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This proposal focuses on the use of multimodal imaging and spectroscopy of post-traumatic soft tissue and bone to assess wound healing. Combining infrared (IR) imaging, near-infrared spectroscopic (NIRS) imaging, and visible reflectance spectroscopic (VRS) imaging with Raman Spectroscopy (RS) will enable the surgeon to probe the tissue with a two-dimensional, real-time approach. This assessment allows optimal determination of the viability of damaged tissue, the suitability of the tissue environment for healing, the potential for wound infection and ectopic bone formation based on the degree of tissue compromise, and development of potential objective indicators for early limb salvage versus amputation. These imaging systems are currently available and readily applicable for clinical use. Combining these technologies in a multimodal system holds great promise in permitting the surgeon to make a better objective assessment of the viability of tissues in ways that have not previously been possible.
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Casualties in Operation Iraqi Freedom (OIF) and Operation Enduring Freedom (OEF) have experienced a high rate of extremity injuries with nearly ubiquitous diffuse tissue damage and compromised local circulation often associated with overt vascular injury. These injuries include traumatic amputations, open fractures, crush injuries, burns, acute vascular disruption, blastwave-associated pressure injuries, air, thrombotic, and fat embolism, and compartment syndrome. In the treatment of such complex traumatic injuries, improved assessment of global and regional perfusion, extent of infection, location and development of necrotic tissue, as well as location and development of early heterotopic ossification would facilitate the resuscitation and definitive treatment of these patients. Noninvasive spectroscopic methods may fulfill such a role, particularly Raman spectroscopy, infrared imaging, near-infrared spectroscopic imaging, and visible reflectance spectroscopic imaging. These technologies are capable of monitoring tissue temperature, perfusion, and associated hypoxia, collagen deposition, and development of calcified tissue.
Aim 2 is comprised of five tasks:

a) Correlate the presence of necrotic tissue with spectroscopic markers.

b) Correlate spectroscopic markers with wound infection.

c) Correlate spectroscopic markers with the development of heterotopic ossification (HO).

d) Correlate spectral parameters and their response with physician and pathologist observations.

Since submission of the proposal, we have received tissue biopsies from wounds for an additional 24 patients and Raman spectra have been collected for these patients. We have also enrolled 50 patients into the 3-CCD study, collecting 3-CCD images of wounds for all of these patients. Briefly, the majority of the patients enrolled in the 3-CCD study to date have been normal healers, so correlation of outcome to 3-CCD data has not yet been possible. We have, however, determined the optimum conditions for obtained quality 3-CCD images and this practice has been implemented by the clinical team.

We are continuing to build our Raman spectral database of bacterial isolates. Preliminary findings were presented at an international conference in January 2012 (Photonics West – see Reportable Outcomes). In addition to the 30 strains of Acinetobacter baumanii, we are adding another 12 species of bacteria with multiple strains of each commonly observed in wounds (including Pseudomonas, Bacillus, Arcanobacterium, Staphylococcus, Enterococcus, Klebsiella, Citrobacter, Stenotrophomonas, Enterbacter, Morganella and Escherichia). These samples have been generously donated by Biomerieux, a manufacturer of in vitro diagnostics for clinical microbiology.

Efforts to collect Raman spectra of colonized wound effluent samples and tissue from patients that develop HO advances.

We are beginning to compile the Raman data into a central database for prediction modeling. We have completed preliminary modeling of a smaller data set (25 wounds) and have presented these results at two national conferences and a workshop (MHSRS, August 2012; SCIX, October 2012; IEEE AIPR, October 2012 – see Reportable Outcomes). Specifically, we have compared various data analysis techniques for predicting wound outcome. Univariate data analysis demonstrates statistically significant differences in the 1004 cm\(^{-1}\), 1040 cm\(^{-1}\), and 1250 cm\(^{-1}\) band areas (Figure 1). Thresholding spectral bands for wound classification (i.e. normal healing or dehisced), however, correctly classifies wound outcome in less than 70% of the test data set. Several multivariate data analysis techniques were probed for predicting wound outcome: naïve Bayesian belief network, support vector machine, and linear discriminant analysis. The Bayesian belief
network model was based on an initial univariate analysis and performed worst (accuracy of ~65% for both normal healing and dehisced wounds – Figure 2A). The addition of clinical data to the data set improved model prediction by almost 10% (Figure 2B). The support vector model provided 80% accuracy for predicting wound dehiscence, while the linear discriminant analysis exhibited 90% accuracy for predicting wound dehiscence. These preliminary results demonstrate great promise for the potential of Raman spectroscopy to predict wound healing.

Aim 3 revolves around the completion of a swine hind limb ischemia protocol.

a) Acquire spectroscopic images of the limb before, during and after limb ischemia using VRS, NIRS and IR imaging.

b) Measure standard systemic assessments of reperfusion injury (creatine kinase (CK), urine myoglobin) in addition to cardiac output, blood pressure, serum lactate, base deficit and hemoglobin levels before, during, and after limb ischemia.

