Equine Wound Management
To our families and friends.

To Aotearoa, for a warm welcome during the preparation of this book – Mauruuru koutou.
Equine Wound Management

Third Edition

Edited by

Christine Theoret, DMV, PhD, Diplomate ACVS
Director, Comparative Veterinary Tissue Healing Laboratory (http://theoretlab.com/index.php/en)
Professor, Département de biomédecine vétérinaire
Faculté de médecine vétérinaire
Université de Montréal
Québec, Canada

Jim Schumacher, DVM, MS, Diplomate ACVS, MRCVS
Professor, Equine Surgery
Department of Large Animal Clinical Sciences
College of Veterinary Medicine
University of Tennessee
Knoxville, Tennessee, USA
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About the Editors

Christine Theoret

Dr. Christine Theoret is a full professor at the University of Montreal where she teaches veterinary anatomy and surgery in the DVM program. She received a DVM degree from the University of Montreal (1991) and completed an internship in equine medicine/surgery at the same institution in 1992. She then undertook a joint residency/MSc program in surgery at the Western College of Veterinary Medicine (1992–1995). She became a diplomate of the American College of Veterinary Surgeons in 1997. In 2000, Dr. Theoret received her PhD degree, studying the molecular mechanisms that govern healing and scarring, from the University of Saskatchewan.

Dr. Theoret founded the Comparative Veterinary Tissue Healing Laboratory in 2002, where she has since trained more than 30 highly qualified personnel, mostly MSc and PhD students. Her research has led to the publication of numerous articles in peer-reviewed, scientific journals. In 2008 she co-edited the second edition of the textbook *Equine Wound Management*. Dr. Theoret has served on the advisory boards of various national and international associations, including a term as President of the Veterinary Wound Management Society.

Jim Schumacher

Jim Schumacher graduated from Kansas State University in 1973. He worked in equine and food animal practices in Texas, California, and Kansas for 5 years after graduation. Most of this time in private practice was spent working in feedyards in Kansas. He completed an MSc degree and a residency in large animal surgery at Texas A&M University in 1980.

He was a member of the faculty at Texas A&M University until 1997. Since then he has worked at the University of London, Auburn University, the University College Dublin, and the University of Tennessee, where he has been a member of the faculty of the Department of Veterinary Clinical Sciences since 2003. He is a diplomate of the American College of Veterinary Surgeons and a member of the Royal College of Veterinary Surgeons. He has lectured students about surgery for the past 35 years.
List of Contributors

Spencer Barber, DVM, Diplomate ACVS
Professor, Equine Surgery
Western College of Veterinary Medicine
University of Saskatchewan
Saskatoon, Saskatchewan
Canada

Gary M. Baxter, VMD, MS, Diplomate ACVS
Associate Dean for Clinical Services
College of Veterinary Medicine
University of Georgia
Athens, Georgia
USA

Christophe Celeste, DrVet, PhD, Diplomate ACVS, Diplomate ECVS
Clinique Vétérinaire Sagamie
Alma, Quebec
Canada

Linda A. Dahlgren, DVM, PhD, Diplomate ACVS
Associate Professor, Large Animal Clinical Sciences
Virginia–Maryland College of Veterinary Medicine
Blacksburg, Virginia
USA

Andrew J. Dart, BVSc, PhD, Diplomate ACVS, ECVS
Professor, Equine Veterinary Science
Director of the Research and Clinical Trials Unit
Veterinary Medical Teaching Hospital, Camden
The University of Sydney
Australia

Yvonne A. Elce, DVM, Diplomate ACVS
Associate Professor, Equine Surgery
Faculté de médecine vétérinaire
Université de Montréal
Montréal, Québec
Canada

R. Reid Hanson, DVM, Diplomate ACVS, ACVECC
Professor, Equine Surgery
College of Veterinary Medicine
Auburn University
Auburn, Alabama
USA

Stine Jacobsen, DVM, PhD, Diplomate ECVS
Professor, Large Animal Surgery
Department of Large Animal Sciences
Faculty of Health and Medical Sciences
University of Copenhagen
Denmark

Derek C. Knottenbelt, OBE, BVM&S, DVMS, Diplomate ECEIM, MRCVS
Emeritus Professor, Equine Internal Medicine
University of Liverpool
Neston, Wirral
United Kingdom

Beth Kraus, DVM, Diplomate ACVS
Chadds Ford, Pennsylvania
USA

Amelia S. Munsterman, DVM, MS, Diplomate ACVS, ACVECC
Clinical Lecturer, Equine Critical Care Medicine and Surgery
College of Veterinary Medicine
Auburn University
Auburn, Alabama
USA

James A. Orsini, DVM, Diplomate ACVS
Associate Professor, Equine Surgery
School of Veterinary Medicine
University of Pennsylvania
Kennett Square, Pennsylvania
USA

Jim Schumacher, DVM, MS, Diplomate ACVS, MRCVS
Professor, Equine Surgery
Department of Large Animal Clinical Sciences
College of Veterinary Medicine
University of Tennessee
Knoxville, Tennessee
USA

John Schumacher, DVM, MS, Diplomate ACVIM
Professor
Department of Clinical Sciences
College of Veterinary Medicine
Auburn University
Auburn, Alabama
USA
Kathryn A. Seabaugh, DVM, MS, Diplomate ACVS, ACVS/SMR
Assistant Professor
College of Veterinary Medicine
University of Georgia
Athens, Georgia
USA

Albert Sole-Guitart, DVM, Diplomate ACVS
Clinician, Equine Surgery
Camden Equine Center
The University of Sydney
Australia

Ted S. Stashak, DVM, MS, Diplomate ACVS
Professor Emeritus, Equine Surgery
Colorado State University
Santa Rosa, California
USA

Christine Theoret, DMV, PhD, Diplomate ACVS
Director, Comparative Veterinary Tissue Healing Laboratory (http://theoretlab.com/index.php/en)
Professor, Département de biomédecine vétérinaire
Faculté de médecine vétérinaire
Université de Montréal
Montréal, Québec
Canada

Ferenc Toth, DVM, PhD, Diplomate ACVS
Assistant Professor
Department of Veterinary Population Medicine
College of Veterinary Medicine
University of Minnesota
St. Paul, Minnesota
USA

Jacintha M. Wilmink, DVM, PhD
Woumarec
Hamsterlaan 4
6705 CT Wageningen
The Netherlands
Preface

Wounds are among the most common medical conditions seen by veterinarians in their equine patients and one of the topics least addressed during the veterinary curriculum or at continuing education meetings. Because the horse's response to wounding differs from that of man, laboratory animals or even other veterinary patients, wound-management textbooks used in the human healthcare field or in small animal practice cannot be relied on to provide appropriate/specific therapeutic guidelines. Moreover, the general-purpose textbook covering equine medicine and surgery cannot possibly address the topic with the required depth because of the abundance of information on wound physiology and management now available. Consequently, the Equine Wound Management textbook is an essential reference for equine veterinarians because it provides readers with state-of-the-art theoretical and practical information, enhanced by an abundance of helpful tables, line drawings, and color figures.

With Dr. Ted Stashak firmly embracing a well-earned retirement, I (Dr. Theoret) was faced with the choice of a new co-editor willing to fill big shoes. Dr. Jim Schumacher courageously accepted the challenge and, due to his vast clinical experience and his familiarity with editorship, he has been an inestimable asset to the smooth execution of the task at hand. Together, we have striven to create a well-balanced book that addresses the needs of students, practitioners, postgraduate veterinarians in training programs (research- or clinically-oriented), and specialists (surgeons). Moreover, with the aim to make the book more reader-friendly, practical information has been highlighted in the text in easy-to-spot, quick-to-read "tips", and a companion interactive website posts text, questions/answers, figures, case series, "how to" videos, etc.

