle stretch causes neuro-ischaemia as evidenced by the new IVU 7 sign.

Patients and methods: All patients presenting with loin pain with or without hematuria during 10 years were entered into a prospective observational study and underwent thorough clinical, laboratory and imaging investigations. Repeated standard imaging was invariably normal, when supine. However, 190 patients demonstrated SN of > 1.5 vertebrae on repeating intravenous urography (IVU) with erect film. Of whom 36 (18.9%) patients developed recurrent episodes of painful hematuria for which no organic pathology was detected on all imaging, when supine – thus fitting the definition of LPHS. The IVU 7 sign, with its horizontal and vertical segments, represents the renal pedicle at supine and erect IVU films, respectively was used for measuring renal pedicle stretch causing renal ischaemia.

Results: Results of 190 with SN on IVU-E, 182 were females and 8 males. The mean age was 28.8; duration of symptoms 15.7 and hospital follow up 6.6 years. Patients showed no abnormality on IVU or ancillary imaging when supine. All patients showed renal drop of >1.5 vertebrae (>5cm) on erect IVU film. Other demonstrable pathology on erect IVP film included: pelviuretric junction kink affecting the right kidney in 116 (61.1%) and bilateral in 19 (10%) of patients. Stretch/rotation of renal pedicle causing neuro-ischaemic pain of LPHS was demonstrable on the right side in 72 (37.9%) and bilaterally in 7 patients.

Complications of SN on IVU erect film included both obstructive and neuro-ischaemic: obstructive complication included ballooned renal pelvis, hydronephrosis and upper pole diverticulum. Neuro-ischaemic complications induced by pedicle stretch and rotation/twist were haematuria of the LPHS affecting 36 (18.9%), auto nephropexy affecting 12 right kidneys, upper pole calyctiasis with extra-vasation affecting 28 (15.8%) right kidney and 2 bilateral that are best shown on RGP. Renal atrophy affected 4 right kidneys. Upper pole infarction affected 2 kidneys. Retrograde pyelography (RGP) also demonstrated upper pole calyctiasis with extra-vasation. Surgical treatment was used in 28 patients; 10 had simple nephropexy and 18 had RSD&N for severe LPHS. Four of patients treated with simple nephropexy had recurrence of LPHS while those who had RSD&N were all cured.

Conclusion: Upright IVU film and RGP are essential for the diagnosis of SN complicating into LPHS. The new IVU 7 sign affirms that pedicle stretch causes neuro-ischaemic nephropathy. Renal sympathetic denervation and nephropexy is curable for LPHS but simple nephropexy is not.

Keywords: LPHS; Nephroptosis; Obstruction; Ischemia; Neuropathy; Autonephropexy; Auto-nephrectomy; Sympathetic nephroplegia; Renal sympathetic denervation; Nephropexy


Introduction
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here is to demonstrate that renal pedicle stretch causes neuro-ischaemia as evidenced by the new IVU 7 sign.

Episodic loin pain (LP) or LPHS that has no detectable pathology is frustrating to both patient and urologist. I inherited 30 cases from my predecessor at King Khaled Hospital, Najran, Saudi Arabia. Repeated investigations and imaging were all normal. A patient showed me a maneuver to bring her own right kidney down to right iliac fossa and keeping it there for examination (Loin grip sign). Ever since I diagnosed 1.7 new cases/m and found the underlying cause of SN and LPHS.

The main management problem of loin pain was the lack of demonstrable pathology on repeated imaging, when supine. The underlying SN though well known [3] was disparaged [4] and LPHS though well documented, its existence may be doubted [5] and both are extremely problematic to manage [5-8].

Demonstrable renal pathology of loin pain and haematuria was invariably lacking on all supine imaging of the received protocol [5-8]. Urinary tract infections (UTI) may affect a few patients during the occasional episodes but UTI, stones and organic causes play no role in the pathogenesis of LPHS. Many complex ramifying management problems of SN and LPHS are well known, but have no solutions. Some of the problems were communicated [9,10] and the illusive overlooked link of SN with LPHS was pointed out recently [1,2].

The objective of this study was to answer the following questions:

a. Is chronic episodic LP/LPHS genuine?
b. Is there an organic pathology? How to reveal it?
c. Why is the Standard Imaging Protocol (SIP) negative?

Defining

b. Standard Imaging Protocol (SIP) as in current use.
c. Upright Imaging Protocol (UIP) in which an IVU with erect film (IVU-E) is done.

