This guideline covers assessing and managing urinary incontinence and pelvic organ prolapse in women aged 18 years and older. It also covers complications associated with mesh surgery for these conditions.

**Who is it for?**

- Healthcare professionals
- Service commissioners
- Women with urinary incontinence, pelvic organ prolapse, or complications associated with mesh surgery for urinary incontinence or pelvic organ prolapse, their families and carers, and the public

This guideline will update NICE guideline CG171 (published September 2013). NICE guideline CG171 updated NICE guideline CG40 (published October 2006).

We have reviewed the evidence in the following areas of the 2013 guideline:

- multidisciplinary teams
- urodynamic testing in urinary incontinence
- absorbent products for urinary incontinence
- medicines for overactive bladder
- botulinum toxin type A for overactive bladder
- surgical procedures for stress urinary incontinence.
We have also reviewed the evidence and added new recommendations in these areas:

- assessing and managing pelvic organ prolapse
- managing coexisting urinary incontinence and pelvic organ prolapse
- assessing and managing complications associated with mesh surgery for urinary incontinence or pelvic organ prolapse.

You are invited to comment on the new and updated recommendations. These are marked as [2019].

We have not reviewed the evidence for the recommendations shaded in grey, and cannot accept comments on them. In some cases, we have made minor wording changes for clarification.

See update information for a full explanation of what is being updated.

This draft guideline contains:

- the draft recommendations
- recommendations for research
- the guideline context.

Information about how the guideline was developed is on the guideline’s page on the NICE website. This includes the evidence reviews, the scope, and details of the committee and any declarations of interest.

Full details of the evidence and the committee’s discussion on the 2019 recommendations are in the evidence reviews. Evidence for the 2013 recommendations is in the full version of the 2013 guideline.

For definitions of medical terms used in the guideline see supplementary material B: glossary and abbreviations.
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1 **Recommendations**

People have the right to be involved in discussions and make informed decisions about their care, as described in your care. *Making decisions using NICE guidelines* explains how we use words to show the strength (or certainty) of our recommendations, and has information about prescribing medicines (including off-label use), professional guidelines, standards and laws (including on consent and mental capacity), and safeguarding.

2 **1.1 Multidisciplinary teams**

3 **Local multidisciplinary teams**

4 **1.1.1 Local multidisciplinary teams (MDTs) for women with primary stress urinary incontinence (UI), overactive bladder (OAB) or primary prolapse**

5 should:

6 - review the proposed treatment for all women offered invasive procedures for primary stress UI, OAB or primary prolapse
7 - review the proposed management for women with primary stress UI, OAB or primary prolapse if input from a wider range of professionals is needed
8 - work within an established clinical network that has access to a regional MDT. [2019]

9 **1.1.2 Local MDTs for women with primary stress UI, OAB or primary prolapse**

10 should include:

11 - 2 urogynaecologists or urologists with expertise in female urology
12 - a urogynaecology, urology or continence specialist nurse
13 - a pelvic floor specialist physiotherapist

14 and may also include:

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1 NHS England is consulting on specialised gynaecology surgery and complex urogynaecology conditions service specifications. The consultation closes on 13 November 2018.
Regional multidisciplinary teams

Regional MDTs that deal with complex pelvic floor dysfunction and mesh-related problems should review the proposed treatment for women if:

- they are having repeat continence surgery
- they are having repeat, same-site prolapse surgery
- their preferred treatment option is not available in the referring hospital
- they have co-existing bowel problems that may need intervention
- vaginal mesh for prolapse is a treatment option for them
- they have mesh complications or unexplained symptoms after mesh surgery for UI or prolapse. [2019]

Regional MDTs that deal with complex pelvic floor dysfunction and mesh-related problems should include:

- a subspecialist in urogynaecology
- a urologist with expertise in female urology
- a urogynaecology, urology or continence specialist nurse
- a pelvic floor specialist physiotherapist
- a radiologist with expertise in pelvic floor imaging
- a colorectal surgeon with expertise in pelvic floor problems
- a pain specialist
- a healthcare professional trained in biofeedback

and may also include:

- a member of the care of the elderly team
- an occupational therapist
- a plastic surgeon. [2019]
1.1.5 Regional MDTs that deal with complex pelvic floor dysfunction and mesh-related problems should have ready access to the following services:

- psychology
- psychosexual counselling
- chronic pain management
- bowel symptom management
- neurology. [2019]

1.2 Collecting data on mesh surgery and mesh-related complications

1.2.1 In women having mesh surgery for stress urinary incontinence or pelvic organ prolapse, or who have mesh-related complications, seek consent to enter the data listed in recommendation 1.2.2 in a national registry and give them a copy of their data.

1.2.2 Ensure that the following data are collected in a national registry of surgery involving mesh insertion to treat urinary incontinence (UI) or pelvic organ prolapse (POP) in women:

- all surgical procedures for urinary incontinence or pelvic organ prolapse that involve the insertion of synthetic polypropylene mesh, including:
  - date and details of the procedure
  - mesh material and type of sutures.
- the woman’s NHS number
- hospital and consultant identifiers
- follow-up information on key short- and long-term (at least 5 years) outcomes, including:
  - symptom improvement or deterioration
  - objective measures of UI or POP
  - adverse events
  - suspected and confirmed mesh-related complications
- date and details of any investigation for mesh-related complications
1. date and details of any surgical or non-surgical intervention for mesh-related complications. [2019]

1.2.3 The national registry of surgery involving mesh insertion to treat urinary incontinence or pelvic organ prolapse in women should report annually and be quality assured. [2019]

1.3 Assessing urinary incontinence

History taking and physical examination

1.3.1 At the initial clinical assessment, categorise the woman's urinary incontinence (UI) as stress UI, mixed UI or urgency UI/overactive bladder (OAB). Start initial treatment on this basis. In mixed UI, direct treatment towards the predominant symptom. [2006]

1.3.2 If stress incontinence is the predominant symptom in mixed UI, discuss with the woman the benefit of non-surgical management and medicines for OAB before offering surgery. [2013, amended 2019]

1.3.3 During the clinical assessment seek to identify relevant predisposing and precipitating factors and other diagnoses that may require referral for additional investigation and treatment. [2006]

Assessing pelvic floor muscles

1.3.4 Undertake routine digital assessment to confirm pelvic floor muscle contraction before the use of supervised pelvic floor muscle training for the treatment of UI. [2006, amended 2013]

Urine testing

1.3.5 Undertake a urine dipstick test in all women presenting with UI to detect the presence of blood, glucose, protein, leucocytes and nitrites in the urine. [2006]

1.3.6 If women have symptoms of urinary tract infection (UTI) and their urine tests positive for both leucocytes and nitrites, send a midstream urine
specimen for culture and analysis of antibiotic sensitivities. Prescribe an appropriate course of antibiotic treatment pending culture results. [2006]

1.3.7 If women have symptoms of UTI and their urine tests negative for either leucocytes or nitrites, send a midstream urine specimen for culture and analysis of antibiotic sensitivities. Consider the prescription of antibiotics pending culture results. [2006]

1.3.8 If women do not have symptoms of UTI, but their urine tests positive for both leucocytes and nitrites, do not offer antibiotics without the results of midstream urine culture. [2006]

1.3.9 If a woman does not have symptoms of UTI and her urine tests negative for either leucocytes or nitrites, do not send a urine sample for culture because she is unlikely to have UTI. [2006]

Assessing residual urine

1.3.10 Measure post-void residual volume by bladder scan or catheterisation in women with symptoms suggestive of voiding dysfunction or recurrent UTI. [2006]

1.3.11 Use a bladder scan in preference to catheterisation on the grounds of acceptability and lower incidence of adverse events. [2006]

Symptom scoring and quality-of-life assessment

1.3.12 Use the following incontinence-specific quality-of-life scales when therapies are being evaluated: ICIQ, BFLUTS, I-QOL, SUIQQ, UISS, SEAPI-QMM, ISI and KHQ. [2006]

