

Commissioning Policy Excluded and Restricted Procedures Version 2.1

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Name of originator/authors:	Head of Planned Care, Cancer & Diagnostics/
	IFR Improvement Manager
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Target audience:	NHS and partnering agencies, all health providers that provide care to Staffordshire and Stoke-on-Trent patients, Primary and Community Care including GPs either via a contract or NCA arrangements, and care professionals; general public within Staffordshire and Stoke-on-Trent Integrated Care Board



CONSULTATION SCHEDULE

Name and Title of Individual	Groups consulted	Date Consulted
ICB Portfolio & Clinical Leads	Internal	2024/2025

GOVERNANCE SCHEDULE

Name of Committee	Role	Date
Clinical Prioritising Advisory Group (CPAG)	Support	November 2024
Quality and Safety Committee	Support	February 2025
Portfolio Boards:	Support	December 2024 and January 2025
Finance and Performance Committee	Endorsement	N/A
Health & Care Senate	Approval	13.02.2025
Health & Care Senate	Approval	04.07.2025

VERSION CONTROL

Version	Version/Description of amendments	Date	Author/amended by
1.0	ERP approved at ICB Board	11/10/2023	Jackie Newman
2.0	Reviewed and updated as per review schedule.	02/10/2024	Vanessa Weaver & Jackie Newman
2.1	Addition of weight management drugs, bariatric surgery for patients over 18 years, simple ear wax, aquatic therapy for children & young people, aquatic therapy for inflammatory arthritis, sequential oral sensory (SOS) therapy. Additional narrative for orthotic provision	April-July 2025	Mel Mahon, Vanessa Weaver, Catherine Lewin, Graeme OMalley & Jackie Newman

IMPACT ASSESSMENTS – available on request				
	Date	Stage	Comments	



Privacy Impact Assessment	N/A	N/A	N/A
Quality Impact Assessment Impact assessments V2	06/02/2025	Gateway 1	QIA24-042 - This QIA sets out clear criteria for health professionals and patients to ensure equity of application.
EHIA – Amendment	29/01/2025	Stage 1	Amendment to the ERP alignment) No changes to original EHIA in 2022 – Satisfied that due regard has been demonstrated)

1.0 Purpose

1.1 The purpose of this policy is to ensure that Staffordshire and Stoke-on-Trent (SSOT) Integrated Care Board, which will be referred to as 'the ICB' throughout this policy only fund treatments that are clinically effective and only offer interventions on the NHS that are evidence-based and appropriate.

The policy details the treatments deemed to have sufficient clinical evidence to justify funding from the ICB's allocated fixed budget. The aim is to improve the quality of care being provided to patients by reducing unnecessary interventions and preventing avoidable harm. This will enable the ICB to reallocate this finite resource and ensures any money saved is spent on other more effective treatments.

This policy provides a collated list of all procedures and services that the ICB have restricted NHS funding for. Patients should only be referred for the interventions listed if they meet the eligibility criteria. The onus is on the treating clinician, and the Provider Organisation, to seek the necessary authorisation from the ICB prior to commencing treatment or procedure. It is the clinician's responsibility to provide relevant, accurate evidence of how the patient meets the criteria.

If a provider undertakes either an excluded or a restricted procedure within this policy without gaining prior approval the ICB will not pay for the procedure, this is in conjunction with the Provider signed contract.

If a provider undertakes one of the listed procedures which is listed as 'excluded' then the ICB will withhold payment. The only route which funding may be considered is on grounds of clinical exceptionality. Please refer to IFR link https://staffsstoke.icb.nhs.uk/contact-us/individual-funding-requests/

2.0 Introduction

2.1 Across the country most, if not all ICBs have a set of policies and procedures for limiting the number of low clinical value treatments and interventions. This Policy supports the decision-making process associated with the allocation of resources for commissioning. The ICB has a prioritisation process which is overseen by the Clinical Advisory Prioritisation Group (CPAG) https://staffsstoke.icb.nhs.uk/your-nhs-integrated-care-board/our-



<u>publications/governance-handbook/all-policies/commissioning/icb-policy-on-the-prioritisation-of-healthcare-resourcesmaster/?layout=default</u>

- 2.2 CPAG use an approved standardised scoring system, which has been supported by the ICB Board, to evaluate clinical evidence that is available for a particular therapeutic treatment, intervention or a diagnostic test.
- 2.3 In addition, the Evidence Based Intervention (EBI) programme, which was launched in 2018 is a clinical initiative led by the Academy of Medical Colleges in partnership with NHS England, NHS Clinical Commissioners, National Institute for Health and Care Excellence (NICE), Getting it Right First Time (GiRFT), providers and the public. The aim of the EBI programme is to improve the quality of care and is designed to reduce the number of medical or surgical interventions as well as some tests and treatments where the evidence shows that it is inappropriate for some patients in some circumstances.

Another focus of the CPAG portfolio, in conjunction with the EBI programme, is to free up valuable resources so they can be put to better use elsewhere in the NHS.

- 2.4 The Policy establishes the framework within which the ICB can demonstrate that its decision-making processes are fair, equitable, ethical, and legally sound. The risk of not having an ERP Policy means that each decision is left solely to the clinician to manage each patients' individual problems without guidance and context of the health needs of the wider population.
- 2.5 This Policy is designed to help the ICB to meet its statutory obligation in providing equitable access to health care. It aims to achieve this by supporting a robust decision-making process that is reasonable and open to scrutiny.
- 2.6 Where patients fall outside of this policy, then the only route which funding may be considered is on grounds of clinical exceptionality. Please refer to IFR link https://staffsstoke.icb.nhs.uk/contact-us/individual-funding-requests/

3.0 Background (Rationale for Policy)

- 3.1 The ICB is required to commission comprehensive, effective, accessible and equitable services which are free to users at the point of entry (except where there are defined charges) within a finite resource. It is, therefore, necessary to make decisions regarding the investment of resources in interventions which achieve the greatest health gain for the population.
- 3.2 This policy supports clinicians in making decisions on when to refer, where the primary reason for the referral is surgical intervention. The policy is also a statement about what the ICB will routinely pay for.



- 3.3 Each year the NHS is subjected to greater pressures. Ensuring that treatment and care is focused where it can make the biggest difference is a vital part of making the best use of finite resource. The ICB receives funding to commission health services for their resident population and make decisions within the context of statutes, statutory instruments, regulations, and guidance. This is a significant challenge for all NHS organisations and is a key focus for the ICB. The ICB has a responsibility to seek the greatest health advantage possible for local populations using the finite resources allocated to them. This policy will help clinicians identify interventions with limited clinical benefit which enables potential reinvestment, where benefits are greater for the population of Staffordshire and Stoke-on-Trent.
- 3.4 This policy covers interventions where there is significant risk that patients undergoing them will gain little health benefit. The procedures have low rather than no clinical value. Some may be effective but may have low value because other treatments could be tried first. Other effective procedures may provide large benefits for some cohorts but less to those with few symptoms, where risks and benefits are closely balanced. There are some interventions which are effective in some but give no clinical value in others. There are those interventions that whilst effective are undertaken for primarily cosmetic reasons which commissioners often consider as providing low clinical value.

4.0 Definition of "Low Priority Treatments"

- 4.1 The term "treatment" describes clinical care and programmes of care that include:
 - Medicines both prescribed and non-prescribed.
 - Surgical procedures
 - Therapeutic and other healthcare interventions
- 4.2 On systematic evaluation, some interventions have been identified as being either marginally effective or ineffective with limited clinical value in the vast majority of cases. Others have been shown to be an inefficient use of resource given their high cost per quality adjusted life year gained.

5.0 Operating Policy for the Development and Implementation of this Policy

5.1 Scope

5.1.1 A number of national organisations, such as National Institute for Heath & Care (NICE) in England, Scottish Intercollegiate Guidelines Network (SIGN) in Scotland, and the Medical Colleges are committed to producing evidence-based commissioning policies. The emphasis is on high value care pathways. In addition, Public Health clinicians from across Staffordshire and the West Midlands have developed evidence-based advice to inform both the prioritisation process and commissioning decisions on low priority treatments. Throughout this Policy these treatments or procedures are categorised as Excluded or Restricted. Excluded treatments or procedures will not be funded by the ICB. Restricted treatments or procedures will only be funded for those patients where an



appropriate threshold for the intervention as stated in this Policy has been met.

The term 'female' and 'male' for the purpose of this policy relates to the biological sex of the patient. The ICB is including those born with internal and external genitalia, chromosomes and/or hormones that relate to that sex and not their gender.

5.1.2 This policy should be used in conjunction with patient symptomatic clinical referral pathways.

5.2 Determining the Evidence Base

5.2.1 Evidence for treatment effectiveness and efficacy is available from many sources, including NICE, the Cochrane Institute, Royal Colleges, other professional guidelines, and sources such as peer reviewed journals or technical notes. Evidence varies in its robustness, ranging from meta-analyses of randomised control trials with large populations of participants, to traditional consensus about best practice. The ICB in arriving at this Policy have taken advice from Public Health locally on the source, extent, and quality of the evidence in reaching their decisions and assessed the relative clinical value of many specific procedures through the local prioritisation process, see section 2.

5.3 Populations

5.3.1 Unless stated otherwise the restrictions within this policy apply to patients of all ages. If the restriction is a paediatric case, then this is defined as a child between the ages of 0 and 18.

Unless stated otherwise the restrictions within this policy apply to all patients registered with a GP Practice in Staffordshire and Stoke-on-Trent Integrated Care System.

5.4 Ethical and Legal Policy for Decision Making

5.4.1 The ICB has Prioritisation Frameworks which are reviewed on an ongoing basis. Utilisation of these prioritisation frameworks informs the review of this policy and the procedures and treatments that it covers.

5.5 Implementation and Compliance

5.5.1 This policy will be reviewed and approved by the Health & Care Senate and will be made publicly available via https://staffsstoke.icb.nhs.uk/your-nhs-integrated-care-board/our-publications/governance-handbook/all-policies/commissioning/icb-excluded-and-restricted-procedures-policy-v3-1/?layout=default



Queries relating to the content of this policy or has difficulty understanding this policy should in the first instance contact the IFR Team IFRTeam@staffsstoke.icb.nhs.uk

- 5.5.2 The schedule showing low priority treatments is set within this policy. This will be formally incorporated into the contracts and service level agreements and will be subject to routine monitoring for compliance. The ICB will require primary, tertiary and secondary care service providers and other organisations acting on behalf of the ICB to embrace and abide by the policy and to advise patients accordingly.
- 5.5.3 This Policy must be implemented by GPs and all Primary Care Health Professionals when advising and referring patients and by providers when considering the treatment options for patients. Those making referrals should refer to any provider for treatment or procedure covered in this policy. In the first instance where there is a Tier 3 service provider pathway in place the Primary Care referrer MUST refer to the Tier 3 service prior to offering Patient Choice. Patient Choice will be offered if the patient requires onward referral from the Tier 3 service or the referral relates to a suspicion of cancer or is an emergency. Providers should not suggest, recommend, or otherwise offer excluded treatments or procedures covered in this Policy to any patient. Providers should only suggest, recommend, or otherwise offer restricted treatments or procedures covered by this Policy to patients who satisfy the appropriate threshold statement for that treatment or procedure.
- 5.5.4 **ALL** restricted procedures contained in this policy are subject to the ICB's requirement for prior approval. The preferred method of prior approval within the ICB is via an automated web-based software system however where providers do not use this facility then formal requests must be emailed to the ICBs secure email address IFRTeam@staffsstoke.icb.nhs.uk and will be processed by the IFR Team and a unique approval code will be issued. Providers will **NOT** be paid if a unique approval code is not issued. Retrospective approval is prohibited.
- 5.5.5 If treatment has been privately financed, it is assumed that the patient has sufficient funds to cover the whole package of care; it would not be reasonable to expect the NHS to pick up the costs associated with private treatment unless there is a medical emergency or the patient has exceptional circumstances, please see https://staffsstoke.icb.nhs.uk/contact-us/individual-funding-requests/
- 5.5.6 The ICB acknowledges that if an individual patient has commenced private treatment that is routinely commissioned by the ICB the patient has a right to transfer back to NHS care with the caveat that the patient is reassessed by an NHS clinician in line with current policy to ensure compliance with the current policy that is in situ. Where the patient meets criteria, the patient will be subject to standard NHS clinical pathways. Any devices that are privately funded the NHS will not pick up the costs of consumables or maintenance unless they meet



NHS criteria.

The continuation of loan device trials will not be routinely funded. Where treatments have been initiated by providers this is done so at the providers' own risk. The provider must manage the patients' expectations and have a clear exit strategy for the trial and/or continuing care.

5.5.7 Within this policy, and as stated against individual treatments or procedures listed appendix 1 treatments and procedures are classified as Excluded or Restricted by the ICB.

