

# Pilot study to assess the feasibility and patient numbers required prior to a randomised controlled trial of the effect of visual distraction for painful procedures in the burns unit

<b>Submission date</b> 30/09/2004	<b>Recruitment status</b> No longer recruiting	<input type="checkbox"/> Prospectively registered
<b>Registration date</b> 30/09/2004	<b>Overall study status</b> Completed	<input type="checkbox"/> Protocol
<b>Last Edited</b> 03/11/2015	<b>Condition category</b> Injury, Occupational Diseases, Poisoning	<input type="checkbox"/> Statistical analysis plan
		<input type="checkbox"/> Results
		<input type="checkbox"/> Individual participant data
		<input type="checkbox"/> Record updated in last year

**Plain English summary of protocol**  
Not provided at time of registration

## Contact information

**Type(s)**  
Scientific

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## Additional identifiers

**Protocol serial number**  
N0060131616

## Study information

**Scientific Title**

Pilot study to assess the feasibility and patient numbers required prior to a randomised controlled trial of the effect of visual distraction for painful procedures in the burns unit

**Study objectives**

Does the use of 2D video glasses with DVD film images and stereo sound during dressing changes for patients with burns decrease the intensity of their experience of pain and anxiety?

**Ethics approval required**

Old ethics approval format

**Ethics approval(s)**

Not provided at time of registration

**Study design**

Randomised controlled trial

**Primary study design**

Interventional

**Study type(s)**

Treatment

**Health condition(s) or problem(s) studied**

Pain

**Interventions**

Short visual analogue scale patient rated scores for pain and anxiety, an appropriate questionnaire and short concluding taped interview after intervention. Quantitative methods will be used.

**Intervention Type**

Other

**Phase**

Not Specified

**Primary outcome(s)**

Improved pain control and toleration of regular dressing change and debridement of burns with reduced anxiety and awareness.

**Key secondary outcome(s)**

Not provided at time of registration

**Completion date**

30/09/2003

**Eligibility****Key inclusion criteria**

Minimum 10 consenting adults suitable to trial criteria and protocols.

**Participant type(s)**

Patient

**Healthy volunteers allowed**

No

**Age group**

Adult

**Sex**

All

**Key exclusion criteria**

Does not meet inclusion criteria

**Date of first enrolment**

01/04/2003

**Date of final enrolment**

30/09/2003

**Locations**

**Countries of recruitment**

United Kingdom

England

**Study participating centre**

Chelsea & Westminster Hospital

London

United Kingdom

SW10 9NH

**Sponsor information**

**Organisation**

Department of Health

**Funder(s)**

**Funder type**

Hospital/treatment centre

**Funder Name**

Chelsea and Westminster Healthcare NHS Trust (UK)

**Results and Publications****Individual participant data (IPD) sharing plan****IPD sharing plan summary**

Not provided at time of registration