Bladder diaries

1.3.13 Use bladder diaries in the initial assessment of women with UI or OAB. Encourage women to complete a minimum of 3 days of the diary covering

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2 NICE is developing guidance on antimicrobial prescribing for lower urinary tract infections (publication expected 12 October 2018).
3 See the 2013 full guideline for details.
Pad testing

1.3.14 Do not use pad tests in the routine assessment of women with UI. [2006]

Urodynamic testing

1.3.15 Do not perform multichannel filling and voiding cystometry before primary surgery if stress UI or stress-predominant mixed UI is diagnosed based on a detailed clinical history and examination. [2019]

1.3.16 After undertaking a detailed clinical history and examination, perform multichannel filling and voiding cystometry before surgery for stress UI in women who have any of the following:

- urge-predominant mixed UI or UI in which the type is unclear
- symptoms suggestive of voiding dysfunction
- anterior or apical prolapse
- a history of previous surgery for stress UI. [2019]

Other tests of urethral competence

1.3.17 Do not use the Q-tip, Bonney, Marshall and Fluid-Bridge tests in the assessment of women with UI. [2006]

Cystoscopy

1.3.18 Do not use cystoscopy in the initial assessment of women with UI alone. [2006]

Imaging

1.3.19 Do not use imaging (MRI, CT, X-ray) for the routine assessment of women with UI. Do not use ultrasound other than for the assessment of residual urine volume. [2006]

Indications for referral to a specialist service

1.3.20 In women with UI, indications for consideration for referral to a specialist service include:
1.3.21 For women aged over 45 who have haematuria, or a recurrent or persistent unexplained UTI, follow the recommendations on referral for bladder cancer in the NICE guideline on suspected cancer. [2006, amended 2019]

1.4 Non-surgical management of urinary incontinence

Lifestyle interventions

1.4.1 Recommend a trial of caffeine reduction to women with overactive bladder (OAB). [2006]

1.4.2 Consider advising women with urinary incontinence (UI) or OAB and a high or low fluid intake to modify their fluid intake. [2006]

1.4.3 Advise women with UI or OAB who have a BMI greater than 30 to lose weight. [2006]

Physical therapies

1.4.4 Offer a trial of supervised pelvic floor muscle training of at least 3 months' duration as first-line treatment to women with stress or mixed UI. [2006]

1.4.5 Pelvic floor muscle training programmes should comprise at least 8 contractions performed 3 times per day. [2006]
1.4.6 Do not use perineometry or pelvic floor electromyography as biofeedback as a routine part of pelvic floor muscle training. [2006]

1.4.7 Continue an exercise programme if pelvic floor muscle training is beneficial. [2006]

1.4.8 Do not routinely use electrical stimulation in the treatment of women with OAB. [2006]

1.4.9 Do not routinely use electrical stimulation in combination with pelvic floor muscle training. [2006]

1.4.10 Electrical stimulation and/or biofeedback should be considered for women who cannot actively contract pelvic floor muscles to aid motivation and adherence to therapy. [2006]

### Behavioural therapies

1.4.11 Offer bladder training lasting for a minimum of 6 weeks as first-line treatment to women with urgency or mixed UI. [2006]

1.4.12 If women do not achieve satisfactory benefit from bladder training programmes, the combination of an OAB medicine with bladder training should be considered if frequency is a troublesome symptom. [2006]

### Neurostimulation

1.4.13 Do not offer transcutaneous sacral nerve stimulation (surface electrodes placed above the sacrum, often known as transcutaneous electrical nerve stimulation [TENS]) to treat OAB in women. [2013]

1.4.14 Explain that there is insufficient evidence to recommend the use of transcutaneous posterior tibial nerve stimulation (surface electrodes placed above the posterior tibial nerve) to treat OAB. [2013]

1.4.15 Do not offer transcutaneous posterior tibial nerve stimulation for OAB. [2013]
1.4.16 Do not offer percutaneous posterior tibial nerve stimulation (needles inserted close to the posterior tibial nerve) for OAB unless:

- there has been a **local** multidisciplinary team (MDT) review **and**
- non-surgical management including OAB medicine treatment has not worked adequately **and**
- the woman does not want botulinum toxin type A\(^4\) or percutaneous sacral nerve stimulation. **[2013, amended 2019]**

1.4.17 Explain that there is insufficient evidence to recommend the use of percutaneous posterior tibial nerve stimulation to routinely treat OAB. **[2013]**

**Absorbent containment products, urinals and toileting aids**

1.4.18 Absorbent containment products, hand held urinals and toileting aids should not be considered as a treatment for UI. Use them only as:

- a coping strategy pending definitive treatment
- an adjunct to ongoing therapy
- long-term management of UI only after treatment options have been explored. **[2006]**

1.4.19 Offer a review at least once a year to women who are using absorbent containment products for long-term management of UI. The review should cover:

- routine assessment of continence
- assessment of skin integrity
- changes to symptoms, comorbidities, lifestyle, mobility, medication, BMI, and social and environmental factors
- the suitability of alternative treatment options
- the efficacy of the absorbent containment product the woman is currently using and the quantities used

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\(^4\) At the time of consultation (October 2018) most botulinum toxin type A preparations did not have a UK marketing authorisation for this indication. Evidence was only available for the licensed botulinum toxin type A preparation (Botox, Allergan).
• long-term management strategies. [2019]

1.4.20 Reviews for women who are using absorbent containment products for long-term management of UI should be carried out by either:

• a registered healthcare professional who is trained in assessing continence and making referrals to specialist services or
• a non-registered healthcare worker, under the supervision of a registered healthcare professional who is trained in assessing continence and making referrals to specialist services.

See indications for referral to a specialist service in this guideline. [2019]

Catheters

1.4.21 Bladder catheterisation (intermittent or indwelling urethral or suprapubic) should be considered for women in whom persistent urinary retention is causing incontinence, symptomatic infections, or renal dysfunction, and in whom this cannot otherwise be corrected. Healthcare professionals should be aware, and explain to women, that the use of indwelling catheters in urgency UI may not result in continence. [2006]

1.4.22 Offer intermittent urethral catheterisation to women with urinary retention who can be taught to self-catheterise or who have a carer who can perform the technique. [2006]

1.4.23 Give careful consideration to the impact of long-term indwelling urethral catheterisation. Discuss the practicalities, benefits and risks with the woman or, if appropriate, her carer. Indications for the use of long-term indwelling urethral catheters for women with UI include:

• chronic urinary retention in women who are unable to manage intermittent self-catheterisation
• skin wounds, pressure ulcers or irritations that are being contaminated by urine
• distress or disruption caused by bed and clothing changes
• where a woman expresses a preference for this form of management. [2006]

1.4.24 Indwelling suprapubic catheters should be considered as an alternative to long-term urethral catheters. Be aware, and explain to women, that they may be associated with lower rates of symptomatic UTI, 'bypassing', and urethral complications than indwelling urethral catheters. [2006]

**Products to prevent leakage**

1.4.25 Do not use intravaginal and intraurethral devices for the routine management of UI in women. Do not advise women to consider such devices other than for occasional use when necessary to prevent leakage, for example during physical exercise. [2006]

**Complementary therapies**

1.4.26 Do not recommend complementary therapies for the treatment of UI or OAB. [2006]

**Medicines**

1.4.27 Before starting treatment with a medicine for OAB, explain to the woman:

- the likelihood of the medicine being successful
- the common adverse effects associated with the medicine
- that some adverse effects of anticholinergic medicines, such as dry mouth and constipation, may indicate that the medicine is starting to have an effect
- that she may not see the full benefits until she has been taking the medicine for 4 weeks
- that the long-term effects of anticholinergic medicines for OAB on cognitive function are uncertain. [2019]

1.4.28 When offering anticholinergic medicines to treat OAB, take account of the woman’s:

- coexisting conditions (such as poor bladder emptying, cognitive impairment or dementia)
• current use of other medicines that affect total anticholinergic load
• risk of adverse effects, including cognitive impairment. [2019]

