Interventional procedure overview of epidermal radiotherapy using rhenium-188 paste for non-melanoma skin cancer

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Table 1 Abbreviations

Abbreviation	Definition
BCC	Basal cell carcinoma
EMPD	Extramammary Paget's disease
EBRT	External beam radiation therapy
NMSC	Non-melanoma skin cancer
SCC	Squamous cell carcinoma
SCT	Skin cancer therapy
SD	Standard deviation

Indications and current treatment

NMSC is the most common type of cancer. It affects the cells in the top layers of the skin. The most common types of NMSC are BCC and SCC. The main symptom is the appearance of lesions (lumps or discoloured patches) on the skin. The lesions are most commonly found on skin that is regularly exposed to the sun.

The current standard of care within the NHS depends on the initial presentation of non-melanoma skin cancer, such as the type, size and location of the lesion. Surgery is the main treatment. Other treatment options include chemical therapies, cryotherapy, brachytherapy, external beam radiotherapy and photodynamic therapy.

What the procedure involves

The procedure is done without the need for anaesthesia or inpatient admission. It uses a beta-emitter radioisotope rhenium-188, which can penetrate the human tissue up 3 mm deep. The rhenium-188 is bound to a matrix to form a paste and is applied using a specially designed applicator. During the treatment, the area to be treated is protected from direct contact with the paste by a cream or sterile

transparent foil. The rhenium-188 paste is then applied on the area of the lesion with a safety margin. The treatment time is calculated based on the applied radioactivity and the area of the region being treated and is typically about 30 to 180 minutes. The paste dries out during the treatment time and turns into a flexible film. The film is removed when the treatment is over. The dead cancer cells are gradually replaced with new healthy cells.

Unmet need

NMSCs are the most common class of skin cancers. The most common treatment is surgery. Surgery may lead to scarring if treating large or multiple lesions. Surgery or alternative therapies (chemical therapies, cryotherapy, brachytherapy, external beam radiotherapy and photodynamic therapy) may also be technically difficult in some common lesion sites, for example on fingers, ears or the nose. This procedure may offer a treatment option when surgery is not advised because of comorbidities or contraindicated, or for lesions which the anatomical position may result in a suboptimal cosmetic result using conventional approaches. Rhenium-188 paste can be used on genital lesions. Avoiding surgery on these lesions may be beneficial for preserving sexual function and quality of life.

Outcome measures

The main efficacy outcome is remission status. Four studies determined this with dermoscopic or histological assessment (Carrozzo, 2013; Carrozzo, 2014; Castellucci, 2021; Cipriani, 2022); 2 studies did not report the method used to determine remission status (Cipriani, 2017; Sedda, 2008).

One study used the Common Terminology Criteria for Adverse Events (CTCAE 5.0) to grade early skin toxicity (Castellucci, 2017). This tool presents sets of criteria against each category of adverse event or symptom that is commonly caused by cancer treatments (for example, erythema). The criteria enable IP overview: Epidermal radiotherapy using rhenium-188 paste for non-melanoma skin cancer

clinicians to make standardised assessments of the severity of the adverse event against descriptions of the event. The descriptions are marked against graded severity ratings from grade 1 (mild) to 5 (death related or due to adverse event).

One study used the Radiation Therapy Oncology Group criteria (RTOG) to evaluate cosmetic results (Castellucci, 2017). This scale describes cosmetic changes to the skin. It ranges from poor (for example, ulceration or necrosis) to excellent (for example, no changes or slight pigment change).

Evidence summary

Population and studies description

This interventional procedures overview is based on publications reporting on the safety and efficacy outcomes of approximately 218 people with NMSC from 1 single-arm trial (Castellucci, 2021) and 5 case series (Carrozzo, 2013; Carrozzo, 2014; Cipriani, 2017; Cipriani, 2022; Sedda, 2008). Additional safety outcomes from 22 people with NMSC reported in 1 conference abstract were also included. All patients had the procedure. This is a rapid review of the literature, and a flow chart of the complete selection process is shown in figure 1. This overview presents 6 published studies and 1 abstract as the key evidence in table 2 and table 3.

The single-arm trial was the only prospective study (Castellucci, 2021). All 6 key studies were done in Italy. Of the key studies, 4 included people from the same hospital. Not all studies reported the dates that participants had treatment. All key studies included at least one author that co-authored at least one other publication included in this overview. There may be some overlap in participants included in these studies.