The Individual Funding Request (IFR) process is to be used in circumstances when an individual patient does not meet the clinical criteria for a procedure as listed in this policy but can demonstrate exceptionality in accordance with the IFR Policy https://staffsstoke.icb.nhs.uk/contact-us/individual-funding-requests/

- 5.5.8 Restricted procedures and treatments are not commissioned by the ICB except where an individual patient satisfies the threshold statement or criteria against a procedure or treatment that have been recommended by the Clinical Prioritisation Advisory Group (CPAG) and approved by the ICB Board. Clinicians considering offering a patient a restricted procedure or treatment should satisfy themselves that the threshold statement or criteria against the procedure or treatment are met. Where a patient satisfies the threshold statement or criteria the procedure or treatment is prior approved and can be undertaken. Where the threshold statement or criteria are not met then the procedure or treatment is excluded for that patient and paragraph 5.5.7 above applies.
 - 5.5.9 This Policy is distributed to all provider organisations, primary care contractors, Primary Care Networks and will be published on the ICB website.

5.6 Monitoring the Policy

- 5.6.1 The ICB will monitor the adherence to this policy through the contractual process, using contractual levers where breaches of the Policy are identified.
- 5.6.2 Referrals to secondary care that are outside of this Policy will be routinely monitored by the Portfolio Management and the Contracts Management Teams of the ICB.
- 5.6.3 The ICB will provide periodic reports to their Boards reporting the number and nature of breaches of the Policy, by provider and by procedure.
- 5.6.4 Where there are defined thresholds, the compliance with the criteria will be subject to regular clinical audits carried out or organised by the ICB. The audit process will require providers to produce patient specific evidence that confirms the threshold criteria for procedures were satisfied at the time the decision to offer the procedure to the patient was taken. Where audit shows that the



evidence is not available or is deficient or fails to satisfy the auditor that the threshold criteria were met at the time the decision to perform the procedure was taken, then the default will be to consider the procedure was excluded and therefore it will not attract payment from the ICB.

- 5.6.5 The ICB reserves the right to reduce the value of all payments for procedures with OPCS codes that match those for Excluded and Restricted procedures (as listed within this policy)
- 5.6.6 Any procedures marked as 'Requires Prior Approval' must be approved by the ICB before the procedure is undertaken using the agreed form/mechanisms. The ICB will not pay for any procedures undertaken without the required approval. It is the responsibility of the requester to provide the ICB with the necessary clinical evidence to demonstrate that the patient meets the criteria. Where a clinical opinion is required, a decision will be made within 30 working days of receipt by the ICB (this may be delayed if insufficient information is provided, and further clarification is required). Please refer to the clinical opinion standard operating procedure <a href="staffsstoke.icb.nhs.uk/your-nhs-integrated-care-board/our-publications/governance-handbook/all-policies/commissioning/sop-clinical-opinions-ifr-final20240604/?layout=default
- 5.6.7 Prior approval is not required if a patient is being referred to secondary care for consultant management other than a procedure listed in this policy (for example diagnostic tests, investigations or treatment options).
 - 5.6.8 If a procedure is not listed in this policy but the Office of Population Censuses & Surveys (OPCS) codes have appeared in the Secondary Uses Services (SUS) data within the last two financial years of the request, this would indicate that both provider and the ICB would classify the activity as commissioned and therefore is chargeable by prior approval. If a procedure has not appeared in the SUS data and is also not covered in this policy this would indicate that both the provider and the ICB acknowledge that this procedure is unlikely to have been actively commissioned and on that basis the provider must seek clarification and appropriate approval before carrying out the procedure. Failure to do so will result in the procedure not been funded.
- 5.6.9 This policy is only applicable to elective/planned activity; non-elective activity is excluded from all the terms and conditions of this policy with no restrictions or exclusions applying. The policy also does not apply to patients who are on a 2ww/cancer pathway or where there is suspicion of malignancy, and this activity will be excluded from any challenge or audit process.
- 5.6.10 Prior approval for NHS funding is not the same as a guarantee of treatment. The final decision around appropriateness and clinical safety will be a clinical one which will be discussed with the patient.



6.0 Audit

- The preferred method of prior approval within the ICB is via an automated webbased software system to be used to pre-authorise procedures where restrictions apply. Due to the time scales obtaining prior approval may not always be feasible; therefore ad hoc audits will be requested by the ICB to ensure that the restricted diagnostics meet the threshold criteria. However where providers do not use this facility then formal requests must be emailed to the ICBs secure email address IFRTeam@staffsstoke.icb.nhs.uk and will be processed by the IFR Team and a unique approval code will be issued. Providers will NOT be paid if a unique approval code is not issued. Retrospective approval is prohibited.
- 6.2 All monitoring will exclude any procedures where Commissioners have identified that the OPCS code is either not applicable or not relevant or a not directly linked to an OPCS code or where the OPCS code is not specified as the Trust is unable to 'track' this activity.

7.0 Maintaining an up-to-date Policy.

- 7.1 The ICB will abide by this policy when making decisions relating to the provision of low priority treatments. Specifically, the role of the ICB is to:
 - Monitor the implementation of the Policy and the impact it has on clinical decision making.
 - Inform referrers of the Policy.
 - Inform all service providers with whom the ICB have formal contractual arrangements of the Policy.
 - Review the policy and the accompanying schedule on an ongoing basis and/or where an urgent consideration of new evidence is justified.

8.0 Managing Expectations

- 8.1 When Clinicians are assessing their patient's, providers should, if necessary, make it clear that the decision by the ICB to consider treatments or procedures to be of low priority under this policy is considered a decision made against their responsibility to seek the greatest health advantage possible for local populations using the resources allocated to them and that it is necessary for the ICB to make decisions regarding the investment of resources in interventions which achieve the greatest health gain for the local population.
- 8.2 Where individual patient circumstances require the escalation of their care providers should refer to the Policy and Procedure for the Authorisation and Management of Individual Funding Request.



Appendix 1

Behavioural/Psychological Therapy

Not routinely commissioned with the exception of children where assessment away from family may be vital to determine the root cause, fully exclude physical illness and allow protection and space for disclosure.

Procedures	Thresholds	Status
Psychotherapy for Borderline Personality Disorder and Dissociative disorders	Not routinely commissioned	Excluded
Behaviour Therapy for Gilles de la Tourette syndrome and tic disorders	Not routinely commissioned	Excluded
Inpatient cognitive behavioural therapy for Chronic fatigue syndrome	Not routinely commissioned	Excluded
Specialist Cognitive Behavioural Therapy (CBT) for Management of Aggressive Behaviour in People with Learning Disabilities	Not routinely commissioned.	Excluded
Specialist Cognitive Behavioural Therapy (CBT) for stuttering	Not routinely commissioned	Excluded
Dyadic Development Psychotherapy (DDP)	Not routinely commissioned	Excluded



Cardiology

Procedures	Thresholds	Status
Closure of Patent Foramen Ovale (PFO) for the prevention of stroke	Not routinely commissioned	Excluded
Closure of Patent Foramen Ovale (PFO) for migraine headache	Not routinely commissioned	Excluded
Angioplasty for Percutaneous Coronary Intervention (PCI) in Stable Angina	PCI is not routinely commissioned for patients with stable angina unless at least ONE indication in criteria 1 is met or at least ONE indication in criteria 2 and lifestyle interventions have been adhered to: Weight management, smoking cessation, cardioprotective Regular physical activity. Criteria 1 • There is ongoing anginal symptoms despite optimal anti-anginal medication* OR • There is ongoing angina symptoms with intolerance of anti-anginal medications* Criteria 2 If agreed at an appropriately constituted myocardial revascularisation cardiac multidisciplinary meeting (MDM), PCI may also be performed in patients with stable angina in the following cases: • In patients with impaired left ventricular systolic function OR • In patients with left main stem disease OR • In patients with significant ischemic burden OR • Where PCI is otherwise considered appropriate by the MDM. AND All patients being considered for elective revascularisation should have documented evidence that a formal shared decision-making process has taken place with informed patient choice. * Optimal risk medical management should be offered and needs to be stated on the Blueteq.	Restricted



Complementary Medicine/Therapies

Procedures	Thresholds	Status
Acupuncture	Acupuncture will only be commissioned as an adjunct to pain management and only through specialist pain clinics	
Complementary Therapies/ Medicines	Including but not restricted to:	Excluded



Procedures	Thresholds	Status
Aquatic Therapy for Children & Young People	Not routinely commissioned	Excluded
Aquatic Therapy for Inflammatory Arthritis	Not routinely commissioned	Excluded
Animal Therapy	Not routinely commissioned	Excluded
Sequential Oral Sensory (SOS) Therapy	Not routinely commissioned	Excluded

Dermatology and Plastic Surgery

Unless stated as an exemption, all cases for patients aged 18 years and over will be referred from primary care to the Tier 3 Community Dermatology provider for clinical triage. Following clinical triage patients will be assessed and treated within the local tier 3 community or if there are any patients who have clinical 'red flags' then they should be referred directly to the acute service. The community service will be responsible for ensuring appropriate conservative and non-surgical treatment options are tried and have failed before referral for surgical opinion.

The assumption is that all cases referred for surgical opinion by the community service will be eligible for surgery in keeping with the criteria set out for various conditions in this policy. It is the responsibility of the receiving Provider to seek approval for all restricted procedures that they undertake in-line with their contracts with the ICB. Direct referrals into an acute setting that are not in line with above criteria should be returned to the referrer.

When a direct referral has been made directly to the Trust there is an expectation that all appropriate conservative and non-surgical treatment options have been tried and have failed prior to referral. Referrals should include appropriate imaging.

Procedures	Thresholds	Status
Excision of sweat gland bearing	Not routinely commissioned	Excluded
skin for hyperhidrosis		
Treatment of Minor Skin Lesions	Will be routinely commissioned under the following circumstances:	Restricted
including but not limited to:		
benign pigmented moles,	Significant functional impairment which is seen as the inability to perform normal working	



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Procedures	Thresholds	Status
comedones, corns/callous, lipoma, milia, molluscum contagiosum, seborrhoeic keratosis, skin tags including anal tags, spider naevus, warts, xanthelasma and neurofibromata, epidermoid/Pilar (sebaceous) cysts	movements of the body with normal excursion, strength, speed, coordination and/or endurance, and which the clinical and patient agree is interfering with or compromising several activities of daily living, restriction of joint movement, lesion causing regular pain requiring long-term medication, the lesion bleeds (more than twice weekly for at least four weeks) in the course of normal everyday activities, the lesion causes pressure symptoms which are unavoidable and cannot be managed conservatively and cause atrophy. Verruca on the feet do not normally meet this criterion as they can be pared back to avoid pressure symptoms.	
	2 or more infections requiring prescribed antibiotic and/or pain with prescribed analgesia use over the past 12 months.	
	Prior approval must be sought via the agreed prior approval route as agreed detailed in the contract.	
	Where the lesion prevents the individual from being able to wear mandatory personal protective equipment (PPE) https://www.hse.gov.uk/ppe/ppe-regulations-2022.htmsafety following an occupational health assessment. The OcH assessment will need to evidence that all reasonable adjustments have been explored and are not suitable. The funding request together with the OcH report will need to be forwarded to the https://www.hse.gov.uk/ppe/ppe-regulations-2022.htmsafety following an occupational health assessment. The OcH assessment will need to evidence that all reasonable adjustments have been explored and are not suitable. The funding request together with the OcH report will need to be forwarded to the https://www.hse.gov.uk/ppe/ppe-regulations-2022.htmsafety following an occupational health assessment. The OcH assessment will need to evidence that all reasonable adjustments have been explored and are not suitable. The funding request together with the OcH report will need to be forwarded to the https://www.hse.gov.uk/ppe/ppe-regulations-2022.htmsafety for manual prior approval following a clinical decision.	
Excision of lesion of the eyelid (including Chalazia)	Incision and curettage (or triamcinolone injection for suitable candidates) of lesions of eyelid including chalazia should only be undertaken if at least ONE of the following criteria have been met:	Restricted
	 Has been present for more than 6 months and has been managed conservatively with warm compress, lid cleaning and massage for 4 weeks. Interferes significantly with vision. Interferes with the protection of the eye by the eyelid due to altered lid closure or lid anatomy. Is a source of infection that has required medical attention twice or more within a 6-month time frame 	



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Procedures	Thresholds	Status
	 Is a source of infection causing an abscess which requires drainage If malignancy (cancer) is suspected E.g. Madarosis/recurrent/other suspicious features in which case the lesion should be removed and sent for histology as for all suspicious lesions. Cosmetic eyelid surgery to correct puffy, hooded, wrinkled tired looking eyes will not be commissioned 	
Laser or surgical treatment for congenital vascular abnormalities (inc. Port Wine Stain, Paediatric Haemangiomas)	Will be routinely commissioned under the following circumstances: - Paediatric haemangiomas can have surgical treatment offered for those which: • Threaten life or function, including compromising eyesight, respiratory, cardiac, or hepatic functions OR • Other internal lesions sited in an area liable to scar OR • Large facial hemangiomas that have failed to regress by school age OR • Show a tendency to bleed or to become infected OR • Kasabach-Merritt syndrome (coagulopathy)	Restricted
Abdominoplasty/Apronectomy/ Panniculectomy	Not routinely commissioned	Excluded
Excision of excessive skin from thigh, leg, hip, buttock, arm, forearm. Buttock/Thigh/Arm lift or body contouring	Not Routinely Commissioned	Excluded



