

Work in progress

The outcome of chronic wounds following hyperbaric oxygen therapy: a prospective cohort study – the first year interim report

Glen C Hawkins, Michael H Bennett, Annelies E van der Hulst

Key words

Chronic wounds, hyperbaric oxygen therapy, research

Abstract

(Hawkins GC, Bennett MH, van der Hulst AE. The outcome of chronic wounds following hyperbaric oxygen therapy: a prospective cohort study – the first year interim report. *Diving and Hyperbaric Medicine*. 2006; 36: 94-98.)

Introduction: The treatment of chronic wounds is a major health cost. This study is an ongoing prospective cohort looking at the effects of hyperbaric oxygen therapy (HBOT) on the healing of chronic wounds.

Methods: Data are being collected from patients presenting to hyperbaric facilities in Australia with chronic (> 3 months' duration) non-irradiated wounds, including details of aetiology, wound characteristics and possible predictors of wound healing. Participants are being enrolled whether or not a decision was made to treat with HBOT. Assessments are performed at the end of the course of HBOT and at one, six and 12 months after hyperbaric treatment. The aim is to quantify the proportion healed and to identify any significant predictors for wound healing.

Results: There are 110 participants included in this analysis with 88 receiving HBOT. Excluding the miscellaneous aetiologies, at the end of treatment 52.3% of patients had a 'good' outcome to the wound, increasing to 64.1%, 91.7% and 78.2% at one, six and 12 months respectively. Logistic regression for participants with diabetic wounds suggests that wound area, chronicity and transcutaneous oxygen readings on room air combine to produce a statistically significant model for prediction of wound healing at one month after treatment.

Conclusions: This ongoing cohort study suggests that HBOT is highly associated with the healing of chronic wounds in the patients in this study. The wound area at presentation, the duration of the wound and the transcutaneous oxygen pressure on air may predict the likelihood of a chronic wound in diabetic patients healing by one month after treatment.

Introduction

Chronic wounds are defined as an interruption in the continuity of the skin where conventional treatment has not achieved healing within a reasonable time (e.g., 3 months).¹ Such wounds are an increasing burden on healthcare systems throughout the world. Studies have shown a prevalence in hospital patients of up to 24%, and 2% of the general population have some form of chronic wound at any one time.²⁻⁵ This creates a significant financial burden on funding agencies with costs exceeding several billion dollars per year.⁶⁻⁸ There are compelling reasons to deliver treatment modalities that are cost effective to individuals with such wounds.

Treatment regimens for chronic wounds are multi-modal but have been traditionally of two types: specific treatments designed to reduce the effect of the underlying disease (such as tight glycaemic control in diabetics and compression bandages in venous insufficiency), and wound environment optimisation dressings (e.g., hydrocolloid gels and antibacterial impregnated dressings).

The rationale for adjunctive hyperbaric oxygen therapy (HBOT) in chronic wound care is the premise that the

underlying problem in many of these wounds is hypoxia. While acute wounds require low oxygen tensions, low pH and a high lactate load to initiate angiogenesis and wound healing,^{9,10} later phases of healing are critically dependent on oxygen, e.g., fibroblastic collagen deposition and macrophage bacteriocidal activity.¹¹⁻¹³ It has been suggested that the stimulus for healing is a rapid drop in the partial pressure of oxygen from surrounding healthy tissue to the wound. In chronic wounds there is a much more gradual drop across the wound margin and this may inhibit healing significantly.¹⁴

The Medicare Services Advisory Committee (MSAC) was established in 1997 to advise the Australian Minister for Health and Ageing on the safety and cost effectiveness of new medical technologies and procedures, and to make recommendations for funding under the Medicare Benefits Scheme.¹⁵ One such review was initiated into the provision of HBOT, and in 2001 MSAC recommended that a properly conducted prospective trial should be undertaken on the treatment of chronic non-irradiated wounds with HBOT. This report presents the first results of a prospective cohort of patients enrolled since June 2004.