The mean age of people with NMSC considered for efficacy outcomes was between 66 and 81; 2 of the published studies and the conference abstract did IP overview: Epidermal radiotherapy using rhenium-188 paste for non-melanoma skin cancer

not report the ages of participants in their study (Cipriani, 2017; Schwarzenbock, 2022; Sedda, 2008). Follow up was a mean of 20 months (range: 3 to 33 months) in the one prospective study (Castellucci 2021). Among the case series, mean follow-up was between 288 days (approximately 9.5 months) and 51 months. Follow-up was 12 months in the conference abstract (Schwarzenbock, 2022). It was unclear whether the reported follow up points were after the last treatment received or if further treatment sessions were done during follow-up in some studies.

Whether lesions had been previously treated, which therapies were applied to lesions if previously treated and the type and size of lesions treated with rhenium-188 paste varied between and within studies. The most common type of lesion was BCC but most studies also included SCCs and sometimes EMPD and Bowen's disease. One study only included people with EMPD (Carrozzo, 2014) and one study only included people with SCC (Carrozzo, 2013). Most studies included some people with lesions that were previously treated with another intervention. Most studies included lesions located anywhere on the body but most were on the head and neck. Two studies only included people with lesions on the genitalia (Carrozzo, 2013; Carrozzo, 2014). Across studies, average lesion size was between 3 and 10 cm² but ranged from 0.3 to 65 cm². Lesion depth was between 0.1 to 2.5 mm across studies. Two studies did not report the size or depth of lesions.

In the single-arm trial, 18 out of 60 lesions had already been treated with other therapies and relapsed (5 lesions had previous surgery; 2 lesions had previous surgery and photodynamic therapy or cryotherapy; 10 lesions had previous cryotherapy, laser and photodynamic therapy; 1 lesion had previous imiquimod) while 42 lesions were new diagnoses at presentation. In this study, 68% of lesions were BCCs, 30% were SCCs and 1 lesion was BCC and SCC. Lesions were mostly on the face, ears, nose and scalp (77%) and the remaining on

extremities (15%) and trunk (8%). The mean surface area of lesions was 7 cm² (1 to 36 cm²) and the mean depth was 1.1 mm (0.2 to 2.5 mm). This was not the largest lesion size among the studies (based on the mean lesion size and the maximum lesion size included) but they did treat the deepest lesions compared to other studies. The authors suggested that the size of lesions treated in their study may explain some of their grade 3 side-effects.

In the case series of 53 people, 28% of lesions had previous surgical treatment (Sedda, 2008). Lesions were diagnosed as BCC (70%) and SCC (30%) and mostly on the head and neck (70%). Other lesions were on the upper and lower limbs (22%) and trunk and back (8%). The mean size of lesion was 7 cm² (SD=8.9). The depth of lesions was not reported.

In the case series of 52 people, 54% of lesions had previous surgery or other intervention (Cipriani, 2022). Most lesions were BCCs (58%) but SCCs (35%) and 4 lesions diagnosed as Bowen's disease (n=2) and EMPD (n=2) were also treated. Most commonly, lesions were on the head and neck (73%) or genitalia (15%). Average lesion size was 10 cm² (range between 0.3 and 61 cm²) and depth was between 0.3 and 0.6 mm.

In the case series of 43 people, 26% had previous surgery (Cipriani, 2017). Lesions were mostly BCCs (67%) and the rest were SCCs. Lesions were mostly located on the head and neck (65%) but lesions on the back, arm, leg, and penis were also treated. Median lesion size was 3 cm² (range 1 to 49 cm²). Two lesions were deeper than 0.5 mm.

All participants in the case series of 15 people (Carrozzo, 2013) had previous topical therapy for SCC lesions located on the penis. One lesion was also previously treated with imiquimod and 5-fluorouracil and 2 with surgery. The size of lesions treated in this study was not reported.

In the case series of 5 people (Carrozzo, 2014), 80% of the participants had previous topical treatments for lesions. All 5 lesions were EMPD located on the genitalia. The size of lesions was not reported.

In the abstract reporting a case series of 22 people (Schwarzenbock, 2022), 58% of lesions were BCC, 30% were Bowen's disease and 13% were SCC. Average lesion size was 3 cm² (range 0.5 to 20 cm²). Lesion depth ranged from 0.1 to 2.1 mm. This study did not report on previous treatment of the lesions.

Table 2 presents study details.

