



Your sexual recovery after stoma surgery

Lars, TRE™ Product User

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Hello

Welcome to 'Your sexual recovery after stoma surgery'

This booklet sets out to discuss how your sexual experience and expression can be affected by stoma formation and aims to offer some hints and tips that will hopefully help if you are feeling a little lost, worried or curious with regards to your sexual recovery.

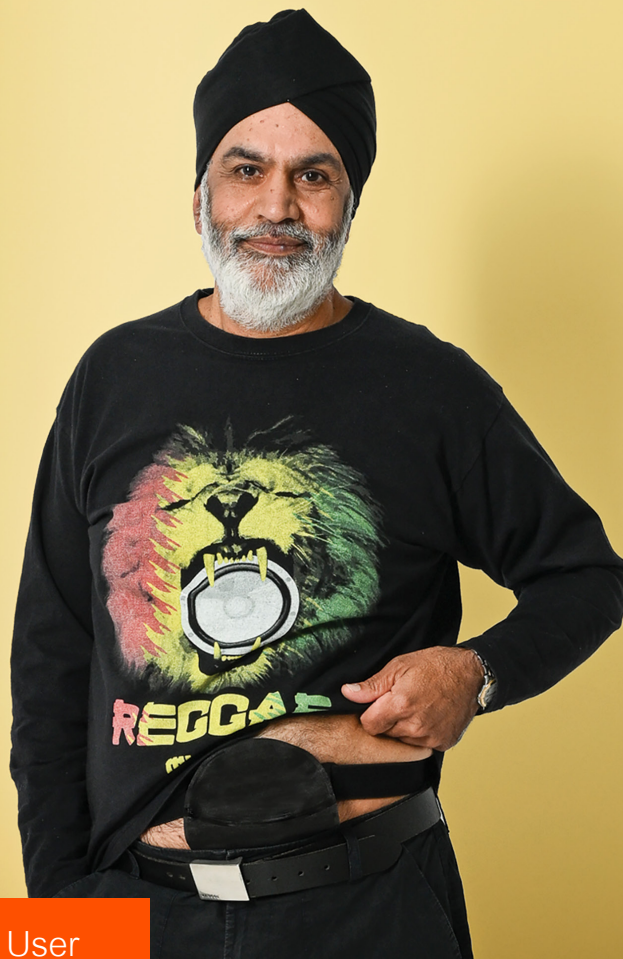
There are many, many aspects to sexuality; we could not possibly address all of them in one handy booklet. Here we focus on the eight topics that most commonly arise when talking to people who have had stoma surgery.

"It is understandable that when you have a stoma, you could feel a little self-conscious about sex"

Helen, TRE™ Product User

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Sapuran, TRE™ Product User

Helping You Discover What Matters Most

A Gentle Guide to Using These Chapters

The chapters use colour coding so you can easily identify those most relevant to you. Because each chapter can be read on its own, some recommendations appear more than once. The key message is to talk with your partner if you have one and to make use of your stoma care nurse, who is often the best source of guidance. If you cannot find the right words, a pull out card at the back of the book can be given to your preferred health care professional to help begin the conversation.

This booklet is not meant to take the place of your professional health care provider. It is provided as an extra educational resource to help guide you with supportive information.



Early Expectations

This chapter sets out to introduce some of the topics people have raised in the early days after surgery, regarding their recovery and what to expect.

Who you are, what is important to you and what you expect from life are very personal things. You are unique – and so is your sex life. How much you express and value your sexuality, celebrate it and enjoy its place in your life is also unique, there is no 'normal' when it comes to sex. Different people have different needs and desires, your 'normal' is what is right for you.

Going through a life-changing illness or injury creates a significant imbalance to your normal world. It affects your body and your mind and has an impact on those you love and who love you. How you respond and how you are supported may be influenced by many factors, including the presence of or absence of a partner, the family you grew up in (and whether they are close to you now), your personality, your religion, community or social network and your earlier life experiences.

Things that positively influence how easily you adapt sexually can include having a relationship that is based on trust and open discussion and having a sex life that values intimacy, love and pleasure above 'performance'. This could be the relationship you have now, or in the future.

There are a number of recommendations that are good for everyone, and which build good foundations for general well-being as well as sexual recovery. These are having a balanced healthy diet, regular exercise and good company. Taking responsibility for your rehabilitation, eating well and building up your fitness can help you to feel good about yourself as you grow in confidence and strength. Sharing this with people who matter to you is important too.

Concerns that are frequently mentioned in the early days and weeks following stoma surgery may not seem directly related to sex and sexuality, but they are a good place to start as they are issues that worry people and worry generally doesn't help you to feel confident or sexy.

Privacy versus sharing

Sharing your thoughts and feelings about what you have been through and what you are still experiencing is usually encouraged as it enables others to support you. Often other people want to help, sometimes in a practical way, sometimes by listening; it makes them feel good to be there for someone they care about.