Procedures	Thresholds	Status
Cosmetic operations on the external ear including Pinnaplasty, split earlobes, excision of lesion of external ear	Not routinely commissioned	Excluded
Cosmetic Operations on Breast (female) Breast augmentation	Will be routinely funded for patients who meet the following criteria: Reconstructive surgery following mastectomy for either suspected or proven malignancy OR Following prophylactic bilateral mastectomy for cancer prevention in high-risk cases.	Restricted
Cosmetic Operations on Breast (female) Mastopexy	Not routinely commissioned	Excluded



Procedures	Thresholds	Status
Cosmetic Operations on Breast (female) Revision of breast augmentation +/- Breast Implant Removal	Breast implant revision surgery will ONLY be supported if one of the following applies: Implants with capsule formation that interferes with mammography Implants complicated by recurrent infection Implants with Baker Class IV contracture associated with pain (to be confirmed by a specialist opinion) Intra or extra capsular rupture of silicone gel filled implants Patient develops Breast Associated Anaplastic Large Lymphoma (BIA-ALCL) Breast implants will ONLY be replaced when the patient meets the acceptance criteria of the current breast augmentation policy. In all other patients faulty or problematic implants will be removed and NOT replaced. Only implant removal should be performed, and no other subsequent cosmetic procedure e.g. mastopexy. *The removal of breast implants due to symptoms termed as Breast Implant Illness (BII) or Autoimmune Syndrome Induced by Adjuvants (ASIA) on social media, or due to the risk of developing Breast Implant Associated Anaplastic Large Cell Lymphoma (BIA-ALCL) is not currently recommended. This criterion does not cover the following: Gender reassignment surgery Implants inserted following surgery for breast cancer or breast cancer prevention performed under the NHS.	Restricted
Cosmetic operations on breast (female) Breast Reduction	Patients are ONLY eligible for surgery to reduce breast size if the following criteria are met: • The patient is suffering from functional problems: neck ache, backache and/or intertrigo, where any possible causes of these conditions have been considered and excluded AND • Symptoms are not relieved by physiotherapy and a professionally fitted brassiere has not relieved symptoms AND • The patient has a body mass index (BMI) within the range 18kg/m2 and 25kg/m2 AND	Restricted



Procedures	Thresholds	Status
	Have a cup size of F+	
	AND	
	Be 21 years of age or over	
	Patients should have an initial assessment prior to an appointment with a consultant plastic surgeon to ensure that these criteria are met. At, or following, this assessment there should be access to a trained bra fitter where it is available. AND	
	There is an expected need to remove at least 500g of tissue from each breast	
Surgery for inverted nipples	Not routinely commissioned	Excluded
Removal of Supernumerary Nipples (Polymastia)	Not routinely commissioned	Excluded
Refashioning of scars/keloids including stretch marks.	Not routinely commissioned	Excluded
Gynaecomastia	Not routinely commissioned however if malignancy (either breast or testicular) is suspected, then normal cancer pathways should be followed. Chronic liver disease, thyroid disease, and renal disease should also be excluded.	Excluded
Silicone Gel Sheeting for Preventing or Treating Hypertrophic Scarring and Keloids.	Not routinely commissioned	Excluded
Cosmetic excision of skin of head or neck – e.g. face lift, brow lifts, rhinoplasty, and blepharoplasty to treat the natural process of aging.	Not routinely commissioned	Excluded
Liposuction of subcutaneous Tissue.	Not routinely commissioned	Excluded



Procedures	Thresholds	Status
Blepharoplasty	Blepharoplasty will be routinely commissioned. only for upper lids in the presence of: • visual field impairment (reducing visual field to 120° laterally and 40° vertically) • Severe congenital ptosis This is available on the NHS for correction of ectropion or entropion or for the removal of lesions of the eyelid skin or lid margin. Note: Excessive skin in the lower lid may cause "eyebags" but does not affect function of the eyelid or vision and therefore does not need correction. Blepharoplasty type procedures may form part of the treatment of pathological conditions of the lid or overlying skin and not for cosmetic reasons. The following procedures will not be funded: • Surgery for cosmetic reasons • Surgery for cyst of moll • Surgery for cyst of zeis • Removal of eyelid papillomas or skin tags • Surgery for pingueculum • Excision of other lid lumps	Restricted
Facial Atrophy – new fill Procedures.	Not routinely commissioned	Excluded



Procedures	Thresholds	Status
Cosmetic Surgery to Genitals	Female genital procedures for cosmetic purposes are not routinely commissioned. Labial surgery should ONLY be offered to patients who fulfil ONE of the following criteria: • Labiaplasty is required secondary to other medical conditions such as cancer OR • Where repair of the labia is required after trauma.	Restricted
Permanent hair removal or reduction techniques for conditions including but not limited to: excess body hair, facial hirsutism, hypertrichosis and as adjunct to surgery for pilonidal sinus	Not routinely commissioned with the exception of electrolysis in the following circumstances: - To manage misdirected lashes causing ocular irritation and corneal injury which would prevent need for further intervention/surgery. OR Where patients have undergone skin transplants using skin from a hair bearing area to a non- hair bearing area.	Excluded
Correction of hair loss including male female pattern baldness and hair transplantation	Surgical and medical treatments are not routinely commissioned. See wig provision. https://www.nhs.uk/using-the-nhs/help-with-health-costs/wigs-and-fabric-supports-on-the-nhs/	Excluded
Laser Treatment for birthmarks and scarring	Laser Treatment for birthmarks and scarring will only be routinely commissioned for large (in excess of 5cm x 5cm) and will be routinely commissioned under the following circumstances: The area to be treated is on the face. AND The patient has been through all other recognised treatments, or it has been considered that the treatment would not be effective due to the size or condition of the area affected.	Restricted
Laser or surgical treatment for rosacea	Not routinely commissioned. Severe cases of rhinophyma may be considered when there is evidence of severe nasal airway obstruction.	Prior Approval



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Procedures	Thresholds	Status
Skin Resurfacing techniques: Dermabrasion, Chemical Peels, Laser treatment and referrals for prescriptions for topical treatments etc.	Not routinely commissioned	Excluded
Botox for excessive sweating	Not routinely commissioned	Excluded
Botox for facial aging or excessive wrinkles.	Not routinely commissioned	Excluded
Tattoo removal.	Not routinely commissioned	Excluded
Gender dysphoria	Not routinely commissioned by ICB. Gender Reassignment is now the responsibility of NHS England and commissioned through specialised commissioning. GPs can refer directly to their contracted services. http://www.nhs.uk/livewell/transhealth/pages/local-gender-identity-clinics.aspx	Excluded
Endoscopic thoracic sympathectomy (ECT) for extreme facial blushing.	Not routinely commissioned	Excluded

Diagnostics

Procedures	Thresholds	Status
Open and Upright MRI and MRI under General Anaesthetic (GA) in adults	Urgent open/upright MRI or MRI under GA requests in cases with red flag symptoms or signs should be made urgently by the referring clinician directly to the commissioned provider and are excluded from this policy.	Prior Approval
	Patients with the following are eligible for funding for open/ upright MRI: • Patients who are unable to tolerate conventional MRI due to claustrophobia despite conservative management of anxiety (including noise-cancelling headphones, visual aids,	
	and scanning feet first) AND the use of sedation (where sedation is not contraindicated).	



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Investigation is clinically required and will directly inform diagnosis or treatment
 OR

• Patients where there is no alternative option (for example, due to obesity, severe contractures, inability to lie flat)

Patients with the following are eligible for funding for MRI under GA:

Sedated and Ventilated on Critical Care-

OR

AND

Confusion and pre-existing agitation

OR

· Adult with Learning Disability

OR

• Unable to lie flat and stationary due to medical reasons or physical impairment AND unable to use an open/upright scanner (assessed using clinical judgement)

Prior approval to be sought in line with current process.

Ear, Nose and Throat and Paediatric surgery

Any patient with clinical 'red flags' or paediatrics should be referred directly to the acute service.

Unless stated as an exemption, all cases for patients aged 18 years and over will be referred from primary care to the Tier 3 Community ENT provider for clinical triage. Following clinical triage patients will be assessed and treated within the local tier 3 community or if there are any patients who have clinical 'red flags' then they should be referred directly to the acute service. The community service will be responsible for ensuring appropriate conservative and non-surgical treatment options are tried and have failed before referral for surgical opinion.

The assumption is that all cases referred for surgical opinion by the community service will be eligible for surgery in keeping with the criteria set out for various conditions in this policy. It is the responsibility of the receiving Provider to seek approval for all restricted procedures that they undertake in-line with their contracts with the ICB. Direct referrals into an acute setting that are not in line with above criteria should be returned to the referrer.

When a direct referral has been made directly to the Trust there is an expectation that all appropriate conservative and non-surgical treatment options have been tried and have failed prior to referral. Referrals should include appropriate imaging.



Procedures	Thresholds	Status
Adenoidectomy	Adenoidectomy will only be funded if undertaken in conjunction with Tonsillectomy and/or Grommets (<i>Please refer to policies for Tonsillectomy and/or Grommets</i>).	Prior Approval
	For adenoidectomy only, prior approval will need to be sought	
Tonsillectomy	To be undertaken in line with the SIGN 2010 guidance: - The following are recommended as indications for consideration of tonsillectomy for recurrent acute sore throat in both children and adults: • Seven or more well documented, clinically significant, adequately treated sore throats in the preceding year, • OR • Five or more such episodes in each of the preceding two years OR • Three or more such episodes in each of the preceding three years AND • Sore throats are due to acute tonsillitis. AND • The episodes of sore throat are disabling and prevent normal functioning. NB A child is under the age of 16 for the purpose of tonsillectomy	Restricted
	An eligible episode must have three of the following criteria:	
	Tonsillar exudates /Tender anterior cervical lymph nodes/ History of fever/ Absence of cough When in doubt as to whether a tonsillectomy would be beneficial, a six-month period of watchful waiting is recommended.	
Tonsillectomy for Peritonsillar	Prior approval will need to be sought.	Prior
Abscess (Quinsy)	Supporting clinic letters MUST accompany request.	Approval
Tonsillectomy for tonsilloliths or tonsil stones	Not routinely commissioned	Excluded



Surgical Treatment for Obstructive Sleep Apnoea (OSA) children	To be undertaken in line with agreed OSA pathway. In the absence of an agreed OSA pathway, these procedures will only be commissioning in the following circumstances: When diagnosis of SDB in children is confirmed based on history, physical examination, audio/video taping, pulse oximetry, and limited or full-night PSG. AHI>5 indicative diagnosis OSA	Restricted
Surgical Treatment for Obstructive Sleep Apnoea (OSA): Adults	 Will be routinely commissioned under the following circumstances: Patients have Epworth Sleepiness Score 15-18 or: Patient sleepy in dangerous situations such as driving. AND Patient has significant sleep disordered breathing (as measured during sleep study, usually by the Apnoea/ Hypopnoea Index: 15- 30/hr. = moderate, >30/hr. = severe AND Patient has already tried CPAP unsuccessfully for 6 months prior to being considered for surgery OR patient has major side effects to CPAP such as significant nose bleeds. AND A clinical decision is that the patient will significantly benefit. AND The patient is fully informed as to the limited effectiveness of procedures, the lack of long-term outcomes and likely adverse effects including pain following surgery. 	Restricted
Treatments for snoring Including, but not restricted to: Uvulopalatopharyngoplasty, Uvulopalatoplasty, Palate Implants and Radiofrequency Ablation of the Soft Palate	Not routinely commissioned	Excluded
Myringotomy with/without grommets for Otitis Media: Children	To be undertaken in line with NICE clinical guideline 60 – Surgical treatment of otitis media with effusion	Restricted



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Myringotomy with/without	Children with persistent bilateral OME documented over a period of 3 months. AND • A hearing level in the better ear of 25-30 dBHL OR • The worse ear averaged at 0.5,1.2 and 4 kHz (or equivalent dBA where dNHL not available) Alternative indications for Grommets. Children should only be considered for grommet insertion if: - • The child has experienced persistent hearing loss for more than a year with deficit estimated to be more than 25 decibels. OR • More than 6 episodes of acute otitis media in previous 12 months OR • The child has developmental impairment (e.g. speech/ language/ cognitive/ behavioural) likely to be due to, or exacerbated by, clinically suspected hearing loss. OR Poor progress at school directly attributable to this condition, the child has proven hearing loss, plus a second disability such as Down's Syndrome or cleft palate. Commissioned for patients where their consultant considers there is development of a	Restricted
grommets for Otitis Media: Adults	retraction pocket and grommet would help prevent cholesteatoma. OR Patient is experiencing persistent hearing loss affecting work or socialization that has persisted for more than a year with deficit estimated to be more than 25 decibels.	Restricted
Surgical treatment of acute otitis media	Not routinely commissioned	Excluded