1.4.29 For women who have a diagnosis of dementia and for whom anticholinergic medicines are an option, follow the recommendations on medicines that may cause cognitive impairment in the NICE guideline on dementia. [2019]

**Choosing medicine**

1.4.30 Do not offer women flavoxate, propantheline or imipramine to treat UI or OAB. [2013]

1.4.31 Do not offer oxybutynin (immediate release) to older women who may be at higher risk of a sudden deterioration in their physical or mental health. [2013, amended 2019]

1.4.32 Offer the anticholinergic medicine with the lowest acquisition cost to treat OAB or mixed UI in women. [2019]

1.4.33 If the first medicine for OAB or mixed UI is not effective or well-tolerated, offer another medicine with a low acquisition cost. [2013]

1.4.34 Offer a transdermal OAB treatment to women unable to tolerate oral medicines. [2013]

1.4.35 For guidance on mirabegron see mirabegron for treating symptoms of overactive bladder (NICE technology appraisal guidance 290). [2013]

1.4.36 The use of desmopressin may be considered specifically to reduce nocturia in women with UI or OAB who find it a troublesome symptom. Use particular caution in women with cystic fibrosis and avoid in those over 65 years with cardiovascular disease or hypertension. [2013]

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5 This could be any medicine with the lowest acquisition cost from any of the medicines reviewed in 2013. The evidence review considered the following medicines: darifenacin, fesoterodine, oxybutynin (immediate release), oxybutynin (extended release), oxybutynin (transdermal), oxybutynin (topical gel), propiverine, propiverine (extended release), solifenacin, tolterodine (immediate release), tolterodine (extended release), trosium and trosium (extended release). See chapter 6 of the 2013 full guideline.
1.4.37 Do not use duloxetine as a first-line treatment for women with predominant stress UI. Do not routinely offer duloxetine as a second-line treatment for women with stress UI, although it may be offered as second-line therapy if women prefer pharmacological to surgical treatment or are not suitable for surgical treatment. If duloxetine is prescribed, counsel women about its adverse effects. [2006]

1.4.38 Do not offer systemic hormone replacement therapy to treat UI. [2006]

1.4.39 Offer intravaginal oestrogens to treat OAB symptoms in postmenopausal women with vaginal atrophy. [2006]

**Reviewing medicine**

1.4.40 Offer a face-to-face or telephone review 4 weeks after starting a new medicine for OAB. Ask the woman if she is satisfied with the treatment and:

- if improvement is optimal, continue treatment
- if there is no or suboptimal improvement, or intolerable adverse effects, change the dose or try an alternative medicine for OAB (see recommendations 1.4.33 and 1.4.34), and review again 4 weeks later. [2013]

1.4.41 Offer a review before 4 weeks if the adverse events of a medicine for OAB are intolerable. [2013]

1.4.42 Refer women who have tried taking medicine for OAB, but for whom it has not been successful or tolerated, to secondary care to consider further treatment. [2013]

1.4.43 Offer a further face-to-face or telephone review if a medicine for OAB or UI stops working after an initial successful 4-week review. [2013]

1.4.44 Offer a review in primary care to women who remain on long-term medicine for OAB or UI every 12 months, or every 6 months if they are aged over 75. [2019]
Invasive procedures for overactive bladder

1.4.45 For women with OAB that has not responded to non-surgical management or treatment with medicine and who wish to discuss further treatment options:

- offer urodynamic investigation to determine whether detrusor overactivity is causing her OAB symptoms and
  - if detrusor overactivity is causing her OAB symptoms, offer an invasive procedure in line with recommendations 1.4.46 to 1.4.59 in this guideline
  - if there is no detrusor overactivity, seek advice on further management from the local MDT. [2013, amended 2019]

Botulinum toxin type A injection

1.4.46 After a local MDT review, offer bladder wall injection with botulinum toxin type A⁶ to women with OAB caused by detrusor overactivity that has not responded to non-surgical management, including pharmacological treatments. [2019]

1.4.47 Consider treatment with botulinum toxin type A⁶ after a local MDT review for women with symptoms of OAB in whom urodynamics has not demonstrated detrusor overactivity, if the symptoms have not responded to non-surgical management and the woman does not wish to have other invasive treatments. [2019]

1.4.48 After a local MDT review, discuss the risks and benefits of treatment with botulinum toxin type A⁶ with women and explain:

- the likelihood of complete or partial symptom relief
- the process of clean intermittent catheterisation, the risks, and how long it might need to be continued

⁶At the time of consultation (October 2018) most botulinum toxin type A preparations did not have a UK marketing authorisation for this indication. Evidence was only available for the licensed botulinum toxin type A preparation (Botox, Allergan).
• the risk of adverse effects, including an increased risk of urinary tract infection
• that there is not much evidence about how long the injections work for, how well they work in the long term and their long-term risks. [2019]

1.4.49 Start treatment with botulinum toxin type A only if the woman is willing, in the event of developing significant voiding dysfunction:

• to perform clean intermittent catheterisation on a regular basis for as long as needed or
• to accept a temporary indwelling catheter if she is unable to perform clean intermittent catheterisation. [2013, amended 2019]

1.4.50 Use 100 units as the initial dose of botulinum toxin type A to treat OAB in women. [2019]

1.4.51 Offer a face-to-face or telephone review within 12 weeks of the first treatment with botulinum toxin type A to assess the response to treatment and adverse effects, and

• if there is good symptom relief, tell the woman how to self-refer for prompt specialist review if symptoms return, and offer repeat treatment as necessary
• if there is inadequate symptom relief, consider increasing subsequent doses of botulinum toxin type A to 200 units and review within 12 weeks
• if there was no effect, discuss with the local MDT. [2019]

7 At the time of consultation (October 2018) most botulinum toxin type A preparations did not have a UK marketing authorisation for this indication. Evidence was only available for the licensed botulinum toxin type A preparation (Botox, Allergan).
8 At the time of consultation (October 2018) the dose of 100 units of botulinum toxin type A has a UK marketing authorisation for overactive bladder. The dose of 200 units has a UK marketing authorisation for urinary incontinence due to neurogenic detrusor overactivity. Note that units of botulinum toxin type A are not interchangeable between preparations. If prescribing outside of the marketing authorisation, the prescriber should follow relevant professional guidance, taking full responsibility for the decision. Informed consent should be obtained and documented. See the General Medical Council’s Prescribing guidance: prescribing unlicensed medicines for further information.
If symptom relief has been adequate after injection of 100 units\textsuperscript{9,10} of botulinum toxin type A but has lasted for less than 6 months, consider increasing subsequent doses of botulinum toxin type A to 200 units\textsuperscript{10} and review within 12 weeks. [2019]

Do not offer botulinum toxin type B to women with OAB. [2019]

**Percutaneous sacral nerve stimulation**

Offer percutaneous sacral nerve stimulation to women after local or regional MDT review if:

- their OAB has not responded to non-surgical management including medicines and
- they are unable to perform clean intermittent catheterisation. [2013, amended 2019]

Consider percutaneous sacral nerve stimulation after local MDT review if a woman's OAB has not responded to non-surgical management (including medicines) and botulinum toxin type A\textsuperscript{9}. [2013, amended 2019]

Discuss the long-term implications of percutaneous sacral nerve stimulation with women including:

- the need for test stimulation and probability of the test's success
- the risk of failure
- the long-term commitment
- the need for surgical revision

\textsuperscript{9} At the time of consultation (October 2018) most botulinum toxin type A preparations did not have a UK marketing authorisation for this indication. Evidence was only available for the licensed botulinum toxin type A preparation (Botox, Allergan).