We have collected images and biological samples from 32 pigs to date, including all sham animals as well as all control animals. As we suspected, imaging data indicates that reperfusion and subsequent oxygenation of the ischemic limb do not proceed at the same rate for severely affected tissues. Calculated 3CCD values at 30 minutes post-reperfusion, indicative of tissue oxygenation, trend in a similar manner to Tarlov scale values (clinical evaluation of mobility) – Figure 3.

c) Optimize resuscitation methods by correlating standard and spectroscopic parameters with porcine model outcomes.

We are in the process of analyzing the data generated by the swine limb ischemia model. Preliminary results of the imaging work have been presented at two national conferences (MHSRS, August 2012 and SCIX, October 2012 – see Reportable Outcomes). Preliminary results of the data collected from clinical samples will be presented at a national conference in November 2012 (AALAS – see Reportable Outcomes).

Starting in 2013, we will begin the experimental arms of the protocol, which involves administering Lifor perfusate after limb ischemia.
In addition to the 30 strains of Acinetobacter baumanii, we are adding another 12 species of bacteria with multiple strains of each commonly observed in wounds.

Efforts to collect Raman spectra of colonized wound effluent samples and tissue from patients that develop HO advances.

We are beginning to compile the Raman data into a central database for prediction modeling.

We have generated preliminary models to predict wound healing and are able to obtain 90% accuracy for prediction of wound dehiscence.

We have collected images and biological samples from 32 pigs to date, including all sham animals as well as all control animals. We are in the process of analyzing the data generated by the swine limb ischemia model.
REPORTABLE OUTCOMES

Poster presentations:


Oral Presentations:


Manuscripts:


CONCLUSION

In this effort we have made progress in all task areas and disseminated our findings through both national presentations and publications. With regards to three of the key outcomes in wounded warriors (the development of HO, wound failure and wound infection) our efforts have begun to demonstrate success. With regards to wound healing, we have enrolled more than 50 patients (each with multiple wounds and time points) for both Raman and 3CCD analysis and are continuing to obtain control specimens from non-injured patients and heterotopic ossification tissue from injured patients. We continue to move closer to an “optical biomarker” which can guide debridement and predict outcomes. Finally, as both HO and wound failure are related to the presence of bioburden our efforts in determining the spectra of common flora in combat patients will not only augment our understanding of the interplay between host and response to injury but help select appropriate anti-microbial approaches. In summary, this multi-faceted approach has laid the foundation for continued advances over the ensuing years. In the future, we hope to apply these efforts to civilian trauma as well as military trauma.
REFERENCES


Figure 1. Band areas and band area ratios calculated for first (A), second (B), and final (C) debridements. Asterisks indicate statistically significant differences. * = $p > 0.05$; ** = $p > 0.01$. 
Figure 2. Receiver operating curve for wound outcome prediction (0 – normal healing wounds, 1 – dehisced wounds). Area under the curves (AUCs) are calculated for both a spectroscopic only dataset (A) and a spectroscopic and clinical dataset (B).
**Figure 3.** Comparison of calculated final 3CCD values (after 30 minutes of reperfusion) and Tarlov scale values. Note that the highest Tarlov scale values correlate with the highest calculated 3CCD values.
Using Multimodal Imaging Techniques to Monitor Limb Ischemia

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INTRODUCTION
• Our goal is to develop technology for real-time monitoring of extremity oxygenation and perfusion.
• Our hypothesis is that changes in tissue oxygenation and perfusion can be accurately monitored using a multimodal imaging system with 3CCD imaging and infrared imaging.
• Thermal imaging provides better data regarding overall perfusion of the extremity.
• 3CCD imaging provides data regarding the surface oxygenation of the extremity and is based on the spectral response of hemoglobin in the visible region of the spectrum.
• By integrating multiple imaging methods, we can quantitatively identify and track regions of tissue where oxygenation and perfusion change in real-time.
• The first step is to ascertain a clinically relevant and objective test to aid clinicians treating critical limb care patients.

STUDY DESIGN
To simulate common vascular injuries sustained during blast-related extremity injuries, two methods are used.

- Direct vascular injury is modeled by occlusion/tamponade of the blast vessel.
- Crush injury is modeled by using a pneumatic hemorrhage simulator, a pressure of 150 mm Hg is used.

IMAGING MODALITIES

- 3CCD (Panasonic P1i AG-344M16B)
  - It is similar in mechanism to a common digital camera, including the lens, image sensor, and image processor.
  - It provides enhanced color and image quality.
- Infrared imaging
  - The IR sensor is placed in the handle of the instrument, and the IR data is recorded using a thermal image acquisition system.
- Thermal imaging
  - Infrared (IR) camera
  - The infrared camera acquires images by detecting the infrared radiation emitted by objects.

RESULTS

3CCD IMAGING

Shown below are 3CCD images at different time points during a technique-induced ischemia.

RESULTS

INFRARED IMAGING

[Diagrams showing thermal images captured at different time points]

DISCUSSION

Here, the combined information of the localized change in limb temperature and skin surface absorption spectra are displayed in a user-friendly visual representation in real-time. The data collected is minimally manipulated using simple matrix arithmetic methods, allowing for easy and rapid on-chip processing.