Surgical Treatment of Meniere's Disease	Not routinely commissioned.	Excluded
Open wound of ear drum - Tympanoplasty	This procedure will ONLY be commissioned in the following circumstances: Chronic discharging ear, with deafness.	Prior Approval
Operations on nose: septoplasty or septorhinoplasty	Septoplasty or septorhinoplasty will be commissioned for the following clinical indications: 1. Continuous nasal airway obstruction that results in mouth breathing reported by the patient and confirmed by the clinician OR 2. Post-traumatic nasal deformity associated with documented sustained interference of the airway OR	Restricted
Surgical Treatment of Chronic Sinusitis	3. As part of the treatment for congenital abnormalities e.g. cleft lip and palate. Referral for specialist secondary care assessment for surgery for chronic rhinosinusitis (CRS) shall only be commissioned where the following criteria and clinical management has been undertaken in Primary Care/interface provider or by a Tier 3 provider prior to patient being referred for surgical opinion: Primary Care or Tier 3 Provider • A clinical diagnosis of CRS has been made (as set out in RCS/ENT-UK Commissioning guidance) in primary care and patient still has moderate / severe symptoms after a 3- month trial of intranasal steroids and nasal saline irrigation. AND • In addition, for patients with bilateral nasal polyps there has been no improvement in symptoms 4 weeks after a trial of 5-10 days of oral steroids (0.5mg/kg to a max of 60 mg). OR • Patient has nasal symptoms with an unclear diagnosis in primary care. OR • Patient has unilateral symptoms or clinical findings, orbital, or neurological features should be referred urgently / via 2-week wait to depend on local pathways.	Restricted



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	No investigations, apart from clinical assessment, should take place in primary care or be a pre-requisite for referral to secondary care (e.g. X-ray, CT scan). There is no role for prolonged courses of antibiotics in primary care. Surgical criteria for chronic rhinosinusitis (CRS) shall only be commissioned where the following criteria are met: • A diagnosis of CRS has been confirmed from clinical history and nasal endoscopy and/or CT scan. AND	
	 Disease-specific symptom patient reported outcome measure confirms moderate to severe symptoms e.g. Sinonasal Outcome Test (SNOT-22) after trial of appropriate medical therapy (including counselling on technique and compliance) as outlined in RCS/ENT-UK commissioning guidance 'Recommended secondary care pathway'. AND Pre-operative CT sinus scan has been performed and confirms presence of CRS. Note: a CT sinus scan does not necessarily need to be repeated if performed sooner in the patient's 	
	 pathway. AND Patient and clinician have undertaken appropriate shared decision-making consultation regarding undergoing surgery including discussion of risks and benefits of surgical intervention. OR Patient with recurrent acute sinusitis, nasal examination is likely to be normal. Ideally, the diagnosis should be confirmed during an acute attack, if possible, by nasal endoscopy and/or CT sinus scan. 	
Simple ear wax removal	For Simple ear wax removal, undertaken in isolation of a clinical pathway, is NOT funded by the NHS / ICB. Patients should follow the NHS self-help guide: Earwax build-up - NHS	Excluded



Endocrinology

Procedures	Thresholds	Status
Continuous Glucose Monitoring and	Please see separate Commissioning Policy for continuous glucose monitoring/integrated insulin	Restricted
Integrated Insulin Pump Therapy	pump therapy.	
Weight Management Injections	Under current NHS guidance, in response to NICE TA1026, patients identified as being in the	Invitation
	highest-risk group and who will benefit the most will be considered and are identified as being	only as per
Access to Tirzepatide (Mounjaro) for	eligible will be invited to participate in the weight-management program.	cohort criteria
Weight Management - cohort 1		
(Aug 2025 – Jul 2026)	For Patients to access treatment they <u>MUST</u> have been initially assessed in the locally	
	commissioned Primary Care Weight-Management Hub <u>AND</u> have: -	
	BMI 40 or higher	
NHSE Eligibility changes in Aug		
2026	AND have	
This policy will be updated in June	FOUR or more of the following weight-related health conditions and receiving treatment	
2026 for cohort 2 inclusion group.	for them and don't have any of the prescribed medication exclusions:	
	> T 0 D' 1	
	> Type 2 Diabetes	
	 High blood pressure Heart disease – Atherosclerotic Cardiovascular Disease (ASCVD) 	
	Obstructive sleep apnoea	
	Abnormal blood fats (dyslipidaemia)	
	7 Abhornal blood fats (dyshpladerina)	
	NOTE: -	
	NO12.	
	Staffordshire and Stoke-on-Trent ICB (SSOT ICB) weight-management program is a Primary	
	Care non-consultant delivered model. Patients will be invited to attend the Weight Management	
	Hub and will be called based on the information recorded within their Primary Care record, i.e.	
	there is a BMI 40 or greater and the patient has four comorbidities.	
	Patients that have been assessed as being eligible must agree to participate in either the	
	locally delivered, nationally funded, weight-management wrap-around care and support	



program or the local Specialist weight-management program. Patients that disengage or don't engage with this will be withdrawn from the treatment.	
Access to this prescribed medication for weight management will ONLY be commissioned for patients that have been through the weight-management Hubs.	

General Surgery

Procedures	Thresholds	Status
Treatment of Minor Skin Lesions including benign pigmented moles, comedones, corns/callous. lipoma, milia, molluscum contagiosum, seborrhoeic keratosis, skin tags including anal tags, spider naevus, warts, xanthelasma and neurofibromata, epidermoid/Pilar (sebaceous) cysts	Will be routinely commissioned under the following circumstances: Significant functional impairment which is seen as the inability to perform normal working movements of the body with normal excursion, strength, speed, coordination and/or endurance, and which the clinical and patient agree is interfering with or compromising several activities of daily living, restriction of joint movement, lesion causing regular pain requiring long-term medication, the lesion bleeds (more than twice weekly for at least four weeks) in the course of normal everyday activities, the lesion causes pressure symptoms which are unavoidable and cannot be managed conservatively and cause atrophy. Verruca on the feet do not normally meet this criterion as they can be pared back to avoid pressure symptoms. OR 2 or more infections requiring prescribed antibiotic and/or pain with prescribed analgesia use over the past 12 months.	Restricted
	Prior approval must be sought via the agreed prior approval route as agreed detailed in the contract. Where the lesion prevents the individual from being able to wear mandatory personal protective equipment (PPE) https://www.hse.gov.uk/ppe/ppe-regulations-2022.htmsafety following an occupational health assessment. The OcH assessment will need to evidence that all reasonable adjustments have been explored and are not suitable. The funding request together with the OcH report will need to be forwarded to the https://www.hse.gov.uk/ppe/ppe-regulations-2022.htmsafety following an occupational health assessment. The OcH assessment will need to evidence that all reasonable adjustments have been explored and are not suitable. The funding request together with the OcH report will need to be forwarded to the https://www.hse.gov.uk/ppe/ppe-regulations-2022.htmsafety following an occupational health assessment. The OcH assessment will need to evidence that all reasonable adjustments have been explored and are not suitable. The funding request together with the OcH report will need to be forwarded to the https://www.hse.gov.uk/ppe/ppe-regulations-2022.htmsafety for manual prior approval following a clinical decision.	



Inguinal Hernia	Surgical repair will be commissioned when.	Restricted
	patients meet one of the following criteria:	
	Incarcerated hernia or not amenable to simple reduction OR	
	Symptomatic inguinal hernia OR	
	Strangulated hernia (emergency surgery)	
	Patients with occult/asymptomatic/minimally symptomatic primary or recurrent inguinal hernias AND who have significant co-morbidity (ASA 3 or 4) AND who do not want to have surgical repair (after appropriate information provided) can be managed conservatively at primary care level.	
	 All children <18 years with inguinal hernia should be referred to a paediatric surgical provider. 	
	Any groin hernia in a woman should be referred urgently to a specialist.	
	• patients who are undergoing or plan to undergo peritoneal dialysis should be referred.	
Umbilical and Para umbilical Hernia	Not routinely commissioned except: • Pain or discomfort sufficient to cause significant functional impairment, significant interference with activities of daily living with prescribed analgesia use OR • If the patient is considered at risk of incarceration or strangulation.	Restricted
	However, patients who are undergoing or plan to undergo peritoneal dialysis should be referred.	
ncisional Hernia	Asymptomatic incisional hernias will not be funded except where peritoneal dialysis is planned.	Restricted



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	Symptomatic incisional hernias will not routinely be commissioned unless the following exceptions are met: -	
	 Incisional hernias with moderate to severe pain with prescribed analgesia will be repaired. OR Risk of strangulation 	
Laparoscopic inguinal or femoral Hernia Repair	 This commissioned ONLY for Bilateral hernia repair (where the patient has bilateral hernias with external swelling on clinical examination) OR Recurrent hernia after previous hernia surgery. 	Restricted
Rectal bleeding	Follow the British Society of Gastroenterology surveillance guidelines for colonoscopy in the management of hereditary colorectal cancer: https://www.bsg.org.uk/wp-content/uploads/2019/12/Guidelines-for-the-management-of-hereditary-colorectal-cancer.fullpdf Follow the British Society of Gastroenterology surveillance guidelines for post-polypectomy and post-colorectal cancer resection: https://www.bsg.org.uk/clinical-resource/bsg-acpgbi-phepost-polypectomy-and-post-colorectal-cancer-resection-surveillanceguidelines/ Colonscopy will not be commissioned as a first line investigation	Restricted
Haemorrhoidectomy	Will be routinely commissioned under the following circumstances: Patient has recurrent grade 3 or grade 4 combined internal/external haemorrhoids with persistent pain or bleeding. OR Irreducible and large external haemorrhoids.	Restricted



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Restricted

Bariatric Surgery for patients	
over 18 years of age.	

For patients with a BMI of 50 or more, surgery should be considered as a first-line treatment intervention, and can be referred for a bariatric assessment, unless a patient has not tried GLP-1 and wishes to try Tirzepatide (Mounjaro) for 12 months prior to surgery.

Patients with a BMI less than 50 should wait to be invited to attend the Primary Care Management Hub. **ALL** patients that have a BMI of less than 50 will need to be assessed by the Primary Care Weight Management Hub prior to being seen for a bariatric surgical assessment.

For a bariatric surgical assessment patients **MUST** meet the following criteria:

- The patient has a BMI of 40kg/m2 or more, or between 35kg/m2 and 40kg/m2 with significant obesity-related complications likely to improve with weight loss and have 4 out of the 5 co-morbidities stated in NHSE Guidance 2025, where GLP-1 have been ineffective in relation to weight loss
- The patient has a BMI of 40kg/m2 or more, or between 35kg/m2 and 40kg/m2 with significant obesity-related complications likely to improve with weight loss and have been assessed by the Primary Care Weight Management Hubs and have been identified as not being suitable to have GLP-1 treatment

ALL patients being considered for bariatric surgery must also meet the following criteria:

- Appropriate non-surgical measures have been tried but the patient has not achieved or maintained adequate, clinically beneficial weight loss over 2 years
 AND
- The patient has been receiving or will receive intensive management in a Tier 3 service or equivalent

AND

The patient is otherwise fit for anaesthesia and surgery

AND

The patient commits to long-term follow-up AND



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	 The patient and clinician have undertaken appropriate share decision-making consultation regarding undergoing surgery including discussion of risks and benefits of surgical intervention and the patient's GP supports the shared care agreement and is willing to continue regular bariatric care when the patient is 2 years post-op and has been discharged. After surgery, the host bariatric surgery unit should follow up with the patient for 2 years. Thereafter the responsibility for follow-up should be handed over to either the local non-surgical Tier 3 service OR the patients GP, who should conduct yearly appointments. Primary Care follow-up should be conducted at least yearly, and these appointments should include weight measurement and a request for nutritional blood tests. See British Obesity & Metabolic Surgery Society (BOMSS) Guidance for more details. 	
Endoscopic radiofrequency ablation for gastro oesophageal Reflux Disease (GORD).	Not routinely commissioned	Excluded
Linx Reflux Management system for Gastro Oesophageal Reflux Disease (GORD).	Not routinely commissioned	Excluded
Cholecystectomy	Cholecystectomy for Asymptomatic Gallstones is not routinely commissioned.	Excluded
	Cholecystectomy for Symptomatic Gallstones is commissioned in line with RCS and NICE Guidance.	Restricted
Sympathectomy for Raynaud's disease	Not routinely commissioned	Excluded
Gastroelectrical stimulation for gastroparesis.	Not routinely commissioned	Excluded



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Implantation of a duodenal-Jejunal Bypass Liner (DJBL) for managing Type 2 Diabetes.	Not routinely commissioned	Excluded
Surgical Treatment of Diverification of Recti (DRAM).	Not routinely commissioned	Excluded
Treatment of Salmonella Enteritis (non-severe) - faecal transplant.	Not routinely commissioned	Excluded
Early Endoscopic Retrograde Cholangiopancreatography (ERCP) Management of Pancreatitis without cholangitis	Not routinely commissioned.	Excluded
*Early ERCP in the treatment of acute gallstone pancreatitis with	Will be routinely commissioned under the following circumstances:	Do odviedo d
cholangitis	 Evidence of cholangitis OR Obstructive jaundice with imaging evidence of a stone in the common bile duct *Refers to being performed on the same admission, ideally within 24 hours 	Restricted
Endoscopic Drainage of Pancreatic Pseudocyst.	Not routinely commissioned	Excluded
Surgical Drainage of Pancreatic Pseudocyst.	Not routinely commissioned	Excluded
Treatment of Atherosclerosis of Renal Artery.	Transluminal balloon angioplasty with or without stent is not routinely commissioned	Excluded



Treatment of non-neonatal achalasia via pneumatic dilation or Heller myotomy and fundoplication (Heller	Will be routinely commissioned for patient with dysphagia and a manometric/radiologic diagnosis of achalasia.	Restricted
Myotomy)	Surgery is for those patients deemed fit for surgery. OR Who choose surgery over dilation? Pneumatic dilation for poor operative candidates or those who refuse surgery.	