Methods

All hyperbaric facilities in Australia and New Zealand were invited to participate in the study. There are currently 13 such chambers treating patients for chronic wounds. Three facilities (Prince of Wales Hospital, Sydney (POW), Wesley Hospital, Brisbane (WES) and Royal Hobart Hospital, Hobart (HOB)) have been able to start in the first year and three other facilities are currently awaiting ethics approval or the conditions of their approval have not permitted submission of data in the first year. No enrolments were undertaken prior to obtaining approval from the relevant local ethics committee. Data were collected on each patient by each facility and an identifying number was included in the data collection sheet that allowed each centre to follow individual patients' progress through the four reporting stages. At the collection centre (POW) each individual patient was given a code number to identify the enrolling centre they were from and order of enrolment. Analysis was performed on the POW code numbered datasheets entered into a computer database.

PATIENT SELECTION

All patients referred to a hyperbaric facility for assessment of one or more chronic wounds (present for more than three months) are eligible for inclusion, regardless of prior therapy. Patients considered unsuitable for HBOT due to the presence of a contra-indication, inadequate prior therapy or anticipated lack of response are therefore also eligible for enrolment. Acute (including extensive debridement within three months) wounds and those due to irradiation tissue injury are excluded from the study. The study authors did not determine assessments or impose HBOT schedules on the study centres as no definitive treatment schedule has been shown to be better than any other. We also feel that this allows the study to reflect 'true practice' in the hyperbaric field, with the variety of equipment available in each centre also influencing clinical practice.

DATA COLLECTION

A standardised datasheet was developed that recorded demographic data, possible factors contributing to poor wound healing, treatment up to the date of assessment, subsequent hyperbaric treatment (if performed) and outcome immediately following HBOT as well as at one month, six months, and twelve months after HBOT.

Data were collected on a Filemaker Pro™ database (v7, Filemaker Inc, Santa Clara, California). Each patient was given a designated identification number for tracking through the four assessment times. Each facility is responsible for data collection on the subjects enrolled at that facility. Units are being encouraged to use all means at their disposal to locate missing subjects, including direct contact and through their local medical services and family members. Each unit was reminded at the appropriate times when a patient was due for re-assessment.

Table 1
Clinical outcome scores for wounds

Clinical description	Category	Outcome class	Outcome
Deceased	1	No benefit	BAD
Nil benefit ± major amputation	2		
Minimal benefit + minor amputation	3	Some benefit	
Improved + minor amputation	4		
Substantially healed	5	Healed	GOOD
Healed	6		

OUTCOMES

Outcomes are scored on a six-point scale originally developed by Dr Harry Oxer (Davis FM, personal communication, 2003) at Fremantle Hospital Hyperbaric Medicine Unit. However, for this interim assessment we categorised all outcomes as either 'good' or 'bad' as shown in Table 1. We have specifically placed amputations of any sort into the 'bad' outcome category because this seems an appropriately conservative approach to assessing the effectiveness of HBOT. While an amputation may indicate a good outcome (e.g., saved limb but lost toes) or poor outcome (e.g., superficial foot ulcer but lost toe) there may be no clear indication which is the case for any individual patient. In addition, any amputation will alter the location and dynamics of the wound – essentially converting a chronic wound into an acute surgical wound. We planned an annual analysis for reporting back to the contributing units.

WOUND CATEGORIES

Wounds were allocated to one of four main aetiological categories for analysis – diabetic (DM), peripheral vascular disease (PVD), venous disease and miscellaneous (including vasculitic and auto-immune diseases). Because the miscellaneous group contains highly diverse aetiologies in very small numbers, no analysis of the fate of this group has been undertaken in this interim report. Similarly, this report does not compare the chance of a good outcome with and without HBOT because of the small numbers in the non-HBOT group.