Demonstrating

a. Features and complications of SN seen on UIP not on SIP
b. Patho-etiology link of SN with LPHS.

defining

a. Reason for disparaging SN, ignoring it by Textbooks and overlooking it in routine urological practice
b. USA Authorities ignores both SN & LPHS. UK Authorities ignores SN and what the leaders choose to ignore, others do not see.

Nephroptosis is differentially diagnosed from ectopic Kidney by blood supply and mobility. In SN the kidney drops >1.5v (>5cm) on erect IVU film causing pain with/out haematuria. In LPHS no detectable anomalies is seen on SIP. Every pair of IVU films, shown here, shows left Supine (S) and right Erect (E) for comparison.

Measuring renal drop, laterality, pelvi-ureteric junction (PUJ) kink and pelvicalyceal dilatation (PC) as evidence of obstruction on IVU E, have established renal pedicle stretch and rotation as evidence of renal neuro-ischaemia. Although SN is known, it was disparaged >60 years ago and omitted from all textbooks. SIP is constantly normal- at supine posture. UIP (IVU-E) is rarely requested, hence chance diagnosis of SN is unlikely and diagnosis is easily missed. Many of SN features and some complications were documented >70 years ago. The link of SN with LPHS is a new discovery explaining its real patho-etiology [1,2].

Patients and Methods

This is based on a prospective observational 10 years study of 236 patients presenting with episodic LP/LPHS. All had NO detectable anomalies or known organic causes of LP/LPHS on SIP. Of presented cases, 190 demonstrated anomalies on UIP of IVU-E and retrograde pyelography (RGP). All 190 SN patients had renal drop >1.5 and up to 5 vertebrae (>5 to 17.5cm) on IVU-E film. Most patients also demonstrated other features and complications on UIP, while SIP is negative as it is routinely done at Supine (S) posture. KUB, US & IVU were normal when S. Ancillary imaging of CT, MRI & Isotope Renography (IR) were also normal at S posture. UIP is the same as SIP after adding erect (E) IVU film. IVU-S was repeated with erect film at 15 min (IVU-E). Isotope renography (IR) was done at S and E (sitting) postures. Retrograde pyelography (RGP) with Cystoscopy was done on patients who consented.

In order to affirm LP/LPHS is genuine we developed the following new methods

a. Loin Grip Sign gripping the loin while the patient in standing position holds the kidney at ptosis position after lying supine, for examination at right iliac fossa.
b. One step jump test asking the patient to jump from one step to the floor causes agony which affirms that pain is real and genuine.

to affirm pedicle stretch causes ischaemia

i. The IVU 7 sign, mapping the renal pedicle on IVU-S and IVU-E demonstrate its degree of stretch
ii. Rubber tube stretch hypothesis shows that stretch narrows the lumen of a tube and induces renal ischaemia.

The kidney was explored by open surgery, placing the kidney at ptosed position caused stretch of renal pedicle causing ischaemic cyanosis of the kidney. This confirmed the IVU 7 sign and tube stretch hypothesis; representing the renal pedicle stretch ischemia and neuropathy. The kidney was explored through loin incision. The renal artery was skeletonised cutting and diathermy all surrounding nerves. Nephropexy was done by the 3 point technique.
Results

TOF 190 patients with SN on IVU-E, 182 were females and 8 males. Of 46 excluded cases, 26 had SN <1.5v, 11 undetermined (allergy) and 9 had urinary stones. A drop of >3 vertebrae (10.5cm) affected 124 (65%). Mean age was 28.7 (12-70), hospital follow up 6.6 (3-16) and symptom duration 15.7 (3-46) y. Age distribution is shown in Figure 1 and ptosis distribution is shown in Figure 2. Ptosis affected right kidney alone (Figure 3) in 107/190 (56.3%) and was bilateral (Figure 4) in 82 (43.2%) patients.

The patients were segregated by presentation into two groups. The Emergency Room (ER) group of patients was mostly young, thin and single females. Their mean body weight 47.7kg, height 155cm and body mass index (BMI) 19.8kg/m2. Severe LP/LPHS episodes required repeated hospital admissions of 97 (51%). Haematuria episodes affected 36 (18.9%) patients. LP affected all right sides of ptosis but may affect the left side as part of bilateral involvement. False diagnosis during an episode of the acute abdomen caused unnecessary surgeries including appendectomy in 28, inter-vertebral disc removal in 3 and cholecystectomy in 4 patients. False label included depression in 18, opiate dependency in 13 and malingering in 3 patients.