Since the second edition of this textbook was published in 2008, hundreds of new, relevant studies have been performed, and summaries of these findings and practical applications thereof have been included in the third edition. The wound-care market for human patients has grown in leaps and bounds over the past few years; consequently, countless new topical medications and interactive dressings have appeared on the market. In many cases, the use of these products in horses has not yet been thoroughly investigated. Despite the aforementioned differences in healing of wounds of horses, veterinarians may be tempted to extrapolate data from human or lab animal trials to the horse. Consequently, Chapters 5 and 6 have been thoroughly updated, and the author presents evidence for the effects of selected products specifically on healing tissues of the horse. Another important change since the previous edition is the increased awareness of antibiotic resistance. Accordingly, a concerted effort has been made by all contributing authors to promote responsible antimicrobial stewardship by better describing the infection continuum and reviewing the premise of antibiotic resistance and biofilms. These topics are particularly well addressed in chapters 3, 4, and 19. Finally, it seemed appropriate to add a section on dog-bite wounds and gunshot wounds to Chapter 20 and to add an entire chapter on innovative adjunctive therapies, discussing the most recent developments.

In closing we wish to express our gratitude to the authors for their willingness in bringing to this textbook all their valuable experience. We are pleased that we were able to include many world-renowned specialists to produce the highest-quality material. We are indebted to these people who generously contributed their clinical insight and current research data.

Acknowledgment

The third edition of Equine Wound Management is here today thanks to Ted Stashak’s vision and commitment to serve the equine veterinary community.
About the Companion Website

This book is accompanied by a companion website:

www.wiley.com/go/theoret/wound

The website includes:
- a webliography from chapter 17 in the book;
- case studies;
- figures from the book as PowerPoint slides, to download;
- interactive multiple choice questions and answers;
- videos.
CHAPTER 1
Physiology of Wound Healing

Christine Theoret, DMV, PhD, Diplomate ACVS

Summary

Prior to undertaking the management of a wound, the underlying biology of wound healing must be understood so that the best approach at the correct time can be selected, and so that problems with healing, if they arise, are recognized.

This chapter aims to provide an update on the physiologic, cellular, biochemical, and molecular aspects of wound repair.

Introduction

A vital trait of living organisms, continually subjected to insults from the environment, is their capacity for self repair. Whether the injury is from surgery or accidental, it generates an attempt by the host to restore continuity to tissue. Two processes are involved in healing: regeneration and repair. Regeneration entails the replacement of damaged tissue with normal cells of the type lost and is only possible in tissues with a sustained population of cells capable of mitosis, such as epithelium, bone, and liver. Conversely, repair is a “stop-gap” reaction designed to re-establish the continuity of interrupted tissues with undifferentiated scar tissue. Repair is, therefore, an inferior method of healing, producing scar tissue that is less biologically useful than the tissue it replaced, and that may adversely affect adjacent normal tissues. When complications of wound healing arise, the final result is even worse.

Accidental wounds occur commonly in horses and exert a significant welfare concern and financial burden on the equine industry. A large study by the United States Department of Agriculture’s National Animal Health Monitoring System found, in 2006, that injury/wound/trauma was the most common medical condition affecting horses, with a prevalence of 4.7% in equids 6 months of age and older. Injury/wound/trauma was the leading cause of death of foals less than 6 months old, accounting for 24% of deaths, while for horses at least 6 months old, it accounted for 16% of deaths and was the leading cause of mortality, after old age.

A study in Mexico conducted specifically on a population of working equids found a prevalence of 20.6% for cutaneous pathologic conditions; among these, skin wounds (abrasions, lacerations, abscesses) were the most prevalent (6.8%).

Figures are also concerning in Europe. A study conducted in the UK found that wounds were the most common type of injury reported by horse owners, accounting for roughly half of all injuries occurring over a 12-month period. Another study found that wounds accounted for 21.6% of veterinary...
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treatments of injured polo ponies in the UK. Horses in the southern hemisphere do not seem to fare any better: wounds ranked as the third most common medical condition encountered by equine practitioners in Australia and New Zealand and ranked second, after colic, as the most common cause of death or euthanasia.

Finally, skin trauma/wounds are a frequent cause of morbidity in athletic horses. A study in Thoroughbred racehorses has shown that 70% of injuries leading to early retirement are the result of a musculoskeletal injury, of which 7% are associated with wounds or lacerations.

The objective of repair is to re-establish an epithelial cover and to recover the integrity, strength, and function of the skin. Partial-thickness cutaneous wounds (e.g., abrasions and erosions) heal primarily by migration and proliferation of epidermal cells from the remaining underlying epithelium, as well as from the adnexal structures (i.e., hair follicles and sweat and sebaceous glands), with little participation of inflammatory or stromal cells. In contrast, second-intention repair of full-thickness cutaneous wounds hinges on four coordinated and interrelated phases (Figure 1.1). Partitioning the process into discrete phases suggests simplicity while, in reality, healing is exquisitely complex. The phases rely on interactions between multiple cellular types, their surrounding matrix, and the soluble mediators that govern the numerous activities required to rebuild the tissue. Moreover, the interactions are not static but rather in a state of constant flux, resulting in a microenvironment that is continually evolving as the wound heals.

Before veterinarians can positively influence wound healing, they must understand its mechanisms so that they select the appropriate techniques of wound management. In fact, Hipppocrates once said, “Healing is a matter of time, but it is sometimes also a matter of opportunity.”

Skin anatomy

The skin is the largest organ and serves key functions including physical protection, sensation, temperature regulation, and insulation. It is composed of two compartments – the epidermis and the dermis (Figure 1.2a). In the horse, the epidermis consists of five layers of keratinocytes: the *stratum basale, stratum spinosum, stratum granulosum, stratum lucidum,* and *stratum corneum* (Figure 1.2b). Additional epidermal components, referred to as skin appendages, include hair follicles, sweat glands, sebaceous glands, and hooves/nails. Although 90–95% of the cells populating the epidermis are keratinocytes, this compartment also includes melanocytes, Langerhans cells, and Merkel cells. Epidermis is attached to the dermis at the level of the basement membrane, a thin, glycoprotein-rich layer composed primarily of laminin and type IV collagen. This attachment is through hemidesmosomes, which physically attach the basal cells of the epidermis to the underlying dermis, as well as by vertically oriented type VII collagen anchoring fibrils, which bind the cytoskeleton.

The dermal compartment consists of two regions, the papillary dermis and the reticular dermis. This compartment is composed of dense, fibroelastic connective tissue and constitutes the bulk of the skin. The epidermis projects into this underlying connective tissue via extensions known as rete pegs or ridges. A network of collagen fibers provides tensile strength to the dermis, and elastin and glycosaminoglycans (GAGs) ensure resilience. Collagen type I is the major collagen of the dermis (~62%) whereas collagen type III comprises ~15% of the dermis. The fibroblast is the principal type of cell found in the dermis; perivascular mast cells and tissue macrophages are also found within the dermis. The connective tissue supports these cells and also a network of nerves, epithelial glands, keratinizing...
appendages, and a microvascular and lymphatic system. Indeed, only the dermal compartment is vascularized; nutrients reach the epidermis by diffusion.

In the horse, the thickness of the skin varies according to body site. For example, in the Dutch warmblood, the skin on the head, neck, and ventral abdomen is relatively thin, measuring between 1.73 mm (±0.16) and 2.03 mm (±0.2) whereas the skin on the limbs is slightly thicker, measuring an average of 2.83 mm (±0.27) for the forelimb and 2.89 mm (±0.24) for the hindlimb, depending on the specific anatomic location (e.g., the skin over the dorsal coffin joint is particularly thick, measuring 4.54 mm [±0.39]).

The subcutaneous tissue (i.e., tissue just deep to the skin) is also known as the hypodermis or superficial fascia and is not considered part of the skin. It is comprised of loose connective tissue; approximately half of the body’s fat stores are located in this region. The hypodermis anchors the skin to the underlying organs and allows the skin to move relatively freely. It also acts as a shock absorber and insulates the deeper body tissues from heat loss.

## Phases of wound repair

### Hemostasis/coagulation

The first phase of wound healing begins immediately upon injury, is completed within hours, and is dedicated to hemostasis and the formation of a provisional wound matrix. Hemostasis was long considered to be a component of the inflammatory phase, and only recently has its individual significance to wound healing been recognized. Many of the processes that occur during the ensuing phases rely on the normal execution of this short, but critical, initial phase.

Wounding traumatizes blood vessels, which results in hemorrhage. The injured endothelial cellular membrane releases phospholipids that are transformed into arachidonic acid and its metabolites that mediate vascular tone and permeability. Peripheral vasoconstriction, lasting 5–10 minutes, limits bleeding but simultaneously starves the surrounding tissues of oxygen and nutrients normally carried by the blood. The resulting transitory hypoxia and increased glycolysis, as well as pH changes, are beneficial because, along with the effects of the original vascular trauma, they enhance the activation, adhesion and aggregation of platelets, thereby initiating the intrinsic coagulation cascade, ultimately leading to the formation of a blood clot that seals the vessel.