\textsuperscript{10} At the time of consultation (October 2018) the dose of 100 units of botulinum toxin type A has a UK marketing authorisation for overactive bladder. The dose of 200 units has a UK marketing authorisation for urinary incontinence due to neurogenic detrusor overactivity. Note that units of botulinum toxin type A are not interchangeable between preparations. If prescribing outside of the marketing authorisation, the prescriber should follow relevant professional guidance, taking full responsibility for the decision. Informed consent should be obtained and documented. See the General Medical Council's Prescribing guidance: prescribing unlicensed medicines for further information.
Tell women how to self-refer for prompt specialist review if symptoms return following a percutaneous sacral nerve stimulation procedure. [2013]

**Augmentation cystoplasty**

Restrict augmentation cystoplasty for the management of idiopathic detrusor overactivity to women whose condition has not responded to non-surgical management and who are willing and able to self-catheterise. Preoperative counselling for the woman or her carer should include common and serious complications: bowel disturbance, metabolic acidosis, mucus production and/or retention in the bladder, UTI and urinary retention. Discuss the small risk of malignancy occurring in the augmented bladder. Provide life-long follow-up. [2006, amended 2013]

**Urinary diversion**

Urinary diversion should be considered for a woman with OAB only when non-surgical management has failed, and if botulinum toxin type A, percutaneous sacral nerve stimulation and augmentation cystoplasty are not appropriate or are unacceptable to her. Provide life-long follow-up. [2006, amended 2013]

**1.5 Surgical management of stress urinary incontinence**

When offering a surgical procedure discuss with the woman the risks and benefits of the different treatment options for stress urinary incontinence (UI). Include information about differences in type of anaesthesia, expected length of hospital stay, surgical incisions and expected recovery period. [2013, amended 2019]

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11 At the time of consultation (October 2018) most botulinum toxin type A preparations did not have a UK marketing authorisation for this indication. Evidence was only available for the licensed botulinum toxin type A preparation (Botox, Allergan).

12 NICE is developing shared decision aids on surgery for stress urinary incontinence and pelvic organ prolapse. They will be published with the final guideline in April 2019.
If non-surgical management for stress UI has failed, offer the woman a choice of:

- colposuspension (open or laparoscopic) or
- a retropubic mid-urethral mesh sling or
- an autologous rectus fascial sling. [2019]

When offering surgery for stress UI, advise the woman that there are long-term complications associated with all procedures and uncertainty about the proportion of women affected. [2019]

If the woman’s chosen procedure is not available from the consulting surgeon, refer her to an alternative surgeon. [2019]

### Mid-urethral mesh sling procedures

When offering a retropubic mid-urethral mesh sling, advise the woman that it is a permanent implant and complete removal might not be possible. [2019]

If a retropubic mid-urethral mesh sling is inserted:

- give the woman written information on the implant including name, manufacturer, date of insertion, and implanting surgeon’s name and contact details;
- ensure that details of the procedure and its subsequent short- and long-term outcomes are collected in a national registry (see collecting data on mesh surgery and mesh-related complications in this guideline). [2019]

When planning a retropubic mid-urethral mesh sling procedure, surgeons should:

- use a device manufactured from type 1 macroporous polypropylene mesh
- consider using a retropubic mid-urethral mesh sling coloured for high visibility, for ease of insertion and revision. [2019]
1.5.8 Do not offer a transobturator approach unless there are specific clinical circumstances (for example, multiple previous abdominal procedures) in which the retropubic approach should be avoided. [2019]

1.5.9 Do not use the ‘top-down’ retropubic mid-urethral mesh sling approach or single-incision sub-urethral short mesh sling insertion except as part of a clinical trial. [2019]

Other procedures

1.5.10 Do not offer women anterior colporrhaphy, needle suspensions, paravaginal defect repair or the Marshall–Marchetti–Krantz procedure to treat stress UI. [2019]

1.5.11 Do not offer women porcine dermis slings to treat stress UI. [2019]

1.5.12 Do not offer women intramural bulking agents to manage stress UI unless alternative surgical procedures are not suitable or acceptable. Explain to women that:

- repeat injections may be needed to achieve efficacy
- efficacy is limited and diminishes with time. [2019]

1.5.13 Do not offer women an artificial urinary sphincter to manage stress UI unless previous surgery has failed. Offer lifelong follow-up to women who have an artificial urinary sphincter. [2019]

Follow-up after surgery

1.5.14 Offer a follow-up appointment within 6 months to all women who have had a surgical procedure to treat stress UI. For women who have had retropubic mid-urethral mesh sling surgery the follow-up appointment should include a vaginal examination to check for exposure or extrusion of the mesh sling. [2019]

1.5.15 For women whose primary surgical procedure for stress UI has failed (including women whose symptoms have returned):
• seek advice on assessment and management from a regional MDT that
deals with complex pelvic floor dysfunction or
• if the woman does not wish to have another surgical procedure, offer
her advice about managing urinary symptoms and explain that if she
changes her mind at a later date she can book a review appointment to
discuss past tests and interventions and reconsider her treatment
options. [2019]

1.6 Assessing pelvic organ prolapse

1.6.1 For women presenting in primary care with symptoms or an incidental
finding of vaginal prolapse:
• take a history to include symptoms of prolapse, urinary, bowel and
sexual function
• do an examination to rule out a pelvic mass or other pathology and to
document the presence of prolapse (see the sections on ovarian
cancer and bladder cancer in the NICE guideline on suspected cancer)
• discuss the woman’s treatment preferences with her, and refer if
needed. [2019]

1.6.2 For women referred to secondary care for an unrelated condition who
have incidental symptoms or an incidental finding of vaginal prolapse,
consider referral to a clinician with expertise in prolapse. [2019]

1.6.3 For women who are referred for specialist evaluation of vaginal prolapse,
perform an examination to:
• assess and record the presence and degree of prolapse of the anterior,
central and posterior vaginal compartments of the pelvic floor, using the
POP-Q (Pelvic Organ Prolapse Quantification System)
• assess the activity of the pelvic floor muscles
• assess for vaginal atrophy
• rule out a pelvic mass or other pathology. [2019]
1.6.4 For women with pelvic organ prolapse consider using a validated pelvic floor symptom questionnaire to aid assessment and decision-making. [2019]

1.6.5 Do not routinely perform imaging to document the presence of vaginal prolapse if a prolapse is detected by physical examination. [2019]

1.6.6 If the woman has symptoms of prolapse that are not explained by findings from a physical examination, consider repeating the examination with the woman standing or squatting, or at a different time of day. [2019]

1.6.7 Consider investigating the following symptoms in women with pelvic organ prolapse:

- urinary symptoms that are bothersome and for which surgical intervention is an option
- symptoms of obstructed defaecation or faecal incontinence (the NICE guideline on faecal incontinence in adults has recommendations on baseline assessment of faecal incontinence)
- pain
- symptoms that are not explained by examination findings. [2019]

1.7 Managing pelvic organ prolapse

1.7.1 Discuss management options with women who have pelvic organ prolapse, including no treatment, non-surgical treatment and all surgical options, taking into account:

- the woman’s preferences
- site of prolapse
- benefits and risks of individual procedures
- comorbidities, including cognitive or physical impairments
- age
- desire for childbearing
- previous abdominal or pelvic floor surgery
- lifestyle factors. [2019]
Lifestyle modification

1.7.2 Consider giving advice on lifestyle to women with pelvic organ prolapse, including information on:

- losing weight, if the woman is obese
- avoiding heavy lifting
- preventing constipation
- exercising and its effect on symptoms. [2019]

Topical oestrogen

1.7.3 Consider vaginal oestrogen for women with pelvic organ prolapse and signs of vaginal atrophy. For recommendations on managing urogenital atrophy see managing short-term menopausal symptoms in the NICE guideline on menopause. [2019]

1.7.4 Consider an oestrogen-releasing ring for women with pelvic organ prolapse and signs of vaginal atrophy who have cognitive or physical impairments that might make vaginal pessaries or creams difficult to use. [2019]

Pelvic floor muscle training

1.7.5 Consider a programme of supervised pelvic floor muscle training for at least 16 weeks as a first option for women with symptomatic POP-Q (Pelvic Organ Prolapse Quantification System) stage 1 or stage 2 pelvic organ prolapse. If the programme is beneficial, advise women to continue pelvic floor muscle training afterwards. [2019]