Gynaecology

Any patient with clinical 'red flags' should be referred directly to the acute service.

Unless stated as an exemption, all cases for patients aged 18 years and over will be referred from primary care to the Tier 3 Community Gynaecology provider for clinical triage. Following clinical triage patients will be assessed and treated within the local tier 3 community or if there are any patients who have clinical 'red flags' then they should be referred directly to the acute service. The community service will be responsible for ensuring appropriate conservative and non-surgical treatment options are tried and have failed before referral for surgical opinion.

The assumption is that all cases referred for surgical opinion by the community service will be eligible for surgery in keeping with the criteria set out for various conditions in this policy. It is the responsibility of the receiving Provider to seek approval for all restricted procedures that they undertake in-line with their contracts with the ICB. Direct referrals into an acute setting that are not in line with above criteria should be returned to the referrer.

When a direct referral has been made directly to the Trust there is an expectation that all appropriate conservative and non-surgical treatment options have been tried and have failed prior to referral. Referrals should include appropriate imaging.

Procedures	Thresholds	Status
Infertility and Assisted	Please see separate Commissioning Policy for Infertility and Assisted Reproduction	Restricted
Reproduction	Assisted Conception - Staffordshire and Stoke-on-Trent, Integrated Care Board	
Intra Uterine Contraceptive Devices (IUCDs) including Mirena coils.	Insertion, removal and checks of IUCDs should not routinely be undertaken within secondary care. It is not commissioned as a stand-alone secondary care service.	Excluded
	Tier 3 Community Gynaecology Providers and/or Primary Care will fit ALL IUCDs unless:-	Restricted



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Patients requiring a fitting within secondary care for clinical reasons where a fitting in Tier 3 Community Gynaecology Provider and/or Primary Care is not possible.	Restricted
Removals of lost or displaced IUCDs will be commissioned within secondary care where circumstances dictate that this cannot be managed within the Tier 3 Community Gynaecology Provider/Primary Care.	Restricted
IUCDs fitted as a secondary procedure/OPCS code will be commissioned within secondary care.	
Insertion and removal of vaginal ring pessaries will only be commissioned within Tier 3 Community Gynaecology Provider. First fitting will be commissioned as part of a first outpatient appointment where clinically necessary and will be commissioned for follow up if complications arise.	Restricted
Insertion and removal of vaginal space occupying pessaries will be commissioned within secondary care but only within an outpatient setting.	Restricted
The original pessary plus subsequent replacements will be routinely commissioned.	
NOTE: vaginal ring pessaries are not commissioned in secondary care as a standalone procedure	
Female sterilisation for the purpose of contraception will be routinely funded in the following circumstances.	Restricted
Documented evidence that the patient is unable to tolerate other forms of contraceptive.	
There is an absolute clinical contraindication to pregnancy	
Patients are aged 18 years and over	
 The patient has given fully informed consent for the permanent sterilisation procedure and have been informed that reversal of sterilisation is NOT available on the NHS and reversal of sterilisation has poor success rates 	
	Community Gynaecology Provider and/or Primary Care is not possible. Removals of lost or displaced IUCDs will be commissioned within secondary care where circumstances dictate that this cannot be managed within the Tier 3 Community Gynaecology Provider/Primary Care. IUCDs fitted as a secondary procedure/OPCS code will be commissioned within secondary care. Insertion and removal of vaginal ring pessaries will only be commissioned within Tier 3 Community Gynaecology Provider. First fitting will be commissioned as part of a first outpatient appointment where clinically necessary and will be commissioned for follow up if complications arise. Insertion and removal of vaginal space occupying pessaries will be commissioned within secondary care but only within an outpatient setting. The original pessary plus subsequent replacements will be routinely commissioned. NOTE: vaginal ring pessaries are not commissioned in secondary care as a standalone procedure Female sterilisation for the purpose of contraception will be routinely funded in the following circumstances. • Documented evidence that the patient is unable to tolerate other forms of contraceptive. OR • There is an absolute clinical contraindication to pregnancy AND • Patients are aged 18 years and over AND • The patient has given fully informed consent for the permanent sterilisation procedure and have been informed that reversal of sterilisation is NOT available on the NHS and reversal



Hysteroscopy	This procedure will be routinely commissioned within an outpatient setting unless clinically indicated.	Restricted (prior approval not required)
	Treatment carried out within an inpatient or day case setting is not routinely commissioned.	
Dilatation and curettage (D&C) in women for menorrhagia.	Not routinely commissioned	Excluded
Hysterectomy for menorrhagia	 Will be routinely commissioned under the following circumstances: There has been an unsuccessful trial with a levonorgestrel intrauterine system (e.g. Mirena®) and it has failed to relieve symptoms unless it is medically inappropriate, or contraindicated. AND At least two of the following treatments have failed, are not appropriate or are contraindicated in line with the National Institute for Health and Clinical Experience (NICE) guidelines: Non-steroidal anti-inflammatory agents Tranexamic acid Other hormone methods (injected progesterones, combined oral contraceptives, Gn-RH analogue) AND Surgical treatments such as endometrial ablation or myomectomy have failed to relieve symptoms, or are not appropriate, or are contra-indicated. 	Restricted



Haematology

Procedures	Thresholds	Status
Home Monitoring of INR	Not routinely commissioned	Excluded

Neurology

Procedures	Thresholds	Status
Trans-magnetic stimulation (TMS) for Migraine.	Not routinely commissioned	Excluded
Percutaneous Electro Neuro Stimulation (PENS) for Neuropathic pain.	Not routinely commissioned	Excluded
Targeted training to improve trunk (postural) control in children with cerebral palsy.	Not routinely commissioned	Excluded
Targeted training to improve trunk (postural) control in severely disabled children (inclusive of all gross motor classifications).	Not routinely commissioned	Excluded
Ketogenic diet for the treatment of epilepsy in children	Not routinely commissioned by ICBs. This is the responsibility of NHS England and commissioned through specialised commissioning. https://www.england.nhs.uk/wp-content/uploads/2018/09/E09-S-b-Paediatric-Neurosciences-Neurology.pro .2013.04.v2.pdf	Excluded
Ketogenic diet for the treatment of epilepsy in adults	Not routinely commissioned	Excluded
Seizure Detection Sensors/Systems for the management of Epilepsy	Not routinely commissioned	Excluded
Sodium Oxybate for treatment of narcolepsy and cataplexy in adults.	Not routinely commissioned	Excluded



Sodium Oxybate for treatment of narcolepsy and cataplexy in children	Not routinely commissioned by ICBs. This is the responsibility of NHS England and commissioned through specialised commissioning. https://www.england.nhs.uk/publication/sodium-oxybate-for-symptom-control-of-narcolepsy-with-cataplexy-children-and-adolescents-aged-7-until-19-years/	Excluded
Sativex for multiple sclerosis (MS)	Not routinely commissioned	Excluded
Cranial Molding Orthosis (Helmet Therapy)	Not routinely commissioned	Excluded

Obstetrics

Procedures	Thresholds	Status
Routine Doppler ultrasound of umbilical and uterine artery in lowrisk pregnancies.	Not routinely commissioned	Excluded
A planned Caesarean Section should NOT be routinely offered to women with:	 With a 'small for gestational age' baby. On the grounds of HIV status to prevent mother-to- child transmission of HIV to: o women on highly active anti-retroviral therapy (HAART) with a viral load of less than 400 copies per ml or; women on any anti-retroviral therapy with a viral load of less than 50 copies per ml. with hepatitis B with hepatitis C with a recurrence of genital herpes simplex virus (HSV) at birth with a body mass index (BMI) of over 50 alone as an indicator. 	Excluded
A planned Caesarean Section should be offered to women with:	 With a singleton breech presentation at term, for whom external cephalic version is contraindicated or has been unsuccessful. In twin pregnancies where the first twin is breach A placenta that partly or completely covers the internal cervical os (minor or major placenta) 	Restricted



praevia) • A previous caesarean section where it is clinically indicated. • With injury/tears to the vagina and/or perineum/rectum • With orthopaedic anomalies impeding the patient's ability of having a vaginal delivery • In patients with HIV who: o are not receiving any anti-retroviral therapy or o is receiving any anti-retroviral therapy and have a viral load of 400 copies per ml or more. • With both hepatitis C virus and HIV • With primary genital herpes simplex virus (HSV) infection occurring in the third trimester of pregnancy	
Pregnant women who may require a planned caesarean section should have consultant involvement in the decision-making process.	

Ophthalmology

Any patient with clinical 'red flags' should be referred directly to the acute service.

Unless stated as an exemption, all cases for patients aged 18 years and over will be referred from community and primary care to the ICBs Choice and Referral Centre for clinical triage, where patient choice will be offered.

The assumption is that all cases referred for ophthalmology opinion by the community service will be eligible for surgery in keeping with the criteria set out for various conditions in this policy. It is the responsibility of the receiving Provider to seek approval for all restricted procedures that they undertake inline with their contracts with the ICB.

When a direct referral has been made directly to the Trust there is an expectation that all appropriate conservative and non-surgical treatment options have been tried and have failed prior to referral. Referrals should include appropriate imaging.

Procedures	Thresholds	Status
Blepharoplasty	Blepharoplasty shall be routinely commissioned only.	Restricted
	for upper lids in the presence of:	
	 visual field impairment (reducing visual field 	
	to 120° laterally and 40° vertically)	
	OR	
	Severe congenital ptosis	



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Excision of lesion of the eyelid (including Chalazia)	This is available on the NHS for correction of ectropion or entropion or for the removal of lesions of the eyelid skin or lid margin. Note: Excessive skin in the lower lid may cause "eyebags" but does not affect function of the eyelid or vision and therefore does not need correction. Blepharoplasty type procedures may form part of the treatment of pathological conditions of the lid or overlying skin and not for cosmetic reasons. The following procedures will not be funded: Surgery for cosmetic reasons Surgery for cyst of moll Surgery for cyst of zeis Removal of eyelid papillomas or skin tags Surgery for pingueculum Excision of other lid lumps Incision and curettage (or triamcinolone injection for suitable candidates) of lesions of eyelid including chalazia should only be undertaken if at least ONE of the following criteria have	Restricted
(including Chalazia)	 Has been present for more than 6 months and has been managed conservatively with warm compress, lid cleaning and massage for 4 weeks. Interferes significantly with vision. Interferes with the protection of the eye by the eyelid due to altered lid closure or lid anatomy. Is a source of infection that has required medical attention twice or more within a 6-month time frame Is a source of infection causing an abscess which requires drainage If malignancy (cancer) is suspected E.g. Madarosis/recurrent/other suspicious features in which case the lesion should be removed and sent for histology as for all suspicious lesions. Cosmetic eyelid surgery to correct puffy, hooded, wrinkled tired looking eyes will not be commissioned. 	



Laser Treatment for myopia (short sightedness).	Not routinely commissioned	Excluded
Screening for diabetic retinopathy by consultant ophthalmologists.	Not routinely commissioned in secondary care.	Excluded
Screening for glaucoma by consultant ophthalmologists.	Not routinely commissioned in secondary care.	Excluded
Cataract Surgery	Commissioned when a patient meets the following criteria for each affected eye: Visual Acuity: • Metres 6/9 or worse • US 20/30 or worse • Decimal 0.66 or worse • LogMar 0.18 or worse • VAS 91 or worse • NAD • Difficulty carrying out everyday tasks such as recognising faces, watching TV, viewing computer/mobile screens, cooking, playing sports/cards etc. • Reduced mobility, unable to drive or experiencing difficulty with steps or uneven ground. • Ability to work, give care or live independently is affected. The Visual Acuity should clearly indicate which eye it refers to and which eye the cataract removal is being requested for. Other indications for cataract surgery include facilitating treatment for one or more of the following: - • Monitoring posterior segment disease e.g. diabetic retinopathy • Correcting anisometropia • Patient with Glaucoma who require cataracts surgery to counteract intraocular pressure Patients with Single Sight (Monocular Vision): The indications for cataract surgery in patients with monocular vision and those with severe reduction in one eye e.g. dense amblyopia are the same as for patients with binocular vision, but the ophthalmologist should explain the possibility of total blindness if severe complications occur.	Restricted



Implantable Intraocular Lens	Not routinely commissioned	Excluded
Systems for Age Related Macular		
Degeneration.		