Data collected from this study so far indicates that this method could allow for a rapid, objective assessment of the severity and extent of ischemic damage. This could provide surgeons with accurate information regarding overall limb viability as well as the appropriate timing of wound closure.
Chronicling Wound Healing with Raman Spectroscopy

Nicole J. Crane Ph.D.¹,³, Rajiv Luthra¹, Emily Valaik¹, Jonathan A. Forsberg M.D.¹,²,³, Eric A. Elster M.D.¹,²,³

¹Naval Medical Research Center
²Walter Reed National Military Medical Center
³Uniformed Services University of the Health Sciences
Acute Combat Wounds
Acute Combat Wounds

• The management of modern traumatic war wounds remains a significant challenge for clinicians.
  • Extensive osseous and soft-tissue damage caused by blasts and high-energy projectiles.

• The ensuing inflammatory response ultimately dictates the pace of wound healing and tissue regeneration.

• The timing of wound closure or definitive coverage is subjectively based.

• Despite the use and application of novel wound-specific treatment modalities, some wounds fail to close, or dehisce.
Surgical debridements are performed every 2-3 days.
- remove devitalized tissue
- decrease bacterial load

Negative pressure wound therapy (NPWT) is applied between debridements. NPWT promotes wound closure.

Wound assessment involves:
- patient’s general condition
- injury location
- adequacy of perfusion
- gross appearance of the wound
Monitor wound healing *in vivo*, i.e. monitor wound healing during surgical debridements.

- Is it the best time to close the wound?
- Is the wound developing HO?
- Is the wound infected? With what?

Develop an objective and predictive model for wound healing.
The Toolbox
Our Toolbox

- Real-time PCR
- Multiplex Protein Assay
- Raman Spectroscopy
- FTIR Imaging
- Thermography
- Visible Reflectance Imaging
- Bayesian Belief Network modeling
Sample Collection

Wound is surgically cleaned

- Collect 1 cm$^3$ tissue biopsy from center of wound bed
- NPWT is applied
- Protein assay
- Raman spectroscopy
- Micro culture

NPWT is applied

- Serum is collected
- Effluent is collected
- Protein assay
- RT-PCR
- Raman spectroscopy
- Micro culture
• Previous study demonstrated the potential of Raman spectroscopic analysis of wound biopsies for classification of wounds as normal or impaired healing from changes in the 1665 cm\(^{-1}\)/1445 cm\(^{-1}\) band area ratio.


• Impaired healing wounds demonstrate a significant decrease in the 1665/1448 cm\(^{-1}\) band area ratio compared to normal healing wounds, as demonstrated by Raman spectroscopic mapping.
Preliminary Study – Real-Time PCR Analysis Results

• Results were corroborated by altered collagen/collagenase gene expression profiles of tissue biopsies.

• Gene expression profiles confirm decreased gene expression of collagen types I and III at the first debridement and collagen type III at the final debridement in impaired wounds.

• COL18A1 mRNA expression remains elevated for impaired healing wounds at almost all time points when compared to normal healing wounds. Continued elevation of endostatin would inhibit neovascularization.

During early wound healing, type III collagen is the most abundant collagen and is gradually replaced by type I collagen.

- delayed deposition of type III collagen = delayed deposition of type I collagen = delayed re-epithelialization

Could this type of analysis be extended to intact wound biopsies and ultimately obviate the need for excisional wound biopsies?
Chronicling Wound Healing with Raman Spectroscopy
Approximately 1 cm² tissue biopsy is excised from the center of the wound bed. Tissue is fixed in 10% neutral buffered formalin for storage. Prior to spectral acquisition, samples are rinsed in 0.9% NaCl saline solution.

Examine multiple spots across the tissue.

40 accumulations, 5s spectrum.
Monitoring a Wound Over Time

- Spectral differences between biopsies collected from normal healing wounds (n=8) and dehisced wounds (n=8) are not readily apparent.
- There is, however, a significant difference in the background intensity (mostly fluorescence) that the Raman signal rides on.
Peakfitting for Spectral Deconvolution

<table>
<thead>
<tr>
<th>Raman Shift (cm$^{-1}$)</th>
<th>Vibrational Band Assignment</th>
<th>Component</th>
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<tr>
<td>860</td>
<td>$\nu$(C-C)</td>
<td>nucleic acids</td>
</tr>
<tr>
<td>920,940</td>
<td>$\nu$(C-N), $\nu$(C-C)</td>
<td>nucleic acids, keratin</td>
</tr>
<tr>
<td>1004</td>
<td>$\nu$(C-C) ring</td>
<td>phenylalanine</td>
</tr>
<tr>
<td>1040</td>
<td>$\nu$(C-C) skeletal</td>
<td>glycogen, keratin</td>
</tr>
<tr>
<td>1125</td>
<td>$\nu$(C-C), $\nu$(C-N)</td>
<td>nucleic acids, protein</td>
</tr>
<tr>
<td>1250</td>
<td>$\nu$ (C-N) and $\delta$(N-H); Amide III</td>
<td>protein</td>
</tr>
<tr>
<td>1320</td>
<td>$\delta$(CH$_2$) twisting</td>
<td>nucleic acids, protein</td>
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<tr>
<td>1445</td>
<td>$\delta$(CH$_3$) and $\delta$(CH$_2$) scissoring</td>
<td>protein</td>
</tr>
<tr>
<td>1555</td>
<td></td>
<td>aromatic amino acids, heme</td>
</tr>
<tr>
<td>1665</td>
<td>$\nu$(C=O); Amide I</td>
<td>protein</td>
</tr>
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Univariate Analysis of Band Areas

Comparison of mean band areas and band area ratios for all debridements for biopsies from normal healing wounds and dehisced wounds.