Pessaries

1.7.6 Consider a vaginal pessary for women with symptomatic pelvic organ prolapse, alone or in conjunction with supervised pelvic floor muscle training. [2019]

1.7.7 Refer women who have chosen a pessary to a urogynaecology service if pessary care is not available locally. [2019]

1.7.8 Before starting pessary treatment:
1. consider treating vaginal atrophy with topical oestrogen
2. explain that more than one pessary fitting may be needed to find a suitable pessary
3. discuss the effect of a pessary on sexual intercourse
4. describe common complications including vaginal discharge, bleeding and pessary expulsion
5. explain that the pessary should be removed at least once every 6 months. [2019]

Surgery

1.7.9 Offer surgery for pelvic organ prolapse to women whose symptoms have not improved with or who have declined non-surgical treatment. [2019]

1.7.10 Do not offer surgery to prevent incontinence in women having surgery for prolapse who do not have incontinence. [2019]

1.7.11 Explain to women considering surgery for anterior or apical prolapse who do not have incontinence that there is a risk of developing postoperative urinary incontinence and further treatment may be needed. [2019]

1.7.12 If a woman has agreed to have a surgical procedure for pelvic organ prolapse, before surgery discuss:

- the risks and benefits of each procedure, including changes in urinary, bowel and sexual function
- the risks of recurrent prolapse
- the role of intraoperative prolapse assessment in finalising the choice of surgical procedure. [2019]

1.7.13 If the woman’s chosen procedure for pelvic organ prolapse is not available from the consulting surgeon, refer her to an alternative surgeon. [2019]

1.7.14 If mesh is to be used in prolapse surgery, explain to the woman:

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13 NICE is developing shared decision aids on surgery for stress urinary incontinence and pelvic organ prolapse. They will be published with the final guideline in April 2019.
• what type of mesh will be used and whether it is permanent.
• the uncertainty about long-term complications associated with mesh and about the proportion of women affected. [2019]

1.7.15 If mesh is to be used in prolapse surgery

• give the woman written information on the implant including name, manufacturer, date of insertion, and implanting surgeon’s name and contact details
• ensure that details of the procedure and its subsequent short- and long-term outcomes are collected in a national registry (see collecting data on mesh surgery and mesh-related complications in this guideline). [2019]

Surgery for uterine prolapse

1.7.16 Discuss the options for surgery with women who have uterine prolapse, including surgery that will preserve the uterus and hysterectomy. [2019]

1.7.17 For women with uterine prolapse who wish to preserve their uterus, offer a choice of:

• vaginal sacrospinous hysteropexy with sutures
• sacro-hysteropexy with mesh (abdominal or laparoscopic)
• Manchester repair, unless the woman is considering a future pregnancy or might become pregnant. [2019]

1.7.18 For women with uterine prolapse who have no preference about preserving their uterus, offer a choice of:

• vaginal hysterectomy, with or without sacrospinous fixation with sutures
• sacro-hysteropexy with mesh (abdominal or laparoscopic)
• vaginal sacrospinous hysteropexy with sutures
• Manchester repair. [2019]

1.7.19 If sacro-hysteropexy with mesh (abdominal or laparoscopic) is used, ensure that details of the procedure and its subsequent short- and long-
term outcomes are collected in a national registry (see collecting data on mesh surgery and mesh-related complications in this guideline). [2019]

### Surgery for vault prolapse

1. Offer women with vault prolapse a choice of:
   - sacrocolpopexy (abdominal or laparoscopic) with mesh
   - vaginal sacrospinous fixation with sutures. [2019]

2. If sacrocolpopexy (abdominal or laparoscopic) with mesh is used, ensure that details of the procedure and its subsequent short- and long-term outcomes are collected in a national registry (see collecting data on mesh surgery and mesh-related complications in this guideline). [2019]

### Colpocleisis for vault or uterine prolapse

3. Consider colpocleisis for women with vault or uterine prolapse who do not intend to have penetrative vaginal sex and who have a physical condition that may put them at increased risk of operative and postoperative complications. [2019]

### Surgery for anterior prolapse

4. Offer anterior repair without mesh to women with anterior vaginal wall prolapse. [2019]

5. Consider synthetic polypropylene or biological mesh insertion for women with recurrent anterior vaginal wall prolapse only after:
   - regional MDT review and
   - discussion with the woman about the risks of mesh insertion

6. and if:
   - apical support is adequate or
   - an abdominal approach is contraindicated. [2019]

7. If a synthetic polypropylene or biological mesh is inserted, ensure that details of the procedure and its subsequent short- and long-term
outcomes are collected in a national registry (see collecting data on mesh surgery and mesh-related complications in this guideline). [2019]

Surgery for posterior prolapse

1.7.25 Offer posterior vaginal repair without mesh to women with a posterior vaginal wall prolapse. [2019]

Follow-up after surgery

1.7.26 Offer women a review 6 months after surgery for prolapse. Ensure that the review includes a vaginal examination and, if mesh was used, check for mesh exposure. [2019]

1.8 Surgery for stress urinary incontinence and pelvic organ prolapse

1.8.1 Consider concurrent surgery for stress urinary incontinence (UI) and prolapse in women with anterior and/or apical prolapse and stress UI. [2019]

1.8.2 When offering concurrent surgery for stress UI and prolapse explain to the woman:

- that there is uncertainty about whether the combined procedure is effective for treating stress UI beyond 1 year, and that stress UI might persist despite surgery
- the risk of complications related to having surgery for stress UI at the same time as prolapse surgery compared with the risk of complications related to having sequential surgery. [2019]

1.9 Assessing complications associated with mesh surgery

1.9.1 For women who report new-onset symptoms after having mesh surgery for urinary incontinence (UI) or pelvic organ prolapse, evaluate whether the symptoms might be caused by a mesh-related complication. These symptoms could include:
• pain or sensory change in the back, abdomen, vagina, pelvis, leg, groin or perineum that is:
  – either unprovoked, or provoked by movement or sexual activity and
  – either generalised, or in the distribution of a specific nerve, such as the obturator nerve
• vaginal problems including discharge, bleeding, painful sexual intercourse, penile trauma or pain
• urinary problems including recurrent infection, incontinence, retention, or difficulty or pain during voiding
• bowel problems including difficulty or pain on defaecation, faecal incontinence, rectal bleeding or passage of mucus
• symptoms of infection, either alone or in combination with any of the symptoms outlined above. [2019]

1.9.2 Refer women with a suspected mesh-related complication to a urogynaecologist, urologist or colorectal surgeon for specialist assessment. [2019]

1.9.3 For women who are referred for specialist evaluation of a suspected mesh complication:

• take a history of all past surgical procedures for prolapse or incontinence using mesh, including the dates, type of mesh and site of mesh placement and the relationship of the symptoms to the surgical procedure(s)
• consider using a validated pelvic floor symptom questionnaire and a pain questionnaire to aid assessment and decision-making
• perform a vaginal examination to:
  – assess whether mesh is palpable, exposed or extruded
  – localise pain and its anatomical relationship to mesh
• consider performing a rectal examination, if indicated, to assess for the presence of mesh perforation or fistula
• consider performing a neurological assessment to assess the
distribution of pain, if present, sensory alteration or muscle weakness.

[2019]

1.9.4 For women with a confirmed mesh-related complication or unexplained
symptoms after a mesh procedure:

• refer to a consultant at a regional centre specialising in the diagnosis
and management of mesh-related complications or
• if the woman has a vaginal exposure of mesh that is smaller than 1 cm²
and no other symptoms, follow recommendation 1.10.3 in this
guideline. [2019]

1.9.5 The responsible consultant should develop an individualised investigation
plan for each woman with suspected or confirmed mesh-related
complications, involving other members of the regional MDT if needed
and using table 1 in this guideline.