STATISTICS

No sample size calculations were performed for this study, as it is an ongoing opportunistic cohort study. We performed a descriptive statistical analysis and a backward stepwise

Table 2
Number of patients and mean ages (with range) of all patients enrolled in the study (whether they had HBOT or not) by aetiology

Aetiology	Number	Average age years (range)	% total wounds
DM	46	66.4 (42–89)	41.8%
PVD	27	73.9 (37–91)	24.5%
Venous	18	69.2 (43–87)	16.4%
Miscellaneous	19	61.7 (11–83)	17.3%
Total	110	67.8 (11–91)	100%

(DM – diabetes mellitus; PVD – peripheral vascular disease; venous – venous insufficiency)

logistic regression analysis on each aetiological group for factors that may predict wound outcome. This was done in order to develop a predictive model for wound healing after HBOT. All calculations were performed using StatsDirect v2.4.5 (StatsDirect Ltd., StatsDirect statistical software. <<http://www.statsdirect.com>> England: 2002).

Results

There were 110 patients enrolled in the study of whom 88 received hyperbaric oxygen treatment. Sixty-seven (61%) were males and 43 (39%) females. The group receiving hyperbaric oxygen had 54 (61%) males with an average age of 67.2 years. The group that did not receive hyperbaric oxygen had 13 (59%) males with an average age of 70.4 years. The breakdown by aetiology is shown in Table 2.

HYPERBARIC TREATMENT

The average number of treatments for the patients receiving HBOT was 24.4 (range 1–70). The average number of treatments, for each aetiology, is given in Table 3, while the overall frequency distribution is shown in Figure 1.

OUTCOME DATA

Figure 2 shows the percentage of people in each aetiological group with a ‘good’ outcome (Scores 5 and 6, Table 1). Overall, immediately after the HBOT course, 52.3% of all aetiological groups combined had a ‘good’ outcome and this proportion increased to 64.1%, 91.7% and 78.2% at one, six and twelve months respectively. These data suggest that diabetic wounds improve most rapidly following HBOT, with venous wounds catching up at one month and arterial wounds at six months. At the time of the final draft of this interim report, we have follow-up data on 60% of those enrolled, and 43% at one year. Because these data sets are substantially incomplete, they will be reported in future annual analyses.

Because of the small numbers enrolled in the study, regression analysis for outcome was possible only for those patients with diabetes mellitus. We performed univariate analysis and a logistic regression for ‘good’ outcome at the end of HBOT and at one month after HBOT with the potential predictors being gender, duration of wound (months), wound area (cm²), transcutaneous partial pressure of oxygen (P_{tc}O₂) in air (mmHg), and P_{tc}O₂ on 100% oxygen at 1 ATA for 10 minutes (mmHg). Neither at the end of HBOT nor at one month follow up were there any significant predictive factors identified on univariate analysis. Stepwise logistic regression for healing at the end of the HBOT course did not produce a useful model. However, analysis at one month follow up suggested the following model was predictive of healing:

$$\text{Log (OR)} = 2.30 - (0.09 * \text{TWA}) - (0.11 * \text{DUR}) + (0.06 * \text{P}_{tc}\text{O}_2)$$

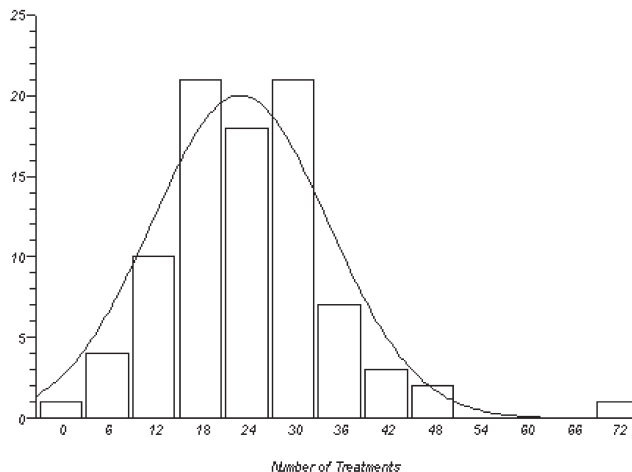
where OR = odds ratio, TWA = total wound area in cm², DUR = duration of wound in months and P_{tc}O₂ = transcutaneous partial pressure of oxygen on air at 1 ATA in mmHg.