Figure 1 Flow chart of study selection

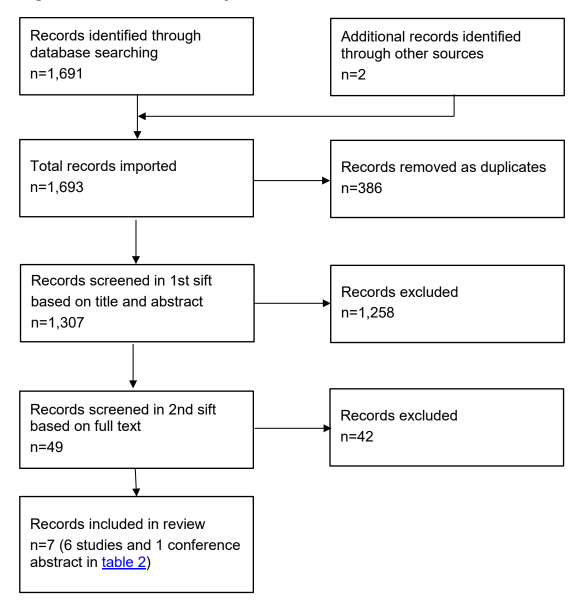


Table 2 Study details

Study no.	First author, date country	Patients (male: female)	Age	Study design	Inclusion criteria	Intervention	Follow up
1	Castellucci, 2021, Italy	n=50 (70% male, 30% female) 60 histologically proven NMSCs (n=41 BCC; n=18 SCC; n=1 BCC and SCC)	Mean=81	Single-arm trial October 2017 to January 2020	1) histologically proven BCC or SCC; (2) lesion thickness invasion less than 2.5 mm; (3) lesions located in the scalp, face, ears, or fingers or other areas in which surgery, EBRT or standard brachytherapy would have been difficult; (4) contraindication or refusal of surgery	Rhenium-188 brachytherapy using Rhenium-SCT (Oncobeta GmbH, Germany) Results were evaluated after 1 treatment session.	Mean 20 months (range 3 to 33)
2	Sedda, 2008, Italy	n=53 (55% male, 45% female) n=37 BCC; n=16 SCC n=9 had multiple BCCs; n=3 had multiple SCCs	Not reported	Case series	Histologically confirmed BCC or SCC; relapse of tumour or surgery considered impossible or aesthetically unacceptable.	Rhenium-188 brachytherapy 43 people had 1 treatment session, 8 people had 2 and 2 people had 3 treatment sessions.	Mean 51 months (range 20 to 72)

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Study no.	First author, date country	Patients (male: female)	Age	Study design	Inclusion criteria	Intervention	Follow up
3	Cipriani, 2022, Italy	n=52 (65% male, 35% female) n=55 lesions which included n=32 BCC, 2 SCC, 2 Bowen's disease, 2 EMPD	Mean=71.7	Retrospective case series 2005 to 2014	Histologically confirmed BCC, SCC, Bowen's disease or EMPD	Rhenium-188 brachytherapy using Rhenium-SCT Results were evaluated after 1 session.	Mean=414 days; Median= 296 days
4	Cipriani, 2017, Italy	Unclear; n=42, 43 and 44 reported in different parts of the full-text (approximately 58% male, 42% female) n=87 histologically confirmed lesions including BCC and SCC. N=1 patient had Bowen's disease.	Not reported	Retrospective case series	Histologically confirmed NMSC; complete histological record, dosimetry information and imaging material	Rhenium-188 brachytherapy Fractionated treatment was planned for people with lesions on lips and genitalia across 2 or 3 sessions. Otherwise responses reported after 1 session.	Mean 288 days (range 35 to 1,150 days)
5	Carrozzo, 2013, Italy	n=15 (100% male)	Mean= 66	Case series June 2005 to April 2010	Confirmed diagnosis of SCC of penis	Rhenium-188 brachytherapy using Rhenium-SCT	Mean 51 months (range 12

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Study no.	First author, date country	Patients (male: female)	Age	Study design	Inclusion criteria	Intervention	Follow up
						(ITM, Germany)	to 84 months)
						Up to 7 sessions. Some people had multiple sessions to treat different lesions.	
6	Carrozzo, 2014, Italy	n=5 (20% male; 80% female) 80% primary EMPD; 20% secondary	Mean=69	Case series First treatment sessions were between 2008 and 2010	Clinical diagnosis of EMPD	Rhenium-188 brachytherapy using Rhenium-SCT (ITM, Germany) 1 session unless relapsed.	Mean 34 months (range 27 to 48)
7	Schwarzenbock, 2022, Germany	n=22 n=37 lesions (n=23 BCC, 12 Bowen's disease, 5 SCC)	Not reported	Conference abstract reporting case series	Histologically confirmed NMSC	Rhenium-188 brachytherapy using Rhenium-SCT	Up to 12 months

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Study no.	First author, date country	Patients (male: female)	Age	Study design	Inclusion criteria	Intervention	Follow up
						Results reported after 1 session.	