This clot, besides providing a small amount of strength to the wound, has multiple roles. It forms a provisional matrix, rich in fibrin, fibronectin, vitronectin, and thrombospondin, that fills the space created by the wound and serves as a scaffold for migrating cells. Special surface receptors (integrins) on inflammatory and stromal cells recognize binding sites on the proteins within the scaffold, ensuring ingrowth of cells involved in healing. This cellular influx is mediated by chemoattractants released by degranulating platelets, among other cells, present in the scaffold. Indeed, activated platelets are amongst the earliest

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**Figure 1.2** (a) Diagram of a cross-section of skin, showing the epidermal and dermal compartments. (b) Diagram of the layers of the epidermis of horse skin. Source: Stashak & Theoret 2014.9 Reproduced with permission of Elsevier.
promoters of inflammation, via the release of potent chemoattractants and mitogens from their storage granules.16 Mediators released not only by platelets but also by mast cells modify vascular tone and permeability, which increase within 5–10 minutes of wounding (apparent clinically as a localized redness, heat and swelling of the wound), and thereby facilitate the aforementioned cellular migration and the diffusion of nutrients and oxygen required to sustain these newly arriving cells.

Over time, the surface clot desiccates to form a scab that protects the wound from infection. This scab is, in turn, lysed by plasmin and sloughs, along with dead inflammatory cells and bacteria, as healing proceeds underneath.

**Inflammation**

The inflammatory phase of the wound healing cascade (also referred to clinically as the debridement phase) is activated during the hemostasis/coagulation phase. It can roughly be divided into an early phase, characterized by recruitment of neutrophils, and a late phase, characterized by the appearance and transformation of monocytes. Inflammation prepares the wound for the subsequent phases of healing. It purges the body of alien substances and disposes of dead tissue, while the participating cellular populations liberate mediators to amplify and sustain the events to follow. The intensity of the inflammatory response is strongly correlated to the severity of trauma and determines the extent of scarring.

Leukocytes are recruited from blood circulating to the site of injury by the numerous vasoactive mediators and chemoattractants supplied by the coagulation and activated complement pathways, by platelets, by mast cells,10 and by injured or activated stromal cells.17 These signals initiate the processes of rolling, activation, tight adhesion, and, finally, transmigration of inflammatory cells through the microvascular endothelium. Chemoattractants also stimulate the release of enzymes by the activated neutrophils, expediting their penetration through vascular basement membranes. Diapedesis of neutrophils is further facilitated by increased capillary permeability brought about by the release of a spectrum of vasodilatory agents. Cellular influx begins within minutes and the concentration of neutrophils at the wound progressively increases to reach a peak 1–2 days after injury. Neutrophils act as a first line of defense in contaminated wounds by destroying debris and bacteria through phagocytosis and subsequent enzymatic and oxygen-radical mechanisms. Neutrophil migration and phagocytosis cease when contaminating particles are cleared from the site of injury. Most cells then become entrapped within the clot, which is sloughed during the later phases of repair. The neutrophils remaining within viable tissue die in a few days and are phagocytized by the tissue macrophages or modified wound fibroblasts, marking the termination of the early inflammatory phase of repair.17 Although neutrophils help create a favorable environment within the wound and serve as a source of pro-inflammatory cytokines, they are not essential to repair of non-infected wounds. Indeed, a classic series of experiments in the 1970s showed that, as long as conditions were kept sterile, depletion of neutrophils in a guinea pig wound model seemed not to perturb tissue repair.18 More recent knockdown experiments in mice support the depletion studies of the 1970s and go further, showing that repair is even more rapid than in wild-type sibling mice so long as conditions are sterile.19

The rapid increase in the number of macrophages in inflamed tissue is predominantly caused by the emigration of monocytes from the vasculature, followed by differentiation of the monocytes into macrophages to assist resident tissue macrophages at the wound site for a period of days to weeks. In this manner, the responsive and adaptable pluripotent monocytes can morph into macrophages whose functional properties are determined by the conditions they encounter at the site of mobilization and that change during healing. Macrophages play a central role in all phases of wound healing and orchestrate the overall process. During the early inflammatory phase, macrophages exert pro-inflammatory functions, such as antigen presentation, phagocytosis, and the production of inflammatory cytokines and growth factors that facilitate wound healing (Figure 1.3). The phenotype of wound macrophages during this phase is probably the classically activated or so-called “M1 phenotype.” Later, during the proliferative phase of healing, macrophages stimulate proliferation of dermal, endothelial, and epithelial tissue to complete formation of the extracellular matrix (ECM), angiogenesis, and epithelialization. Macrophages can then change the composition of the ECM during the remodeling phase by releasing degradative enzymes (Figure 1.4). This suggests an important role for alternatively activated macrophages (also known as “M2 phenotype”) in this phase of wound healing.20.21

More recent studies using genetically modified neonatal mice have shown that, like neutrophils, macrophages might not be essential for tissue repair.22 Nevertheless, they probably play an important role in the regulation of fibrosis and scarring by degrading matrix.23

In spite of this new, somewhat conflicting evidence, acute inflammation is still considered crucial to the normal repair of wounds in adult mammals exposed to infectious agents. When inflammation fails to resolve, however, and a chronic inflammatory response is established, the process can become dysregulated, resulting in pathologic wound repair and the accumulation of permanent fibrotic scar tissue at the site of injury. This fibrosis is characterized by the excessive accumulation of ECM components, including collagens, fibronectin, and hyaluronic acid at the site of injury, leading to a decrease in organ function and, in some cases, organ failure and death. In humans, an estimated 45% of deaths in the western world are now attributed to diseases in which fibrosis plays a major etiologic role.24 One such “fibroproliferative disorder” encountered in full-thickness cutaneous wounds of the horse and that leads to increased scarring is the development of exuberant granulation tissue (the reader is referred to Chapter 15).

After injury, once infection has been countered and repair completed, all the inflammatory cells disperse from the wound.
For inflammation to resolve, each of the events that initiated it must be halted or even reversed. Apoptosis, or programmed cellular death, is the universal pathway for eliminating unneeded cells in a phagocytic process that does not elicit additional inflammation. This mechanism prevails during all phases of wound repair because each phase relies on rapid increases in specific cellular populations that prepare the wound for repair (inflammatory cells) or deposit new matrices and mature the...
The main objective of the proliferative phase (also referred to clinically as the repair phase) is to achieve protection of the wound's surface via the formation of granulation tissue and a new epithelial cover and to restore the vascular network to nourish the new tissues.

**Fibroplasia**

The proliferative phase of repair comes about as inflammation subsides and is characterized by the appearance of red, fleshy granulation tissue, which ultimately fills the defect. Although the earliest part of this phase is very active on a cellular level, this activity does not immediately translate into a gain in the wound's strength. Indeed, during the first 3–5 days following injury, fibroblasts and endothelial and epithelial cells rapidly invade the wound in preparation for synthesis and maturation of the matrix or for wound coverage; however, these latter reinforcing mechanisms lag somewhat. Granulation tissue is formed by three elements that simultaneously move into the defect created by the wound: macrophages, which debride and produce mediators, such as cytokines and growth factors, that stimulate angiogenesis and fibroplasia; fibroblasts, which proliferate and synthesize new components of the ECM; and new blood vessels, which carry oxygen and nutrients necessary for the metabolism and growth of cells, and confer to the granulation tissue its characteristic red, granular appearance.

This stroma, rich in fibronectin and hyaluronan, replaces the fibrin clot to provide a physical barrier to infection and, importantly, provides a surface across which cells can migrate. A number of matrix molecules as well as cytokines and growth factors released by inflammatory cells are believed to stimulate fibroblasts from adjacent uninjured dermis and subcutaneous tissue to proliferate and express integrin receptors to assist their migration into the defect. Integrins are transmembrane proteins that act as the major cell-surface receptors for ECM molecules and thus mediate interactions between cells and their environment. They are particularly critical to the migratory movements exhibited by cells involved in wound healing, such as epithelial and endothelial cells and fibroblasts. Migration of fibroblasts immediately precedes advancing capillary endothelial buds but follows macrophages that have cleared a path by phagocytizing debris. Fibroblasts themselves also possess an active proteolytic system, comprising proteinases, such as plasminogen activator (PA), various collagenases, gelatinase, and stromelysin, to aid their migration into the fibrin blood clot.