![Graph showing comparison of mean band areas and band area ratios for normal and dehisced wounds.](image)
Univariate Analysis of Band Areas

Comparison of mean band areas and band area ratios for first debridements for biopsies from normal healing wounds and dehisced wounds.

![Graph showing comparison of band areas and ratios for normal and dehisced wounds.](image)
Univariate Analysis of Band Areas

Comparison of mean band areas and band area ratios for final debridements for biopsies from normal healing wounds and dehisced wounds.

Band Area (a.u.)

- **Normal**
- **Dehisced**

*p < 0.02*
Monitoring a Wound Over Time – Normal Healing Wounds vs. Dehisced Wounds

• Overall, there is an increase in the mean 1040 cm$^{-1}$ band area (BA) for all normal healing wound biopsies compared to all dehisced wound biopsies in this preliminary study.

• When just comparing the first debridement biopsies, protein related band areas and band area ratios are significantly different for normal healing wound biopsies and dehisced wound biopsies – 1445 cm$^{-1}$ BA, 1665/1445 cm$^{-1}$ band area ratio (BAR), and 1004 cm$^{-1}$ BA.

• At the final debridement, the 1040 cm$^{-1}$ BA for all normal healing wound biopsies is decreased compared to the dehisced wound biopsies.

• We have over 250 tissue biopsies from over 50 patients, most of which we have collected spectra in triplicate (over 4 years).

• Peak fitting needs to be performed over the entire spectrum to determine which vibrational bands provide optimal sensitivity and specificity. Some regions of the spectrum require a more in depth peak fitting (Amide I and III bands).

• Other aspects of the spectra, such as the fluorescence background, may also provide indications of the status of the tissue overall.
Project Goals

• Build our spectral biopsy database for Bayesian Belief Network modeling and other multivariate analysis techniques.

• Determine threshold values (band area ratios, fluorescence background, etc.) that are capable of differentiating normal healing wounds and wounds that are susceptible to dehiscence.

• Validate spectroscopic wound healing prediction models in a military and civilian population.
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http://www.med.navy.mil/sites/nmrc/Pages/ccc_regenmed.htm
The multidisciplinary care of these patients would not have been possible without the dedicated efforts of everyone at WRAMC and NNMC. Both civilian and military personnel have rendered skilled and compassionate care for these casualties. All of our efforts are dedicated to those who have been placed in harm’s way for the good of our nation.

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This study was approved by the Walter Reed National Military Medical Center Institutional Review Board in compliance with all Federal regulations governing the protection of human subjects.
Using Multimodal Imaging Techniques to Monitor Limb Ischemia: A Rapid, Non-Invasive Method for Assessing Extremity Wounds

Rajiv Luthra¹, Nicole J. Crane Ph.D.¹,³, Jonathan A. Forsberg M.D.¹,²,³, Eric A. Elster M.D.¹,²,³

¹Naval Medical Research Center
²Walter Reed National Military Medical Center
³Uniformed Services University of the Health Sciences
Acute Combat Wounds

- Approximately 70% of military casualties from Operation Iraqi Freedom and Operation Enduring freedom were recorded as involving major limb injury.
- Roughly 90% of which were caused by blast injuries from improvised explosive devices (IEDs).
- The management of modern traumatic war wounds remains a significant challenge for clinicians.
The timing of wound closure is subjectively based.

Surgical debridements are performed every 2-3 days to assess wound condition and remove devitalized tissue.

Wound assessment involves combination of:

- patient’s general condition
- injury location
- adequacy of perfusion
- gross appearance of the wound
Acute Combat Wounds - Challenge

• Need for an objective method to measure limb viability.

• Aim to develop technology for real-time monitoring of extremity oxygenation and perfusion.

• Ideally the technology would be:
  - Low-cost
  - Non-invasive
  - Rapid and Reliable
Potential Solution

- Multimodal imaging system with 3CCD and infrared imaging.
- Accurately monitors changes in tissue oxygenation and perfusion.
  - Real-time
  - Non-invasive
  - Uses existing commercially available cameras
Imaging Modalities

- **3CCD:**
  - Provides data for surface oxygenation.
  - Based on the spectral response of hemoglobin in the visible region of the spectrum.

- **IR:**
  - LWIR: 7.5 - 13.5 μm.
  - Provides data for overall tissue perfusion.
  - Based on relationship of tissue temperature with perfusion.
Understanding 3CCD Cameras

• Advantages:
  • Technology similar to common digital camera.
  • Commercially and easily available.
  • Inexpensive & easy to use.
  • Provides enhanced color and image quality.