Table 3 Study outcomes

First author, date	Efficacy outcomes	Safety outcomes
Castellucci, 2021	Complete response at 6 months=98% (53/54 evaluable lesions; total sample included 60 lesions) Partial response in 1/54 lesions (there was a small residual lesion, which was surgically excised with good cosmetic results).	No notable pain during or after treatment. 93% (56/60) of lesions showed early localised side-effects (grades 1 to 2) including skin erythema, faint or moderate oedema and ulcerations. These resolved in a mean of 4 weeks.
	Lesions free from relapse at 12 months=100% (41/41 evaluable lesions) Lesions free from relapse at 24 months=96% (23/24 evaluable lesions) 1 lesion required retreatment within 24 months.	7% (4/60) of lesions showed more severe (grade 3) early side effects which resolved within 90 days (mean=10 weeks). Late side effects (3 to 33 months) included dyschromia, slight skin atrophy, hair loss. These were not considered significant.
	Cosmetic results 12 to 33 months after 1 session • Excellent= 73% (30/41) • Good= 27% (11/41)	

First author, date	Efficacy outcomes	Safety outcomes
Sedda, 2008	Complete clinical remission= 100% (53/53) 3 to 5 months after treatment (43 people had 1	Erythema in irradiated area which disappeared after 2 to 7 days.
	treatment session, 8 people had 2 and 2 people had 3 treatment sessions)	Bleeding in large lesions which stopped and formed a scab 10 to 30 days after treatment.
		No pain.
	There were no relapses within 20 to 72 month follow up.	No systemic or topical side-effects. No measurable radioactive contamination.
	Complete healing - 1000/ (52/52) ofter 20 to 72	No longer term side effects.
	Complete healing= 100% (53/53) after 20 to 72 months (mean= 51 months) and up to 3 sessions. No disfiguring scarring.	
	Histological analysis, available for about 60% of patients, confirmed complete tumour regression after the treatment (timing not reported).	
Cipriani, 2022	Complete remission= 100% (52/52) of people after 1 session.	10% (5/52) died after 1 year of treatment. Cause of death was not reported.
	Remission was sustained throughout follow up (median= 296 days; 27% (14/52) people were lost	Slight depigmentation of the treated region.
	to follow up and may not have been included in this figure).	No other complications or post-interventional problems.
		No radioactivity contamination of patients, staff or equipment

First author, date	Efficacy outcomes	Safety outcomes
Cipriani, 2017	Complete remission at follow up=100% (mean follow up 288 days (range 35 to 1,150 days, median 212 days; 1 person was lost to follow up).	Some lesions produced clear serum for 1 to 2 weeks after treatment but stopped without further intervention.
	5% (2/43) required unplanned retreatment because the security margin was too tight, resulting in a recurrence at the border of the treated area. All achieved full remission after this. Radiation wounds healed within 30 to 154 days (median=53 days).	No other side effects or adverse events during or between 30 and 154 days after treatment. No pain during or after treatment. No haematological toxicity observed. No radioactivity contamination.
Carrozzo, 2013	Complete remission at follow up=80% (12/15) (mean 51 months; 1 person lost to follow up). 13% (2/15) required surgical salvage therapy.	Faint redness in irradiated area immediately after treatment. Erythema present after a few days. Serum secretion which formed a crust. Visual clinical healing in 3 to 4 months. The authors report no pain, discomfort or other side effects.
Carrozzo, 2014	Complete remission at the end of treatment=100% (5/5) Of the 5 patients,1 had 1 treatment session and 4 had 2 sessions; 2 of the 5 patients had relapse inside the treated area and 2 had relapse at the periphery of the previously treated area. Clinical healing in a mean of 34 months.	Faint redness in irradiated area immediately after treatment. Burning sensation and superficial erosions in days after treatment requiring topical or analgesic treatment. Erythema Serum secretion from wound which formed a scab and resolved in 2 to 3 weeks.

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First author, date	Efficacy outcomes	Safety outcomes
		One patient, who had secondary EMPD, died from metastatic bladder cancer.
Schwarzenbock, 2022	Efficacy data is not extracted from conference	Not otherwise reported
	abstracts.	<u>Day 14</u>
		Most lesions increased in size up to 900% because of inflammation.
		Itching reported in 18% (7/40)
		Scaling in 8% (3/40)
		Mild pain in 13% (5/40)
		4 months
		8% (3/40) reported itching
		5% (2/40) reported pain (these were people with persistent NMSC and ulceration)
		12 months
		48% (13/27) showed hypopigmentation
		67% (18/27) showed mild scarring

Procedure technique

Four studies and the conference abstract named the device used to apply rhenium-188 paste to the skin (Carrozzo, 2013; Carrozzo, 2014; Castellucci, 2021; Cipriani, 2022; Schwarzenbock, 2022). All used the Rhenium-SCT kit. This device contains the radioactive material in an applicator and has a shield to protect the person applying the substance to the skin. All treatments were done within a nuclear medicine department. Cipriani (2017) used a device that looks like this kit but did not name it. Sedda (2008) gave no information on device or application tool.