After fibroblasts have arrived within the defect created by the wound, they proliferate then switch their function to protein synthesis and commence to gradually replace the provisional matrix with a collagen-rich one, probably under the influence of various cytokines and growth factors. As the wound matures, the ratio of type I (mature) to type III (immature) collagen markedly increases; proteoglycans also become abundant within the mature matrix. The greatest rate of accumulation of connective tissue within the wound occurs 7–14 days after injury, at least in the laboratory rodent, which translates into the period of most rapid gain in tensile strength (see Figure 1.1). Thereafter, the collagen content within the wound levels off as fibroblasts down-regulate their synthetic activities; this corresponds to a much slower gain in tensile strength as the wound remolds. The fibroblast-rich granulation tissue is subsequently replaced by relatively avascular and acellular scar tissue, as the capillaries within the wound regress and fibroblasts either undergo apoptosis or acquire characteristics of smooth muscle and transform into myofibroblasts that participate in wound contraction. The latter phenomena are regulated by the physiologic needs and/or the

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**Figure 1.5** This metatarsal wound failed to heal for 7 months as a result of chronic low-grade inflammation due to exposure as well as superficial and deep infection. The wound in fact became larger rather than smaller, illustrating suspension of the healing process. Courtesy of Dr. Derek Knottenbelt.

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**Cellular proliferation**

Clinicians have the greatest influence on the acute inflammatory phase of healing: proper surgical debridement and irrigation, good hemostasis, and adequate drainage can greatly hasten wound healing.
microenvironmental stimuli present at the wound. It appears that if the signal to down-regulate fibroblast activity is delayed beyond a specific time point, apoptosis is permanently impaired, ultimately leading to an imbalance between collagen synthesis and degradation and the formation of excessive scar tissue.

**Angiogenesis**

Besides initiating the inflammatory response through interactions with leukocytes, microvascular endothelial cells play a key role in the proliferative phase of repair. The formation of new capillaries from pre-existing ones (angiogenesis) is necessary to restore oxygenation and to provide required nutrients to the granulation tissue newly formed within the wound. Angiogenesis, which occurs in response to tissue injury and hypoxia, is a complex and dynamic process mediated by diverse soluble factors provided by the serum and the surrounding ECM. These factors are, in particular, angiogenic inducers, including growth factors [most notably vascular endothelial growth factor (VEGF), fibroblast growth factor (FGF)-2, platelet-derived growth factor (PDGF), and members of the transforming growth factor (TGF)-β family], chemokines and angiogenic enzymes (notably the serine proteinase thrombin), endothelial-specific receptors, and adhesion molecules, such as integrins, many of which are released during the inflammatory phase of repair.

Construction of a vascular network requires sequential steps that include augmented microvascular permeability, the release of proteinases from activated endothelial cells with subsequent local degradation of the basement membrane surrounding the existing vessel, migration and sprouting of endothelial cells into the interstitial space, endothelial cellular proliferation and differentiation into mature blood vessels (i.e., arterioles and venules), eventually followed by regression and involution of the newly formed vasculature as tissue remodels. Angiogenesis depends not only on the cells and cytokines present, but also on the production and organization of components of the ECM that provide a scaffold through which endothelial cells can migrate and a reservoir and modulator for growth factors. Thus, endothelial cells at the tip of capillaries begin their migration into the wound in response to angiogenic stimuli and the absence of neighboring cells on the second day after injury. Cytoplasmic pseudopodia extend through fragmented basement membranes; subsequently, the entire endothelial cell migrates into the perivascular space. Cells remaining in the parent vessel near the tip of the angiogenic sprouts begin to proliferate, providing a continuous source of microvascular endothelial cells for angiogenesis. A new capillary sprout has no lumen when it first develops; after it fuses with a neighboring sprout to form an arcade, it canalizes, allowing erythrocytes to pass into and through it. Formation of a lumen probably involves the joining of plasma membranes of individual and/or adjacent cells, as well as extensive intracellular vacuolization followed by fusion of the vacuoles to form “ring cells,” which ultimately fuse to form seamless capillaries. Capillaries then become stable as endothelial cells interact with the new basement membrane within 24 hours of new vessel formation and via the recruitment of pericytes and smooth muscle cells. In healing wounds, this vigorous angiogenic response results in a density of vessels that far exceeds that of capillaries in normal, uninjured tissue and provides the granulation tissue with its red, granular appearance.

After the stroma has been completely reconstituted, a rich vascular supply to the wound is no longer needed. Pro-angiogenic stimuli are down-regulated and/or the local concentration of anti-angiogenic factors (thrombospondin, interferon gamma-induced protein 10/CXC motif chemokine 10, and Sprouty-2) increases, and most of the recently formed capillary network quickly involutes through the activity of matrix metalloproteinases (MMPs), in particular MMP-1 and MMP-10, and selective apoptosis of endothelial cells. The color of the wound pales as the rich capillary bed disappears from the granulation tissue.

**Epithelialization**

Epithelialization is defined as the process of covering denuded epithelial surfaces and is essential for successful closure of the wound. In addition to the aforementioned hemostatic activities that establish a temporary barrier, the residual epithelium beneath the clot moves centripetally to participate in closure of the wound. Even though epithelial migration commences 24–48 hours after wounding, the characteristic pink rim of new epithelium (Figure 1.6) is not macroscopically visible until some time later. This “lag” varies according to the species of animal and the site, size, and substrate of the wound. For example, epithelialization is accelerated in a partial-thickness wound because migrating cells arise not only from the residual epithelium at the wound’s periphery but also from remaining epidermal appendages. Furthermore, the basement membrane is intact in this type of injury, precluding a lengthy regeneration. On the other hand, during second-intention healing of a full-thickness wound,
epithelialization cannot proceed until a bed of granulation tissue has formed. In full-thickness, 7–9 cm² wounds created experimentally in horses and ponies and left to heal by second intention, after an initial lag phase, epithelialization progressed over the wound surface at a rate of 0.63 mm/week for metatarsal wounds in ponies, 0.48 mm/week for metatarsal wounds in horses, 0.75 mm/week for buttock wounds in ponies, and 0.62 mm/week for buttock wounds in horses between the third and the seventh week of healing.38

The regenerative capacity of the epidermis relies on keratinocyte stem cells (KSC) that reside within specific microenvironments referred to as stem cell niches. The following three niches have been identified: the bulge of the hair follicle (HF), the base of the sebaceous gland, and the basal layer of the interfollicular epidermis (IFE).39 In response to epidermal injury, the HF and IFE niches participate in epithelialization of the defect.40

To close the defect in the epidermis, keratinocytes at the wound’s edge must first loosen their adhesions to each other (desmosomes) and to the basal lamina (hemidesmosomes) and develop the flexibility required to migrate over the new matrix. Numerous regulators play a critical role in modulating the proliferation and migration of keratinocytes during epithelialization; these include chemokines, cytokines, integrins, keratins, ECM molecules, and MMPs, among others. A discussion of these is beyond the scope of the text; the reader is referred to a recent review of the subject.41

Additionally, various degradative enzymes necessary for the proteolysis of components of the ECM are up-regulated within cells at the leading edge of neo-epithelium, facilitating ingestion of the clot and debris found along the migratory route. This route is determined by the array of integrin receptors for various ECM proteins, expressed on the surface of migrating epithelial cells. Once the surface of the wound is covered by epithelial cells contacting one another, further migration is inhibited by the expression, within the ECM, of laminin, a major factor responsible for adhesion of epithelial cells.

Although initial cellular migration does not require an increase in cellular multiplication, basal keratinocytes at the wound’s margin do begin to proliferate 1–2 days after injury to replenish the migratory front; this corresponds histologically to epithelial hyperplasia (Figure 1.7). The new cells leap-frog over those at the wound margin and adhere to the substratum, only to be replaced in turn by other cells coming from above and behind. The newly adherent monolayer subsequently restratifies to restore the original multi-layered epidermis.