But it is also true that how much you want to share, who with and when, is entirely up to you. One thing you can be certain of is that anything you discuss with the stoma care nurse will be dealt with sensitively and confidentially. If your stoma care nurse feels that you would benefit from a referral to another health care professional, this will always be discussed with you first.



Expect to have good days and bad days in the weeks following surgery.

Managing early recovery

Sometimes it seems that everyone is in a hurry to see you getting better from the weeks before your operation, through your in-patient stay and then afterwards from the community teams as well as visitors and well-wishers. It can start out feeling very safe, but can become rather stifling, so that privacy is hard to find. It is often a good idea to set a regular rest time in the early weeks and let everyone know when that is, so you can have some time for yourself, your partner – and for a nap.

Expect to have good days and bad days in the weeks following surgery. Both physically and emotionally you are likely to be 'up and down'. If you are culturally different from your stoma care nurse, it may be useful to you both to talk through what having a stoma means to you and those who

share your background or faith. Talking about it will mean that you get truly inclusive care and that you can be directed to resources and information that are helpful. There are information sheets on so many aspects of stoma care and if there isn't one that directly suits your needs, you'll still be amazed at how resourceful your stoma nurse can be!

When is the right time for sex?

This depends what you mean by 'sex'. In small and simple ways in the early days you can concentrate on how you look and feel while building your strength and confidence and reconnecting with 'you'. Go slowly, you've a lot to learn - finding the right appliance, caring for your skin, getting to know your stoma.

Perhaps it's best to start by standing in front of a mirror and looking at and touching your body.

Recognise your thoughts. These may begin with sadness or anger but over time things will improve as you adjust. The next step may be to invite your sexual partner to stand with you as you look and touch together and you can explain if anything hurts, or you have areas of numbness.

Not everyone wants to share this intimacy of course and that's okay, for lovemaking you can choose to wear a top or t-shirt, stoma belt, pouch cover or stay in your underwear.

Expectations of your partner

Partners often request to be shown how to care for the stoma 'just in case'. You need to decide how comfortable you are that your partner is involved - this is your body and your choice. With regard to your partner's expectations of lovemaking or sexual intercourse, again this can create anxiety. There is some adjustment for you both and you both need to feel comfortable and ready to experiment in order to improve the chances of a happy outcome. If you have lost your libido (sex drive), which commonly

happens in the early weeks, talk about it... and agree to review the situation as you recover. Your partner probably wants to hear that this is common and temporary. Try to offer and to accept hugs and affection and if you have always shared a bed, try and stay in the same bed.

It's advisable for you to both read Chapter 7 'Sexual function and resuming an active sex life', once you are ready to move on to more active lovemaking.

If you are single

Those going through stoma surgery without a partner suffer all the same sexual side effects and worries as those people with a partner and sometimes more. Besides there is no-one to experiment with along the way. The worry about whether and when to tell a new potential partner about the presence of the stoma can feel very confusing. Try not to let this worry grow.

If the stoma is a big issue for a potential partner – remember, it's not you, it's them! Maybe they need time to adjust, or maybe they are just not 'the one'.

Be who you are, with friends you trust. Enjoy hobbies and activities that please you.

Big decisions

Sometimes having a life-threatening illness or injury is a springboard for re-evaluating life and making big decisions. Perhaps there is a lost dream that it now feels extremely important to grab hold of again, a 'bucket list' that has to be emptied, a relationship that has floundered for some while and needs to be worked on, or let go. People have been known to do all sorts of things in the aftermath of illness; sell up, buy a camper-van and 'hit the road', leave their partners, emigrate, start a university degree. Many will say that they had to do this very soon after their illness, because they felt that if they waited, they wouldn't do it at all. If you find you are now re-evaluating your life, try not to 'jump' too soon.

Taking time to recover, having people around you that you love and trust and allowing time to let life settle will not mean you lose your motivation for change. It will mean that you can enter into that change in a stronger body and more organised manner – it will let you 'look before you leap'.

Early Expectations

Hints & Tips

- Confidence comes from getting to know your body again and accepting the changes, as well as having faith in your appliances.
- If you are worried about baring your body entirely, wearing a top, tee-shirt or stoma belt, or choosing a pouch cover may help you to feel more comfortable when it comes to lovemaking.
- Accept compliments, go people watching – how many people truly look at each other? No-one except those you tell knows that you have a stoma.
- Do something that you are good at, enjoy your success and share it with someone who matters to you.
- If you are not ready to make love, talk about it, suggest a time for reviewing the situation and start courting again – have fun with cuddling and kissing or even just holding hands.
- Take your time.



"Initially the thought of sex when you have a stoma seems like an odd combination. However, I found that once I had my colostomy and began to irrigate, thereby necessitating the need only to wear a stoma cap, it was all very possible. I did not in any way feel different from how I had done pre-stoma and my husband frankly didn't seem to notice!

It could promote humour whenever the stoma cap filled with air and produced its usual inelegant noises – I certainly found it funny – nothing wrong with a bit of schoolboy humour.

