Pelvic Floor Dysfunction, the Role of Imaging and Reconstructive Surgery
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ABSTRACT
We are putting forward three novel concepts describing the pathophysiology concerning:
• Micturition, factors that control urinary continence and different types of urinary incontinence.
• Genital organs support and genital prolapse.
• Defecation, causes of fecal incontinence (FI)

I. Urinary continence depends on high urethral pressure (Pura) which depends upon two factors: One inherent and one acquire.

1. The inherent factor is the tough strong collagen layer constituent of the internal urethral sphincter (IUS), that creates the high wall tension necessary for keeping high urethral pressure (Pura). The IUS is a collagen-muscle tissue cylinder that extends from the bladder neck to the perineal membrane in both sexes.

2. The acquired factor, which is high alpha-sympathetic tone at the IUS gained from learning and training in early childhood, keeps it contracted and the urethra closes all the time until there is a need or a desire to void as social circumstances allow. Injury to one or both factors leads to urinary incontinence.

II. The vagina is a cylinder of collagen-elastic-muscle tissues. The strong tough collagen sheet is responsible for the upright position of the vagina. The main function of the pelvic ligaments is to assign the pelvic organs to their anatomical site and keeps the pelvic organs in situ. Childbirth trauma damages the collagen layer due to overstretching of the vagina and leads to flabby and redundant vaginal walls with subsequent vaginal prolapse. When the pelvic ligaments suffer most of the trauma, the insult will lead to weakness of the pelvic ligaments, leading to vault and uterine prolapse.

III. The integrity of both anal sphincters, internal anal sphincter (IAS) and external anal sphincter (EAS) is an essential factor in keeping fecal continence. Fecal continence also depends on strong pelvic floor muscles which keep an angle between the rectum and the anal canal. In addition, it depends on an acquired behavior, gained by learning and training in early childhood of maintaining high alpha-sympathetic tone at the IAS keeping
the anal canal empty and closed all the time until there is a desire and/or a need to pass flatus and/or stool and there are favorable social circumstances.

The intimate relation of the IUS with the anterior vaginal wall and the IAS with the posterior vaginal wall exposes them to the childbirth trauma with subsequent damage. This will lead to stress urinary incontinence (SUI) and FI in addition to vaginal prolapse.

Therefore, we have innovated an operation to treat SUI, FI and vaginal prolapse. ‘Urethro-ano-vaginoplasty’ repair operation. It consists of anterior and posterior sections. In the anterior section, we have corrected the SUI and the anterior vaginal wall descent through the following steps:

1. Expose the IUS and mend its torn wall.

2. Strengthen the anterior vaginal wall by overlapping the two vaginal flaps, and hence we can add extra support to the mended IUS and preserve the body collagen.

In the posterior section, we have the following:
1. Exposed the IAS and mended the torn sphincter.
2. We have approximated the two-levator ani muscles.
3. Strengthened the posterior vaginal wall by overlapping the two vaginal flaps; as such, we would have also added extra support to the mended IAS and kept the natural body collagen.
4. We repaired the perineum.

Keywords: Micturition, Urinary continence, Urinary Incontinence, Stress urinary incontinence, Genital prolapse and fecal incontinence.

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INTRODUCTION
The female pelvis contains three major tracts that lie on and traverse the pelvic floor. These are the urinary bladder (UB) and the urethra anteriorly, the female genital tract in the middle, and the rectum and anal canal posterior. The pelvic floor consists of the pelvic floor muscles, mainly the levator ani muscles, and connective tissues and ligaments.

The relative contributions to the structural support of the pelvic floor and its functions have been the subject of controversy. With increasing age, women can develop voiding troubles as urgency, overactive bladder, frequency, nocturia and stress urinary incontinence (SUI). Other concomitant troubles, which occur, are genital prolapse, fecal incontinence (FI) and pelvic pain. All of these symptoms can be associated, to a greater or lesser extent, with pelvic floor defects.