All studies used materials to prevent direct contact between the radioactive material and the skin. One study (Sedda, 2008) used protective cream, 4 studies used an adhesive plastic foil (Carrozzo, 2013; Carrozzo, 2014; Castellucci, 2021; Cipriani, 2017) and 1 study used the cream and/or the foil (Cipriani, 2022). The conference abstract did not report what method was used (Schwarzenbock, 2022).

All studies tailored the dose according to initial radioactivity, isotope emission energy, contact time and the size and depth of the lesion. All studies used at least one algorithm to calculate the dose required. Castellucci (2021) used a progressive reduction of delivered doses because of the incidence of early side-effects.

Studies applied the radioactive substance to an area which included the lesion and a border of healthy tissue. Carrozza (2013; 2014) used a margin of 2 to 4 mm. Cipriani (2017) used a margin of 3 to 5 mm. Castellucci (2021) used a margin of 3mm. Cipriani (2022) used a margin of 5 mm.

Irradiation time varied within and between studies. Sessions lasted between 15 minutes and 2 hours in Sedda (2008) and Cipriani (2017). Carrozzo (2013; 2014) both reported sessions lasted between 30 minutes and 1 hour. Castellucci IP overview: Epidermal radiotherapy using rhenium-188 paste for non-melanoma skin cancer

(2021) reported sessions lasted between 21 and 285 minutes and Cipriani (2022) reported sessions lasted between 8 and 240 minutes. In the conference abstract, Schwarzenbock (2022) reported sessions lasted between 38 to 175 minutes.

The number of sessions and reasons for multiple sessions varied between studies. Details are presented under efficacy outcomes in terms of relapse and reintervention rates.

Efficacy

Complete remission

All 6 studies reported data on remission. Complete remission ranged from 80% to 100%.

In the single-arm trial of 50 people with 60 lesions, 54 lesions were followed up 6 months after 1 treatment session and 98% showed complete response to treatment (Castellucci, 2021). Partial response was observed in 1 lesion.

In the case series of 53 people, 100% (53/53) were in apparent complete clinical remission after 3 to 5 months (Sedda, 2008). 81% (43/53) were treated with 1 session, 15% (8/53) with 2 sessions and 4% (2/53) with 3 sessions.

In the case series of 52 people, 100% (52/52) showed complete remission after 1 session (Cipriani, 2022).

In the case series of 43 people, 100% (43/43) showed complete remission between 35 and 1,150 days after treatment (median=212 days; Cipriani, 2017). This included 2 people (5%) whose lesions needed unplanned retreatment because of recurrence at the border of the treated area.

In the case series of 15 people with SCC of the penis, complete remission was observed in 80% (12/15) of people, 13% (2/15) of lesions did not respond to

therapy and 1 person was lost to follow-up (Carrozzo, 2013). Lesions were treated in up to 7 sessions and mean length of follow-up was 51 months.

In the case series of 5 people, all 5 showed complete remission after a mean follow up period of 34 months (Carrozzo, 2014). Lesions were treated with 1 session in 1 person, and with 2 sessions in 4 people. Retreatment was triggered by partial response in these 4 people.

Relapse

In the single-arm trial of 50 people including 60 lesions treated in 1 session, 100% of people who were followed up (41/41) were still in remission at 12 months (Castellucci, 2021) and 96% of those who were followed up (23/24) were still in remission at 24 months. One lesion needed retreatment by 24 month follow up.

In the case series of 53 people, there were no relapsed cases after 20 to 72 months (mean=51; Sedda, 2008). Lesions were treated in up to 3 sessions. The authors did not report whether additional sessions were planned or triggered by recurrence.

In the case series of 52 people, there were no relapsed cases after a median follow-up of 296 days (Cipriani, 2022). This figure may not include 14 patients that were lost to follow-up. They reported 10% (5/53) of people died after the first post-interventional year and reason for death was not reported.

In the case series of 43 people, fractionation of treatment across 2 or 3 sessions was planned for people with lesions on their lips or genitals. Further treatment was triggered in some cases by recurrence at follow up (Cipriani, 2017). They reported that 5% (2/43) of people needed unplanned retreatment with rhenium-188 paste. All had full remission after retreatment.

In the case series of 15 people with SCC of the penis, lesions were treated in up to 7 sessions (Carrozzo, 2013). It was unclear whether retreatment was triggered by partial response or recurrence which required further intervention or if this was planned. Some people with SCC of the penis were treated in different locations in different sessions. They reported that 13% (2/15) required surgical salvage therapy because of non-response. Results were reported at a mean follow-up of 51 months. One participant was lost to follow-up.