It is understandable that when you have a stoma, you could feel a little self-conscious about sex but I just reminded myself that the stoma is not my persona but just a bit of slightly more unusual anatomy which isn't pinned onto my face. I don't believe that it has changed who I am and therefore hasn't changed the relationship I have with my husband."





Helen, TRE™ Product User





Jen, TRE™ Product User

Body Image

This chapter sets out to discuss some of the difficulties people have raised when trying to come to terms with a changed body.

Body image by most definitions has two elements. Firstly, there is the picture in your mind of how your body looks (its size, shape and appearance for example). Secondly there is what you believe, think and feel about your body. How you view your body has many influences including your culture, the media, fashion, age and wellbeing amongst others. It is common for body image to change through the years, though this 'natural' change generally happens gradually and you adapt without really noticing. Stoma formation imposes big and sudden changes to body image, both physically and emotionally and therefore can have an impact sexually.

Some people feel this very positively, while for others the process of adjustment and acceptance takes time.

Give yourself the time you need.

Try to explain how you feel about your body to your partner or supportive friend.

Maybe you have a positive view – your stoma has saved your life, has given you freedom from incontinence or bowel disease or pain, but maybe your feelings are very negative, affecting every aspect of life.

Give your family and friends the best chance to understand and be there for you. It can be really hard for them too, watching the person they love in distress. You can be assured that you are not the only person who ever felt this way or who needed emotional support. Stoma care nurses want to help you to heal, inside and out. If you would like your partner or a close friend to be present while you talk with your stoma care nurse, that will be fine.

Some hospitals, support groups and associations have a 'buddy' scheme where you can be introduced to someone who has experienced the same surgery as you or who has overcome the same difficulties. Other people like to use websites or read blogs and prefer the anonymity of this option. Again, your stoma nurse can often advise on the best place to look.

Please remember though that some websites do not offer research-based information and blogs are usually posts about someone else's opinion and may not be medically sound. Check with your health care team before taking any such 'advice'.

It may also be useful to read Chapter 3, 'Feelings and emotions'.

Body Image

Hints & Tips

- Try not to avoid going to new places or seeing new people. Avoidance can reinforce the idea that you will be stared at and that you won't be able to manage. Perhaps start with a walk in the park, everyone is busy doing their own thing, they are not even looking in your direction and there are toilets there, just in case you need them. You've a high chance of a successful outing!
- Remember what originally attracted your partner to you – it probably wasn't your tummy!
- Rather than worry about someone being too curious, rehearse some one-line statements that you can use to deflect curiosity. Try looking them in the eye, stand tall, smile and say something like 'I want to forget about that for now – let's talk about you'.
- Consider what is positive about you and your appearance, say it out loud so that you hear it as well as think it.
- Remember, if it is a positive feeling that you have, or a positive feeling that has slowly grown, perhaps your stoma comes to represent well-being and being rid of illness – it is good to share that too!





Lars, TRE™ Product User

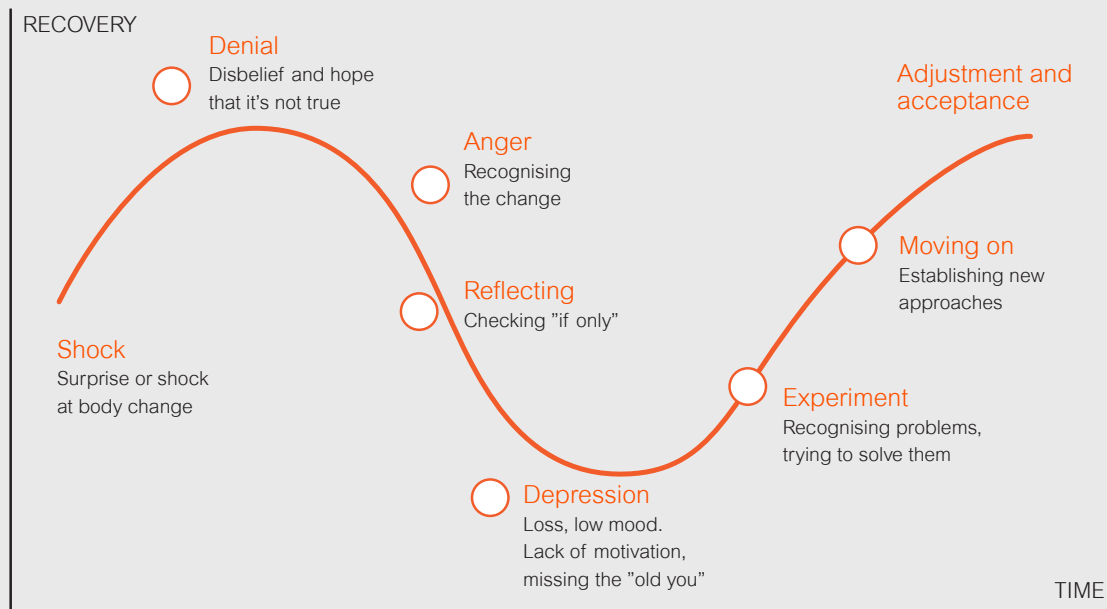
Feelings and Emotions

Bewilderment and loss

For many people who have stoma surgery, a long and healthy life lies ahead. Despite this, a deep sense of grief can be present after stoma surgery. This loss can be multi-faceted and associated with 'loss' not only of a bodily function – the control of bowel movements or bladder function, but potentially loss of the healthy person they used to be; loss of body confidence, loss of social activities, loss of privacy, loss of ability to do what they did before. This is part of a grief process that can be very profound.