Table 3
Number and treatment averages of those patients who had HBOT by aetiology

Aetiology	Number of patients	Mean number of treatments (SD)
DM	40	23.4 (10.2)
PVD	20	24.7 (8.46)
Venous	13	24.2 (9.49)
Miscellaneous	15	27.0 (14.63)
Total	88	24.4 (10.53)

(DM – diabetes mellitus; PVD – peripheral vascular disease; venous – venous insufficiency)

Figure 1
Frequency distribution (with curve of best fit) of number of treatments for patients receiving HBOT



This model suggests that, at presentation, wound healing is negatively impacted by increased wound area, duration of wound and a lower $P_{tc}O_2$ in air. For example, using this model we would predict that for a wound 7 cm² in area and of 10 months' duration with a resting $P_{tc}O_2$ (in air) of 30 mmHg, the odds of healing at one month after completion of a course of HBOT are nearly 11 to one (odds ratio 10.7).

Discussion

This study suggests that we can expect 50% of chronic wounds to heal by the end of a course of HBOT and up to 90% of wounds to be healed at six-month follow up. These wounds have all persisted for at least three months at presentation despite comprehensive wound care, and we believe this represents a real and important clinical benefit. Although 50% may not seem a particularly large proportion, given the population of Australia (20,404,617)¹⁶ and an assumed prevalence for chronic wounds of 1%, this represents over 100,000 people who could potentially have a good outcome from HBOT.

Hyperbaric facilities have been treating chronic wounds for several years but there has been very little high-quality clinical research evidence to demonstrate the effectiveness of HBOT. A recent Cochrane meta-analysis on the efficacy of HBOT for chronic wounds included four randomised controlled trials (RCTs).¹ Three of these studies enrolled diabetic foot-ulcer patients and one enrolled patients with venous ulcers. There were no RCTs on the effects of HBOT on arterial ulcers. These RCTs suggest that there was a benefit in having HBOT for diabetic and venous ulcers but a larger, multi-centre study is required.

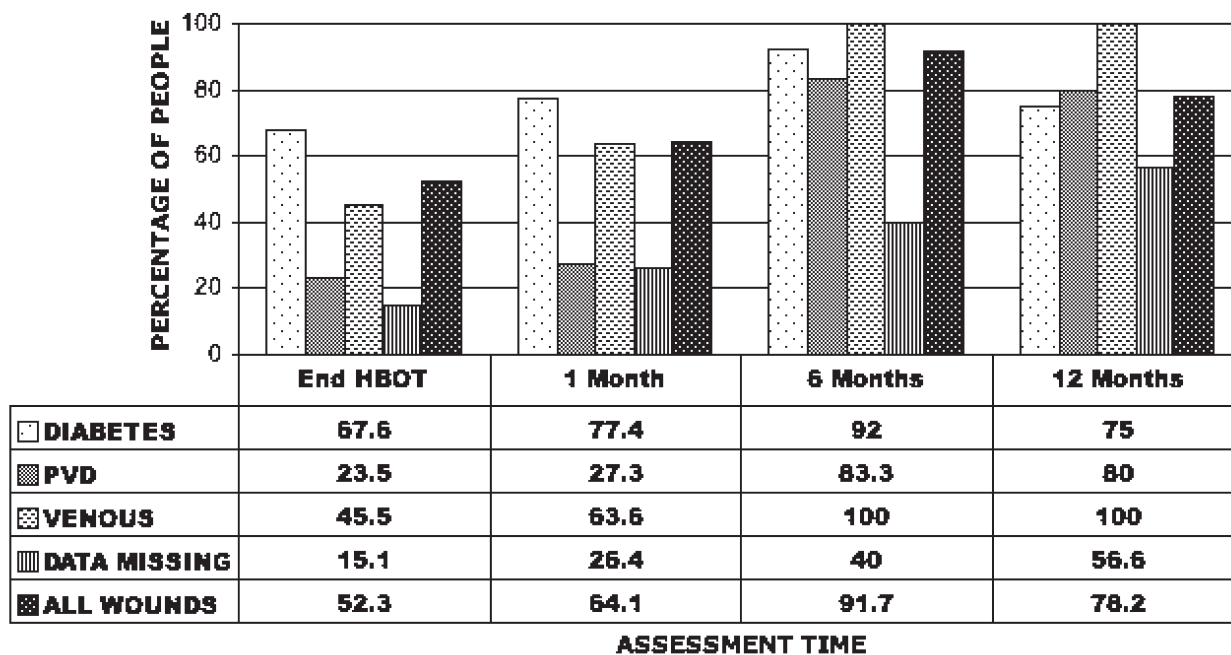
Because of the small number of patients enrolled in this study who did not receive HBOT (n = 22), we have not reported the fate of ulcers in that subgroup of patients in this analysis. While the reasons they were thought unsuitable for HBOT were not always clear, we hypothesise most of them had normoxic $P_{tc}O_2$ levels, failed to adequately respond to oxygen challenge with an increase in $P_{tc}O_2$ or declined to undergo therapy. We intend to more fully address this group in our next report.

