The Role of Oxygen in Wound Healing

HBOT, TO, and TCOT all play critical treatment roles.

BY WINDY COLE, DPM

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Without oxygen, there can be no life. The attainability of oxygen, the most capable electron acceptor on our planet, has allowed organisms to develop mechanisms of highly efficient energy production, which has led to the evolutionary development of aerobic life forms from the very first multicellular organisms to early vertebrates to the patients we care for today. You may recall memorizing the steps of the Krebs cycle during your general biology studies in high school. It is through this process of aerobic respiration that most living things generate energy. In aerobic organisms, the mitochondria in cells utilize oxygen (O₂) as the final electron acceptor to synthesize high-energy adenosine triphosphate (ATP) from adenosine diphosphate (ADP) (Figure 1). ATP is the fuel needed for all human cellular processes to occur. In the absence of adequate levels of O₂, cells convert to anaerobic metabolism, which is less advantageous. Hypoxia, or lack of adequate oxygen levels, will ultimately result in inadequate ATP production, tissue acidosis, and cellular dysfunction.

Understanding how wounds heal enables clinicians to apply the appropriate treatment and management techniques at each phase to support the healing process. The wound healing cascade is composed of four overlapping phases: hemostasis, inflammation, repair, and...
remodeling. Although the stages of wound healing are linear, wounds can progress backward or forward depending on internal and external patient conditions. Throughout these stages wounded tissues exhibit an increased energy demand leading to a hypermetabolic state.\(^1\) Therefore, wound healing is heavily reliant on the presence of adequate oxygen levels within the injured tissues. Oxygen is essential to multiple wound healing processes including oxidative killing of bacteria, cellular signaling and proliferation, collagen deposition, and angiogenesis.\(^3\)

**The Antimicrobial Effects of Oxygen**

Controlled inflammation is beneficial to wound healing. When flesh is injured (Figure 2) chemical signals such as histamine are released triggering tissue inflammation. Local blood vessels dilate and become porous to allow phagocytic cells such as macrophages and neutrophils to migrate to the wounded tissue.\(^2\) These phagocytic cells help to break down denatured extracellular matrix components, debris, and consume harmful bacteria.

Increased oxygen levels can speed the rate of collagen deposition.

Oxygen plays a vital part in controlling bacterial burden during the inflammatory phase of wound healing. Within the membranes of neutrophils, NADPH oxidase generates superoxide (\(2O_2\)). Through the process of respiratory burst, \(2O_2\) is eventually converted to hypochlorous acid (HOCL). Reactive oxygen species (ROS) such as HOCL are responsible for the oxidative killing of bacteria and serve as the body’s natural protection against infection.\(^2\) Without adequate local tissue oxygenation, the respiratory burst is impaired, resulting in increased tissue susceptibility to infection. Certain disease conditions can also impair the formation of ROS. Chronic granulomatous disease (CGD), a primary immunodeficiency that affects phagocytes, is one such condition. CGD is caused by a mutation in NADPH oxidase.\(^4\) Lack of ROS formation will predispose patients to an increased risk of bacterial infection.\(^4\) Breathing air with reduced oxygen levels can even affect bacterial levels in wounds.

Hohn et al. conducted a study in which they determined that skin wounds of rabbits exposed to air containing low oxygen concentrations had more elevated levels of Staphylococcus aureus than skin.
Oxygen (from page 94)

wounds of rabbits exposed to air containing high levels of oxygen. Local tissue oxygen levels are a pivotal determinant in the overall wound microbial burden and probability of infection. Support of the body’s innate immunogenic response to bacterial contamination with additional oxygen supplementation may decrease clinically significant wound infections, abscess formation, osteomyelitis, and amputations.

Growth Factor Regulation

The transition between wound healing phases is managed and regulated by biologically active substances called growth factors. These polypeptides control the growth, differentiation, and metabolism of cells. Fibroblasts migrate into the wound and replicate in response to growth factors released during inflammation. The essential role of fibroblasts during the reparative phase of wound healing is the secretion of the polysaccharide gel that makes up the extracellular matrix of newly formed connective tissue. ROS in low levels play a role in growth factor release. Acute hypoxia stimulates growth factor production, but chronic hypoxia inhibits or eliminates it.

Siddiqui and colleagues investigated the proliferation of fibroblasts in varying environments. They compared cells grown in a hypoxic environment (1% O2) vs. standard culture conditions (20% O2). The results of the study showed cell proliferation was three times slower with exposure to 1% oxygen versus 20% oxygen. The investigators concluded that chronically low levels of local oxygen were detrimental to growth factor production and led to decreases in fibroblast proliferation.