• Mechanism:
  • Utilizes a trichroic prism to split incoming light into three channels (Red, Green and Blue).
  • Each channel has its own detector a charge-coupled device (CCD).
  • Higher color sensitivity and dynamic range.
Absorption Spectra of Hb\textsubscript{O2} and Hb

Peak Differences between Hb\textsubscript{O2} and Hb
Relating $P_{O_2}$ values with 3CCD

Each well initially contains the same concentration of oxygenated blood. Varying amounts of sodium dithionite, a reducing agent that scavenges the available $O_2$, are added to change the final $P_{O_2}$. 

0 mg/ml  
1.2 mg/ml  
1.9 mg/ml  
2.5 mg/ml
Relating $P_{O_2}$ values with 3CCD

1. Image imported into Matlab.
2. Split into its component channels.
Relating $P_{O_2}$ values with 3CCD

3CCD Image

Red Channel

Red-Blue Intensity

Mean blue channel intensity in a ROI is subtracted from the mean red channel intensity to give the R-B 3CCD value.
Relating $P_{O_2}$ values with 3CCD

Measured $P_{O_2}$ values against the 3-CCD values for each well show a very clear correlation.
Swine Model of Limb Ischemia
Swine Limb Ischemia Protocol

- Goal: Characterize extremity injury caused by ischemia and reperfusion.

- To score the severity of the extremity injury the modified Tarlov scale is used.

  0 – Complete paralysis
  1 – Minimal movement
  2 – Stands with assistance
  3 – Stands alone
  4 – Weak walk
  5 – Normal gait
Limb Ischemia

- Study extremity ischemia and reperfusion in 2 cases.
- To simulate a vascular injury, we use direct vessel occlusion by isolating and tying the femoral artery and vein.
- To simulate a crush type injury, such as one sustained by an IED blast, we use a pneumatic tourniquet inflated to 250 mm Hg.
Swine Limb Ischemia Protocol - Tourniquet

- Study tourniquet induced extremity ischemia in 2 cases.
  - Case I:
    - 2 hours ischemia followed by 30 minutes of reperfusion.
    - We predicted the damage caused to be mostly “reversible”.
  - Case II:
    - 3.5 hours ischemia followed by 30 minutes of reperfusion.
    - After 3 hours of ischemia, muscle cells are much more likely to become hypoxic and die.
    - Here we predicted the damage caused to be “irreversible”.
3CCD Results
2 hr Tourniquet Ischemia and Reperfusion 3CCD Images

“Reversible Ischemia”

0 minutes: Start Ischemia

120 minutes: Maximal Ischemia

125 minutes: Start Reperfusion

150 minutes: Maximal Reperfusion
3.5 hr Tourniquet Ischemia and Reperfusion 3CCD Images

“Irreversible Ischemia”

0 minutes: Start Ischemia

210 minutes: Maximal Ischemia

215 minutes: Start Reperfusion

240 minutes: Maximal Reperfusion
• All R-B values were normalized to peak hypovolemic values.

• Within the 2 hour ischemia study we see 2 cases appear to be a milder ischemia than the other 5 cases.
Peak tissue oxygenation values for all 5 cases is ~ 6 minutes after reperfusion starts.
The final R-B value correlates with the severity of the extremity injury.
3.5 hr Tourniquet Ischemia and Reperfusion: Mean 3CCD R-B Values

- Again all R-B values were normalized to peak hypovolemic values.
- Here we see 2 cases appear to be a milder ischemia than the other 3 cases.
Peak tissue oxygenation values are at different times for each case.
• Here we see a very good agreement between final R-B value (240 minutes) and the modified Tarlov score.

• We may be able to predict severity of tissue damage using this method.
Different Peak Times due to Delayed Reperfusion?

• One explanation would be:
  
  • Different rates of extremity reperfusion cause different peak oxygenation values.

• To look at reperfusion we turn to IR imaging.
IR imaging Results
IR images during 2 hour Ischemia and Reperfusion

0 minutes: Start Ischemia

120 minutes: Maximal Ischemia

125 minutes: Start Reperfusion

150 minutes: Maximal Reperfusion
IR images during 3.5 hour Ischemia and Reperfusion

0 minutes: Start Ischemia

210 minutes: Maximal Ischemia

215 minutes: Start Reperfusion

240 minutes: Maximal Reperfusion
Average extremity temperature during Ischemia and Reperfusion

- Temperature drop greater in 3.5 hr.
- No clear deviation in trends for either 2 hour or 3.5 hour cases.
- Limb temperature returns to pre-ischemic values.
Comparing Mild and Strong Ischemic cases during Reperfusion

- Looking at 3.5 hr tourniquet cases.
- Reperfusion rates are different between the two cases.
Comparing Tissue Oxygenation and Reperfusion – Milder Ischemic cases

- IR values do not directly correspond to 3CCD values.
- The delay in 3CCD value reaching its peak value is not explained well by only the reperfusion state.
Comparing Tissue Oxygenation and Reperfusion – Stronger Ischemic cases

- Here, the fastest reperfusion case has the maximum delay to peak oxygenation.
- Possible tissue damage: reperfusion but reduced oxygenation?
- Need to monitor both reperfusion and tissue oxygenation.
Conclusions

• We have developed a multimodal imaging system with 3CCD and infrared imaging.