The first vaginal birth is especially associated with the development of prolapse, whereas additional vaginal births do not show significant increases in the odds of prolapse. We are putting forward three novel concepts describing the pathophysiology, anatomy, morphology of the structures, functions and pathology of these three tracts.

1. Micturition, factors that control urinary continence, causes of urinary incontinence and voiding troubles.
2. Genital organs support and genital prolapse.
3. Defecation, causes of FI and description of the anatomy morphology of the structures and functions of the internal anal sphincter (IAS).

The UB stores the urine, which we void through the urethra. Voiding has two stages:

First stage, before training in infancy and early childhood:
As soon as the UB is full, sensation of fullness of the UB travels along the pelvic parasympathetic nerves to the spinal cord. Through spinal cord’s centers, reflex parasympathetic activity (S2, 3 and 4) leads to contraction of the detrusor muscle voiding the urine through an open urethra irrespective of neither time nor place.

Second stage after training: The mother starts to teach and train her infant-child how to hold up until proper social circumstances are available. The teaching and training leads to learning how to maintain high alpha-sympathetic tone (T10-L2) at the internal urethral sphincter (IUS) keeping it contracted and the urethra closed all the time until favorable social circumstances allow.
Sensations of bladder fullness travel along the pelvic parasympathetic (S2, 3 and 4) to the CNS. This allows the person, according to the social circumstances available, to choose either to retain the urine to a later time until favorable social circumstances allow, or to void. If she chooses to retain, three neuromuscular actions take place:
1. Increase of the alpha-sympathetic tone to the IUS confirming closure of the urethra.
2. The second action is inhibition of the parasympathetic impulses to the detrusor muscle inhibiting its contractions.
3. The third action is increase of the tone of the external urethral sphincter (EUS) which is a skeletal muscle innervated by voluntary NS.

When appropriate time and place are available then, controlled by the CNS, synergistic actions between the somatic and the autonomic nervous systems four neuromuscular actions take place:
1. Lowering of the high alpha-sympathetic tone at the IUS relaxing the sphincter and opening the urethra,
2. Relaxing the EUS which is a striated muscle innervated by somatic nerve supply,
3. Activating pelvic parasympathetic nerves to contract the detrusor muscle and empty the UB,
4. The EUS (compressor urethrae) acts to propagate and propel the stream of urine and at the end to squeeze the urethra to expel the last drops of urine8-13,15 (Fig. 1).

High urethral pressure (Pura) that is much higher than vesical pressure (Pves) is the main factor that keeps urinary continence.

The main causes of high Pura are as follows:

1. Structural or inherent factor, which is the tough strong collagen tissue layer constituent of the IUS’ cylinder, which is the skeleton of the IUS, that creates the high wall tension necessary for keeping high Pura.

2. The acquired or behavior factor which is the high alphasympathetic tone at the IUS that keeps it contracted and the urethra closed all the time until there is a need or a desire to void and social circumstances allow.
Fig. 1: The steps followed in the second stage of micturition and its CNS control. Sensations of bladder filling travels along the pelvic parasympathetic nerves S2, 3 & 4. Controlled by the CNS, depending on the social circumstances, synergistic neuromuscular actions take place. If time and place do not allow voiding, the woman will increase the alpha-sympathetic tone at the IUS. She will also inhibit the pelvic parasympathetic preventing detrusor contractions. In addition, she will confirm closure of the EUS. When social circumstances allow, she will inhibit the high alpha-sympathetic tone at the IUS, thus opening the urethra. She will activate the pelvic parasympathetic inducing detrusor contractions. She will relax the EUS thus allowing voiding. The EUS tone increases to allow propulsion and ejection of the stream of urine and at the end of micturition to squeeze the urethra from the last few amount of urine.