Augmentation of local tissue oxygen levels may therefore enable increased fibroblast migration and proliferation in wounded tissue to help speed healing rates and prevent wound chronicity.

The formation of the stable collagen fiber triple helix is O2 dependent (Figure 3). Without oxygen, the hydroxylation of the proline and lysine side chains do not allow for the proper assembly of the triple helix structure and the resulting procollagen is non-functional. It is these extracellular cross-linkages that are ultimately responsible for the tensile strength needed in prolonged wound healing.

Angiogenesis levels of local oxygen were detrimental to growth factor production and led to decreases in fibroblast proliferation. Augmentation of local tissue oxygen levels may therefore enable increased fibroblast migration and proliferation in wounded tissue to help speed healing rates and prevent wound chronicity. Oxygen levels can also affect the rate of collagen deposition and possibly decrease the overall time to wound healing. Hunt and colleagues used a rabbit model to track the rate and density of collagen formation with changes in oxygen levels. The results demonstrated that exposure to hyperoxic environments accelerated collagen synthesis. Oxygen continues to be an important element throughout the remodeling stage of wound healing as continued collagen deposition during this time is key. The caliber of the collagen fibers will ultimately determine the quality of scar tissue formation. The more robust the collagen, the better the long-term healing rates of the injured tissue will be.

Angiogenesis

The formation of new blood vessels, or angiogenesis, is another critical step in the process of tissue repair. Angiogenesis begins as growth factors stimulate endothelial cells to migrate and proliferate through a healing wound. This neovascularization begins through budding of the existing capillary network to provide new channels for active cells, nutrients, and oxygenated hemoglobin to travel to the wounded tissues (Figure 4). Vascular endothelial growth factor (VEG-F) is a major angiogenic stimulus. Initially, hypoxia acts as a stimulus to

Figure 3: The formation of the stable collagen fiber triple helix.

Collagen Synthesis

As the neutrophil count decreases in wounded tissue, the number of fibroblasts increases. After the extracellular matrix is laid down, fibroblasts begin to secrete collagen fibers. Collagen is the most abundant protein produced in the human body. The overall structure and integrity of human skin is proportionally related to the amount and quality of collagen found within the tissues. Therefore, deposition of collagen is a fundamental step in the wound healing process. The formation of this natural scaffold is essential for new tissue growth and repair. Strong collagen tendrils develop through the aggregation of mature collagen fibers with a tight triple helical structure.
Oxygen (from page 95)

VEG-F. However, prolonged hypoxia inhibits VEG-F formation and function and obstructs neovascularization. Supplementary oxygen wound therapy has been shown to increase VEG-F levels, thus stimulating angiogenesis. The most common forms of supplementary oxygen therapy available in the wound care space are hyperbaric oxygen therapy (HBOT) and topical oxygen therapy (TOT). The mechanism of action of these therapies varies greatly.

The effects of HBOT are based on the gas laws. Patients enter a chamber where they breathe 100% O₂ while exposed to increased atmospheric pressure, typically two to three times atmospheric absolute (ATA). It is theorized that HBOT improves wound healing rates by amplifying oxygen gradients along the periphery of ischemic wounds. In contrast, TOT is a low-pressure treatment. A device delivering normoxic oxygen is directly applied to the wound. There is no increased inspiration of oxygen and the effects of TOT are localized in comparison to HBOT.

A 2008 study by Gordillo, et al. evaluated outcomes in 1,854 outpatient wound clinic patients who were screened for non-randomized enrollment into either hyperbaric oxygen therapy treatment or topical oxygen for the treatment of their chronic wounds. The investigators determined that there were no significant changes in wound measurements in the HBOT patient group. However, the TO treatment group did exhibit a noticeable decrease in overall wound dimensions. Tissue biopsies were obtained from wounds in both treatment arms. There was an increase in VEG-F found within the TO group. Overall, the investigators concluded that TO showed better wound healing benefits compared to HBOT.

Acute hypoxia stimulates growth factor production, but chronic hypoxia inhibits or eliminates it.

Increasing O₂ Levels in Wounded Tissue

From inflammation through tissue remodeling, oxygen plays a key role in the essential stages of wound healing. Despite this critical need for oxygen, levels are frequently insufficient in patients with chronic wounds due to a variety of systemic disease states causing poor circulation, inactivation of growth factors, and cellular senescence. Low levels of oxygen in the wounded tissues will prolong healing. Therefore, it stands to reason that supplying additional oxygen to wounds may promote healing.