There are a number of stages to the process of coming to terms with grief and loss and these can happen in any order, moving back and forth in waves, gradually moving nearer to adjustment and acceptance. Knowing about the stages of grief can help you to understand your feelings and emotions and to recognise waves of grief when you are suddenly having a bad day, after a period of doing well.

Like other losses you may have experienced in life, you can overcome this, one step at a time, with support along the way.



Based on seminal work by Kubler Ross^{1&2}

Many people suffering illness experience anger at some point. Some people are angry with the disease, asking questions like 'Why me?'

Anger

Many people suffering illness experience anger at some point. Some people are angry with the disease, asking questions like 'Why me?' It can feel unfair and undeserved. This is a natural part of the grieving process and of adjustment, it can give you strength to fight, to find a solution, to be determined, or to help you to tell others what is bothering you. Anger however can turn into a very negative emotion when it stops you thinking clearly, simmers away constantly – taking up energy that could be better used, or leading to aggression aimed at those who have no responsibility for triggering the anger. This anger is no longer useful and can damage relationships. Try not to let your partner have all of your anger, just because they are 'there' but take steps to manage this unhelpful anger. Talking through your "anger triggers" with someone who knows you well is a great place to start. Are there lessons to be learned, is there something you need to change? Take constructive action.

Find a way to diffuse this anger – through physical exercise, or deep breathing or relaxation exercises.

Worry about illness returning is a common theme reported by patients. Sadly, this worry may be a very real possibility for some.

Worry, fear and anxiety

Facing an uncertain future is hard. Worry about what is to come, or of the unknown can feel utterly consuming and you may find yourself withdrawing from the very people who love you the most and who can help you the best. As always, talk. Talk to your partner, your friend, your surgeon, your oncologist, your stoma care nurse, or your GP about your fears.

Try not to assume this is a permanent situation. If you can deal with the anxiety, you have a greater chance of a good sexual experience.

Not knowing what to expect can lead to feelings that are very difficult to deal with and can build to become persistent fear and anxiety. As you leave hospital, there are so many new experiences to be faced for the first time, with your stoma as company. Feeling a bit nervous is common but persistent fear and anxiety can have an impact not only on how you feel but on how you think, your ability to get on with day-to-day life, how your body behaves and how you relate to others.

Fear, when it is sudden, for instance in response to a threat such as an angry dog attack, releases a burst of adrenaline into the body, a chemical sometimes referred to as the 'fight or flight' hormone.

It raises the heart rate, gives a surge of energy and helps to either run away from the dog or to fight off the dog. It is designed to keep us safe. When this same hormonal response is released for a long time because of constant anxiety and there is no end to the feeling of 'threat', it can feel truly dreadful.

The heart rate stays high, sometimes feeling like palpitations, the shakes develop, there may be light-headedness, lack of concentration and a feeling of unrest as that energy surge has nowhere to go.

It can also trigger a short temper and snappiness with others close at hand, sparking arguments (fight), blocking intimacy and pleasure seeking, self-confidence and of course, sexual warmth.

The alternative risk is that the adrenaline surge will tempt you to withdraw from what you feel to be threatening situations and those who can help you (flight), to 'hide' and find a sense of safety that way. This situation is not good for your sex life, especially if the perceived 'threat' is sex itself.

High levels of anxiety and adrenaline in men can cause reduced libido (sex drive) and difficulty getting and maintaining an erection; women can also find it harder to have any feelings of sexual desire and vaginal lubrication may be reduced.³

This can lead people to believe that the surgery itself has caused 'sexual dysfunction', when it may be the emotional response to the surgery which is causing the problem. Try not to assume this is a permanent situation. If you can deal with the anxiety, you have a greater chance of a good sexual experience.

If things are not improving over time, or you have worrying thoughts about your ability to carry on, please call your stoma care nurse or GP. Medication can help and there are often counsellors and psychologists attached to your GP surgery or your health care team who can help you to move forward. Reaching out is a positive step. Please remember that.



If you are struggling emotionally, take a look at: **www.nhs.uk/mental-health/self-help/** where lots of great resources are available.



Feelings and Emotions

Hints & Tips

- Share your feelings...talk if you can, or write them down. Writing down your feelings gives you something to look back on over time, so you can see how you have progressed or perhaps use them to support someone else in the future.
- If you are feeling depressed every day, with fewer or no good days, seek help. If you do nothing else, call your stoma care nurse and tell her, or ask a family member to do it for you.
- If you are very anxious about starting lovemaking again, share that with your partner. Together try and work out how you can make it easier, set small manageable goals and aim for achievable success.
- If you have always shared a bed with your partner, try to stay in the same bed.