The out-patient department (OPD) group regularly attended clinics with episodes of LP and the multiple associated splanchnic symptoms (MASS). They were married multi-parous women with a mean body weight of 66kg, height 155cm and BMI 28 kg/m2. They regularly attended many clinics of medical, gastro-intestinal tract (GIT), Nephrology, UTI, Backache and Pain clinic.

MASS indicated sympathetic over-activity and included:

a. GIT symptoms of acid peptic disease (APD) and irritable bowel syndrome (IBS) affected 94 (49%) patients.

b. Backache affected 120 (63%) of whom 24 (12.6%) patients had sciatica-like pain (normal CT spine).

c. Cystitis without UTI affected 96 (51%) patients, of whom 12 had bladder biopsy and showed interstitial cystitis.

Presentation

i. Recurrent attacks of severe LP or LPHS to OPD or ER

ii. ER Group had the differential diagnosis of right acute abdomen.

iii. OPD Group had LP and MASS of GIT of APD / IBS, Backache and Cystitis. All specialist investigations were normal.

Kink at PUJ was demonstrable on IVU-E of 135/190 (71.1%), 116 (61.1%) right and 19 (10%) bilateral. PUJ kink with renal pelvis dilatation on IVU-E affected 46 (24.2%) patients, 37 right and 9 bilateral kidneys. Renal pelvis dilatation became evident on follow up later on IVU-S of 18 (9.5%) and classical PUJ obstruction developed in 7 cases (Figure 5).

Ischaemia due to pedicle stretch and renal twist torsion rotation around pedicle demonstrated in 124 (65.3%) patients. The renal ischaemia is evidenced by the IVU 7 sign and tube stretch
hypothesis (Figure 6 & 7). Ischaemia caused erosion of medullary papillae best shown on RGP (Figures 8 & 9).

The 99mTc-DTPA IR at supine and seated posture showing blood flow, GFR and washout curves. Heterogeneous causes of LP and LPHS in SN demonstrable on IR but are posture dependant. Split kidney function was shown on changing posture from S to E (sitting). The pathology shown on IR may show ischaemia (Blood Flow & GFR) and or obstruction (Drainage) (Figure 10).

Figure 5: Shows IVU with nephroptosis complicated with PUJ obstruction.

Figure 6: Shows renal pedicle mapped on a supine IVU film (Horizontal) and erect film (Vertical) where the renal pedicle is stretched to 3 times its normal length.

Figure 7: Compares the rubber tube stretch with IVU 7 sign which demonstrate that stretch of a tube or artery causes stenosis of the lumen.

Figure 8: Shows RGP with multiple pyelocalyctasis with erosion of medullary papillae and contrast leakage into veins (papillary venous fistula).

Figure 9: Compares IVU with RGP of the same patient, IVU looks normal while RGP reveals extensive bilateral renal damage (eroded papillae of pyelocalyctasis).

Figure 10: Shows isotope renography at supine and sitting postures with split renal function.

Figure 11: Shows agony on a patient’s face who suffered from LPHS after jumping from one step to the floor.
Open surgical treatment was used in 28 patients suffering from severe LPHS; 10 had simple nephropexy and 18 had RSD&N. Four of those treated with simple nephropexy had recurrence of LPHS while those who had RSD&N were all cured of LPHS. Nephropexy alone is inadequate treatment. The evidence that repeated stretch of renal pedicle causes ischaemia and focal infarctions of the kidney is shown in Figure 11 as a result of the severe vascular stenosis affecting renal Pedicle (Figure 12). The new physical signs that demonstrate LP and LPHS are genuine are shown in (Figures 13 & 14).

Figure 12: Shows the ioin grip sign applied when the patient is standing then keep it as she lies supine (Top) to allow for kidney examination at the right iliac fosse (bottom).

Figure 13: Shows an explored right kidney of LPHS patient which is cyanotic with focal infarction.

Figure 14: Shows an explored right renal pedicle after denervation of LPHS patient shown in Figure 13 with severely stenosed artery and vein (effect of chronic repeated pedicle stretch). The Nelaton tube is approximately the size of a normal renal artery.