The use of supplemental oxygen for wound healing can be traced back to the 1960s to the clinical implementation of hyperbaric oxygen (HBO). Around this same time topical oxygen (TO) was also introduced for the treatment of chronic wounds. By way of research and clinical studies, HBO gained wide acceptance within the medical community as a viable adjunct to wound care therapy. TO has only gained ground as a potential wound healing therapy in the last decade. Local delivery of oxygen to treat wounds has been used with varying success. In a 2015 article by Dissemond and colleagues, evidence of the effectiveness of topical oxygen therapies for wound healing was reviewed. The group concluded that although no singular topical oxygen therapy approach is widely used universally throughout the wound healing community, there is a growing level of evidence suggesting its effectiveness as an adjunctive therapy to speed wound healing outcomes.

It has been the author’s experience that continuous topical oxygen therapy (TCOT) offers an effective non-invasive chronic wound treat-
of skin breakdown and complicates wound healing while elevating the likelihood of recurrence.

Previous wound management had included multilayer compression bandaging, antimicrobial creams, silver alginates, collagen, and amniotic tissue grafts. Unfortunately, the wound failed to show significant signs of healing. A full wound assessment was performed including ABI which was 1.01. Upon initial intake, the wound measured 3.1 cm x 3.2 cm x 0.1 cm (Figure 8). Significant psoriatic plaques and hemosiderin staining were evident on the patient’s legs.

Near-infrared spectroscopy imaging was performed to determine the functional perfusion and oxygen saturation of the wound tissues. At baseline, the wound NIRS oxygenated hemoglobin level was 58% (Figure 9). TCOT was initiated at this visit along with standard of care consisting of a secondary foam dressing covered with a multilayer compression bandage.

Weekly wound assessments, including wound measurement, were performed in clinic. After three weeks of therapy, there were active signs of robust new granulation tissue development as well as new islands of epithelial tissue throughout the wound base (Figure 10). At the three-week mark, the wound NIRS oxygenated hemoglobin level increased to 78% (Figure 11). The wound continued to progress in a healing trajectory over the next few weeks with a continuous reduction in documented wound size. Complete wound closure was achieved by week 9 of therapy (Figure 12). The patient reported no issues with managing the device in between clinic visits and was thrilled that after a year of suffering from a VLU, he had healed in nine weeks of oxygen therapy.

**Discussion**

The process of wound healing begins at the very moment of tissue injury. As detailed in this article, oxygen
is essential throughout the phases of wound healing. It is an important biomarker in determining the healing potential of a wound. The oxygen gradient in wounded tissue is unequal. Supply may not meet demand. This is especially true in patients suffering from systemic conditions such as diabetes and venous disease. Although the etiology of non-healing chronic wounds is multi-factorial, hypoxia is a common component in a vast majority of cases.

Healing wounds characteristically have a limited inflammatory phase, high cellular mitogenic activity, robust neovascularization, intact functional extracellular matrix, strong new collagen deposition, and increased tissue tensile strength; whereas chronic wounds display a prolonged inflammatory phase, low mitogenic activity, a dysfunctional matrix, senescent cells, and excessive scarring and contracture. Oxygen levels in the tissue are a major contributory factor in determining the wound’s clinical pathway. Hypoxia is a common limitation to healing in chronic wounds of varying etiologies. Maintaining adequate levels of oxygen in wounded tissues continues to be a challenge.

For many years, hyperbaric oxygen therapy (HBOT) has been employed to deliver high-pressure 100% oxygen to the tissue in the hopes of increasing wound oxygenation. HBOT relies on adequate arterial perfusion in order to transfer oxygen to the wounded tissue. The advent of newer technologies can now enable small, portable battery-powered oxygen generators to allow continuous oxygen therapy (TCOT) to be applied directly to the wound base at normospheric pressure. TCOT is a novel treatment that may potentially be applied to a broader cross-section of patients to improve long-term wound healing outcomes. Each of these modalities has both pros and cons. It is imperative that clinicians choose the appropriate treatment course based on the needs of the patient. Most commonly, successful wound healing occurs as a culmination of multiple evidence-based therapeutic modalities working in conjunction with each other.

It has been the author’s experience that utilizing NIRS to monitor and track tissue oxygenation has proven to be an effective predictor of wound healing. By incorporating this near-infrared imaging technology, the limitations typically found with standard NIVS have been eliminated. In the patient cases presented in this manuscript, the NIRS was able to track changes in the oxygen levels of wound tissues with more precision than ABI alone. The changes noted in wound tissue oxygenation exhibited in these case studies also had better correlation with the weekly clinical appearance of the wound. Using treatments that maintain oxygen levels can decrease median time to heal in chronic recalcitrant wounds of multiple etiologies. Using non-invasive devices such as NIRS to track wound oxygen levels will help clinicians determine the appropriate therapeutic pathway for each individual patient week to week.