Respiratory

Procedures	Thresholds	Status
Sinus X ray	Not routinely commissioned	Excluded
Insertion of Endobronchial Nitinol coils to Improve Lung Function in Emphysema	Not routinely commissioned	Excluded
Non-Invasive Ventilation	This intervention will ONLY be routinely commissioned in the following circumstances: • Patient has Type 2 respiratory failure. AND • Shows improvement in blood gases, Oximetry or symptoms and demonstrates compliance with equipment at review after at least 6 weeks	Restricted
Cough assist Therapy	Mechanical insufflation-exsufflation (MI-E) therapy, or Cough Assist, for neuromuscular disorders and cervical spinal cord injury patients is commissioned in the following circumstances: 1. An established diagnosis as paralytic/restrictive disorder including but not exclusively: - spinal cord injuries (SCI) - neuromuscular diseases such as ALS - Guillain-Barre Syndrome - myasthenia gravis - muscular dystrophy - multiple sclerosis - post-polio - kypho-scoliosis - syringomyelia 2. Patient is unable to cough or clear secretions effectively with a - PCF (Peak Cough Flow) less than 160 L/min - VC (vital capacity) below 1.1L in general respiratory muscle weakness, or voluntary	Restricted



Home monitoring to prevent sudden	Not routinely commissioned	Excluded
infant death syndrome (SIDS)		

Specialist Therapies

Procedures	Thresholds	Status
Themed Bimanual Training for Cerebral Palsy.	Not routinely commissioned	Excluded
Specialist Sensory Integration Therapy (SIT) for children with autism spectrum disorder and other diagnosed developmental disorders.	Not routinely commissioned	Excluded

Trauma and Orthopaedics

Unless stated as an exemption, all cases for patients aged 16 years and over will be referred from Primary Care to the ICBs Choice and Referral Centre for clinical triage into the local tier 3 community service or directly into local tier 3 community services for clinical triage where a Choice and Referral Centre is not available.

Any case with clinical 'red flags' or paediatrics should be referred directly into the acute service. The community service will be responsible for ensuring appropriate conservative and non-surgical treatment options are tried and have failed before referral for a surgical opinion.

The assumption is that all cases referred for surgical opinion by the community service will be eligible for surgery in keeping with the criteria set out for various conditions in this policy. Direct referrals into acute setting that are not in line with above criteria should be returned to the referrer.

<u>ONLY</u> East Staffordshire PCN may directly refer to the Trust however there is an expectation that all appropriate conservative and non-surgical treatment options are tried and have failed prior to referral. Referrals should include appropriate imaging.



Procedures	Thresholds	Status
Low Back Pain & Radicular Back	ck Pain Procedures/Treatments	
X-rays and MRI of the lumbar spine for non-specific pain	Do not routinely commission imaging in a non-specialist setting for people with low back pain with or without sciatica in the absence of red flags or suspected serious underlying pathology following medical history and examination.	Monitored via Audit
	As part of a diagnostic pathway MRI for Chronic lumbar back-pain (>6 weeks) with no clinical or serological indicators of infection or neoplasia or other red flags to be used in specialist care only where management will be altered.	
	Imaging in low back pain should be offered if serious underlying pathology is suspected. Serious underlying pathology includes but is not limited to: cancer, infection, trauma, spinal cord injury (full or partial loss of sensation and/or movement of part(s) of the body) or inflammatory disease.	
Spinal injections, not limited to – Could include:	Not routinely funded for the treatment of non-specific low back pain.	Excluded
Intraarticular Facet joint injectionsIntradiscal therapy	Medial branch blocks (spinal injections) can be used diagnostically for patients with isolated lower back pain who have not responded to rehabilitation.	
ProlotherapyPlatelet rich plasma	Medial branch blocks should not be used therapeutically for patients with isolated lower back pain. (See Medial Branch Block criteria).	
 Stem cell therapy Trigger point injections with any agent, including botulinum toxin. 	Radiofrequency denervation should be offered for patients with isolated lower back pain who meet the criteria. (See Radiofrequency denervation criteria).	
 Epidural steroid injections for chronic low back pain or for neurogenic claudication in patients with central spine stenosis 		
Alternative therapy such as acupuncture and Tens	Are not funded within the NHS	Excluded



Medial nerve branch blocks as a
diagnostic prior to the
Radiofrequency denervation
(rhizolysis)

Patients **must** have been triaged or seen in the Musculoskeletal MSK Service or Chronic Pain Management Service and exhausted all appropriate non-surgical options within current episode. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics.

Restricted

- Patients have received a biopsychosocial assessment within a specialist pain service (ideally multidisciplinary)
- Back pain severity on a scale of ≥6/10 which has been assessed using a validated pain score questionnaire such as VAS (https://www.physio-pedia.com/Visual Analogue Scale)
- Patients must be actively involved in shared decision making in respect of their treatment and demonstrated commitment to their long-term treatment plan
- Patients must have a commitment in taking responsibility for managing their condition by demonstrating lifestyle changes which may include weight loss, increased fitness through exercise and physiotherapy; diet control, avoidance of illicit drugs and alcohol, and improvement in sleep patterns, managing mood and mental health; and improved engagement in activities of daily living and purposeful occupation where appropriate
- Back pain has persisted for at least 12 months and all clinically appropriate conservative management options, including medication, physiotherapy, and exercise, have already been tried without success
- Back pain causes significant impact on daily functioning which has been assessed using the MSK HQ tool

Radiofrequency denervation for chronic non-specific low back pain will only be funded in accordance with the criteria below:

 Moderate or severe localised back pain (rated as 6 or more on a visual analogue scale or equivalent).

AND



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Radiofrequency denervation (rhizolysis)	 The main source of pain is thought to come from structures supplied by the medial branch nerve as evidenced by a previous positive response to one or two diagnostic medial branch blocks. AND Patient is being treated in the context of a specialist (ideally multidisciplinary) Chronic Pain Management Service https://www.nice.org.uk/guidance/ng59 (See Page 55) 	Restricted
Spinal fusion	Spinal fusion is only commissioned for back pain in the presence of ONE or more of the following: • Spondylolisthesis or spondylolysis • Spinal deformity • Post discectomy or decompression. Neurological compression with associated neural compression symptoms	Restricted
Disc replacement	Disc replacement will not routinely be funded for patients with non-specific low back pain.	Excluded
Vertebral Augmentation (vertebroplasty or kyphoplasty) for Painful Osteoporotic Vertebral Fractures	Patient with vertebral augmentation (vertebroplasty or kyphoplasty) for painful osteoporotic vertebral fractures when clinically indicated and they meet the following criteria. Will be funded: Patient has severe (7/10 or greater on VAS scale) ongoing pain after a recent, unhealed	Restricted
	vertebral fracture despite optimal pain management.	
	AND	
	The acute vertebral fracture has been proven on imaging and correlates with the site of maximal pain on clinical examination AND	
	Multidisciplinary team discussions have taken place AND	
	The procedure will take place at a facility with access to spinal surgery services AND	
	Processes for audit and clinical governance are in place AND	
	• Vertebroplasty must be performed in conjunction with additional measures to improve bone- health	



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	NOTE: Older patients (>60 years old) with fractures at most 6 weeks old and with severe pain despite optimal pain management benefit most from the procedure.	
Lumbar Radicular Pain		
Diagnostic nerve root block	Diagnostic nerve root blocks are only funded after surgical review when decompressive surgery is being considered for nerve root compression. Repeated diagnostic nerve root blocks are not routinely funded for the same level of injection for the same nerve root	Restricted
Epidural injections and therapeutic spinal nerve blocks	Epidural injections and therapeutic nerve blocks: Not routinely funded for the treatment of low back pain without root compression.	Restricted
	There should be evidence that a patient with low back pain and/or sciatica has been assessed in line with NICE guidance NG59.	
	 Patients should have been triaged or seen by a Musculoskeletal (MSK) Service. Where MSK is not available (East Staffordshire PCN) all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics) and have been referred on with evidence available for: 	
	MRI (or similar diagnostic test) which shows radicular compression with documented clinical correlation (MRI scans should be relevant to the presenting symptoms and be at least within 6-12 months of the current episode)	
	AND Symptoms persisting despite non-operative treatment for at least 6 weeks (conservative management including exercise, medication and education)	
	 Referral for injection for acute back pain (less than 6 weeks after onset) is only funded for patients with root compression symptoms (leg pain) that are severe, and debilitating and patient is immobile (despite a documented trial of a full analgesic ladder and physiotherapy without success). 	
	3. Patients may receive one further injection after a documented improvement for a period of 6 months after the first injection. A maximum of 2 injections will be funded for that episode of care. Further injections will require prior approval from the ICB.	



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	 All requests for repeated therapeutic nerve injections for chronic radicular pain, on the grounds of clinical need, need to be made via the ICB Prior Approval Process. The ICB would only consider repeated spinal injections for patients with chronic radicular pain (duration >12 weeks) after prior approval has been sought in cases where there is documented clinical improvement from previous injections. Lumbar facet joint injections will not be routinely commissioned. 	
Spinal decompression and discectomy (lumbar) for radicular pain/spinal claudication	Spinal decompression (laminectomy / laminotomy / foraminotomy) and discectomy will only be funded for patients with sciatica (radicular pain) or stenotic symptoms (neurogenic claudication) in accordance with the following criteria:	Restricted
	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics and referred on with evidence of:	
	MRI (or similar diagnostic test) showing radicular compression with documented clinical correlation (MRI scans should be relevant to the presenting symptoms and be at least within 6-12 months of the current episode)	
	 Symptoms need to be persisting despite non-operative treatment for at least 12 weeks (conservative management including exercise, medication and education). Patient consents for surgical treatment 	
	Patient is aged 19 years and over (as per Evidence Based Interventions guidance on AOMRC website) This policy restriction does not apply to patients with red flag symptoms.	
Lower Limb Foot & Ankle	The part of the pa	



Surgical Referral for symptomatic	Surgical referral has not been made for cosmetic purposes alone.	Restricted
Hallux Valgus (Bunion) Surgical correction for hallux valgus using minimal access techniques (IPG332) is not commissioned. Evidence on safety is inadequate therefore procedure should only be used within special arrangements for clinical governance, consent and audit or research. Hallux Valgus in patients with diabetes should be treated in line with the diabetes pathway and referred to podiatry for an urgent assessment.	 Referral for surgical consideration of hallux valgus shall only be considered where a patient meets ALL of the following: Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. Patients have persistent symptoms despite at least 6 months of conservative management. Prior conservative management must include ALL of the following: Reasonable modification of footwear – avoidance of high-heeled shoes, narrow fitting shoes. Wear wide fitting shoes which will naturally stretch and breathe. Non-surgical treatments Simple analgesia Significant persistent pain preventing patients from fulfilling vital activities of daily living. 	
	Severe deformity which prevents the patient from wearing suitable footwear.	
Autologous Chondrocyte Implantation in the Ankle	Not routinely commissioned	Excluded
Common Foot and Ankle Procedures	Treatment of and not exclusive: ganglion excision, claw toe, hallux rigidus, hammer toe, in growing toenail, metatarsalgia, Morton's neuroma, plantar fasciitis, metatarsal damage, achilles tendon disorders, tibialis posterior dysfunction, arthritis are not routinely commissioned by the ICB except if the following criteria are met: -	Restricted



	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. • Have persistent symptoms despite at least 6 months of conservative management excluding in growing toenail. • Significant persistent pain preventing patients from fulfilling vital activities of daily living. OR • Have recurrent ulcers and infections. Prior conservative management must include ALL of the following: - • Reasonable modification of footwear – avoidance of high-heeled shoes, narrow fitting shoes. Wear wide fitting shoes which will naturally stretch and breathe. • Simple analgesia • Foot/ankle exercises	
Acquired Flat Foot Stage I Disease – Debridement, this may be supplemented with the use of an arthrodesis screw	Referral for surgical consideration of flat foot shall only be considered where a patient meets ALL of the following: • Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service	Restricted
Stage II Disease - a Flexor Digitorum Longus transfer, Calcaneal Osteotomy and Spring ligament	and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics.	
Reefing. Adjunctive procedures include gastrocnemius recession, midfoot fusion or osteotomy and subtalar arthrodesis.	If the patient is unresponsive to conservative treatment or there is significant persistent pain or loss of function impacting on daily living	
Stage III Disease - Triple arthrodesis of the subtalar, calcaneocuboid and	Surgery will be considered for patients with flatfeet (who have pre-existing or recent onset of symptomatic flat feet) in the following circumstances:	
talonavicular joints.	If the deformity is recent onset or deteriorating this should be made a priority.	