• We have demonstrated that we can track changes in tissue oxygenation and perfusion in a limb ischemia model.

• A combination of R-B and IR values will likely enable prediction of limb viability.

• This technology can be easily visualized in real-time.
We can imagine a clinical imaging system for visualization of ischemia and reperfusion…….
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Disclaimer

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Predicting Wound Outcome from Raman Spectroscopic Data: Univariate versus Multivariate Techniques

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Acute Combat Wounds
Acute Combat Wounds

- The management of modern traumatic war wounds remains a significant challenge for clinicians.
  - Extensive osseous and soft-tissue damage caused by blasts and high-energy projectiles.
- The ensuing inflammatory response ultimately dictates the pace of wound healing and tissue regeneration.
- The timing of wound closure or definitive coverage is subjectively based.
- Despite the use and application of novel wound-specific treatment modalities, some wounds fail to close, or dehisce.
Surgical debridements are performed every 2-3 days.

- remove devitalized tissue
- decrease bacterial load

Negative pressure wound therapy (NPWT) is applied between debridements. NPWT promotes wound closure.

Wound assessment involves:

- patient’s general condition
- injury location
- adequacy of perfusion
- gross appearance of the wound
Monitor wound healing *in vivo*, i.e. monitor wound healing during surgical debridements.

- Is it the best time to close the wound?
- Is the wound developing HO?
- Is the wound infected? With what?

Develop an objective and predictive model for wound healing.
The Toolbox
Our Toolbox

- Real-time PCR
- Multiplex Protein Assay
- Raman Spectroscopy
- FTIR Imaging
- Thermography
- Visible Reflectance Imaging
- Bayesian Belief Network modeling
Sample Collection

Wound is surgically cleaned

Collect 1 cm³ tissue biopsy from center of wound bed

RT-PCR
Raman spectroscopy
Micro culture

Protein assay
Raman spectroscopy
Micro culture

Effluent is collected

NPWT is applied

Serum is collected

Protein assay
Preliminary Study – Raman Mapping Results

- Previous study demonstrated the potential of Raman spectroscopic analysis of wound biopsies for classification of wounds as normal or impaired healing.


- Impaired healing wounds demonstrate a significant decrease in the 1665/1445 cm\(^{-1}\) band area ratio compared to normal healing wounds - these results were corroborated with collagen gene expression in the same samples.

Could this type of analysis be extended to intact wound biopsies and ultimately obviate the need for excisional wound biopsies?
Chronicling Wound Healing with Raman Spectroscopy
Raman Fiber Probe Data Collection

Approximately 1 cm² tissue biopsy is excised from the center of the wound bed.

Tissue is fixed in 10% neutral buffered formalin for storage.

Prior to spectral acquisition, samples are rinsed in 0.9% NaCl saline solution.

Examine multiple spots across the tissue.

14 accumulations, 5s spectrum
7 accumulations, 10s spectrum
Monitoring a Wound Over Time

- Spectral differences between biopsies collected from normal healing wounds (n=12) and dehisced wounds (n=13) are not readily apparent.
Peakfitting for Spectral Deconvolution

<table>
<thead>
<tr>
<th>Raman Shift (cm(^{-1}))</th>
<th>Vibrational Band Assignment</th>
<th>Component</th>
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<tbody>
<tr>
<td>860</td>
<td>(\nu(C-C))</td>
<td>nucleic acids</td>
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<tr>
<td>920,940</td>
<td>(\nu(C-N), \nu(C-C))</td>
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<tr>
<td>1250</td>
<td>(\nu(C-N)) and (\delta(N-H)); Amide III</td>
<td>protein</td>
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<td>(\delta(CH_3)) and (\delta(CH_2)) scissoring</td>
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<tr>
<td>1665</td>
<td></td>
<td>protein</td>
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</tbody>
</table>
Univariate Analysis of Band Areas

First Debridement

- heeled
- dehisced

Graph showing band areas at different wavenumbers (1665 cm⁻¹, 1555 cm⁻¹, 1455 cm⁻¹, 1445 cm⁻¹, 1320 cm⁻¹, 1250 cm⁻¹, 1040 cm⁻¹, 1004 cm⁻¹, 1665 cm⁻¹, 1004 cm⁻¹, 940 cm⁻¹) for healed and dehisced conditions.
Univariate Analysis of Band Areas

Final Debridement

- Healed
- Dehisced

* denotes significant difference.
Univariate Analysis of Band Areas

• Based on a univariate analysis, there are only three band areas that differ based on outcome (healed or dehisced) - 1004 cm\(^{-1}\), 1040 cm\(^{-1}\), and the band area ratio 1665/1004 cm\(^{-1}\).

• We can also look for changes in the band area profiles of healed and dehisced wounds over time, i.e. first debridement versus final debridement.

For wounds that heal, there are no statistically significant difference between calculated band areas chronologically.

For wounds that dehisce, the band area at 1250 cm\(^{-1}\) demonstrates a statistically significant difference between the first and final debridement (\(p < 0.03\)).