Injury to one or both factors leads to drop of Pura and leakage of urine. The drop in Pura happens:

a. Physiologically when she, voluntary, inhibits the high alpha-sympathetic tone at the IUS that leads to opening the urethra and voiding occurs.

b. Pathologically, because of rupture, atrophy and degeneration of the collagen sheet of the IUS. The weak IUS will not stand against sudden rise of abdominal pressure on physical activity, as on cough, sneezing, jumping, coitus or sometimes changing postural position and urine will leak (Figs 2 to 9). The woman will be embarrassed, and a quick reflex reactive sympathetic activity will increase the existing acquired high alphasymathetic tone at the IUS, closing the urethra and preventing further urinary leakage.16-19
Childbirth trauma causes rupture of the collagen layer constituent of the
IUS leading to its weakness. Drop in the estrogen level causes atrophy of
the pelvic collagen and subsequent weakness.

Chronic and or repeated genitourinary infections causes degeneration of
the collagen of the IUS and of the vagina that leads to its weakness
causing SUI and vaginal prolapse (Figs 2 to 12).
Imaging with 3D US and MRI can demonstrate the rupture in the IUS
clearly.
The level of the rupture along the cylinder of the IUS and its extent will
determine the type of SUI(DO,genuine SUI or mixed type of
incontinence)and the morphological shape of the urethra seen on imaging.
If the rupture affects mainly the upper part of the IUS, detrusor
overactivity (DO) ensues, and funneling of the bladder neck with loss of
urethrovessical angle appears on imaging. When the main damage is in the
lower part of the IUS, it will lead to genuine SUI and a ‘flask-shape’
appearance on imaging. If the damage affects the entire length of the IUS,
then mixed type of urinary incontinence is the result, and a collapsed,
apparent short urethra with irregular walls is seen on imaging (Figs 4 to 8).
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Figs 2A to D: A diagram of the IUS as a cylinder of collagen-muscle lined with urothelium;
and an image (A) with three-dimension ultrasound (3DUS) shows the IUS as a thick tissue
cylinder that extends from the bladder neck down to the perineal membrane with muscle lying
on top of collagen. The cross section in (B), shows the muscle lying on top of muscle. In (C),
you can kindly notice the connection of the muscle layer above with the detrusor muscle. In
(D), an MRI image of a patient with full distended UB with an intact thick IUS when
compared to the normal healthy vaginal wall thickness.
Intact wall compact sheet of collagenous tissue with muscle fibers lie on and intermingle with the collagen fibers in the middle part of the cylinder.

The torn collagenous tissue cylinder, with the muscle layer intact and seen connected with the detrusor muscle in a patient with SUI.
Figs 6A and B: MRI images, sagittal section, of a patient with Mullerian duct agenesis (A) with absent uterus and vagina, the IUS is seen as a tissue cylinder that extends from the bladder neck to the perineal membrane with a closed urethra. In (B), the IUS is torn especially in its upper part causing funneling of the bladder neck and leading to DI alone or mixed-type of urinary incontinence. The vagina is torn and prolapsed as indicated by the yellow arrow.
Cases with severe SUI, where you can see lacerations that affect the IUS

**Fig. 7:** Patients who suffer severe SUI scanned with 3D US. The IUS is torn, seen clearly in the cross section, and the urethra is open and flask shape.

**Figs 8A to D:** Images with 3D US, (A) is a coronal section of a normal continent woman with intact IUS, (B) is a coronal section showing lacerations of the IUS affecting the whole length with irregular walls and an open dilated urethra, (C) is a coronal section showing the lacerations in the IUS mainly in the lower part giving a flask-shape appearance to the urethra, (D) is a sagittal section that shows loss of the posterior urethrovesical angle; it also shows an intact IAS with a closed and empty anal canal.
Figs 9A and B: Images with MRI, (A) is a cross section of a normal nulliparous woman with H-shape vagina and normal cross section of an intact IUS, (B) is a cross section, which shows lacerated prolapsed vagina and lacerated IUS.