There are some simple strategies that can help you to manage anxiety when you recognise it building up:

1. Cut back on caffeine in tea, coffee and soft drinks. Caffeine can contribute to and magnify feelings of anxiety.
2. Try relaxation exercises and practice them. Use the techniques before you expect to make love.
3. Adopt a message to yourself that you can say in your head to remind yourself that this feeling will only last a short time.
4. Communicate. Apologise if you have been hurtful and take steps not to let anger or anxiety lead to snappiness or aggression.
5. Take your time with lovemaking. If you know you are tense, explain to your partner and both of you make allowances. Try and keep the focus on pleasure giving and taking, rather than on 'performance'. Hug, caress, reassure.
- 6 Try not to assume that a lost erection, or pain, or a lack of lubrication is a permanent state.



Nils, TRE™ Product User

"I have always been very open as a person and my tactic with the stoma has always been to show everybody and make it be as normal as possible. All my friends and family, and all my colleagues at work, everybody knows about it and everyone has seen it. I am very comfortable with the stoma and I am comfortable with myself, the biggest problem for me was the operation in general. My life after has been a hundred times better than the times before the operation.

I know there is a stigma around talking about having a stoma, but I think if you are open and if you can talk about it, people accept you as you are."





Darren, TRE™ Product User

Relationships

This chapter sets out to explore the impact of relationship stress and some simple ideas that may help.

Relationships are complex, even in good health and everyday life, relationships are complex. Relationships are put under considerable strain when life is not straightforward or a crisis arises, when change is forced upon it, or when someone is ill. Illness can mean reduced household income, one partner absorbing the day-to-day chores and trying to continue with work, as well as being the emotional support for children and the 'patient'. Considerable strain for everyone can be the result. Stress, worry, anger, anxiety and fear are not good for you, your relationships or sex life, as this can potentially affect your ability to relax, and impact your confidence, sexual desire (libido), and sexual function. Dealing with stress in your relationship is important, it affects you both and like ripples in a pond, can touch others around you.

Understanding the causes of your stress is necessary. You and your partner can work through this together, recognising that neither of you can fix a relationship under strain by yourself! Try to write down all of the things that are stressing you both, then go back and put them in order, the most stressful and pressing issue first, down to the little niggles. Next collaborate to make a plan to tackle one big issues and perhaps one or two of the little niggles for starters (a quick 'win' is good for you). Remind each other that this exercise is not about criticism and stay focussed on improving things. Doing this together can really help not only to reduce the stress, but can bring you closer, as can expressing gratitude for the support you each offer- both positive steps for intimacy and sexual recovery.



If you are single

Being single often means that you turn to family members or close friends to help with your long-term recovery, the stresses of life, managing your household and your illness. Don't be afraid to ask and to take up offers of help – especially in the early days. There are no prizes for doing this all by yourself.

When it comes to your sexual recovery, this can be another stress all of its own.

It may also be useful to read Chapter 1, 'Early expectations' talks about 'the time to tell' a potential new partner about your stoma – but there are ways that close friends can help you here too.



Good friends don't exclude someone who is not able to do a particular activity – they change the activity.

But they need to know what your ability is, your energy levels and your fitness – so they can choose wisely and by including you, help you grow your confidence.

Adjusting to your body changes can be helped by showing your tummy to someone you trust (and is curious to see), so that when you show it to a new partner, it won't be the first time you have ever done that. It's ok to take your time when a potential partner comes along - enjoy romance, affection, intimacy and foreplay.

Communicate

There are physical changes for both men and women that may influence your sexual ability, comfort and pleasure. Being able to openly discuss those changes with a new partner may reduce worry and increase understanding.

If you are concerned about your ability to get and keep your erection, or to be comfortably penetrated, then trying things out on your own first can be a good idea!

Physical changes are further discussed in Chapter 7 'Sexual function and resuming an active sex life'.

For everyone, whether you have a partner or not, it is often easier to talk about deeply private matters as you walk side-by-side with someone who you are close to, sharing what is on your mind. It's also a good way to make space for hearing about their life and worries as well – it is healthy communication and support becomes mutual. Hopefully, you'll also get some positive feedback about your progress – and if they don't volunteer it, then ask for it!



Lauren, TRE™ Product User

Orientation

This chapter acknowledges some of the concerns faced by people who do not identify as heterosexual (straight).

As was discussed at the beginning of this book, we are all different from each other.

If you are trans and have had surgery or use hormone therapy, there is a fair chance that the team looking after you will know already, from your medical history and notes.

What your health care team cannot know without you sharing, is your gender identity and sexual orientation. They cannot know the sexual role you play or how your particular surgery is likely to worry or potentially change things for you sexually, without you telling them.

You are the expert on 'you' and health care professionals are often very glad if you can explain to them what it means to be 'you'.