1.9.6 The responsible consultant should ensure that details of any confirmed
mesh-related complications are:

• collected in a national registry (see collecting data on mesh surgery
and mesh-related complications in this guideline) and
• reported to the Medicines and Healthcare Products Regulatory Agency.

[2019]
### Table 1 Investigations for assessing suspected mesh-related complications

Individualised investigation plans may include, but are not limited to, one or more of these investigations.

<table>
<thead>
<tr>
<th>Investigation</th>
<th>Type of mesh</th>
<th>Indications</th>
<th>Benefits and risks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Examination under anaesthesia</td>
<td>All types of mesh</td>
<td>Pain, or suspected:</td>
<td>Benefits:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• vaginal or rectal exposure or extrusion</td>
<td><strong>Allows diagnosis when not revealed by awake examination or when an awake examination is not tolerated</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• sinus tract, urinary or bowel fistula</td>
<td><strong>Risks</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>Anaesthetic risk</strong></td>
</tr>
<tr>
<td>Cystourethroscopy</td>
<td>All types of mesh</td>
<td>Suspected:</td>
<td>Benefits:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• urethral perforation</td>
<td><strong>allows diagnosis by direct visualisation</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• bladder perforation</td>
<td><strong>aids management planning</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• fistula</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• calculus on suture or mesh material</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>Risks</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>Anaesthetic risk and risk of urinary tract infection</strong></td>
</tr>
<tr>
<td>Sigmoidoscopy</td>
<td>Abdominally, laparoscopically or vaginally placed mesh for pelvic organ prolapse</td>
<td>Suspected bowel perforation by mesh</td>
<td>Benefits:</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>allows diagnosis by direct visualisation</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>aids management planning</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>Risks</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>anaesthetic risk if carried out under anaesthetic</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td><strong>risk of bowel perforation</strong></td>
</tr>
<tr>
<td>Laparoscopy</td>
<td>Abdominally or laparoscopically placed mesh for pelvic organ prolapse</td>
<td>• Pain</td>
<td>Benefits:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Suspected bowel entrapment around mesh</td>
<td><strong>allows diagnosis by direct visualisation</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Suspected adhesions secondary to</td>
<td><strong>aids management planning</strong></td>
</tr>
</tbody>
</table>

Urinary incontinence and pelvic organ prolapse in women NICE guideline DRAFT (October 2018)
<table>
<thead>
<tr>
<th>MRI. protocolled and reported by a clinician with experience in interpreting mesh complications</th>
<th>All types of mesh</th>
<th><strong>Risks</strong></th>
<th><strong>Benefits</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>- anaesthetic risk</td>
<td>- show implanted material and complications nearby</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- risks of laparoscopy, including bowel injury</td>
<td>- show location of mesh in relation to the vaginal wall and sacrum</td>
</tr>
</tbody>
</table>

**Risks**
Generally regarded as safe, with a low risk of short- and long-term harms

**Benefits**

**Ultrasound scan (transperineal, transvaginal or translabial, or 3D), performed and reported by a clinician with experience in interpreting mesh complications**

<table>
<thead>
<tr>
<th>Vaginally placed mesh to treat incontinence</th>
<th>Pain</th>
<th><strong>Benefits</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Voiding dysfunction</td>
<td>- show implanted material and local complications</td>
</tr>
<tr>
<td></td>
<td>Suspected infection</td>
<td>- identify mid-urethral slings</td>
</tr>
<tr>
<td></td>
<td>Suspected urethral mesh perforation</td>
<td>- show location of mesh in relation to the vaginal wall and urethra</td>
</tr>
<tr>
<td></td>
<td>Anatomical mapping to guide excision surgery</td>
<td></td>
</tr>
</tbody>
</table>

**Risks**
Discomfort

**Benefits**

**CT**

<table>
<thead>
<tr>
<th>All types of mesh, although CT is not commonly used to show implanted material</th>
<th><strong>Benefits</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>May be useful in assessing for urinary fistulae or bowel injury</td>
</tr>
</tbody>
</table>
### Managing complications associated with mesh surgery

#### 2.10 General considerations before removing mesh

1.10.1 The decision to remove mesh for any indication should be made only after a discussion with the woman and a regional MDT review. [2019]

1.10.2 When discussing surgery to remove mesh inserted to treat urinary incontinence or pelvic organ prolapse, explain to the woman that:

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Mesh Types</th>
<th>Suspected:</th>
<th>Benefits</th>
<th>Risks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluoroscopic studies (cystography or contrast enema)*</td>
<td>All types of mesh</td>
<td>Suspected urinary or bowel fistula</td>
<td><strong>Benefits</strong></td>
<td><strong>Risks</strong></td>
</tr>
<tr>
<td>*Perform with water-soluble contrast media. Fluoroscopic studies and CT may be used according to local preference and expertise.</td>
<td></td>
<td></td>
<td>Aids management planning</td>
<td>Potential radiation-related harms</td>
</tr>
<tr>
<td>Urinary flow studies and post-void residual volume assessment or cystometry</td>
<td>All types of mesh</td>
<td>• Voiding dysfunction</td>
<td><strong>Benefits</strong></td>
<td><strong>Risks</strong></td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Urinary incontinence</td>
<td>Aids management planning</td>
<td>Urinary tract infection and radiation risks if fluoroscopy is used</td>
</tr>
<tr>
<td>Neurophysiology, including nerve conduction studies and quantitative sensory testing*</td>
<td>All types of mesh</td>
<td>Suspected nerve injury</td>
<td><strong>Benefits</strong></td>
<td><strong>Risks</strong></td>
</tr>
<tr>
<td>*Quantitative sensory testing is very specialised and not available in all centres</td>
<td></td>
<td></td>
<td>Allows diagnosis of impaired nerve function</td>
<td>Nerve conduction studies are difficult to perform and can induce more pain</td>
</tr>
</tbody>
</table>
surgery to remove mesh can have significant complications including
organ injury, worsening pain, and urinary, bowel and sexual dysfunction

- it is not certain that removing the mesh will relieve symptoms
- it might not be possible to remove all of the mesh
- removing only part of the mesh might be just as effective at improving
  symptoms as removing all of it
- urinary incontinence or prolapse can recur after the mesh has been
  removed. [2019]

Managing vaginal complications

1.10.3 Consider non-surgical treatment with topical oestrogen cream for women
with a single area of vaginal mesh exposure or extrusion that is smaller
than 1 cm². [2019]

1.10.4 Consider partial or complete surgical removal of the vaginal portion of
mesh if:

- the area of vaginal mesh sling exposure or extrusion is 1 cm² or larger
  or
- there has been no response to non-surgical treatment after a period of
  3 months. [2019]

1.10.5 Offer imaging and further treatment to women who have signs of infection
in addition to vaginal mesh exposure or extrusion. [2019]

1.10.6 Explain to women who have vaginal complications after mesh sling
surgery for stress UI that:

- complete removal of the vaginal portion of mesh sling is associated
  with a greater risk of recurrence of stress UI than partial removal
- partial removal is associated with a higher rate of further mesh sling
  extrusion
- complete removal might not be possible. [2019]

1.10.7 Explain to women who have vaginal complications after mesh surgery for
pelvic organ prolapse that:
Urinary incontinence and pelvic organ prolapse in women NICE guideline DRAFT (October 2018)

1.10.8 Explain to women who have vaginal complications after abdominally placed mesh that:

- removal is associated with a risk of urinary tract and bowel injury
- there is a risk of recurrent prolapse
- they might need abdominal surgery to remove the mesh
- complete removal might not be possible. [2019]

Managing pain and sexual dysfunction

1.10.9 For women who have pain or painful sexual intercourse suspected to be related to previous mesh surgery:

- if specialist assessment indicates a mesh-related complication, seek advice from a regional MDT
- if assessment and investigation do not show a mesh abnormality such as vaginal extrusion or exposure, or an infection, offer non-surgical treatments such as pain management, vaginal oestrogen, dilators, psychosexual counselling and physiotherapy
- if pain does not respond to initial management, seek advice from a regional MDT. [2019]