In the case series of 5 people, relapse triggered a second session of therapy in 4 of 5 people (Carrozzo, 2014). After retreatment, all 5 were in complete remission at a mean of 34 months follow-up.

Healing and cosmesis

In the single-arm trial of 50 people with 60 BCC and SCC lesions, Radiation Therapy Oncology Group criteria were used to evaluate cosmetic results 12 to 33 months after 1 session (Castellucci, 2021). Of 41 evaluable lesions (68% of original sample), 73% (30/41) were rated excellent and 27% (11/41) were rated good.

In the case series of 53 people, the authors reported no disfiguring scarring after 20 to 72 months (mean=51 months) and that all lesions had completely healed after up to 3 sessions (Sedda, 2008).

In the case series of 52 people with lesions treated in 1 session, slight depigmentation was observed (Cipriani, 2022).

In the case series of 43 people, radiation wounds were observed between 30 and 154 days after treatment (Cipriani, 2017). These healed without medical intervention. The authors reported that treatment leaves no scar but in some cases faint discolouration of the skin is present.

In the case series of 15 people with SCC of the penis, visual and clinical healing was reported 3 to 4 months after treatment (Carrozzo, 2013).

In the case series of 5 people, complete healing was observed after a mean of 34 months follow up (Carrozzo, 2014).

Safety

In the single-arm trial of 50 people with 60 BCC and SCC lesions, Common Terminology Criteria for Adverse Events (5.0) was used to assess early skin toxicity within the first 30 days of 1 session of treatment in all 60 lesions (Castellucci, 2021). 93% (56/60) showed early side-effects (grades 1 to 2) including erythema, faint or moderate oedema and ulcerations. These resolved in a mean of 4 weeks. 7% (4/60) lesions showed more severe (grade 3) early side effects which resolved within 90 days (mean=10 weeks). No notable pain was reported during or after treatment. Late side-effects (3 to 33 months) included dyschromia, slight skin atrophy, hair loss. The authors did not consider these to be significant side-effects.

In the case series of 53 people, erythema was reported in the irradiated area which disappeared after 2 to 7 days (Sedda, 2008). Bleeding was reported in large lesions which stopped and formed a scab 10 to 30 days after treatment. No pain, systemic or topical side-effects, measurable contamination or other longer-term side effects were reported.

In the case series of 52 people, the authors reported no other complications, post-interventional problems or contamination (Cipriani, 2022). They reported 9% (5/53) of people died after the first post-interventional year and reason for death was not reported.

In the case series of 43 people, the authors reported that some lesions produced clear serum for 1 to 2 weeks after treatment but stopped without further

intervention (Cipriani, 2017). The authors reported that no pain, haematological toxicity, contamination or other side effects were observed during or after treatment.

In the case series of 15 people with SCC of the penis, faint redness in the irradiated area was observed immediately after treatment (Carrozzo, 2013). Erythema was present after a few days. There were 3 cases of serum secretion from the treated area which then formed a crust. The authors reported no pain, discomfort or other side effects.

In the case series of 5 people, there was faint redness in the irradiated area immediately after treatment (Carrozzo, 2014). The authors reported a burning sensation and superficial erosions in the days after treatment, requiring topical or analgesic treatment. Erythema was observed and some lesions secreted serum which later formed a scab.

Other adverse events were reported in the conference abstract. At day 14, Schwarzenbock (2022) reported up to 900% increase in lesion size because of inflammation, itching in 18%, mild pain in 13% and scaling in 8%. At 4 months, 8% of participants reported itching and 5% reported pain. The people who reported pain had either persistent NMSC or ulceration of the lesion. At 12 months, 48% showed hypopigmentation and 67% showed mild scarring.

Anecdotal and theoretical adverse events

The authors of the single-arm trial (Castellucci et al, 2021) stated that a longer observation period is needed to rule out the theoretical possibility of developing another (secondary) cancer in the local skin due to the radiation treatment.

Expert advice was sought from consultants who have been nominated or ratified by their professional society or royal college. They were asked if they knew of any other adverse events for this procedure that they had heard about

(anecdotal), which were not reported in the literature. They were also asked if they thought there were other adverse events that might possibly occur, even if they had never happened (theoretical).

They listed the following anecdotal and theoretical adverse events:

- infection
- skin burns
- dry skin
- alopecia
- skin induration
- hyperpigmentation
- telangiectasia
- secondary/ late malignancy in treated region (up to decades later)
- delayed wound healing
- unintended exposure to other parts of the body
- keloids
- necrosis.

Four professional expert questionnaires for this procedure were submitted. Find full details of what the professional experts said about the procedure in the specialist advice questionnaires for this procedure.

