1 Guidance

1.1 The evidence on extracorporeal shockwave therapy (ESWT) for refractory tennis elbow raises no major safety concerns; however, current evidence on its efficacy is inconsistent. Therefore, this procedure should only be used with special arrangements for clinical governance, consent and audit or research.

1.2 Clinicians wishing to undertake ESWT for refractory tennis elbow should take the following actions.

- Inform the clinical governance leads in their Trusts.
1.3 NICE encourages further research into ESWT for refractory tennis elbow. Future research should take the form of clinical studies with clearly described patient selection and treatment protocols, including a description of local anaesthesia use and the type of energy applied (see section 2.5). The studies should include validated outcome measures and be based on a minimum of 1-year follow-up. NICE may review the procedure on publication of further evidence.

2 The procedure

2.1 Indications and current treatments

2.1.1 Tennis elbow (also known as lateral epicondylitis) is characterised by chronic degeneration at the origin of the extensor carpi radialis brevis muscle on the lateral epicondyle of the humerus. It is usually caused by injury or overuse. Symptoms include pain, weakness and stiffness of the outer elbow.

2.1.2 Conservative treatments include rest, application of ice, analgesic medication, non-steroidal anti-inflammatory drugs, orthotic devices, physiotherapy, eccentric training/stretching and corticosteroid injection.

2.2 Outline of the procedure

2.2.1 Extracorporeal shockwave therapy is a non-invasive treatment in which a device is used to pass acoustic shockwaves through the skin to the affected area. Ultrasound guidance can be used to assist with positioning of the device.

2.2.2 Extracorporeal shockwave therapy may be applied in one or several
sessions. Local anaesthesia may be used because high-energy ESWT can be painful. Different energies can be used and there is evidence that local anaesthesia may influence the outcome of ESWT.

2.2.3 The mechanism by which this therapy might have an effect on tendinopathy is unknown.

Sections 2.3 and 2.4 describe efficacy and safety outcomes from the published literature that the Committee considered as part of the evidence about this procedure. For more detailed information on the evidence, see the overview.

2.3 Efficacy

2.3.1 In a randomised controlled trial (RCT) comparing 272 patients treated by ESWT or sham ESWT, success was reported in 26% (32/124) and 25% (31/122) of patients, respectively, at 3-month follow-up (success defined as a Roles and Maudsley score of 1 or 2 out of 4 and no requirement for additional treatment). An RCT of 93 patients treated by ESWT or steroid injection reported treatment success (at least 50% improvement in Visual Analogue Scale [VAS] score) in 60% (29/48) and 84% (21/25) of patients, respectively, at 3-month follow-up (p < 0.05).

2.3.2 An RCT of 75 patients treated by ESWT or sham ESWT reported that at 3 months, 35% (14/40) and 37% (13/35) of patients, respectively, had at least a 50% improvement in VAS score for pain during the day (p < 0.01 in both treatment groups); and 30% (12/40) and 43% (15/35) of patients, respectively, had at least a 50% improvement in VAS score for pain at night (p value stated as ‘non-significant’ and p < 0.05, respectively).

2.3.3 The Specialist Advisers listed key efficacy outcomes as relief of symptoms and functional improvement.

2.4 Safety

2.4.1 Two RCTs of 272 and 114 patients reported pain in 11% (15/134) and 50% (28/56) of ESWT patients, and 4% (6/136) and 22% (13/58) of sham
2.4.2 In the RCT of 272 patients, transient skin reddening occurred in 31% (42/134) of ESWT patients and 8% (11/136) of sham ESWT patients, and transient swelling occurred in 7% (9/134) and 6% (8/136) of patients, respectively.

2.4.3 The RCT of 114 patients treated by ESWT or sham ESWT reported a local reaction (not otherwise described) in 11% (6/56) and 9% (5/58) of patients, respectively. In this study 18% (10/56) of patients in the ESWT group experienced nausea compared with none in the sham ESWT group.

2.4.4 In the RCT of 75 patients, 2 patients in the ESWT group had worsened symptoms after 2 treatment sessions and withdrew from the study.

2.4.5 The Specialist Advisers listed adverse events as bruising, transient skin reddening and local skin damage. Theoretical adverse events include rupture of the common extensor tendon.

2.5 Other comments

2.5.1 The Committee found interpretation of the data difficult because of the diversity of treatment protocols and comparators used, varying reported end points and inconsistencies in terms of the use of local anaesthesia and energy type. The results of studies conflicted and there was evidence of a substantial placebo response. Previous guidance on this procedure published in 2005 had found the evidence on efficacy inadequate, and new evidence has not been published to alter that view.

2.5.2 Tennis elbow is a common condition and many patients who have it are refractory to other treatments. If the procedure is efficacious in selected patients, it has the potential for a high impact. This makes provision of robust data particularly important.
3 Further information

3.1 This guidance requires that clinicians undertaking the procedure make special arrangements for audit. NICE has identified relevant audit criteria and developed an audit tool (which is for use at local discretion).

3.2 For related NICE guidance see our website.

Information for patients

NICE has produced information on this procedure for patients and carers ('Understanding NICE guidance'). It explains the nature of the procedure and the guidance issued by NICE, and has been written with patient consent in mind.

4 Other NICE recommendations on extracorporeal shockwave therapy

Guidance has also been issued on Extracorporeal shockwave therapy for refractory Achilles tendinopathy and Extracorporeal shockwave therapy for refractory plantar fasciitis. It replaces the previous guidance on Extracorporeal shockwave therapy for refractory tendinopathies (plantar fasciitis and tennis elbow) (IPG139, November 2005).

5 About this guidance

NICE interventional procedure guidance makes recommendations on the safety and efficacy of the procedure. It does not cover whether or not the NHS should fund a procedure. Funding decisions are taken by local NHS bodies after considering the clinical effectiveness of the procedure and whether it represents value for money for the NHS. It is for healthcare professionals and people using the NHS in England, Wales, Scotland and Northern Ireland, and is endorsed by Healthcare Improvement Scotland for implementation by NHSScotland.

This guidance was developed using the NICE interventional procedure guidance process.

We have produced a summary of this guidance for patients and carers. Tools to help you
put the guidance into practice and information about the evidence it is based on are also available.

Changes since publication

6 January 2012: minor maintenance.

Your responsibility

This guidance represents the views of NICE and was arrived at after careful consideration of the available evidence. Healthcare professionals are expected to take it fully into account when exercising their clinical judgement. This guidance does not, however, override the individual responsibility of healthcare professionals to make appropriate decisions in the circumstances of the individual patient, in consultation with the patient and/or guardian or carer.

Implementation of this guidance is the responsibility of local commissioners and/or providers. Commissioners and providers are reminded that it is their responsibility to implement the guidance, in their local context, in light of their duties to avoid unlawful discrimination and to have regard to promoting equality of opportunity. Nothing in this guidance should be interpreted in a way which would be inconsistent with compliance with those duties.

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Contact NICE

National Institute for Health and Clinical Excellence
Level 1A, City Tower, Piccadilly Plaza, Manchester M1 4BT

www.nice.org.uk
nice@nice.org.uk
Endorsing organisation

This guidance has been endorsed by Healthcare Improvement Scotland.

Accreditation

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