Figs 10A to C: Images with 3D US (A) and MRI (B and C) of the vagina that show lacerations and prolapse of the vagina, the transverse axis of the vagina suffers most of the trauma. The IUS, which is intimately related to the anterior vaginal wall, is also lacerated and the urethra is open. Damage affects the entire length of the IUS, then mixed type of urinary incontinence is the result, and a collapsed, apparent short urethra with irregular walls is seen on imaging (Figs 4 to 8).
Figs 11A to C: Images with MRI, (A) is an image of a normal woman showing the IUS as a tissue cylinder that extends from the bladder neck down to the perineal membrane. The vagina is standing up thanks to its tough collagen layer, which is shown in (B). In (C), the IUS cylinder is torn more marked in its upper part that shows funneling of the bladder neck. The vagina is also torn and prolapsed.

Figs 12A and B: Images with MRI, sagittal section, they are comparison between normal IUS and vagina in (A); in contrast to torn IUS and vagina in (B). In (A), the lower arrow points at normal vagina, which is standing up because of its tough collagen sheet.
Thus, by understanding this new concept, we can explain all voiding troubles.8-19 Functional disturbances, and/or structural damage of the IUS will lead to urinary incontinence, and voiding troubles.

**Functional Damage of the Nervous Control of the IUS**

1. Failure to acquire the second stage of micturition leads to nocturnal enuresis.13-15 These failures can be complete failure or partial failure. Complete failure: Here there is a stop at the first stage of micturition, as the UB fills it empties irrespective to neither time nor place, leading to day and night enuresis. This occurs in about 10% of nocturnal enuresis patients.

The failure can be partial, as the bladder is full; the patient will be embarrassed of wetting herself, so she increases the alpha-sympathetic tone contracting the IUS and closing the urethra, preventing involuntary urination until she reaches the toilet. However, on sleeping this weak partial alpha-sympathetic tone will be lost and nocturnal enuresis will occur. This occurs in 90% of nocturnal enuresis patients. Therefore, the treatment of nocturnal enuresis is not by giving anticholinergic drugs, but by giving alpha-sympathomimetic drugs.

2. Sympathetic overactivity, e.g. painful stimuli (e.g. episiotomy, abdominal or pelvic surgery), leads to retention of urine. Spinal cord injury below the second lumbar neural level or spinal anesthesia, leads to loss of the pelvic parasympathetic sparing the thoracolumbar sympathetic supply will cause retention of urine or retention with overflow.

3. Sympathetic failure, like severe fear leads to transient urinary incontinence. In addition, alcohol, getting drunk may lead to transient UI.

**Structural damage of the IUS**

The damage can affect the whole thickness of the urethra and this will lead to true urinary incontinence. The damage and rupture can be from inside, as after urethral instrumentation, this will lead to urethral diverticulum and post voiding dribbling. Damage from outside due to over distension of the vagina in difficult, prolonged or multiple frequent labors will tear the collagen layer of the IUS causing SUI. In a review article de Boer et al found an association between vaginal prolapse and over active bladder.7 Other authors tried ultrasonic assessment of the bladder, urethra and pelvic organs with different results.20-23
GENITAL PROLAPSE

The vagina is a cylinder of collagen elastic muscle tissues. The strong tough collagen sheet is the one responsible for the upright position of the vagina. Childbirth trauma injures the collagen layer due to overstretching of the vagina and leads to flabby and redundant vaginal walls with subsequent vaginal prolapse (Figs 9 to 12). The injury affects the transverse axis of the vagina much more than the longitudinal axis (Figs 10A to C). The vagina becomes wider and more redundant. Vaginal deliveries, specially prolonged difficult labor, multiple frequent labors and rough instrumental labors cause drastic trauma to the pelvic collagen with lacerations and subsequent atrophy and weakness. Several other factors contribute to exaggeration of the weakness of the pelvic collagen by atrophy and/or degeneration, e.g. hormone deficiency, especially after menopause, chronic or repeated genitourinary and rectoanal infections; in addition to congenital collagen weakness.