In full-thickness wounds healing by second intention, such as those commonly managed in equine practice, provisional matrix is eventually replaced by a mature basement membrane. Repairing epidermis reassembles its constituents from the margin towards the center of the wound.17 Epidermal cells then revert to a quiescent phenotype and become attached to this new basement membrane through hemidesmosomes and to the underlying neodermis through type VII collagen fibrils. This particular aspect of epithelialization is time consuming, occurring long after total coverage of the wound by epithelium is apparent, which may explain the continued fragility of neoeplidermis for extended periods after repair is macroscopically complete. This is particularly evident in large wounds of the limb, where new epidermis is often thin and easily traumatized (Figure 1.8).

### Matrix synthesis and remodeling (also referred to clinically as the maturation phase)

Mature ECM is a non-cellular scaffold composed of proteins, glycosaminoglycans, polysaccharides, and water, that facilitates bidirectional communication between cells and their biochemical/biophysical environment.43 During the remodeling phase (occurring from 3 weeks to up to 1 year after injury), the components of the ECM undergo certain changes to ensure strength, integrity, and function of the replacement tissue.

In addition to epithelialization, contraction contributes to the successful closure of full-thickness wounds. Contraction is defined as a process whereby both dermis and epidermis bordering a full-thickness skin deficit are drawn from all sides centripetally over the exposed wound.44 This usually occurs during the second week after injury. Wound contraction not only accelerates closure, it also enhances the cosmetic appearance and strength of the scar because proportionally less area of the wound must be covered by newly formed epithelium of inferior quality, which is fragile and lacks normal nervous, glandular, follicular, and vascular components (Figure 1.8b). For this reason, a high degree of contraction is a desired feature of wound repair, at least in the horse.

Wound contraction is thought to result from a combination of tractional forces generated by migrating fibroblasts and the action of a specialized fibroblast phenotype, the myofibroblast. As fibroblasts migrate into the provisional matrix of the wound under the influence of various cytokines, tension within the wound reaches the threshold required, along with the action of TGF-β1 and the ED-A splice variant of fibronectin,45 to trigger fibroblasts to differentiate into myofibroblasts.46 The latter are the most abundant cellular elements of healthy granulation.
tissue and are aligned within the wound along the lines of contraction. The most striking feature of the myofibroblast is a well-developed alpha smooth muscle actin (α-SMA) microfilamentous system, arranged parallel to the cell's long axis and in continuity with the components of the ECM via various integrins. In addition to these cell–substratum links, intercellular connections, such as gap junctions and hemidesmosomes, ensure that neighboring cells exert tension on one another.

Wound contraction is divided into three phases. An initial lag phase (wherein skin edges retract, and the area of the wound increases temporarily for 1–2 weeks, depending on its anatomic location) occurs prior to substantial fibroblastic invasion of the wound, as a prerequisite for contraction. Subsequently, a period of rapid contraction is followed by one of slow contraction as the wound approaches complete closure. The number of myofibroblasts found in a wound appears proportional to the need for contraction, and thus, as repair progresses and the rate of contraction slows, this number decreases accordingly. During wound contraction, the surrounding skin stretches by intussusceptive growth, and the wound takes on a stellate appearance. Contraction ceases in response to one of three events: the wound's edges meet, causing contact inhibition to halt the processes of epithelialization and contraction; tension in the surrounding skin becomes equal to or greater than the contractile force generated by the α-SMA of the myofibroblasts; or, in the case of chronic wounds, a low number of myofibroblasts in the granulation tissue may result in failure of the wound to contract, despite laxity in the surrounding skin. As contraction concludes, myofibroblasts disappear, either by reverting to a quiescent fibroblastic phenotype or by apoptosis, primarily in response to reduced tension within the ECM. The myofibroblast persists in fibrotic lesions where it may be involved in accumulation of more ECM and pathologic contracture, a condition leading to substantial morbidity, particularly when it involves joints or orifices, but rarely encountered in the horse.

The conversion of ECM from granulation to scar tissue constitutes the final phase of wound repair, also referred to as maturation, and consists of synthesis of connective tissue, lysis, and remodeling. Proteoglycans replace hyaluronan during the second week of repair, support the deposition and aggregation of collagen fibers, and make the mature matrix more resilient. Collagen macromolecules provide the tensile strength to the wound as their deposition peaks within the initial week, when healing occurs by first intention, and between 7 and 14 days, when healing occurs by second intention, in the laboratory rodent.

Although this corresponds to the period of most rapid gain in strength, only 20% of the final strength of the skin wound is

Wound contraction was measured in full-thickness, 7–9 cm² experimental wounds left to heal by second intention on the limbs and hindquarters of horses and ponies. The wounds were bandaged for the first 3 weeks then left unbanded. Following an initial lag phase of 1 week, wound contraction became apparent in buttock wounds of horses and ponies and in metatarsal wounds of ponies. The percentage decrease in wound surface area attributable to wound contraction between the second and the fourth week of healing was 47% for the body wounds of ponies vs. 35% for the limb wounds of ponies, and 38% for the body wounds of horses vs. 0% for the limb wounds of horses. After week 4, the rate of wound contraction slowed to less than 5% per week for these wounds, up to complete healing. The metatarsal wounds of horses showed a different pattern: the lag phase of healing lasted 4 weeks and this was followed by an average rate of contraction that did not exceed 2.5% per week.

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Figure 1.8 (a) A 5-day-old, full-thickness, experimentally created wound over the dorsal surface of the fetlock. Granulation tissue is beginning to fill the wound. (b) The same wound, 75 days after creation. Neopidermis is thin, dry, and hairless, and could be easily traumatized. Courtesy of Dr. Ted Stashak.
achieved within the first 3 weeks of repair. At this time, collagen synthesis is balanced by collagen lysis, which normally prevents accumulation of excessive amounts of collagen and formation of a pathologic scar. The balance between synthesis and degradation determines the overall strength of a healing wound at a particular time. The first newly deposited collagen tends to be oriented randomly and, therefore, provides little tensile strength, whereas during remodeling, the fibers reform along lines of stress and, therefore, resist dehiscence more effectively. Cross-linking of the later-formed collagen is also more effective, but never occurs to the same extent as in the original tissue. The new collagen weaves into that which pre-existed and also appears to bond to the ends of old collagen fibers. These welds are points of weakness, which may rupture when stressed.

Remodeling of ECM within a wound depends on the presence of various proteolytic enzymes (proteinases) released from inflammatory and mesenchymal cells, such as MMPs and serine and cysteine proteinases (cathepsins). Those of the MMP family are collectively capable of degrading virtually all components of the ECM. Although MMPs are not constitutively expressed in skin, up-regulation occurs whenever proteolysis is required, such as during cellular migration and remodeling of the matrix. Inactive precursors of the MMPs (pro MMPs) are cleaved in the extracellular space by proteinases, such as plasmin and trypsin, left over from the inflammatory and proliferative phases, but also by other MMPs. To date, two dozen different MMPs, all distinct gene products, have been characterized.27

Based on domain organization and preference for substrate, MMPs may be divided into the following groups: 1 - collagenases; 2 - gelatinases; 3 - stromelysins; 4 - matrilysins; 5 - metalloelastases; 6 - membrane-type MMPs; 7 - other MMPs.49 Although the major function of most MMPs is probably to process bioactive molecules, such as chemokines and cytokines, as well as their respective receptors, their ability to degrade ECM proteins, as demonstrated by some members of the MMP family (MMP1, MMP3, MMP13, and MMP14) capable of cleaving collagen; MMP7 can process syndecan-1 and elastin), suggests they have a role in remodeling during wound healing.27 Comprehensive lists of MMPs, including their physiologic and in vitro substrates, can be found in proteinase databases.50,51

Homeostasis between collagen synthesis and degradation during the remodeling phase depends on the simultaneous presence of MMPs and non-specific inhibitors, such as α2-macroglobulin and α1-antiprotease, as well as natural specific inhibitors, the tissue inhibitors of metalloproteinases (TIMPs). TIMPs are a gene family of four structurally related members, TIMP-1 through -4, that inhibit conversion of MMPs from a zymogen to an activated state and that irreversibly bind to the catalytic site of active MMPs.19

Under most circumstances, a balance between MMPs and TIMPs leads to abnormal resolution and delayed repair. Indeed, although the presence of MMPs is essential for the wound to mature, it may also be responsible for the inability of chronic wounds to heal. For example, fluid found in chronic wounds is characterized by elevated concentrations of proteinases, particularly MMP-9 and serine proteinases, which lead to excessive degradation of proteins and the inactivation of critical growth factors. Chronic wounds also contain reduced concentrations of TIMPs, in particular TIMP-1.52

Wound remodeling continues for up to 2 years, during which time there is no net increase in collagen content, but instead, the collagen fibers rearrange into a more organized lattice-like structure, under the influence of local mechanical factors, progressively increasing the tensile strength of scar tissue. The majority of type III collagen fibers laid down early in healing are replaced by collagen type I, fibers become increasingly cross-linked, and the normal skin ratio of 4:1 type I to type III collagen is re-established. Glycosaminoglycans are steadily degraded until they reach concentrations found in normal dermis. The duration of the maturation phase depends on a variety of factors, including the patient's genetic makeup, age, location of the wound, type of injury, and duration of inflammation.