If you can share, it means less assumption, less confusion and often better experiences and outcomes.

Of course, it is up to you what you share and who you share it with. Perhaps you are not out at all or perhaps are not out to everyone.

Perhaps you are just discovering your sexuality - or you may have a long-term partnership, or you may have many lovers, you may identify as straight but have same sex lovers occasionally?

Whatever your unique situation, your health care team just want to work with you as a partner in your own care; listen to your worries, support you and those important to you, and ensure that you have access to relevant information.

Whatever your gender and whatever the gender of your partner, whether you define as gay or straight, this is important - If receiving anal penetration has been an important part of your sex life, we would strongly encourage you to talk to your health care team about your future sexual activity, following ano-rectal surgery.

Above all, your stoma should not be penetrated nor should you ever push anything into your stoma as this can cause significant damage which may lead to further surgery.

Being open about your sexual orientation and worries can help your health care professional provide the best care possible.





Natalie, TRE™ Product User

Roles

This chapter is about sexual roles and how these may change following illness or injury.

Sexual roles are, like everything, influenced by many factors, but in partnership terms tend to settle into a pattern that works for this particular relationship. Perhaps one person is always the one who reaches out to the other and starts a sexual encounter, or maybe you are both able to initiate sex. It may be that one partner is more active and one more passive during lovemaking, possibly one person suggests position changes or is more adventurous or creative than the other. How it all happens varies from couple to couple, and sometimes from one time to the next. Initiation can happen perhaps by talking about it and agreeing, perhaps by making a 'date' in the diary, perhaps by routine 'Saturday night' sex, or maybe by giving clues, kissing and caressing and hoping for a positive response.

What happens next can be influenced by the roles taken in the general relationship, experience, personality and sometimes habit.

If the person who does all the initiating gets ill or injured, they are not necessarily in a position to start things off.

That doesn't always mean that they don't want sex. Perhaps the person who is normally passive is simply unable to fill that now empty role – they have never practiced being the initiator and feel awkward and uncertain. That doesn't necessarily mean that they don't want sex either. Changing roles can be really difficult. Many couples find it impossible to talk about their needs, to ask for what they want and at that point allow assumption and feelings of rejection to grow.

Further to this, other changed roles such as not being able to work – to be 'the breadwinner' or for that matter, 'the homemaker' can erode sense of self, identity and confidence. Equally it can be a direct journey from partner to patient, or lover to carer, or even a shift to a 'parent and child' type relationship – but a hard journey back to a mutually fulfilling sexual partnership.

There are people who can help. Psychosexual therapists are trained to help people to talk, to identify their needs, to learn to initiate sex – or to politely decline – to communicate and to practice. More information about psychosexual therapy can be found in the next chapter.

"I had just met my now husband and we had only been together a few months. After surgery I was quite worried about – well, quite bluntly - whether he would dump me. But he's been fantastic and he just wanted to see me well. When I'm with my husband we have a very functional relationship."





Jim, TRE™ Product User

Sexual function and resuming an active sex life



Sexual function and resuming an active sex life

This chapter discusses in very direct language the potential impact of stoma surgery on sexual function as well as some of the medications and resources available to both men and women.

Resuming an active sex life does not generally mean leaping into action with full penetrative intercourse the first week after your operation. In fact, this is absolutely not recommended – as gentle rehabilitation is usually required for even the most menial of tasks, such as making the bed in the first place! Most surgeons will recommend a period of taking it easy, lasting about 6 weeks.

By this time, you are likely to be moving more easily and more comfortably and at least beginning to resume your usual activities, hobbies and generally ‘finding yourself’ again.

What is clear is that there is no right time to resume intercourse!

This is down to you, your relationship(s), how you feel, what sex meant to you before your surgery, your desired sex life now, all of those things we have mentioned in the other chapters. It is a very individual thing. It is always suggested that talking with your sexual partner is very likely to help when it comes to 'doing it'.

Being worried about your partner and how they will react, about how your body will feel and respond, and about how your stoma/product will behave creates too much tension.

Remember to talk and reduce that tension. Talking about sex with your partner may not be something that you have ever done. If that is the case, then having to renegotiate your sex life or invent a new way of doing things can be really tricky. There is help out there if you simply cannot talk about it together.

Your stoma care nurse may be able to advise you or may suggest a sex therapy referral.

Psychosexual therapists are trained to help people negotiate difficulties in their sexual lives and relationships, including those caused by illness or injury. For some, this is just about talking it through once – with the therapist as a facilitator, while for others a programme of therapy is desirable.

Sexual confidence

Getting back to the things that you are good at or that make you feel good is a great start when energy levels allow. Dressing in your usual style, rather than to 'hide' your middle can go a long way to re-establishing your identity. Taking compliments (and even writing them down) is good too.

Perhaps you have been chronically ill for a long time and having your stoma represents the end of a long road and the start of a new healthy you. Maybe you are only at the beginning of your journey and are facing other treatments, feeling depleted and constantly tired – do what you can to stay in touch with who you are, and who you will be again.