There are differences in response to HBOT between aetiologies. Diabetic wounds have a faster resolution (higher percentage of those with a good outcome at the end of treatment) and appear to have a higher chance of a good outcome for the first few months after treatment. The estimated difference narrows rapidly and at six months there is very little between the three main aetiologies.

Logistic regression suggests that even with this small data set, we are able to show that features such as total wound area, duration of wound and $P_{tc}O_2$ at the wound site breathing air at 1 ATA are significant predictors of the proportion of wounds that will heal. We hope that as the data set grows, the regression model will become increasingly predictive of those wounds that can be expected to heal. This would have useful clinical applications for the selection of candidates for HBOT.

There are several limitations to the interpretation and applicability of this study. First among them is the loss of data as the study progresses. This is largely due to inability to contact some patients for follow up, despite considerable efforts to do so. Currently 56.6% of patients' data are lost

Figure 2
Number of people with 'good' outcome after HBOT as a percentage of total people receiving HBOT



at the 12-month assessment time reflecting difficulties in following up patients out to this time period. We have attempted to address this with better patient tracking and by co-ordinating the follow up of patients with active reminders to the collection centres involved. Some apparent loss of data is in fact due to significant numbers of participants who have not yet reached the final assessment time.

Another significant limitation for this cohort study is the relatively small number of participants who had chronic wounds but did not receive HBOT. Financial considerations have made it impractical to improve the methodology of this study by the active recruitment of a comparison cohort of participants for whom hyperbaric referral has not been considered. Such a study is beyond our means at this time but remains highly desirable.

In conclusion, we have reported the first 110 patients of an ongoing prospective study. Our results suggest that a clinically important proportion of patients can expect a good outcome by one month after the completion of hyperbaric therapy. We continue to collect data prospectively and hope to generate a useful predictive model by which to identify those patients in whom HBOT is appropriate. We believe that this study is important in helping to better define the role of hyperbaric oxygen in these patients.

Acknowledgements

The authors wish to thank the staff at Royal Hobart Hospital, Wesley Hospital and Prince of Wales Hospital for the collection of the wound care data.