Managing urinary complications

1.10.10 Refer women who have mesh perforating the lower urinary tract to a centre for mesh complications for further assessment or management. [2019]

1.10.11 For women with urinary symptoms after mesh surgery for stress UI or pelvic organ prolapse who are considering mesh removal surgery, explain that:

- complete removal might not be possible
- complete removal has a higher risk of urinary tract or bowel injury than partial removal
- there may be a risk of recurrent prolapse. [2019]
• urinary symptoms might not improve and new symptoms might occur after complete or partial removal of the mesh
• stress UI might recur after mesh removal, and the risk of this happening is higher with complete than with partial mesh removal
• complete removal of the mesh might not be possible
• further treatment might be needed for mesh complications, or recurrent or persistent urinary symptoms
• there is a risk of adverse events such as urinary tract fistula. [2019]

1.10.12 Consider division of mesh sling for women with voiding dysfunction caused by mesh sling surgery. [2019]

1.10.13 Refer women considering excision of mesh sling for persistent voiding dysfunction to a centre specialising in the diagnosis and management of mesh-related complications for assessment and management. [2019]

1.10.14 For women considering surgery to alleviate voiding symptoms caused by mesh surgery, explain that:

• the risk of recurrent stress incontinence is higher after mesh excision than mesh division
• further surgery might be needed. [2019]

Managing bowel symptoms
1.10.15 For women who present with functional bowel disorders after mesh surgery for pelvic organ prolapse, follow the recommendations in the NICE guideline on faecal incontinence in adults for women with faecal incontinence or locally agreed protocols for women with obstructed defecation. [2019]

1.10.16 Discuss bowel complications in women that are directly related to mesh placement, such as erosion, stricture, or fistula, with a regional MDT that deals with complex pelvic floor dysfunction and mesh-related problems to formulate an individualised treatment plan. [2019]
1.10.17 Explain to women with bowel complications directly related to mesh placement that:

- complete removal might not be possible
- bowel symptoms might persist or recur after mesh removal
- they might need a temporary or permanent stoma after mesh removal.

[2019]

Recommendations for research

The guideline committee has made the following recommendations for research.

Key recommendations for research

1 Anticholinergic medicines

What is the effectiveness and safety of anticholinergic medicines for overactive bladder in older women?

To find out why the committee made this research recommendation see evidence review C: risks to cognitive function for women taking anticholinergic drugs for overactive bladder.

2 Colpocleisis compared with sacrospinous fixation for pelvic organ prolapse

What is the effectiveness of colpocleisis compared with sacrospinous fixation for pelvic organ prolapse in elderly women?

To find out why the committee made this research recommendation see evidence review I: surgical management of pelvic organ prolapse.

3 Assessing complications associated with mesh surgery

What is the effectiveness of ultrasound-guided visualisation compared with clinical assessment to identify complications after mesh surgery for stress urinary incontinence or pelvic organ prolapse in women?

To find out why the committee made this research recommendation see evidence review K: assessing mesh complications after pelvic floor mesh surgery.
4 Pessaries or surgery for pelvic organ prolapse
What is the long-term patient satisfaction with pessaries compared with surgery for pelvic organ prolapse in women?

To find out why the committee made this research recommendation see evidence review I: surgical management of pelvic organ prolapse.

5 Long-term risks of surgery with and without mesh
What are the long-term risks of mesh surgery compared with non-mesh surgery for pelvic organ prolapse in women?

To find out why the committee made this research recommendation see evidence review I: surgical management of pelvic organ prolapse.

Other recommendations for research

Long-term effectiveness of botulinum toxin type A for overactive bladder
What is the long-term effectiveness of bladder wall injection with botulinum toxin type A for overactive bladder in women?

Lifestyle interventions for pelvic organ prolapse
Which lifestyle interventions improve the symptoms and prevent the progression of pelvic organ prolapse in women?

Surgery for stress urinary incontinence and pelvic organ prolapse
What is the most effective surgical management for women with both stress urinary incontinence and pelvic organ prolapse, including the sequence of interventions?

Pain management after mesh surgery
What is the effectiveness of pain management for women who present with chronic pain 3 months after mesh surgery for stress urinary incontinence or pelvic organ prolapse?

Context
Urinary incontinence (UI) is a common symptom that can affect women of all ages, with a wide range of severity and nature. Although it is rarely life-threatening, UI can
be very detrimental to the physical, psychological and social wellbeing of the women it affects. The impact on families and carers can also be profound, and the resource implications for the health service are considerable. UI is defined by the International Continence Society as ‘the complaint of any involuntary leakage of urine’. UI can be a result of functional abnormalities in the lower urinary tract or of illnesses. Stress UI is involuntary urine leakage on effort or exertion or on sneezing or coughing. Urgency UI is involuntary urine leakage accompanied or immediately preceded by urgency (a sudden compelling desire to urinate that is difficult to delay). Mixed UI is involuntary urine leakage associated with both urgency and exertion, effort, sneezing or coughing. Overactive bladder (OAB) is defined as urgency that occurs with or without urgency UI and usually with frequency and nocturia. OAB that occurs with incontinence is known as 'OAB wet'. OAB that occurs without incontinence is known as 'OAB dry'. These combinations of symptoms are suggestive of the urodynamic finding of detrusor overactivity, but can be the result of other forms of urethrovesical dysfunction.

Pelvic organ prolapse is defined as symptomatic descent of one or more of: the anterior vaginal wall, the posterior vaginal wall, the cervix or uterus, or the apex of the vagina (vault or cuff). Symptoms include a vaginal bulge or sensation of something coming down, urinary, bowel and sexual symptoms, and pelvic and back pain. These symptoms affect women’s quality of life.

The prevalence of pelvic organ prolapse is high; in primary care in the UK, 8.4% of women reported vaginal bulge or lump and on examination prolapse is present in up to 50% of women. One in 10 women will need at least one surgical procedure, and the rate of re-operation is as high as 19%. There is likely to be an increasing need for surgery for urinary incontinence and pelvic organ prolapse because of the ageing population.

The NHS England Mesh Working Group report published in December 2015 raised a number of concerns about the safety and efficacy of surgery for stress urinary incontinence and pelvic organ prolapse using mesh devices. The report made the following recommendations for NICE:
• to produce a clinical guideline that describes, holistically, care for women with pelvic organ prolapse

• to review the current clinical guideline on urinary incontinence in women (CG171)

• to review evidence on complications arising from surgery for stress urinary incontinence and pelvic organ prolapse.

NICE accepted these recommendations and commissioned an update of the existing urinary incontinence guideline to review the evidence on complications arising from surgery for stress urinary incontinence and managing pelvic organ prolapse.

Finding more information and resources

To find out what NICE has said on topics related to this guideline, see our web pages on gynaecological conditions and urological conditions.

Update information

October 2018

This guideline is an update of NICE guideline CG171 (published September 2013) and will replace it. NICE guideline CG171 updated NICE guideline CG40 (published October 2006).

We have reviewed the evidence on assessment and management for women with urinary incontinence, pelvic organ prolapse and complications suspected to be associated with mesh surgery.

Recommendations are marked [2019] if the evidence has been reviewed.

Recommendations that have been deleted or changed

We propose to delete some recommendations from the 2013 guideline. Table 2 sets out these recommendations and includes details of replacement recommendations. If there is no replacement recommendation, an explanation for the proposed deletion is given.

In recommendations shaded in grey and ending [2006, amended 2019] or [2013, amended 2019], we have made changes that could affect the intent without
reviewing the evidence. Yellow shading is used to highlight these changes, and reasons for the changes are given in table 3.

In recommendations shaded in grey and ending [2006], [2006, amended 2013] or [2013], we have not reviewed the evidence. In some cases minor changes have been made – for example, to update links, or bring the language and style up to date – without changing the intent of the recommendation. Minor changes are listed in table 4.