Sexual confidence is also helped by stoma confidence. There are many different stoma products on the market and some will suit you more than others. There are two-piece, one piece, small and super discreet, clear, opaque and black pouches, covers for pouches, support belts etc. Find out, get samples and settle on the one that you are happiest with. If you are confident with the practical matters to do with your stoma, such as pouch security, capacity and bowel activity (or urine production for those with a urostomy) and take some simple steps, such as emptying or changing your pouch (30 mins before-hand to get a good seal), things are likely to be more relaxed.



If you want to know more about sex therapy you can go **to www.cosrt.org.uk** where you can also use their 'find a therapist' facility.



It's usually a good idea to negotiate and agree the first time you are going to make love.

You can make some practical preparations as well as set some goals and ground rules. It helps you both to be clear about what you expect to happen, what you are willing to try and what you are not going to try this time - and that can help reduce anxiety and boost confidence. Fatigue and loss of libido are common features in the early weeks and months.

Remember that sex doesn't have to be energetic, or penetrative, nor does it have to start with deep desire.³ Negotiate tender, gentle, connected lovemaking.

Generally, it is better to start slowly, holding each other, touching, kissing and caressing. Discover together if there are any altered sensations, numb skin patches, or if it hurts anywhere. You can 'map' each other's bodies... where and how do you like to be touched? What do you like less? Talk as you touch – it can work both ways and can be great fun! Grow your confidence and share the experience, as well as how you each feel about it. It is always better to set an easily achievable goal and be successful, than to have high expectations and suffer disappointment.

Build up slowly, keep negotiating what you want to achieve each time, think about positions and what is most likely to be physically comfortable and practical. When (or if) it comes to penetrative intercourse, think ahead.

Stress alone can cause some men to struggle to get or keep their erection and women may need additional lubrication. Please remember that neither are necessarily permanent or due to changes caused by surgery.

Try not to make assumptions - this is almost certainly not because your partner doesn't want to have sex with you.

You can read more about stress in Chapter 4 'Relationships'.

For women

Pain during penetrative intercourse is known as dyspareunia. This is the most commonly reported side effect for women following stoma surgery. It is thought to be caused by anatomical changes following surgery, reduced sexual desire and arousal, and sometimes made worse by stress and reduced lubrication⁴. These problems often improve over time but meanwhile you can take some steps to help matters. Talk to your partner about what you feel and keep them informed about your comfort levels, try to manage your stress and buy some lubricant and be ready to use it. There are many different lubricants on the market – water based, silicone, or oil based⁵, in different colours, flavours and sensations. They are available in the supermarket, pharmacy and online. Some are available on prescription. (NOTE - Oil based lube should not be used with latex condoms as it can reduce reliability.)⁶ Silicone based lubes can cause deterioration of silicone dilators and sex toys.⁵

For women who could experience orgasm before surgery, be aware that some drugs, particularly those for nerve pain or depression can stop or delay orgasm – loss of female orgasm is not a common side effect of the stoma surgery itself.

Some women choose to take a painkiller before intercourse, while others choose not to have penetrative sex until things are much more comfortable. If this is the case explain and find alternative pleasures with your partner.

Many women are offered vaginal dilators or vaginal 'trainers' following low bowel surgery or radiotherapy. They often come with video or written instructions for use. The idea is to stop your vagina from becoming tight and scarred by treatment. Practicing with dilators can also be a good way to know how to relax and position yourself. If you are offered dilators, you do not have to accept them.

For men

Treatment options for erectile dysfunction

Some men have problems with erection prior to their bowel disease, while others have problems caused by their surgery or other treatments. Either way, it may become desirable to have some treatment for erectile dysfunction. The treatment options are many and varied and may include psychosexual therapy and / or tablets, injections, vacuum pumps, creams and implants, depending on how motivated you are to restore your erection and which of the options is acceptable to you.

The first step is to talk to your stoma care nurse, surgeon or GP. Tablet treatment (sildenafil and tadalafil for example) is not medically advisable for some men, but if it is safe to try, often GP's will start tablet treatment as a first step. There are erectile dysfunction clinics (sometimes known as andrology clinics) in most hospitals and this would be the likely next step if tablet treatment is not possible or doesn't have the desired outcome. Remember however that sex can be great without erection or penetration – there are lots of other pleasures to be enjoyed!

For everyone

If receiving anal penetration has been an important part of your sex life, we would strongly encourage you to talk to your health care team about your future sexual activity, following ano-rectal surgery. Above all, your stoma should not be penetrated nor should you ever push anything into your stoma as this can cause significant damage which may lead to further surgery.