Oxygen is vital throughout the wound healing process, especially in the inflammatory and proliferative phases. Research suggests that patient supplementation with oxygen could enhance bacterial killing, increase growth factors, expedite collagen deposition, and improve angiogenesis. Long-term studies are needed to determine if supplemental oxygen therapy can also reduce surgical site infection rates and increase wound tensile strength. The importance of oxygen is often overlooked. Adequate supply does not just come from the air we breathe. Many different supplemental oxygen therapeutic modalities are...
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commercially available and more are entering into the market. Clinicians should keep up on the ever-changing data regarding supplemental oxygen therapy in wound care. PM

References

14. Dr. Cole is an Adjunct Professor and Director of Wound Care Research at Kent State University College of Podiatric Medicine. She also serves as Director of Wound Care Services for Cleveland Regency East Hospital and is the Medical Director at University Hospitals Ahuja Wound Care Center. She is board certified by the American Board of Podiatric Surgery. Her practice focus is on advanced wound care modalities and regenerative medicine. She has published on these topics and speaks nationally and internationally on limb preservation and wound care.

CME EXAMINATION

SEE ANSWER SHEET ON PAGE 101.

1) Which is a true statement about the Krebs cycle?
   A) It is an anerobic process.
   B) ADP is the fuel needed for all human cellular processes to occur.
   C) Oxygen is the final electron acceptor in the synthesis of ATP.
   D) Hypoxia does not affect this process.

2) Why is oxygen important for bacteria management in wounded tissue?
   A) Without oxygenation the respiratory burst is impaired.
   B) Devitalized tissue may remain in the wound bed making bacterial colonization more likely.
   C) Lack of oxygen has been shown to increase tissue susceptibility to infection.
   D) All of the above.

3) All statements about growth factor regulation are true except:
   A) Acute hypoxia decreases growth factor production, but chronic hypoxia stimulates it.
   B) Transition between wound healing phases is managed by growth factors.
   C) Fibroblast migration is in direct response to growth factor release.
   D) Low levels of local oxygen are detrimental to growth factor production.

4) Collagen and procollagen formation differ in what way?
   A) Unlike procollagen, collagen fibers exhibit a tight triple helical structure.
   B) Fibroblasts secrete collagen but not procollagen.
   C) Hydroxylation of the proline and lysine side chains is not required to form procollagen.
   D) Both a and c.

5) How does the level of local oxygen influence the rate of wound healing?
   A) Increased oxygen levels can speed the rate of collagen deposition.
   B) Hyperoxic environments decrease collagen synthesis.
   C) The Hunt and colleagues rabbit study showed that breathing air with increased oxygen had little effect on wound healing.
   D) Oxygen levels do not impact the rate of wound healing.

6) All statements are true regarding angiogenesis except:
   A) Initially hypoxia acts as a stimulus to angiogenesis.
   B) Angiogenesis begins through budding of the existing capillary network.

Continued on page 100
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**CME EXAMINATION**

7) The Gordillo study comparing hyperbaric oxygen therapy to topical oxygen concluded all of the following except:
   A) No significant changes in wound measurements were seen in the HBOT patient group, but the TO treatment group had a noticeable decrease in wound dimensions.
   B) There was an increase in VEG-F found within the wounds of the TO group.
   C) The TO group exhibited better overall wound healing compared to the HBOT group.
   D) HBOT and TO had similar effects on wound healing.

8) How do systemic disease states contribute to decreased rates of wound healing?
   A) Patients with co-morbidities are more non-compliant.
   B) Diabetes and PVD do not complicate wound healing.
   C) Chronic disease states commonly cause decreased circulation, inactivation of growth factors, and cellular senescence.
   D) Co-morbid states increase patients’ physical fitness.

9) Which statement is correct regarding HBOT?
   A) The origins of HBOT can be traced back to 1860.
   B) HBOT has gained acceptance within the medical community as a viable adjunct wound care therapy.
   C) HBO is inexpensive and can be used in a broad cross-section of patients.
   D) There are no known side-effects of HBOT.

10) Chronic wounds display which of the following?
    A) A prolonged inflammatory phase.
    B) Low mitogenic activity.
    C) A dysfunctional matrix.
    D) All of the above.

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