Validity and generalisability

- The 6 studies included in the key evidence had a collective sample of 218 people. Each study had 53 participants or less.
- All 6 key studies were done in Italy.
- Complete response was between 80% and 100%, indicating a consistent efficacy signal.

- The single-arm trial was the only prospective study (Castellucci, 2021) which
 included 50 patients and had a mean follow-up of 20 months. The authors
 stated larger patient population and longer follow-up are needed to confirm the
 preliminary data and find the optimal dose needed to achieve complete
 response without significant side effects.
- The 5 retrospective case series did not have standardised follow up points and length of follow up varied widely within each sample (range: 288 days to 51 months). No studies reported reasons for loss to follow up.
- All 6 studies were done in Italy and had overlaps in the research groups. Four
 of the 6 studies were conducted in the same hospital and all 6 publications
 shared at least one author. The dates that people received treatment were not
 reported in some studies. This may mean that there is some overlap in
 participants.
- Five of the 6 study designs were case series and did not report a clear strategy for patient selection. Cases presented in these reports may have been subject to selection bias. The single-arm trial reported inclusion and exclusion criteria and a clearer protocol for patient identification (Castellucci, 2021).
- Five studies included participants whose lesions were on different locations on the body. One study only included participants with lesions on the penis (Carrozzo, 2013). Four studies reported the surface area of the lesion and these appear to be similar. All studies included some participants whose lesions had previously been treated with another intervention before rhenium-188 paste.
- Studies used a mixture of histological, dermoscopic and 'clinical assessment' (not otherwise described) methods to determine response status. The single-

arm trial was the only study that reported how they classified response status (Castellucci, 2021). They used dermoscopy for all lesions and biopsy was used if needed. This was the only study that used standardised measures to assess cosmetic outcomes and severity of safety events. It was the only study that reported the frequency of safety events. All 5 other studies described the types of adverse events that were observed but did not report the frequency. No studies used patient reported outcome measures.

- There was some evolution of the procedure between the earliest (Sedda, 2008) and subsequent publications. The Rhenium-SCT kit was used to apply the radioactive resin in at least 4 studies. Different materials were used to protect the skin (cream only compared to foil or combination) and several versions of the software used to calculate how long to irradiate the lesion were used.
- One study pre-planned to fraction the dose across multiple sessions for lesions on the lips and genitals or for thicker lesions (Cipriani, 2017). Two studies delivered the intervention and reported outcomes after 1 session (Cipriani, 2022; Castellucci, 2021). Three studies delivered multiple sessions for some participants (Carrozzo, 2013; Carrozzo, 2014; Sedda, 2008) but 2 did not state when they happened or whether they were pre-planned or triggered by partial or non-response to previous sessions (Carrozzo, 2013; Sedda 2008). Interpretation of treatment success and relapse is difficult in these studies.
- Some studies successfully treated large lesions but 5 studies commented that some recurrences related to size, depth of lesion and the success of the security margin mapped around the lesion.

- Castellucci (2021) conducted a statistical analysis that showed that grade 3
 acute toxicity was significantly related to the surface area treated and the
 dose.
- Multiple authors on the Cipriani (2022) publication were employees, former employees or medical consultants for OncoBeta. OncoBeta were funded to publish the paper. Cipriani (2017) acknowledges support from OncoBeta, MAVIG GmbH and GBN Systems Gmb but do not state this is a conflict of interest.

Ongoing trials:

Efficacy of Personalised Irradiation With Rhenium-Skin Cancer Therapy
 (SCT) for the Treatment of Non-Melanoma Skin Cancer: A Phase IV,
 Multi-Centre, International, Open-Label, Single Arm Study (EPIC-Skin;
 NCT05135052); Australia; n=210; study completion date May 2024.

Related NICE guidance

Interventional procedures

- NICE's interventional procedures guidance on electrochemotherapy for primary basal cell carcinoma and primary squamous cell carcinoma.
 (Recommendation: special arrangements).
- NICE's interventional procedures guidance on photodynamic therapy for non-melanoma skin tumours (including premalignant and primary non-metastatic skin lesions). (Recommendation: standard arrangements for BCC, Bowen's disease and actinic (solar) keratosis; special arrangements for invasive squamous cell carcinoma).

Technology appraisals

NICE's technology appraisal guidance on cemiplimab for treating advanced cutaneous squamous cell carcinoma.

NICE's technology appraisal on vismodegib for treating basal cell carcinoma

Medical technologies

NICE's medical technologies guidance on Ambulight PDT for the treatment of non-melanoma skin cancer.

Diagnostics

NICE's diagnostics guidance on VivaScope 1500 and 3000 imaging systems for detecting skin cancer lesions

NICE guidelines

NICE guideline on suspected cancer: recognition and referral.