Sexual function and resuming an active sex life

Hints & Tips

- Fatigue and loss of libido are common after surgery. Take the time you need to recover or make allowances in your sex life.
- Try body mapping. Take it in turns to allow your partner to touch your body from top to toe, front and back. Talk to each other about what feels nice – soft touch, kissing, blowing on the skin for instance, as well as what areas are less pleasant, sore or numb. Include the head and face in the exercise. Say what you like about your partner's body too. Revisit this exercise over time, to see if anything has changed.
- You are usually encouraged to empty or change your bag before lovemaking. If you need to change your base plate, or if you are using a one piece bag, allow at least 30 minutes for a good seal to be achieved before commencing lovemaking. Minimise worry. Whilst it is not acceptable to everyone, trying things out on your own (masturbation) is often a good idea. When it comes to lovemaking, talk about what you want to do and what you don't want to do.

- Talk to your partner about how things may be different and make a plan. Do you need to take a more passive role than usual? Do you need to consider alternative sexual positions? What else might be helpful to achieve a happy outcome?
- Remember that 'lovemaking' doesn't have to mean 'intercourse'. Many people have very satisfying sex lives without penetration.

This is important;

- If receiving anal penetration has been an important part of your sexual repertoire, we would strongly encourage you to talk to your health care team about your future sexual activity following bowel surgery.
- Above all, your stoma should not be penetrated nor should you ever push anything into your stoma as this can cause significant damage which may lead to further surgery.





Darren, TRE™ Product User

Reproduction & reproductive decisions

Reproduction and decisions about fertility preservation are discussed in this chapter.

Whatever your gender, however you identify, it is really important to have a discussion with your health care team about the risks of your surgery on your sexual function, fertility and ability to have children before your operation, if you hope to make babies in the future.

The presence of a stoma should not, on its own, affect your fertility but it remains strongly recommended that you have that discussion, so it is clear in your mind.

If your surgery has already happened and you are not sure about your fertility status, it remains important to check, if it is a concern to you.

Talk to your surgeon at your next consultation or ask your stoma care nurse for advice.

Patients having stoma formation for cancer may have had or now require additional treatment such as radiotherapy or chemotherapy as part of their whole treatment.

At each stage in your care, you will be told about the side effects of each treatment and complete a consent form. You should at this time be told about any additional impact on your ability to have children, as well as about contraception whilst on treatment. Again, if you are unclear, go back and ask.

It may be that you've simply forgotten what you were told, it is a very chaotic time and there is only so much your brain can take in – health care professionals understand that and will be glad to go over things with you.

There will sadly be people who want more children but whose fertility is likely to be at risk. For some people illness can be managed and surgery postponed until later, when babies are born and families are complete. For others, particularly when dealing with cancer, treatment has to start very quickly. The urgency to treat can rule out some fertility preservation options for women, as completing certain procedures can take

a number of weeks, delaying cancer treatment too long. There are still some possibilities to be explored. Following discussion and agreement with your cancer care team, this is most likely to begin with an urgent referral to a specialist team at your local fertility clinic. These are sometimes known as 'assisted conception units'.

Fertility issues are not only experienced by women. Men undergoing stoma surgery, particularly those having surgery on the lower bowel, or having bladder and prostate removal with formation of urostomy will also be counselled regarding the possible impact on sexual function (ability to get and maintain an erection and to ejaculate semen) and fertility.

Sperm storage or 'banking' can be an option and again this would usually require referral to the local fertility clinic or assisted conception unit.

Learning that you will no longer be able to have the family that you dreamed of or that there will be long term sexual consequences from your stoma surgery can be devastating. Your stoma care nurses want to help and to support you. Please talk to them so that they can help.



More information regarding the options for both men and women can often be found on the website of your local fertility centre, or alternatively from the Human Fertilisation and Embryology Authority – along with statistics regarding successful pregnancy outcomes;
www.hfea.gov.uk





Natalie, TRE™ Product User

What next?



What next?

Despite everything that has been said, sexual worries are not inevitable.

Many people going through stoma formation recover sexually, in their own time, without additional support, or referral or therapy. They find their own way. Some reinvent their sex lives, finding greater closeness, better communication, shared pleasures and more satisfaction. Others choose to let go of their sex lives and feel liberated and are glad.

If you do want to recover and are struggling to make progress in any aspect of your sexual rehabilitation, please speak up. Talk to the health care professional you feel most comfortable with, and will help you best.



Useful Links

Hints & Tips

More patient education from your stoma care company.

www.dansac.co.uk

The Lesbian & Gay Foundation:

<http://lgbt.foundation>

College of Sex and Relationship Therapists

www.cosrt.org.uk

Human Fertility and Embryology Authority

www.hfea.gov.uk

The Sexual Advice Association:

Factsheets about sexual function and treatment options.

[http://sexualadviceassociation.co.uk/factsheets/
booklets/booklets-factsheets/](http://sexualadviceassociation.co.uk/factsheets/booklets/booklets-factsheets/)

NHS Mental Health

[www.nhs.uk/mentalhealth/self-help/guides-tools-and-
activities/mental-wellbeing-audio-guides](http://www.nhs.uk/mentalhealth/self-help/guides-tools-and-activities/mental-wellbeing-audio-guides)

Your sexual recovery after stoma surgery

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