Table 2 Recommendations that have been deleted

<table>
<thead>
<tr>
<th>Recommendation in 2013 guideline</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.1.5 Refer women with UI who have symptomatic prolapse that is visible at or below the vaginal introitus to a specialist. [2006]</td>
<td>Replaced by new sections on assessing and managing pelvic organ prolapse.</td>
</tr>
<tr>
<td>1.7.3 Prescribe the lowest recommended dose when starting a new OAB drug treatment. [new 2013]</td>
<td>This is covered in the NICE guideline on medicines optimisation and in the summaries of product characteristics for the individual medicines.</td>
</tr>
<tr>
<td>1.7.4 If a woman's OAB drug treatment is effective and well-tolerated, do not change the dose or drug. [new 2013]</td>
<td>This is covered in the NICE guideline on medicines optimisation and in the summaries of product characteristics for the individual medicines.</td>
</tr>
<tr>
<td>1.7.7 Offer one of the following choices first to women with OAB or mixed UI: • oxybutynin (immediate release), or • tolterodine (immediate release), or • darifenacin (once daily preparation). [new 2013]</td>
<td>The cost of these medicines has changed since 2013 and the efficacy of all anticholinergics for overactive bladder is now similar.</td>
</tr>
</tbody>
</table>

Table 3 Amended recommendation wording (change to intent) without an evidence review
<table>
<thead>
<tr>
<th>Recommendation in 2013 guideline</th>
<th>Recommendation in 2019 guideline</th>
<th>Reason for change</th>
</tr>
</thead>
<tbody>
<tr>
<td>All recommendations that refer to ‘conservative management’</td>
<td>All recommendations except those labelled [2019]</td>
<td>‘Conservative management’ has been changed to ‘non-surgical management’ for clarity and consistency with the 2019 recommendations.</td>
</tr>
<tr>
<td>1.1.14 This recommendation has been replaced by recommendations 1.6.4 and 1.6.5 in the NICE guideline on suspected cancer: recognition and referral.</td>
<td>1.3.21 For women aged over 45 who have haematuria, or a recurrent or persistent unexplained UTI, follow the recommendations on referral for bladder cancer in the NICE guideline on suspected cancer. [2006, amended 2019]</td>
<td>To clarify the cross-reference to the NICE guideline on suspected cancer.</td>
</tr>
</tbody>
</table>
| 1.5.4 Do not offer percutaneous posterior tibial nerve stimulation for OAB unless:  
  • there has been a multidisciplinary team (MDT) review, and  
  • conservative management including OAB drug treatment has not worked adequately, and  
  • the woman does not want botulinum toxin A or percutaneous sacral nerve stimulation. [new 2013] | 1.4.16 Do not offer percutaneous posterior tibial nerve stimulation (needles inserted close to the posterior tibial nerve) for OAB unless:  
  • there has been a local multidisciplinary team (MDT) review and  
  • non-surgical management including OAB medicine treatment has not worked adequately and  
  • the woman does not want botulinum toxin type A or percutaneous sacral nerve stimulation. [2013, amended 2019] | ‘Local’ has been added to differentiate this MDT from the regional MDT, as described in section 1.1. |
<p>| 1.7.6 Do not offer oxybutynin (immediate release) to frail older women. [new 2013] | 1.4.31 Do not offer oxybutynin (immediate release) to older women who are at higher risk of a sudden deterioration in their physical or mental health. [2013, amended 2019] | The wording has been clarified to specify that it refers to cognitive as well as physical impairment. |</p>
<table>
<thead>
<tr>
<th>1.7.17</th>
<th>1.4.45</th>
<th>Initial referral to the MDT has been removed because it is now covered in recommendation 1.1.1. ‘Local’ has been added to clarify that further advice should be sought from the local MDT.</th>
</tr>
</thead>
</table>
| If the woman wishes to discuss the options for further management (non-therapeutic interventions and invasive therapy) refer to the MDT and arrange urodynamic investigation to determine whether detrusor overactivity is present and responsible for her OAB symptoms:  
  - If detrusor overactivity is present and responsible for the OAB symptoms offer invasive therapy (see recommendations in section 1.9).  
  - If detrusor overactivity is present but the woman does not wish to have invasive therapy, offer advice as described in recommendation 1.6.9.  
  - If detrusor overactivity is not present refer back to the MDT for further discussion concerning future management. [new 2013] | For women with OAB that has not responded to non-surgical management or treatment with medicine and who wish to discuss further treatment options:  
  - offer urodynamic investigation to determine whether detrusor overactivity is causing her OAB symptoms and  
    - if detrusor overactivity is causing her OAB symptoms, offer an invasive procedure in line with recommendations 1.4.46 to 1.4.59 in this guideline  
    - if there is no detrusor overactivity, seek advice on further management from the local MDT. [2013, amended 2019] | 1.4.49 Start treatment with botulinum toxin type A only if the woman is willing, in the event of developing significant voiding dysfunction:  
  - to perform clean intermittent catheterisation on a regular basis for as long as needed or  
  - to accept a temporary indwelling catheter if she is unable to perform clean intermittent catheterisation. [2013, amended 2019] |
| 1.9.3 | 1.4.49 | Amended because the likelihood of needing catheterisation is lower with the dose of 100 units of botulinum toxin type A that is recommended in the 2019 guideline, and to remove restrictions on women being offered this treatment. |
| Start treatment with botulinum toxin A only if women:  
  - have been trained in clean intermittent catheterisation and have performed the technique successfully, and  
  - are able and willing to perform clean intermittent catheterisation on a regular basis for as long as needed. [new 2013] |  

Amended because the likelihood of needing catheterisation is lower with the dose of 100 units of botulinum toxin type A that is recommended in the 2019 guideline, and to remove restrictions on women being offered this treatment. |
1.9.10 Offer percutaneous sacral nerve stimulation to women after MDT review if:
- their OAB has not responded to conservative management including drugs, and
- they are unable to perform clean intermittent catheterisation. [new 2013]

1.4.54 Offer percutaneous sacral nerve stimulation to women after local or regional MDT review if:
- their OAB has not responded to non-surgical management including medicines and
- they are unable to perform clean intermittent catheterisation. [2013, amended 2019]

‘Local or regional’ has been added to clarify that this refers to either the local or the regional MDT.

1.9.11 Consider percutaneous sacral nerve stimulation after MDT review if a woman’s OAB has not responded to conservative management (including drugs) and botulinum toxin A. [new 2013]

1.4.55 Consider percutaneous sacral nerve stimulation after local MDT review if a woman's OAB has not responded to non-surgical management (including medicines) and botulinum toxin type A. [2013, amended 2019]

‘Local’ has been added to differentiate this MDT from the regional MDT, as described in section 1.1.

1.10.1 When offering a surgical procedure discuss with the woman the risks and benefits of the different treatment options for SUI using the information in information to facilitate discussion of risks and benefits of treatments for women with stress urinary incontinence. [new 2013]

1.5.1 When offering a surgical procedure discuss with the woman the risks and benefits of the different treatment options for stress urinary incontinence (UI). Include information about differences in type of anaesthesia, expected length of hospital stay, surgical incisions and expected recovery period. [2013, amended 2019]

The table of information referred to in the 2013 recommendation is to be replaced by shared decision aids on surgery for stress urinary incontinence and pelvic organ prolapse.

Table 4 Minor changes to recommendation wording (no change to intent)

<table>
<thead>
<tr>
<th>Recommendation numbers in current guideline</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>All recommendations except those labelled [2019]</td>
<td>The term ‘drug’ has been replaced with ‘medicine’ (in line with current NICE style for recommendations in guidelines). Yellow highlighting has not been applied to these changes.</td>
</tr>
<tr>
<td>All recommendations except those labelled [2019]</td>
<td>Minor changes to language and style have been made (in line with current NICE style for recommendations in guidelines) where possible. Yellow highlighting has not been applied to these changes.</td>
</tr>
</tbody>
</table>

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