Though some authors believe that the first vaginal delivery does the insult that affects the vagina and the intimately overlying urethra.6 The first vaginal birth is especially associated with the development of prolapse, whereas additional vaginal births do not show significant increases in the odds of prolapse.6 Yet cumulative affect of frequent multiple childbirth traumas and hormone deficiency delays the complaint till near menopause.

The weakness and rupture of the vaginal collagen sheet will manifest itself mostly in the transverse axis of the vagina. We can demonstrate this clinically and on imaging (Figs 9 and 10).

1. At first, there will be loss of the nulliparous H-shape vagina, which changes into a transverse slit in parous women.
2. Then, further weakness, will lead to loss of vaginal rugae; the vaginal wall will be smooth without folds as can be seen clinically.
3. Further weakness and rupture of the vaginal collagen will induce vaginal wall redundancy and descent.

At the same time, the same childbirth trauma will injure the intimately closed IUS leading to SUI. In addition, the same trauma, in a backward direction, will injure the IAS causing FI.

The main function of the pelvic ligaments (uterosacral, great transverse cardinal cervical and pubocervical ligaments) is to assign the pelvic organs to the anatomical site and keep the pelvic organs in situ supported. The pelvic ligaments (which are composed of collagen bundles), when they suffer most of the childbirth trauma, the insult will lead to weakness.
of the pelvic ligaments, leading to their laxity and lengthening. This will induce drop of the pelvic organs beyond their anatomical site and produces vault and uterine prolapse.

**DEFECATION AND FECAL INCONTINENCE**

We are putting forward a recent concept on the pathophysiology of defecation.24,25

**Defecation (Fig. 13)**

The mechanism of defecation has two stages.

*First Stage of Defecation*

In infancy and early childhood, before training: Through stretch receptors in the rectum, impulses of rectal fullness travel along pelvic parasympathetic, S2, 3 and 4 to spinal sacral centers leads to:

1. Reflex contraction of the rectal muscles.
2. The anal canal opens and the external anal sphincter (EAS) relaxes allowing defecation to occur.

![Diagram showing CNS control of defecation](image)

**Fig. 13: Diagram, which shows the CNS control the mechanism of defecation.**

**The IAS:** The IAS is a collagen muscle tissue cylinder that surrounds the anal canal; surrounded externally by the EAS. Its nerve supply is thoracolumbar alpha-sympathetic nerves from the hypogastric plexus (T10-L2).
Its functions are as follows:
1. On contraction, is to keep the anal canal closed and empty.
2. On relaxation, is to open the anal canal to allow passage of flatus and or stool.

Second Stage after Training
The mother starts to teach her child how to control himself.
Gaining control is by maintaining high alpha-sympathetic tone at the IAS keeping it closed all the time until appropriate place and time are available.

On rectal distension, stretch receptors are stimulated.
The sensation of rectal distension travels along the pelvic parasympathetic nerves to S2, 3 and 4. In the rectoanal junction, there are specialized sensory end organs, e.g. Krause end-bulbs, Golgi-Mazzoni bodies. Specialized afferent nerves for tension, temperature, texture, touch and friction subserve these organized nerve endings.
An intact sampling reflex allows the individual to choose whether to:
a. Retain the rectal contents or,
b. Discharge flatus and or stool.
According to the available social circumstances, if the woman chooses to retain, then she will do the following steps:
1. She increases the acquired high alpha-sympathetic at the IAS, confirming its closure.
2. She confirms the closure of the EAS, which is a skeletal muscle, innervated with somatic nerve supply.
3. She increases the contraction of the levator ani muscles to exaggerate the angle between the rectum and anal canal.
4. She inhibits the pelvic parasympathetic activity to the rectum and anal canal preventing their contractions.
Discharge is by relaxing the IAS and EAS for a moment only to release flatus, or for longer time to pass stool.