At maximum strength, cutaneous wounds in mice remain 15–20% weaker than the normal surrounding tissue,55 but a study in horses showed that cutaneous wounds, fully healed by second intention, withstood a maximum breaking load equivalent to only 60% of the breaking force of normal, intact skin.44 Importantly, skin appendages are usually not regenerated after wounding, resulting in a loss of other functions of skin.

Non-invasive methods to monitor healing

There are objective, non-invasive methods to monitor the progress and assess the quality of healing of cutaneous wounds, focusing on anatomic, mechanical, physiologic, and esthetic properties.12 The goal of these technologies is to provide detailed information regarding skin components imperceptible to visual inspection and that enable the clinician to assess the severity of a wound and its healing potential, thereby guiding therapeutic decisions. Optical technologies that have, to date, been applied to wound assessment in humans include: near infrared imaging, thermal imaging, optical coherence tomography, orthogonal polarization spectral imaging, fluorescence imaging, laser Doppler imaging, microscopy, spatial frequency domain imaging, photoacoustic detection, and spectral/hyperspectral imaging.56 Both infrared thermography (IRT) and near infrared spectroscopy (NIRS) have been used in the author’s laboratory to study the healing of experimental wounds in horses.57,58 Near infrared spectroscopy provides quantitative information on the structural and chemical components of cutaneous tissue, specifically oxygen saturation, hemoglobin, and water content. This tool enabled the identification of hypoxia in limb wounds relative to body wounds during the early phase of healing in horses.57 Concomitantly, cutaneous wound temperature, as measured by IRT, and by extension skin blood flow, was found to be significantly lower in limb wounds than in body wounds.58 Because low oxygen levels may promote a feeble yet prolonged inflammatory response to wounding, which could interfere with and retard the subsequent phases of healing, these experimental data aid our understanding of the impaired healing that commonly afflicts horses, particularly where limb wounds are concerned.

Other tools allow the measurement of wound dimensions to improve objective monitoring during healing, in view of adjusting the treatment plan, if required, and facilitating communication with the horse owner. A couple of recent studies have evaluated the accuracy and reliability of some of these methods when used in horse wounds.59,60 Two-dimensional (2-D) measurements of the area of the scar/scar can be taken using manual planimetry, digital planimetry or digital imaging combined with computer-aided analysis. With manual planimetry, the borders of the scar are traced on to sterile acetate film placed over the wound.
Mediators of wound repair

Wound repair relies on a complex amalgamation of interactive processes involving formed elements of blood (e.g., erythrocytes, platelets, leukocytes), ECM, and mesenchymal cells. Although histologic and morphometric observations have lead to a detailed description of the kinetics of cellular and macromolecular components involved in repair, much remains to be learned about the regulation of such activities. Restoration of structural integrity and partial functional properties appears to rely on soluble mediators, synthesized by cells present within the wound or within the surrounding tissue, that coordinate migration, proliferation, and synthesis of proteins by the various cellular populations involved in the repair process.

Cytokines, defined as 4–60 kDa signaling glycoproteins released by most nucleated cells, are among the most important soluble mediators regulating wound repair. They act in concentrations of $10^9$–$10^{12}$ M in an autocrine (same cell), paracrine (adjacent cell), or endocrine (distant cell) fashion, via cell surface receptors. Receptors are proteins with an extracellular site to bind the cytokine and a transmembrane component to transmit the signal to the intracellular site where it must reach nuclear DNA for a specific response to occur. Cells may have different numbers of receptors for different factors; the concentration of factors in the area and the number of receptors that are bound determine the response generated. Growth factors are cytokines that exert primarily mitogenic influences. Both in vivo and in vitro studies analyzing non-healing wounds have shown deregulation of various cytokines, suggesting a potential target for therapy that has led to a robust interest in using exogenous cytokines in the clinical setting to improve outcomes of healing wounds.

The cytokines that play a significant role in wound repair are summarized in Table 1.1. For more detail, the reader is referred to a comprehensive review of the topic.

Table 1.1 Cytokines involved in wound repair.

<table>
<thead>
<tr>
<th>Name</th>
<th>Abbreviation</th>
<th>Source</th>
<th>Major function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Granulocyte macrophage colony stimulating factor</td>
<td>GM-CSF</td>
<td>Macrophage, lymphocyte, fibroblast, endothelial cell</td>
<td>Differentiation and maturation of hematopoietic stem cells; recruitment of inflammatory cells; mediation of epidermal proliferation; promotion of myofibroblast differentiation and wound contraction</td>
</tr>
<tr>
<td>Interferon</td>
<td>IFN</td>
<td>Monocyte and macrophage, lymphocyte, mesenchymal cell</td>
<td>Proinflammatory; release of other cytokines; inhibits fibrosis</td>
</tr>
<tr>
<td>Interleukin</td>
<td>IL</td>
<td>All nucleated cells, in particular macrophage and lymphocyte</td>
<td>Proinflammatory; enhances epithelialization, angiogenesis, and remodeling</td>
</tr>
<tr>
<td>Tumor necrosis factor</td>
<td>TNF</td>
<td>Macrophage, lymphocyte, mast cell</td>
<td>Proinflammatory; enhances angiogenesis, epithelialization, and remodeling</td>
</tr>
<tr>
<td>Connective tissue growth factor</td>
<td>CTGF</td>
<td>Fibroblast</td>
<td>Mediator of TGF-β activity (cell proliferation and ECM accumulation)</td>
</tr>
<tr>
<td>Epidermal growth factor</td>
<td>EGF</td>
<td>Platelet, saliva</td>
<td>Epithelialization; chemotactic and mitogenic to fibroblast</td>
</tr>
<tr>
<td>Transforming growth factor-α</td>
<td>TGF-α</td>
<td>Macrophage, epithelial cell</td>
<td>MMP synthesis (remodeling); angiogenesis</td>
</tr>
<tr>
<td>Fibroblast growth factor (basic)</td>
<td>FGF-2</td>
<td>Inflammatory cell, fibroblast, endothelial cell, keratinocyte</td>
<td>Chemotactic and mitogenic to fibroblast and epithelial cell; protein synthesis; angiogenesis</td>
</tr>
<tr>
<td>Insulin-like growth factor</td>
<td>IGF</td>
<td>Liver, platelet</td>
<td>Chemotactic and mitogenic to endothelial cell; migration of epithelial cell; fibroblast proliferation; protein and GAG synthesis</td>
</tr>
<tr>
<td>Keratinocyte growth factor</td>
<td>KGF (FGF-7)</td>
<td>Fibroblast</td>
<td>Chemotactic and mitogenic to epithelial cell; mitogenic to endothelial cell</td>
</tr>
<tr>
<td>Platelet-derived growth factor</td>
<td>PDGF</td>
<td>Platelet</td>
<td>Chemotactic to inflammatory cell and smooth muscle cell; increases keratinocyte motility; mitogenic to fibroblasts; enhances protein synthesis; induction of myofibroblast phenotype</td>
</tr>
<tr>
<td>Transforming growth factor-β</td>
<td>TGF-β</td>
<td>Platelet, lymphocyte, mast cell, macrophage and macrophage, endothelial cell, epithelial cell, fibroblast</td>
<td>Chemotactic to inflammatory and mesenchymal cell; fibroblast proliferation; protein synthesis; ECM deposition (inhibition of MMP; induction of TIMP); wound contraction</td>
</tr>
<tr>
<td>Vascular endothelial growth factor</td>
<td>VEGF</td>
<td>Macrophage; fibroblast; endothelial cell; epithelial cell</td>
<td>Angiogenesis</td>
</tr>
</tbody>
</table>
Conclusion

The equine practitioner, when presented with a wounded horse, should fully understand the physiologic mechanisms underlying repair to enable design of an appropriate treatment plan. In the following chapters of this book, experienced authors share their opinion on how to best manage specific injuries. The reader benefits from a detailed understanding of the different phases of healing, as well as thorough knowledge of the mediators governing them, because these dictate the approach to follow, particularly when the wound is complicated by chronic inflammation and/or an excessive proliferative response.