References

- 1 Kranke P, Bennett M, Roeckl-Wiedmann I, Debus S. Hyperbaric oxygen therapy for chronic wounds. *Cochrane Database of Systematic Reviews*. 2004 (2): CD004123.
- 2 Stausberg J, Kroger K, Maier I, Schneider H, Niebel W, Interdisciplinary Decubitus Project. Pressure ulcers in secondary care: incidence, prevalence, and relevance. *Adv Skin Wound Care*. 2005; 18: 140-5.
- 3 Lahmann NA, Halfens RJ, Dassen T. Prevalence of pressure ulcers in Germany. *J Clin Nurs*. 2005; 14: 165-72.
- 4 Chauhan VS, Goel S, Kumar P, Srivastava S, Shukla VK. The prevalence of pressure ulcers in hospitalised patients in a university hospital in India. *J Wound Care*. 2005; 14: 36-7.
- 5 Graham ID, Harrison MB, Nelson EA, Lorimer K, Fisher A. Prevalence of lower-limb ulceration: a systematic review of prevalence studies. *Adv Skin Wound Care*. 2003; 16: 305-16.
- 6 Albert S. Cost-effective management of recalcitrant diabetic foot ulcers. *Clin Podiatr Med Surg*. 2002; 19: 483-91.
- 7 Bennett G, Dealey C, Posnett J. The cost of pressure ulcers in the UK. [see comment]. *Age Ageing*. 2004; 33: 230-5.
- 8 Gordois A, Scuffham P, Shearer A, Oglesby A, Tobian JA. The health care costs of diabetic peripheral neuropathy in the US. *Diabetes Care*. 2003; 26: 1790-5.
- 9 Knighton DR, Hunt TK, Scheuenstuhl H, Halliday BJ, Werb Z, Banda MJ. Oxygen tension regulates the expression of angiogenesis factor by macrophages. *Science*. 1983; 221: 1283-5.
- 10 Jensen JA, Hunt TK, Scheuenstuhl H, Banda MJ. Effect of lactate, pyruvate, and pH on secretion of angiogenesis and mitogenesis factors by macrophages. *Lab Invest*. 1986; 54: 574-8.
- 11 Hunt TK, Pai MP. The effect of varying ambient oxygen tensions on wound metabolism and collagen synthesis. *Surg Gynecol Obstet*. 1972; 135: 561-7.
- 12 Niinikoski J, Grisliis G, Hunt TK. Respiratory gas tensions and collagen in infected wounds. *Ann Surg*. 1972; 175: 588-93.
- 13 Hohn DC, MacKay RD, Halliday B, Hunt TK. Effect of O₂ tension on microbicidal function of leukocytes in wounds and in vitro. *Surg Forum*. 1976; 27: 18-20.
- 14 Davis JC, Hunt TK, editors. *Problem wounds: the role of oxygen*. New York: Elsevier Science Publishing Co.; 1988.
- 15 Committee MSA. MSAC-About Us. 2005 7/10/2003 [cited 2005 28/8/2005]; Available from: <<http://www.msac.gov.au/bckgrd.htm>>
- 16 Australian Bureau of Statistics: Projected population. 2005 [cited 2005 18/9/2005]; Available from: <www.abs.gov.au>

Glen C Hawkins, MBChB, BmedSc, FANZCA, is Associate Lecturer at the University of New South Wales, and Hyperbaric Fellow, Department of Diving and Hyperbaric Medicine, Prince of Wales Hospital, Sydney, Australia
Michael H Bennett, MBBS, DA, FANZCA, MM (ClinEpi), DipDHM, is Senior Lecturer at the University of New South Wales, and Hyperbaric Consultant, Department of Diving and Hyperbaric Medicine, Prince of Wales Hospital, Sydney, Australia
Annelies E van der Hulst, is a medical student at the Medical Facility, University of Groningen, Groningen, The Netherlands

This paper is based on Dr Hawkins' thesis for his SPUMS Diploma, awarded earlier this year.

Address for correspondence:

*Dr Glen Hawkins
 Department of Diving and Hyperbaric Medicine
 Prince of Wales Hospital
 Barker Street, Randwick, NSW 2031
 Australia
 Phone: +61-(0)2-9382-3880
 Fax: +61-(0)2-9382-3882
 E-mail: <hawkeye@swiftdsl.com.au>*