Wherever, appropriate place is available and there is a desire, under the control of the high CNS centers, through synergistic nervous actions between the autonomic and the voluntary nervous systems, six neuromuscular actions will occur:
1. The woman will lower the acquired high alphasymathetic tone at the IAS relaxing it, opening the anal canal.
2. Through the voluntary NS, she will relax the pelvic floor muscles thus widening the acute anorectal angle (changing it from acute to obtuse), to bring the anal canal and the rectum on one axis. She does so, through relaxing the pelvic floor muscles.
3. Through, voluntary NS, she will also relax the EAS, which is a skeletal muscle innervated by the pudendal nerve. Then two synergistic actions between the voluntary and autonomic nervous system will occur.
4. The abdominal muscles and the diaphragm contract to increase the intra-abdominal pressure and forcing the feces through the anal canal (The voluntary nervous system controls this action).
5. The smooth muscles of the distal colon and rectum contract, propelling the feces into the anal canal (The autonomic nervous system controls this action).
6. Followed by sequential contractions of the three parts of the EAS, deep, then superficial then the subcutaneous parts that will squeeze the anal canal propelling any residual contents and emptying the anal canal completely.

An intact IAS, through the acquired high alphasymmpathetic tone, keeps the anal canal closed and empty. A torn weak IAS will lead to an open anal canal that is easy to discharge flatus or stool on rise of abdominal pressure (Figs 14 to 17).

Vaginal delivery causes lacerations of the posterior vaginal wall leading to posterior vaginal wall prolapse and causes lacerations of the collagen layer of the IAS, which is intimately related to the posterior vaginal wall.

Therefore, FI is prevalent in posterior vaginal wall prolapse (rectocele). Concomitant troubles, which occur, are vaginal prolapse, SUI and FI. 1, 5 and 6 (Fig. 17).

Childbirth trauma causes the major insult, but aging and hormone deficiency (menopause) causes collagen atrophy and adds to weakness of the IUS, IAS and the vagina.

Figs 14A and B: Images with 3D US of the rectum and anal canal in normal continent woman (A) with healthy, intact IAS and a closed empty anal canal. In contrast, in (B) the IAS is torn leading to a widely open anal canal in a patient with FI.
**Figs 15A to D:** Images with 3D US of the rectum and anal canal in normal continent woman (B) with healthy, intact IAS and a closed empty anal canal. In contrast, in (A, C and D) the IAS is torn leading to a widely open anal canal in a patient with FI.

**Figs 16A to D:** Images with 3D US of the rectum and anal canal in patients with FI. Images in C and D are of complete perineal tear (fourth degree), the EAS is torn and appears as horseshoe; in addition, and the IAS is torn as well. Images in A and B are of the IAS, which are torn leading to an open dilated anal canal. The IAS appears like a horseshoe.
Fig. 17: Images by 3D US of patients who suffer from pelvic organs dysfunction, SUI, FI and vaginal prolapse simultaneously. The images show torn IUS and IAS

RECONSTRUCTIVE SURGERY

According to these novel concepts, a weak, torn IUS will lead to SUI. Vaginal prolapse is a consequence of weak, flabby, redundant torn vaginal walls. A weak, torn IAS will lead to FI. Therefore, we have innovated an operation to treat SUI by exposing the rupture in the IUS and mending the torn sphincter. At the same time, we have treated vaginal prolapse and FI by this novel operation26-30 (Figs 18 to 22).

Urethro-ano-vaginoplasty’ Repair Operation

It consists of anterior and posterior sections.
In the anterior section, we correct the SUI and the anterior vaginal wall descent through the following steps (Figs 18 and 19).
We grasp the cervix with two pairs of cervical volsela.
We inject about 10 to 20 ml normal saline with adrenaline (2 per 1,000 concentration), beneath the vaginal wall to act as hydrodissection and vasoconstrictor. This separates the anterior vaginal wall from the posterior wall of the IUS.
Figs 18A to E: Surgical photos of a patient with anterior vaginal wall prolapse, posterior vaginal wall prolapse, SUI and FI. The metal catheter is directed forward and upward (A), which means loss of posterior urethrovesical angle. We dissect the IUS from the anterior vaginal wall (B and C) and mend the torn IUS with simple interrupted sutures (D and E).