Is this reflective of wound “stability”? 
Univariate Analysis of Band Areas: Thresholds

Predicting outcome based on univariate analysis of band areas would misclassify 31% of dehisced wounds and 33% of healed wounds.
Multivariate Analysis: BBNs

• Bayesian Belief Networks fall under the umbrella of machine learning.

  Input = empirical data → output = patterns or predictions that are believed to be responsible for generating the data

• Applies the Bayes probabilistic theory for analyzing data sets.

• Bayesian Belief Networks (BBNs) demonstrate probabilistic relationships – relationships in which knowledge of the value of a variable affects the belief about the likelihood of other variables taking certain values.

  ➢ Data mining
  ➢ Prediction
BBN Analysis of Band Areas: Data Mining

Bayesian Belief Network (BBN) machine learning can elucidate some of the relationships between band area changes.

- $1555 \text{ cm}^{-1} \rightarrow 1250 \text{ cm}^{-1} \rightarrow 1320 \text{ cm}^{-1} \& 1665 \text{ cm}^{-1} \rightarrow 1665/1445 \text{ cm}^{-1} \rightarrow 1004 \text{ cm}^{-1}$

- $1040 \text{ cm}^{-1} \rightarrow 1250 \text{ cm}^{-1} \rightarrow 1665/1004 \text{ cm}^{-1} \rightarrow 1004 \text{ and } 1455 \text{ cm}^{-1} \rightarrow 1665/1004 \text{ cm}^{-1}$
A naïve Bayes model for outcome as the target reveals that outcome is largely dependent on mean 1040 cm\(^{-1}\) and mean 1455 cm\(^{-1}\) band area ratios.

Note, this data set incorporates different collection parameters, operators and regions of the biopsy.
Here, we generated a naïve Bayes model based on a training data set of all debridements except final debridements and used it to predict the wound healing outcome for final debridements.

Unfortunately, the accuracy of the prediction model is only 68% for normal healing wounds and 64% for dehisced wounds.
Multivariate Analysis: Support Vector Machine

Supervised classification method based on a non-probabailistic binary linear classifier.

- assumes that groups are separated by a wide boundaries
- places samples in the two classes as far apart as possible
- able to handle non-linear data
- can be susceptible to overfitting
- may perform better on data sets with more examples of training and fewer labels
Support Vector Machine

Developed a subset of spectra for a training data set (N=61), randomly selected from the whole data set. Used the entire spectrum with no pre-processing (derivatization, smoothing, normalization).

<table>
<thead>
<tr>
<th>Actual</th>
<th>Predicted</th>
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<tbody>
<tr>
<td></td>
<td>Healed</td>
</tr>
<tr>
<td>Healed</td>
<td>36</td>
</tr>
<tr>
<td>Dehisced</td>
<td>0</td>
</tr>
</tbody>
</table>

For a test data set (N=20), 80% of the wound spectra were classified correctly as dehisced.
Multivariate Analysis: Linear Discriminant Analysis

Supervised classification method based on Bayes’ formula.

- assumes a normal distribution
- assumes that the variability between each group has the same structure
- uses PCA to reduce dimensionality
- places samples in the same class close to one another but samples not in the same class far apart
- may perform better than SVM in cases of many labels and little training data
Developed a subset of spectra for a training data set (N=61), randomly selected from the whole data set. Used the entire spectrum with no pre-processing (derivatization, smoothing, normalization).

For a test data set (N=20), 90% of the spectra were classified correctly as dehisced.
Which Model to Choose?

- The models shown require significant optimization.

- We need to ensure complete data sets for training (include different acquisition parameters, multiple acquisitions, different instrument operators – i.e. build robustness into our model).

- The final evaluation of the prediction model will require both the highest sensitivity and specificity.

- Preliminary results demonstrate promise for the use of Raman spectroscopy to predict wound outcome.
Ongoing Projects

• Currently, we are continuing to build our biopsy database for Bayesian Belief Network modeling and other multivariate analysis techniques.

• We have over 250 tissue biopsies from over 50 patients, most of which we have collected spectra in triplicate (over 4 years).

• Peak fitting needs to be performed over the entire spectrum to determine which vibrational bands provide optimal sensitivity and specificity. Some regions of the spectrum require a more in depth peak fitting (Amide I and III bands).

• Other aspects of the spectra, such as the fluorescence background, may also provide indications of the status of the tissue overall.
Project Goals

- Determine threshold values (band area ratios, fluorescence background, etc.) that are capable of differentiating normal healing wounds and wounds that are susceptible to dehiscence.

- Validate spectroscopic wound healing prediction models in a military and civilian population.
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General Surgery, Orthopaedics and Rehabilitation

http://www.med.navy.mil/sites/nmrc/Pages/ccc_regenmed.htm
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This study was approved by the Walter Reed National Military Medical Center Institutional Review Board in compliance with all Federal regulations governing the protection of human subjects.
Profiling wound healing with wound effluent: Raman spectroscopic indicators of infection

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ABSTRACT

The care of modern traumatic war wounds remains a significant challenge for clinicians. Many of the extremity wounds inflicted during Operation Enduring Freedom and Operation Iraqi Freedom are colonized or infected with multi-drug resistant organisms, particularly Acinetobacter baumannii. Biofilm formation and resistance to current treatments can significantly confound the wound healing process. Accurate strain identification and targeted drug administration for the treatment of wound bioburden has become a priority for combat casualty care. In this study, we use vibrational spectroscopy to examine wound exudates for bacterial load. Inherent chemical differences in different bacterial species and strains make possible the high specificity of vibrational spectroscopy.