References

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CHAPTER 2

Differences in Wound Healing between Horses and Ponies

Jacintha M. Wilmink, DVM, PhD

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Summary

Observing differences in wound healing between horses and ponies has provided valuable information about the intrinsic process of wound healing and the common complications encountered when managing traumatic wounds in the equid. Ponies heal faster and with fewer complications than do horses. To a large extent, these differences can be explained by disparity in the local inflammatory response, which, in turn, relates to differences in the functional capacity of leukocytes. Research data indicate that a maximal effect of treatment will be obtained in clinical practice if a differential approach is used, optimizing conditions for each successive phase of wound healing. In particular, the effect of treatment on the inflammatory response is of paramount importance to the other phases of healing and should, therefore, always be considered when managing a wound. When treating wounds healing by second intention, inflammation should be stimulated until the wound has filled with granulation tissue, and thereafter it should be inhibited.

Horses and ponies: same species, different healing characteristics

The horse evolved over the course of millions of years from a small forest dweller to a large ungulate that inhabited the vast open plains of the temperate zone. It became a “flight” animal whose instinctive reaction to danger is to run. Evolution took place as a response to various environmental and climatic challenges, and the development of special features resulted from selection by mankind. Both evolution and selection led to the large variety of breeds known today.

The equine species can be roughly subdivided into horses and ponies, and this division is determined by the height of an adult at the withers (ponies are <1.48 m). Whether ponies are just small horses has been debated for decades. The discussion of this topic, with respect to wound healing, started in the 1980s when Bertone et al. found, in a study examining second-intention healing, that wounds of ponies heal faster than those of horses and without the formation of exuberant granulation tissue (EGT). Other authors, however, reported that wounds of ponies do develop EGT, and the faster healing of wounds in ponies could not be confirmed. Because a difference in wound healing between horses and ponies might provide information about the basic biology of equine wound healing and about the complications commonly associated with wound repair in this species, Wilmink et al. performed a series of experiments to investigate differences in wound healing between horses and ponies. They proved that skin wounds in ponies heal faster and with fewer complications than do similar wounds in horses and demonstrated that these differences were based on the
efficiency and capacity of the leukocyte to produce various mediators.\textsuperscript{8,9} Differences in healing are, however, apparently not related to body size. This is confirmed by a study in donkeys and Caspian miniature horses, equine subspecies of similar size, which showed that wound healing in donkeys is faster than healing in Caspian miniature horses.\textsuperscript{10}

What remains to be elucidated is the cause of the observed differences in healing between horses and ponies. The longer period of domestication of the horse may have precluded natural selection against poor healing, because the wounded horse was/is tended to by man. Additionally, the artificial selection of various features, such as height, athletic capacity, and appearance, might have favored the development of some diseases and undesirable characteristics, including the reduced efficiency of leukocytes. Moreover, horses with wounds so poorly healed that athletic ability is impaired are often used for breeding, thus perpetuating a hereditary tendency toward poor healing. In contrast, ponies have been domesticated for a much shorter time, and, consequently, a poor capacity for healing may have been eliminated by natural selection. Moreover, artificial selection within pony breeds has been less intense, and many pony breeds maintain subpopulations (and genetic reserves) in the wild. The hypothesis that natural and artificial selection have an influence on wound healing is supported by the faster healing of donkeys compared to Caspian miniature horses,\textsuperscript{11} the latter being selected based on a special feature, namely their speed.

The physiologic and pathophysiologic differences in healing observed between horses and ponies with experimentally induced wounds have since been corroborated by other research\textsuperscript{7–9,11,12} and by clinical observation. These observations have resulted in improvements in guiding the management of wounds of equids and led to faster healing and the prevention of complications.

**First-intention healing (primary wound closure)**

Primary closure of traumatic wounds of horses is preferred over healing by second intention because healing by primary intention is usually faster, and the cosmetic and functional results are better. Unfortunately, either partial or complete wound dehiscence may ensue after primary closure. Whether or not a wound dehisces depends on many factors, including those relating to the wound itself, to the horse, to the treatment, and to the environment.\textsuperscript{13,14}

The success of primary closure of traumatic wounds was evaluated retrospectively by Wilmink \textit{et al.} in more than 500 equine patients admitted to a referral clinic.\textsuperscript{7} The study revealed that primary closure is significantly more often successful in ponies than in horses (Figure 2.1), and that bone sequestra form significantly less often in ponies when bone is exposed during

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure2_1.png}
\caption{(a) Wound on the elbow of a horse, which suffered dehiscence following primary closure, as a result of infection. (b) Wound on the distal aspect of the limb of a 4-year-old pony with an open fetlock joint and damage to the lateral collateral ligament. The wound was sutured approximately 8 hours after it occurred and it healed successfully, without dehiscence. Source: (b) Wilmink 2004.\textsuperscript{15} Reproduced with permission of Elsevier.}
\end{figure}
injury. The predominant cause of wound dehiscence and of sequestration of bone is infection. Other factors contributing to dehiscence are tension on the wound’s margins, excessive movement of the sutured region, and involvement of certain structures, such as tendons, ligaments, and synovial cavities. Other factors besides infection contributing to formation of a sequestrum include exposure of cortical bone and extensive trauma. The risk of infection is influenced by many factors, including the time elapsed since injury, the degree of contamination, the degree of tissue damage, and the thoroughness of debridement and irrigation. Wound debridement determines the concentration of bacteria left in the wound and is one factor that can be controlled clinically. The concentration of bacteria, in combination with factors at the wound that facilitate bacterial colonization, such as the presence of dead space, a hematoma, and devitalized tissue, is critical to whether or not a wound becomes infected (the reader is referred to Chapter 4 for more information about factors involved in the development of infection). Although bacterial colonization and development of infection are greatly influenced by the administration of antibiotics, the effectiveness of the patient’s own local defense, its acute inflammatory response, is as least as important to the prevention of infection.

In the aforementioned retrospective study by Wilmink et al., all these factors relating to wound infection were evaluated and compared between horses and ponies. Ponies and horses did not differ with respect to age and gender, nor with respect to wound characteristics, such as location, duration, contamination, and treatment. The wounds of ponies, however, were generally deeper, and ponies were more apt to suffer laceration of an extensor tendon, to incur damage to the periosteum, to have exposed bone, and to have their wound closed while standing, rather than while anesthetized. Additionally, ponies were significantly less likely to receive antimicrobial therapy and to be administered a non-steroidal anti-inflammatory drug (NSAID). Ponies were, therefore, at greater risk of bacterial challenge than were horses because their wounds were deeper and debridement was less complete since it was performed with the ponies standing, and because they were less likely to receive antimicrobial therapy. In spite of this greater risk, wound infection occurred less often, implying that the acute inflammatory response in ponies is more effective than that of horses at reducing bacterial contamination. The more frequent use of NSAIDs in horses, however, may have contributed to a reduced effectiveness of the inflammatory response.

**Second-intention healing**

Wounds are allowed to heal by second intention when closure is not feasible or affordable, or when a wound dehisces after being closed primarily. Second-intention healing was investigated in horses and ponies by creating excisional wounds on the metatarsi and buttocks that extended through the periosteum of the metatarsal bone or 18 mm into the muscle on the buttocks, in imitation of clinical wounds. This investigation found that wounds of ponies heal significantly faster than do wounds of horses (Figure 2.2) and that the speed and efficiency of healing seemed related to remarkable differences between horses and ponies in the phases of healing. Subsequent studies showed that these differences can be attributed to variations in the function of leukocytes.