We do 2 to 4 cm transverse incision about 3 cm above the external cervical os. With a pair of dissecting scissors, we separate the anterior vaginal wall from the IUS. We cut the anterior vaginal wall longitudinally from the transverse cut we made up to the submeatal sulcus, which correspond to the perineal membrane. We grasp each vaginal flap with three pairs of Kocher’s forceps. The defect in the IUS will be apparent and on each side, we can clearly see two clear edges. One edge is of the anterior vaginal wall and the other is of the torn posterior wall of the IUS.

1. Expose the IUS (we dissect the IUS clear from the anterior vaginal wall).
2. Mend the torn posterior wall of the IUS by several (6-8) simple interrupted sutures using number 0 polyglycan thread sutures (Figs 18A to E).
3. Strengthen the anterior vaginal wall by overlapping the two vaginal flaps, using a novel dragging sutures, dragging the right vaginal flap underneath the left vaginal flap. Then we do suturing the free edge of the left vaginal flap as far lateral on the right side of the vagina. So we strengthen the anterior vaginal wall and decrease its width, we also add extra support to the mended IUS, and preserve the body collagen.

In the posterior section, we do as shown in Figures 20 to 22.
We do hydrodissection between the posterior vaginal wall and the anal canal and the rectum; and in the perineum as done anteriorly.
We do a V-shape incision at the line between the posterior vaginal wall and the perineal skin down to the perineum. Then we try to create a space between the posterior vaginal wall and the anal canal by sharp and blunt
dissection. Next with a pair of dissecting scissors, we separate the posterior vaginal wall from the rectum and anal canal. Then we cut the posterior vaginal wall longitudinally in the midline to beyond the apex of the prolapse protrusion.

Figs 19A to D: After mending the IUS, we do overlapping of the two vaginal flaps as seen in the photos. We bring the right vaginal flap underneath the left vaginal flap with this novel dragging suture as seen in (A) and (B), repeating it 4 to 6 times. Then, we suture the free edge of the left vaginal flap as far laterally in the vagina on the right as seen in C and D. Thus, we strengthen the anterior vaginal wall and add extra strength to the mended IUS.

Figs 20A and B: The posterior vaginal wall when seen without straining, this means posterior vaginal wall prolapse as seen in (A). Image (B) is 3D US showing rectocele of the same patient who suffers FI.

We hold each vaginal flap with three pairs of Kocher’s forceps. We will clearly see two different edges on each side, one is the vaginal edge, and the other is the anterior wall of the torn IAS.
1. We dissect the torn IAS clear from the posterior vaginal wall.

2. Mend the torn wall of the sphincter by serial interrupted simple sutures with number 0 polyglycan thread.
3. Approximate the two-levator ani muscles.

4. Strengthen the posterior vaginal wall by overlapping the two vaginal flaps; thus, we also add extra support to the mended IAS and keeping the natural body collagen.

5. Repair the perineum. We put a Foley’s catheter and vagina pack for 24 hours.

*Fig. 21: Images with 3D US of a patient with FI that show torn IAS. The anal canal is wide and open and the IAS is horseshoe*
Figs 22A to D: Surgical steps of posterior repair. We dissect the IAS from the posterior vaginal wall (A). We mend the sphincter (B and C), in addition, we approximate the two levator ani muscles by two stitches, but we do not tie them till we finish overlapping the posterior vaginal wall (D) dysfunction. The new repair operation is a simple successful vaginal operation with no use of synthetic materials with its complications. It corrects the anatomy of the pelvic organs in order to restore their normal physiology.

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