Discussion

The IVU 7 sign and tube stretch shows that stretch to twice the normal length reduces artery diameter to \( \frac{1}{2} \), the cross section area to \( \frac{1}{4} \) and flow to \( \frac{1}{16}(6.25\%) \). Artery twist/torsion induces further ischaemia. The renal pedicle contains renal artery, vein and sympathetic nerves. Thus SN initially causes functional stretch neuropathy and ischaemia explaining pain and haematuria of LPHS. It later causes ischemic damage to renal papillae.

This causes sympathetic neuropathy and nephropathy inducing ischaemia of SN which is initially intermittent and functional causing LP/LPHS. LPHS takes 3-4 years to complicate SN. Ischaemia organic permanent calyctaisis becomes evident on RGP: It starts at upper right calyx, common site of haematuria, spreads to other calyx of the same right kidney and affects the medullary papillae long before cortex. Auto-nephropexy detected in 12 (63%) patients and auto-nephrectomy (Renal atrophy) in 4% of patients. Focal or segmental infarctions may occur. Retrograde pyelography demonstrates the organic pyelocalyctaisis (Figure 8 & 9) as the patho-etiology of painful haematuria, caused initially by intermittent renal ischemia of pedicle ptotic stretch or torsion [11,12]. Also PUJ kink obstruction may occur with nephroptosis which is initially functional intermittent and later cause organic PC dilatation.

The reported patho-etiology of SN pain is heterogeneous with obstruction, ischemia and neuropathy components that have reversible and organic stages. Although, these causes, and rarely UTI, may contribute to the establishment of pyelocalyctaisis, [13-15] the lesion seems primarily ischemic [10-12]. The link of SN with LPHS was illusive and overlooked, particularly when the ischemic complications of auto-nephropexy, auto-nephrectomy and sympathetic nephropregia involving the normal contra-lateral kidney occurred insidiously over many years [1]. Auto-nephropexy, made the link of SN with LPHS most elusive by erasing the evidence on renal mobility, but demonstrated that simple nephropexy [14-16] or spontaneous nephroptosis alone may neither cure the loin pain of SN nor abort its complication into LPHS.

The demonstrable pathology on upright IVU, [2,3,9-16] arteriography [11,12] and IR [2,13] are well documented on SN that was discarded long ago [4]. Upright imaging is currently undone and has not been reported previously in LPHS. Retrograde pyelography findings (Figure 8 & 9) have not previously been documented in either condition. The use of IVU started early in the 20th century while clinical evidence on the genuineness of SN pain dated back to the 15th century [15-17]. Loin pain hematuria syndrome was reported in 19675 while Dietl’s crisis is known for centuries. Organic reno-vascular complications demonstrated on conventional arteriography of SN [11,12] and LPHS [5-8] are of advanced cases. The demonstrable link of SN with LPHS, other ischemic complications of infarction and atrophy “auto-nephrectomy” and the most elusive “autonephropexy” and “sympathetic nephroplegia” are reported here. Hahn [18] reported nephropexy for SN in 1881 and Blacklock [19] reported renal sympathetic denervation in 1989. The IVU 7 sign (Figure 6 & 7) and deformity of renal calyxes seen on erect IVU film demonstrate the...
renal pedicle stretch and torsion, respectively, which are important in explaining the neuro-ischaemic pathology of LPHS.

**Conclusion**

LP and LPHS are genuine symptoms of SN. SN is overlooked on SIP but demonstrable on UIP. The features and complications of SN cause genuine incapacitating episodic LP/LPHS. Pain of SN has heterogeneous causes of intermittent (functional) or permanent (organic) and of obstructive and neuro-ischaemic origin. PUJ kink causes Obstruction. Ishaemic stretch and/or twist torsion of renal pedicle causes neuro-ischaemic nephropathy via sympathetic over-stimulation and neuropathy. SN may complicate into LPHS causing ischaemic papillary necrosis calyctaisis and veno-calyceal shunt which is evident on RGP. The neuro-ischemia due to pedicle stretch/rotation explains the LPHS.

The IVU-E and RGP demonstrate these features of SN while all ancillary imaging remain normal being supine. The IVU 7 sign and calyx deformity are used for detecting stretch neuro-ischaemia and obstructive pathology, respectively. The condition goes through two stages; a reversible stage which is curable and an irreversible stage which is incurable. IVU-E film and RGP are indicated in all LPHS cases. Renal sympathetic denervation and nephropexy (RSD&N) offer the best chance of curing SN and LPHS. In view of the new evidence presented here, the authorities should reconsider SN with link to LPHS and bring it back to current textbooks.

**References**