Stage IV Disease - Pantalar fusion or a triple fusion and ankle replacement.	If the patient is unable to go up onto tiptoe unaided and standing only on the affected foot or if the foot is not correctable when assessed on the couch.	
 Hind foot – Treatment for arthritis Image guided/targeted injections can be used as a diagnostic and also therapeutic tool. Hind foot fusions of one, two or three of the hind foot joints is a surgical procedure to relieve severe pain from arthritis or correct painful deformity. Double and triple fusion (involving the talonavicular and calcaneocuboid joints) 	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. AND • Exhausted appropriate non operative interventions as specified in the OA pathway • Appropriate analgesia	Restricted
Symptomatic ankle arthritis: Refer to a Consultant Orthopaedic Foot & Ankle Surgeon for consideration of surgery:	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. AND Be unresponsive to conservative treatment AND Have symptomatic Osteoarthritis	Restricted



Integrated Care Board Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and **Arthroscopy & Debridement** Restricted (ankle) onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. AND either/or Unresponsive to conservative treatment Clinical examination or MRI scan has demonstrated clear evidence that there is an internal joint derangement e.g. removal of ankle/subtalar loose bodies, debridement of osteochondral defects and resection of scar tissue Ankle Replacement - Arthroplasty Commissioned by NHS England **Excluded** is the responsibility of NHS England PRN00115-prescribed-specialisedservices-manual-v6.pdf (england.nhs.uk) **Extracorporeal Shockwave** Not routinely commissioned **Excluded** Therapy (ESWT) for Achilles **Tendinopathy** https://www.nice.org.uk/guidance/ip q571 **Extracorporeal Shockwave** Not routinely commissioned **Excluded Therapy for Refractory Plantar Fasciitis** https://www.nice.org.uk/guidance/ipg 311 **Functional Electrical Stimulation** FES using skin surface electrodes will be commissioned for patients who meet the following Restricted (FES) for Foot Drop criteria (Please see separate policy for FES)



	This is exempt from the requirement for Musculoskeletal (MSK) Service referral as the problem is neurological in origin.	
	<u>ALL</u> patients must have had a successful trial of FES and demonstrate improved gait.	
	The patient has foot drop caused by upper-level nerve damage.	
	The patient has been assessed by a specialist in foot drop of neurological origin and all treatment options have been considered.	
	There is evidence that foot drop has caused trips or falls, or gait issues.	
	The patient can walk a minimum of 10 metres independently (+/- aids)	
	The patient can physically manage a FES (+/- minimal assistance)	
	The patient's cognitive ability is such that they can manage a FES independently.	
	The patient does not have co morbidities which would affect their capacity to benefit from FES.	
	The patient does not have any of the known clinical contraindications to FES.	
	Clear FES treatment goals and expectations of benefit are outlined, this is in relation to the effectiveness, and these are assessed annually, outlined	
Other types of FES (implanted or	Not routinely commissioned	Excluded
wireless) are not commissioned. Lower Limb – Hip Procedures		
Diagnostic Arthroscopy of Hip	The ICB does not commission diagnostic arthroscopy	Excluded
Therapeutic Arthroscopy of the	This is not routinely commissioned, and patients must have: -	Restricted
Hip	Been triaged or seen in the Musculoskeletal (MSK) Service. Where MSK is not available	
	(East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics.	



	Have exhausted all appropriate non-surgical interventions.	
Arthritic hip with severe acetabular bone loss, abnormal anatomy (such that non-standard implants may be necessary), prior fusion and cases secondary to infection should be undertaken in specialised centres.	The ICB will consider referral for hip replacement if all the criteria below have been met, for each group: - 1. Been triaged or seen in the Musculoskeletal (MSK) Service. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. Has followed the STP agreed MSK OA Hip Pathway or has had previous surgery in same joint. Has exhausted all appropriate non-surgical interventions. Patient still has a painful irritable and stiff hip interfering with sleep, activities of daily living, work or leisure which has not been controlled with measures above There is narrowing of the joint space on radiograph Patients with a BMI of 35 or more must be actively supported to engage in lifestyle modifications including with weight management to reduce their BMI. 2. OR Is a young adult (<40) with persistent hip pain which affects activities of daily living, work, or leisure 3. OR Where joint destruction is rapid and where a delay in surgery may cause total loss of mobility and independence	Restricted



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Hip Impingement – these operations should be undertaken by surgeons with a special interest and expertise in young adult hip problems	Patients must have been triaged or seen in a Musculoskeletal (MSK) Service and onward referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. The ICB will fund open or arthroscopic hip surgery for the treatment of femoro-acetabular impingement (FAI) ONLY when patients fulfil all of the following criteria: Diagnosis of definite femoro-acetabular impingement defined by appropriate investigations, X-rays, MRI and CT scans. An Orthopaedic surgeon who specialises in young adult hip surgery has made the diagnosis. This should include discussion of each case with a specialist musculoskeletal radiologist. Severe symptoms typical of FAI with duration of at least six months where diagnosis of FAI has been made as above. Failure to respond to all available conservative treatment options including activity modification, pharmacological intervention and specialist physiotherapy. Compromised function, which requires treatment within a 6-8 months' time frame, or where failure to treat early is likely to significantly compromise surgical options at a future date. Treatment with more established surgical procedures is not clinically viable.	Restricted
Femoral/pelvic osteotomy These operations should be undertaken in specialised centres such as University Hospital of North Midlands (UHNM), Royal Orthopaedic Hospital (ROH) Coventry & Warwickshire Hospital, Royal National Orthopaedic Hospital (RNOH), or in a DGH by a surgeon with expertise.	 May be considered in: patients aged <50 years with persistent hip symptoms with abnormalities of femoral and/or acetabular anatomy Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. 	Restricted



Lower Limb – Knee Procedures		
Bespoke Knee Prosthetic	Not routinely commissioned	Excluded
Diagnostic Arthroscopy of Knee	 The ICB will not commission diagnostic arthroscopy unless: - For assessment of severe knee pain (based on a recognised pain scale score) following arthroplasty AND Where a detailed understanding of the degree of compartment damage within the knee is required, above that demonstrated by imaging, when considering patients for certain surgical interventions (e.g. high tibial osteotomy) 	Restricted
Hyaluronic Acid Injections into the Knees	Not routinely commissioned	Excluded
Implantation of a Shock or Load Absorber for mild to moderate symptomatic Medial Knee Osteoarthritis.	Not routinely commissioned	Excluded
Therapeutic Arthroscopy of OA knee	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. Knee arthroscopy, lavage and/or debridement for patients with non-mechanical symptoms is not commissioned Knee arthroscopy, lavage and/or debridement will not be routinely funded unless the person has knee osteoarthritis with a clear history of mechanical locking (as opposed to morning joint stiffness, 'giving way' or X-ray evidence of loose bodies) Clear history of intermittent mechanical symptoms e.g. locking that have not responded to non-surgical treatment (if the knee does not unlock then refer urgently to the appropriate clinic)	Restricted



Primary Knee replacement	The ICB will consider referral for knee replacement if all the criteria below have been met: -	Restricted
	 Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. Have exhausted all appropriate non-surgical options. 	
	Have moderate or severe knee pain not adequately controlled after commencement of treatment and appropriate non-surgical management following NICE guidance (NICE cg177)	
	 Patients with a BMI of 35 or more will be actively supported to engage in lifestyle modifications including with weight management to reduce their BMI. 	
	All patients must have engaged in a shared decision-making process about alternatives, with a view to fully involve them in decisions and their care.	
	Patients who do not meet all of the criteria above may be considered in the following circumstances: -	
	Functional disability in the presence of end stage cartilage disease	
	Progressive deformity of the knee (varus/valgus) with functional disability.	
Partial Knee replacement – This involves the replacement of only one compartment of the arthritic knee	As well as the criteria for primary knee replacement patients should have: - • Symptomatic Osteoarthritis predominantly confined to a single joint compartment	Restricted
Partial knee replacement is less		



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common, and it is more appropriately commissioned and delivered by specialised units, with experienced surgeons, performing around 10 such procedures within a unit per year (NJR 2017)		
Knee Revision Surgery	Commissioners do not routinely fund Modular Rotating Hinge knees	Excluded
Upper Limb - Hand Procedures		•
Carpal Tunnel Syndrome (CTS)	All patients referred into secondary care must have been through the Musculoskeletal (MSK) Service to optimise access to conservative treatment unless CTS is severe. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. The ICB will fund carpal tunnel surgery where: • Patient has acute, severe symptoms that persist for more than three months after conservative therapy in MSK/AQP service, treated with either local corticosteroid injection and/or nocturnal splinting OR • There is neurological deficit or median nerve denervation for example sensory blunting, muscle wasting or weakness of thenar abduction AND • Severe symptoms significantly interfering with daily activities and sleep which have been assessed	Restricted
Dupuytren's Disease – palmar fasciectomy	All patients referred into secondary care must have been through the Musculoskeletal (MSK) Service. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community diagnostics. Treatment is not indicated in cases where there is no contracture and in patients with a mild	Restricted



	(less than 20°) contractures, or one which is not progressing and does not impair function.	
	Surgical intervention (collagenase injections, needle fasciotomy, fasciectomy and dermofasciectomy) will only be routinely commissioned in the following circumstance: • Finger contractures causing loss of finger extension of 30° or more at the metacarpophalangeal joint or 20°at the proximal interphalangeal joint. OR • Severe thumb contractures which interfere with function	
Surgical removal of ganglion (Including surgical removal of	Not routinely commissioned for the treatment of asymptomatic ganglia	Restricted
ganglion on wrist surgical removal of seed ganglia at base of digits,	Surgical removal of ganglion on hands will ONLY be routinely commissioned where there is evidence of:	
surgical removal of mucoid cysts at	Aspiration failed to resolve pain or tingling/numbness	
DIP joint)	• Restricted hand function	
	Removal of seed ganglia at base digits Surgical treatment will ONLY be routinely commissioned in the following circumstances: • If ganglion persists or recurs after puncture/aspiration	
	Surgical Removal of Mucoid cysts at DIP joint:	
	Surgical treatment will ONLY be routinely commissioned in the following circumstances:	
	Recurrent spontaneous discharge of fluid OR	
	Significant nail deformity	
Trigger Finger - surgical treatment	Mild cases which cause no loss of function require no treatment or avoidance of activities which precipitate triggering and may resolve spontaneously. Cases interfering with activities or causing pain should be treated with:	Restricted
	1 or 2 steroid injections OR	
	Splinting of the affected finger for 3-12 weeks	



	 Surgical treatment will ONLY be routinely commissioned in the following circumstances: The triggering persists or recurs after one of the above measures (particularly the steroid injections have not worked) OR The finger is permanently locked into the palm. OR The patient has had two other trigger digits unsuccessfully treated with appropriate non-operative methods. OR Diabetics 	
Upper Limb – Shoulder Procedure		
Allograft reconstruction for glenoid bone loss in glenohumeral instability	Not routinely commissioned	Excluded
Arthroscopic Capsular Release (ARC) for Adhesive Capsulitis (Frozen Shoulder)	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. Consideration will be given for those patients who have not responded to a maximum of 3 months of conservative management as listed below: - Appropriate medicines management Physiotherapy (Minimum of 6 weeks, continue for a further 6 weeks if patients function and symptoms have improved) May include advice, exercises, manual therapy, thermotherapy, electrotherapy and steroid injection. Have relevant patient information leaflets/support. Patients must have significant persistent pain preventing them from fulfilling vital activities of daily living. Disturbance of sleep AND	Restricted



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	Have failed to respond to a steroid injection in conjunction with physiotherapy.	
Hydrodilatation for Adhesive Capsulitis (frozen shoulder)	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. Consideration will be given for those patients who have not responded to maximum of 3 months of conservative management as listed below: - • Appropriate medicines management • Have relevant patient information leaflets/support. • Physiotherapy • Patients must have significant persistent pain > 3months preventing them from fulfilling vital activities of daily living. • Disturbance of sleep Due consideration to be given MUA & injection as valid alternative to Hydrodilation.	Restricted
Ultrasound and/or MRI for soft tissue shoulder (if rotator cuff injury is suspected)	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. Consideration will be given for those patients who have not responded to 6 weeks of conservative management as listed below: - • Appropriate medicines management • Have relevant patient information leaflets/support. • Patients must have significant persistent pain preventing them from fulfilling vital activities of daily living.	Restricted



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D' 4 4 4	Disturbance of sleep The IOD shows the state of the	
Diagnostic Arthroscopy	The ICB does not support the use of arthroscopy for diagnostic purposes. Alternatives should be used such as X-ray, MRI, Ultrasound.	Excluded
Therapeutic Shoulder Arthroscopy	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics.	Restricted
	Consideration will be given for those patients who have not responded to conservative management between 3-6 months as listed below: - • Activity modification • Physiotherapy Programme • Appropriate analgesia • Steroid injections where clinically appropriate AND • Full thickness rotator cuff tear as demonstrated by clinical symptoms and radiological imaging. OR • Significant superior labrum anterior posterior (SLAP) tear as demonstrated by clinical symptoms and radiological imaging. OR • Partial thickness rotator cuff tear as demonstrated by clinical symptoms and radiological imaging which has not responded to 3 months of conservative management.	
	 OR Adhesive capsulitis demonstrated by clinical symptoms which has not responded to 6 months of conservative management. OR Adhesive capsulitis demonstrated by clinical symptoms and in the view of the treating consultant is having an extraordinarily severe impact on quality of life, and which has not responded to conservative management including corticosteroid injection where clinically appropriate. OR 	