**Clinically apparent phases during wound healing**

Wound healing is often divided into general phases of hemostasis, inflammation, proliferation, and synthesis and remodeling of matrix (see Chapter 1). Because these phases overlap and occur simultaneously in all tissue components, distinguishing them from one another is difficult. Consequently, this division is somewhat academic. In a clinical setting, dividing healing into the following macroscopically apparent events may be more practical: inflammation, formation of granulation tissue, wound contraction, and epithelialization. Although these events also overlap, they largely succeed one another and occur more or less chronologically. Moreover, they are clearly visible to the practitioner observing second-intention healing.

**Inflammation**

The inflammatory response to wounding is more prompt in ponies, as demonstrated by comparing the leukocytic infiltration of experimental wounds of ponies and horses. High numbers of polymorphonuclear cells (PMNs) are found during the first 3 weeks of healing, after which the PMNs disappear rapidly from the wounds of ponies. In contrast, the influx of PMNs is sluggish and weak in horses, but PMNs persist so that, beginning 4 weeks after wounding, the number of PMNs exceeds those in the wounds of ponies. The leukocytes of ponies are more efficient than those of horses. An in vitro study showed that the leukocytes of ponies produce more reactive oxygen species, such as H$_2$O$_2$. 

![Figure 2.2](source: Wilmink 1999)
or O\textsuperscript{-} radicals, necessary for bacterial killing after phagocytosis.\textsuperscript{9} Within tissue cages implanted subcutaneously and in newly formed granulation tissue, the leukocytes of ponies also produce higher concentrations of other inflammatory mediators, including tumor necrosis factor (TNF)-\(\alpha\), interleukin (IL)-1, and transforming growth factor (TGF)-\(\beta\), essential to the reinforcement of the inflammatory response, to the induction of fibroplasia, and to wound contraction.\textsuperscript{5,9} The superior production of these mediators can thus explain the higher initial influx of leukocytes into wounds of ponies.

Migrated leukocytes, in turn, release more biologically active substances, thus creating a positive feedback that enhances the inflammatory response.\textsuperscript{19,20} This feedback may account for the faster debridement of non-viable tissue and fibrin in wounds of ponies, leading to the faster development of a healthy bed of granulation tissue. In contrast, granulation tissue in the wounds of horses retains an irregular and purulent appearance, along with persistent deposits of fibrin.\textsuperscript{5,6} An enhanced inflammatory response observed in ponies can, likewise, translate into a more efficient local defense against contaminating bacteria, leading to a superior control of wound infection.\textsuperscript{7} A stronger acute inflammatory response, thus, averts the development of chronic, low-grade inflammation and leads to a faster and more thorough preparation of the wound for repair. Indeed, chronic inflammation, such as that seen in horses,\textsuperscript{4} perpetuates the release of tissue-damaging lysosomal enzymes, as well as mediators, such as TGF-\(\beta\), which in turn, over-stimulate fibroplasia and lead to the formation of EGT, which inhibits contraction.\textsuperscript{19,21–23}

In summary, the inflammatory response to wounding in ponies is stronger and more succinct; this response appears to be more efficient for healing. In contrast, the inflammatory response in horses is weak in onset but persists over time, and this poor but persistent response may be due to the lower initial production of inflammatory mediators.

Formation of granulation tissue

In a study by Wilming et al., granulation tissue formed faster in experimental wounds of horses than in those of ponies.\textsuperscript{5} The abundant granulation tissue in the wounds of the limbs of horses was observed to push the edges of the wound apart, which may explain why experimental wounds on the limbs of horses enlarged so dramatically during the first 2 weeks after they were created (Figure 2.2). Between weeks 2 and 3 of healing, the granulation tissue of all wounds on the limbs and buttocks of horses and ponies protruded above the level of the surrounding skin and was characterized as excessive (clinically referred to as EGT or “proud flesh”); protrusion of granulation tissue was most prominent in wounds on the limbs of horses (Figure 2.3). The wounds were left unbandedaged after week 3, and during week 4 EGT disappeared spontaneously from wounds, except those on the limbs of horses, where the excess tissue had to be trimmed to promote healing.\textsuperscript{5} The granulation tissue in horses was traversed by grooves and clefts for a much longer period, and presented a yellowish purulent surface up to week 5 after wounding. The appearance of the granulation tissue on the limbs may have been due to the weak and delayed onset followed by prolongation of the inflammatory phase. In contrast, the granulation tissue in the wounds of the ponies was smooth, regular, and achieved a healthy pink color significantly sooner (Figure 2.4).\textsuperscript{5} It was microscopically apparent that fibroblasts continued to proliferate in horse wounds even after granulation tissue filled the wound bed, whereas fibroblasts in pony wounds ceased proliferating at this time. Granulation tissue in wounds of horses presented a chaotic cellular pattern and appeared persistently inflamed, whereas it was organized regularly in pony wounds (Figure 2.5).\textsuperscript{6}

There may be a causal relationship between persistent inflammation and the continuous proliferation of fibroblasts and the synthesis of granulation tissue, via the activity of mediators, such as TNF\(\alpha\), IL-1, IL-6, platelet-derived growth factor (PDGF), TGF-\(\beta\), and basic fibroblast growth factor (bFGF), which are known to induce fibrosis.\textsuperscript{22–24} However, the formation of granulation tissue was less extensive in the wounds of ponies than in those of horses, particularly those located on the limb, even though the granulation tissue of ponies initially contained higher concentrations of TNF\(\alpha\), IL-1, and TGF-\(\beta\),\textsuperscript{5,8,9} which mediate the migration and proliferation of fibroblasts and endothelial cells.\textsuperscript{22} Furthermore, fibroblasts from limbs of ponies are known to proliferate faster \textit{in vitro} than those of horses,\textsuperscript{11,25} even though fibroplasia seems more rapid, \textit{in vivo}, in horses. Apparently, the balance of mediators, the interaction of mediators with other factors, and time-scale of the presence of mediators \textit{in vivo} are more important than the absolute concentration of mediators in determining the rate of cellular growth, stressing once more the significance of the overall course of the inflammatory response.

The formation of granulation tissue in horses is excessively fast, compared to that of other species\textsuperscript{26} and to what is observed in ponies.\textsuperscript{5} The fast formation and persistent proliferation of granulation tissue, caused by an unrelenting inflammatory response, clearly contribute to the formation of EGT.\textsuperscript{5,6}

Wound contraction

Contraction of wounds of ponies is faster and significantly more pronounced than is contraction of wounds of horses, and contraction of wounds on the body is faster and significantly more pronounced than is contraction of wounds on the limb (Figure 2.6).\textsuperscript{5} As a result, second-intention healing is significantly faster in ponies than in horses and significantly faster in wounds on the body than in wounds on the limb (Figure 2.2).\textsuperscript{5}

Histologic examination of wounds showed that the myofibroblasts in newly formed granulation tissue of wounds of ponies are organized into a regular pattern within 2 weeks after wounding; the cells are oriented perpendicular to the vessels and parallel to the wound’s surface. This pattern is thought to enhance contraction because a good bond between fibroblasts and the surrounding ECM is required for the contractile forces exerted by smooth muscle actin filaments within the fibroblasts.
to pull the margin of the wound centripetally (i.e., towards the center of the wound). Although the number of fibroblasts and the amounts of smooth muscle actin and collagen do not differ between wounds of horses and those of ponies, the alignment of myofibroblasts of horses is delayed (Figure 2.5).6

Wound contraction occurs when the forces exerted by myofibroblasts exceed centrifugal (i.e., outward) forces and the local resistance to displacement. Centrifugal forces present in the skin of horses and ponies are similar, as evidenced by the identical enlargement that occurs immediately after the creation of experimental wounds (Figure 2.5). Moreover, there is no reason to believe that the local resistance to contraction in horses and ponies should differ. Variations in wound contraction are, therefore, most likely related to differences in the contractile force generated by myofibroblasts within the granulation tissue.

Surprisingly, the inherent capacity of fibroblasts of ponies to contract, in vitro, is similar to that of horses,11 suggesting that factors in the wound, such as the presence of inflammatory mediators, determine the contractile forces exerted by myofibroblasts and hence the extent of contraction. Indeed, inflammatory mediators, in particular TGF-β, wield major effects on contraction. Interestingly, the concentration of TGF-β...