Cancer service guidelines

Improving outcomes for people with skin tumours including melanoma.

Professional societies

Specialist Societies to approach for PEs:

- Association of Cancer Physicians
- British Association of Dermatologists
- British Association of Plastic Reconstructive and Aesthetic Surgeons
- British Nuclear Medicine Society
- Royal College of Radiologists.

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Company engagement

NICE asked companies who manufacture a device potentially relevant to this procedure for information on it. NICE received 1 completed submission. This was considered by the interventional procedures team and any relevant points have been taken into consideration when preparing this overview.

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Methods

NICE identified studies and reviews relevant to epidermal radiotherapy using rhenium-188 paste from the medical literature. The following databases were searched between the date they started to 28th April 2023: MEDLINE, PREMEDLINE, EMBASE, Cochrane Library and other databases. Trial registries and the internet were also searched (see the <u>literature search strategy</u>). Relevant published studies identified during consultation or resolution that are published after this date may also be considered for inclusion.

The following inclusion criteria were applied to the abstracts identified by the literature search.

- Publication type: clinical studies were included with emphasis on identifying good quality studies. Abstracts were excluded if they did not report clinical outcomes. Reviews, editorials, and laboratory or animal studies, were also excluded and so were conference abstracts, because of the difficulty of appraising study methodology, unless they reported specific adverse events that not available in the published literature.
- Patients with non-melanoma skin cancer.
- Intervention or test: Rhenium-188 paste.
- Outcome: articles were retrieved if the abstract contained information relevant to the safety, efficacy, or both.

If selection criteria could not be determined from the abstracts the full paper was retrieved.

Potentially relevant studies not included in the main evidence summary are listed in the section on <u>other relevant studies</u>.

Find out more about how NICE selects the evidence for the committee.

Table 4 literature search strategy

Databases	Date searched	Version/files
MEDLINE (Ovid)	28/04/2023	1946 to April 27, 2023
MEDLINE In-Process	28/04/2023	1946 to April 27, 2023
(Ovid)		
MEDLINE Epubs ahead of	28/04/2023	April 27, 2023
print (Ovid)		
EMBASE (Ovid)	28/04/2023	1974 to April 27, 2023
EMBASE Conference	28/04/2023	1974 to April 27, 2023
(Ovid)		
Cochrane Database of	28/04/2023	Issue 4 of 12, April 2023
Systematic Reviews –		
CDSR (Cochrane Library)		
Cochrane Central	28/04/2023	Issue 4 of 12, April 2023
Database of Controlled		
Trials – CENTRAL		
(Cochrane Library)		
International HTA	28/04/2023	-
database (INAHTA)		

Trial sources searched

- Clinicaltrials.gov
- ISRCTN
- WHO International Clinical Trials Registry

Websites searched

- National Institute for Health and Care Excellence (NICE)
- NHS England
- Food and Drug Administration (FDA) MAUDE database
- Australian Safety and Efficacy Register of New Interventional Procedures Surgical (ASERNIP – S)
- Australia and New Zealand Horizon Scanning Network (ANZHSN)
- General internet search

The following search strategy was used to identify papers in MEDLINE. A similar strategy was used to identify papers in other databases.

MEDLINE search strategy

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21 or 22

23 not 24

(("non melanoma*" or non-melanoma* or nonmelanoma*) adj4 skin adj4 (cancer* 1 or tumo?r* or neoplasm* or carcinoma*)).tw. 5401 NMSC.tw. 1407 2 3 Carcinoma, Basal Cell/ 18639 4 (Basal Cell adj4 (Carcinoma* or Epithelioma*)).tw. 14713 5 (Rodent adj4 Ulcer*).tw. 6 Carcinoma, Squamous Cell/ 140111 7 ((Squamous or Epidermoid or Planocellular) adj4 carcinoma*).tw. 112747 8 (BCC or SCC).tw. 25717 9 (keratinocyte adj4 (cancer* or tumo?r* or neoplasm* or carcinoma*)).tw. 868 10 or/1-9 198634 11 Rhenium/ 2170 12 ((Rhenium adj2 "188") or "Rhenium-188").tw. 283 13 Brachytherapy/ 21695 14 (brachytherap* or curietherap*).tw. 17481 15 13 or 14 25127 16 ((High adj4 dose) or HDR or (Dermo adj4 beta)).tw. 118264 5090 17 15 and 16 (((beta adj4 emitter) or epiderm*) adj4 radioisotope*).tw. 18 19 (radioactive adj4 resin*).tw. 40 20 11 or 12 or 17 or 18 or 19 21 10 and 20 620 22 (Oncobeta or Rhenium-SCT).tw. 2

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Animals/ not Humans/5081619