	 Subcromial shoulder pain demonstrated by clinical symptoms which has not responded to 6 months of conservative management. Non-traumatic shoulder joint instability that has not responded to 6 months of conservative management OR Traumatic shoulder joint instability alongside relevant conservative management as clinically appropriate. 	
Therapeutic Arthroscopy for Minor (type I*) SLAP tear repair	The ICB does not routinely fund this procedure	Excluded
Management of recurrent anterior dislocation of the shoulder. Open and arthroscopic Bankart repair. (exclude young recurrent anterior dislocation)	 Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. The following criteria apply:- Young anterior dislocation (less than 25) whether first time or recurrent, should be assessed in secondary care for consideration of early stabilisation surgery (BESS guidelines). 25-45 years can be treated by physiotherapy following initial instability, but recurrent dislocation should be seen and assessed in secondary care. According to BESS guidelines, first time dislocation in patients over the age of 45 should be seen and assessed in secondary care to pick up cases of associated acute cuff tear. Recurrent atraumatic structural instability in the absence of muscle patterning deemed suitable for surgical intervention. 	Restricted
Bristow Latarjet Procedure	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and	Restricted
The Bristow-Laterjet must not be used for multi-directional instability	received 12 weeks of conservative treatment. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics.	



or psychogenic habitual dislocations.	And	
The Bristow-Laterjet Repair for shoulder instability must only be performed by an experienced Orthopaedic surgeon with advanced arthroscopic skills and familiarity with anatomy encountered during the open Bristow-Laterjet Procedure	 have an Instability Severity Index Score (ISIS) of greater than 3. And Have a severe glenoid bone defect (>20% of the glenoid surface as measured on a preoperative CT scan) 	
Shoulder Replacement Surgery Elderly patients with intact but poorly functional cuff can be considered for reverse total shoulder replacement in line with the criteria for reverse shoulder surgery with rotator cuff pathology.	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. Consideration will be given for those patients who have not responded to conservative management as listed below: - • Activity modification • Appropriate analgesia • Severe pain and functional disability that significantly interferes with activities of daily living from injury e.g. osteoarthritis, post traumatic arthritis of shoulder for at least 6 months duration. AND • has severe limited range of motion of the glenohumeral joint on physical examination. AND • Radiographic evidence of destructive degenerative joint disease (as evidence by 2 or more of the following: irregular joint surfaces, glenoid sclerosis, osteophyte changes, flattened	Restricted



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	glenoid, cystic changes in the humeral head, or joint space narrowing of the shoulder joint)	
Reverse shoulder surgery with rotator cuff pathology	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. • Appropriate analgesia	Restricted
	Steroid injection if clinically appropriate (injection should be no less than 3 months before referral)	
	Patient information leaflets/support	
	Patients must have significant persistent pain preventing them from fulfilling vital activities of daily living.	
	Disturbance of sleep	
	Physiotherapy programme	
	Lifestyle modification	
	Surgical intervention may be considered if patient has failed to benefit from ALL conservative treatments and patient remains in significant pain and activities of daily living are greatly affected AND a total shoulder replacement has been considered AND there is evidence of rotator cuff dysfunction.	
	FOR ANY OTHER INDICATION CLINICIANS MUST APPLY FOR FUNDING VIA THE IFR DEPARTMENT.	
Subcromial Decompression for shoulder pain	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics.	Restricted



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	Patient's must have undergone 6-8 weeks of conservative management: - • Appropriate analgesia • Steroid injection if clinically appropriate • Patient information leaflets/support • Patients must have significant persistent pain preventing them from fulfilling vital activities of daily living. • Disturbance of sleep. • Physiotherapy programme • Lifestyle modification • Referral supported by appropriate imaging. Surgical intervention may be considered if patient has failed to respond to ALL conservative treatment and patient remains in significant pain and activities of daily living are greatly affected.	
Rotator Cuff Disorders	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate conservative management must have been undertaken including any relevant community physiotherapy and diagnostics. AND have received: - Appropriate analgesia Steroid injection if clinically appropriate Patient information leaflets/support Patients must have significant persistent pain preventing them from fulfilling vital activities of daily living. Abduction of shoulder between 60 and 120 degrees. Disturbance of sleep. Physiotherapy programme Lifestyle modification Surgical intervention may be considered if patient has failed: Evidenced base conservative treatment and patient remains in significant pain and activities of daily living are greatly affected	Restricted



Supra-scapula nerve block – May	Inclusion criteria:	Restricted
offer temporary benefit in reducing		
symptoms and facilitating	Adults with shoulder pain secondary to musculoskeletal (MSK) disorders commonly treated	
engagement with	in MSK practice.	
physiotherapy/exercise programme.	Osteoarthritis (GHJ or ACJ)	
	Adhesive capsulitis / frozen shoulder	
Policy is not applicable to:	Rotator cuff arthropathy	
Visceral pain	Single or 'one off' suprascapular nerve blocks	
Cancer pain	Either guided using radiology, or via bony landmarks.	
Hemiplegic shoulder pain		
Peri or post-operative pain		
Continuous nerve block via indwelling	Patients must have been triaged or seen in a community Musculoskeletal (MSK) Service and	
catheter	onwardly referred. Where MSK is not available (East Staffordshire PCN), all appropriate	
	conservative management must have been undertaken including any relevant community	
	physiotherapy and diagnostics.	
	AND I II CIII I	
	AND have the following: -	
	Persistent shoulder pain, i.e. chronic shoulder pain (more than three month's duration)	
	which has failed to, or only partially responded to more traditional therapies (other shoulder	
	injections/ physio etc.)	
	injections, project story	
	The shoulder pain is being generated by more than one site in the shoulder e.g. AC OA	
	with GH OA with supraspinatus cuff pathology	
	Nerve block should be offered conjunction with a prescribed/personalised exercise	
	programme	
Upper Limb - Other Procedures		
Lycra splinting for the prevention	All patients must undergo an assessment with the commissioned Orthotics Provider and	Manual Prior
and correction of upper limb	demonstrate, with clinical evidence from this Provider, that their orthotic needs cannot be met	Approval only
contractures for patients with	before any prior approval request. Prior approval requests will not be accepted by the ICB if this	
neurological dysfunction.	assessment has not been undertaken or where clinical evidence has not been provided. Prior	
	approval requests MUST be made to (<u>IFRTeam@staffsstoke.icb.nhs.uk</u>). Should approval be	
	granted the ICB shall require clinical reports which include evidence of outcomes achieved from	



	<u> </u>	
	the orthoses and patient concordance. The ICB shall not approve prior approval requests on the	
	basis of a historical or previously agreed prior approvals.	
Orthotics (defined as a device used to compensate for impairments of the structure and function of the human body. Not limited to and sometimes referred to as splints, braces, collars, calipers, trusses, specialist footwear and insoles).	All patients must undergo an assessment with the commissioned Orthotics Provider and demonstrate, with clinical evidence from this Provider, that their orthotic needs cannot be met before any prior approval request. Prior approval requests will not be accepted by the ICB if this assessment has not been undertaken or where clinical evidence has not been provided. Prior approval requests MUST be made to (IFRTeam@staffsstoke.icb.nhs.uk). Should approval be granted the ICB shall require clinical reports which include evidence of outcomes achieved from the orthoses and patient concordance. The ICB shall not approve prior approval requests on the basis of a historical or previously agreed prior approvals.	Manual Prior Approval only
Orthotic Treatment for Pectus Excavatum	Not routinely commissioned	Excluded
Surgical Treatment for Pectus Excavatum	Not routinely commissioned by ICBs – NHS England commissioning responsibility https://www.england.nhs.uk/wp-content/uploads/2019/02/1675-Policy Surgery-for-pectus-deformity.pdf	Excluded
Intramedullary Nail in Lower Limb Length Discrepancy	Not routinely commissioned	Excluded
Orthopaedics – Other		
Phonophoresis	Phonophoresis (The use of ultrasound to enhance the delivery of topically applied drugs	Excluded
Prolotherapy	Prolotherapy (Injection of e.g. dextrose into tissues to try to promote healing), Platelet Rich Plasma	Excluded
Cryotherapy	Cryotherapy (use of cooling to promote healing) Viscosupplementation (this has gone through clinical priority advisory group (CPAG) and	Excluded
Viscosupplementation	scored as low	Excluded



Surgical treatment of synovitis and	Not routinely commissioned.	Excluded
tenosynovitis		
General Joint Injections for pain (peripheral joints)	Not commissioned in a sterile theatre unless x-ray screening or general anaesthesia is required and where they are performed with other procedures i.e. nerve blocks or manipulation. Not commissioned when a patient could be a candidate for joint replacement in the next 6-12 months except as a diagnostic tool prior to joint replacement in order to confirm the joint as the major source of pain/ symptoms and for patients who are currently unfit or unsuitable for surgery or who do not wish to proceed to surgery.	Restricted
Trigger Point Injections for Pain	Not commissioned in a sterile theatre unless x-ray screening or general anaesthesia is required and where they are performed with other procedures i.e. nerve blocks or manipulation. Trigger point injections of therapeutic substances into peripheral nerves for persistent non-specific neck/back pain are not routinely commissioned. It is important to note this policy does not cover pain due to malignancy, infection, fracture, ankylosing spondylitis and other inflammatory conditions, radicular pain resulting from nerve root compression or cauda equine syndrome.	Excluded
Therapeutic ultrasound in Physiotherapy	Not routinely commissioned	Excluded
Low Intensity Ultrasound (Exogen) for the Healing of Fractures	See separate commissioning policy	Restricted

Urology

Procedures Thresholds	Status
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Circumcision	All ages: Male Circumcision for cosmetic, social, cultural, and religious reasons is not routinely commissioned. Circumcision will be funded in the following medical circumstances for adults and children: - Pathological Phimosis	Restricted
	OR - Three documented episodes of balanoposthitis	
	Relative indications for circumcision or other foreskin surgery include the following: - Prevention of urinary tract infection in patients with an abnormal urinary tract <u>OR</u> - Recurrent Paraphimosis <u>OR</u> - Balanitis Xerotica Obliterans <u>OR</u> - Congenital abnormalities	
Reversal of Sterilisation: reversal of vasectomy or reversal of tubal ligation (Male and Female)	Not routinely commissioned	Excluded
Surgery for protatism	 Will be routinely commissioned where there is evidence of one of the below: - International prostate symptom score >7. Dysuria. Post voided residual vol >150ml. Recurrent UTI. Deranged renal function. PSA > Age adjusted normal values. Retention of urine Failed medical management 	Restricted
Stress incontinence surgery	The ICB will only agree to fund the first episode of primary surgical treatment of urinary incontinence as NHS England is now responsible for the commissioning of Stress Incontinence Surgery where previous surgery has failed. NHS commissioning » Search Results » incontinence	Restricted



Sacral Nerve Stimulation for	Not routinely commissioned – NHSE responsibility	N/A
Jrinary or Faecal Incontinence	https://www.england.nhs.uk/wp-content/uploads/2013/08/a08-p-b.pdf	
Sacral Nerve Stimulation for Constipation	Not routinely commissioned	Excluded
Injection of therapeutic substance into penis for erectile dysfunction	Not routinely commissioned except as 3rd-line treatment in diabetes after lifestyle & phosphodiesterase inhibitors. Patients must have tried at least 2 oral PDE5 inhibitors prior to referral	Restricted
Erectile dysfunction – medical management	Referrals to secondary care should not be made for the purposes of NHS prescribed medication. Medical management of erectile dysfunction should only be undertaken in accordance with current national restrictions reflected in GMS/PMS contract.	Excluded
Penile Implants	Not routinely commissioned by ICBs – NHSE Responsibility https://www.england.nhs.uk/publication/penile-prosthesis-surgery/	Excluded
Vasectomy	Vasectomy is not routinely commissioned within secondary care. All referrals MUST be sent to a Community Provider. Following an assessment if it deemed clinically appropriate by the Community Surgeon a referral can be made into secondary care. (Consideration for referral into secondary care: cryptorchidism, evidence of allergy to local anaesthetic, inability to locate the vas deferens, previous scrotal surgery resulting in scarring that prevents the vas deferens been palpable, Large hydrocoele/Inguinoscrotal hernia, masses, cysts that prevents the vas deferens from being identified, significant bleeding that cannot be managed) – Prior Approval must be sought.	Restricted
Surgery for hydrocele	Patient is over 2 years of age. AND Discomfort and/or disfigurement have resulted in significant functional impairment, significant interference with activities of daily living, recurrent infections requiring prescribed antibiotic and/or pain with prescribed analgesia use.	Restricted



Vascular Surgery

Procedure	Thresholds	Status
Surgical Treatment of uncomplicated varicose veins and reticular veins or telangiectasia	Surgery (stripping of veins) is routinely commissioned. when the patient meets one or more of the following criteria: • Varicose veins which have bled and are at risk of bleeding again. OR • A history of varicose ulceration OR • Signs of prolonged venous hypertension (haemasiderin, pigmentation, eczema, induration lipodermatosclerosis), or significant oedema associated with skin changes) OR • Superficial thrombophlebitis in association with varicose veins OR • Significant symptoms attributable to chronic venous insufficiency which are resulting in significant functional impairment.	Restricted
Carotid endarterectomy	Not routinely commissioned for asymptomatic or mild to moderate carotid stenosis.	Restricted
Geko device for reducing the risk of venous thromboembolism	Not routinely commissioned	Excluded