Keywords: combat wounds; wound effluent; Raman spectroscopy; bacteria; Acinetobacter baumannii

1. INTRODUCTION

Infections are common complications of combat wounds and affect not only quality of life but also wound outcome (healing or non-healing). At the beginning of the twentieth century, improvements in military hygiene and disease control significantly reduced the number of war-time deaths due to pestilence.[1] While deaths from “war-time” pestilence are not common in recent conflicts such as Operation Iraqi Freedom (OIF) and Operation Enduring Freedom (OEF), infection control of multi-drug resistant organisms such as Acinetobacter, Klebsiella, and Pseudomonas has presented a challenge.[2] Acinetobacter isolates were the most predominant microorganisms found in a recent study of combat wounds, accounting for over 60% of all bacterial isolates.[3] Multi-drug resistant Acinetobacter infections can be problematic due to the small number of effective drugs for treatment – carbapenems and tigecycline.[4] Thus, accurate identification of the species and strain of the infecting organism becomes important. Currently, we are evaluating wound effluent from combat-wounded soldiers for bacterial and correlating wound colonization to wound outcome.

Wound effluent is the fluid that is exudated from the wound during the wound healing process. The composition of wound effluent changes over the course of wound healing and is a complex milieu of blood, plasma, cells, immunoglobulins, other various proteins such as enzymes, cytokines and chemokines, and bacteria, in the case of infection. In a previous study, Brown and coworkers have shown that inflammatory cytokine and chemokine profiles extracted from wound effluent are associated with the extent of wound colonization.[5] In this study, we explore the use of Raman spectroscopy to probe wound effluent for bacterial infection.

Raman spectroscopy is a molecularly specific technique that is capable of probing samples noninvasively and nondestructively. It has been used to assess tissues at the molecular level with diverse clinical and diagnostic applications to include the analysis of cellular structure and the determination of tumor grade and type.[6-22] This makes Raman spectroscopy an ideal technology for evaluating wound effluent, particularly for detecting bioburden. There have been numerous Raman spectroscopic studies of microorganisms, many focusing on rapid identification.
of the microorganisms.[23-31] By creating a Raman spectral database of microorganisms, it is possible to identify bacteria at the strain level. We hypothesized that Raman spectroscopy could evaluate bioburden in wound effluent and differentiate strains of the same species of bacteria, namely *Acinetobacter baumannii*.

## 2. MATERIALS AND METHODS

### 2.1 Clinical Studies and Sample Collection

The clinical studies were approved by the institutional review boards of the National Naval Medical Center (NNMC) and the Walter Reed Army Medical Center (WRAMC). All study participants were recruited from wounded Operation Iraqi Freedom and Operation Enduring Freedom U.S. service members evacuated to the National Capital Area. Informed consent was obtained from all participating patients.

For the treatment of combat wounds, surgical debridement and pulse lavage were performed in the operating room every 48-72 hours until definitive wound closure or coverage. Negative pressure wound therapy (NPWT) was applied to the wounds between surgical debridements, as per current standard practice at NNMC and WRAMC.[32] All wounds were examined once daily following wound closure or coverage until the sutures were removed. All patients were followed clinically for 30 days. Wound effluent was collected from the NPWT canister (without gel pack; Kinetic Concepts, Inc., San Antonio, TX) two hours following the first surgical debridement and over a 12 hour period prior to each subsequent wound debridement. Samples were stored at 4°C prior to spectral acquisition.

### 2.2 Culturing *Acinetobacter baumannii*

Thirty *Acinetobacter baumannii* isolates were streaked onto lysogeny broth agar (LBA) plates and placed in a 37°C incubator. After approximately three days of growth, at least 10 μL of each isolate was available for analysis by Raman spectroscopy. Additionally, microorganisms were cultured from the wound effluent itself by plating 50–100 μL of effluent onto a blood agar plate. Bacteria counts are reported as CFU/mL by plate.

### 2.3 Raman Spectroscopy

Raman spectra of reference standards (plasma, whole blood, cells, bacteria and immunoglobulin G) were transferred to an aluminum foil covered weighing dish for spectral acquisition. Uncentrifuged, unfiltered wound effluent samples were placed in a 1 cm³ quartz cuvette for spectral acquisition. Bacterial isolates were transferred to an aluminum foil covered weighing dish with a 10 μL inoculating loop for spectral acquisition. A 785 nm Raman PhAT system (Kaiser Optical Systems, Inc., Ann Arbor, MI) was used to collect spectra of the effluent and bacteria. Final spectra were the accumulation of twenty 5 second spectra (for bacteria) and thirty 5 second spectra (for effluent), acquired using the 3 mm spot size. For some bacteria isolates, fluorescence signal overwhelmed the Raman scatter. To reduce the fluorescence, the bacteria isolates were transferred to a 0.5 mL centrifuge vial and rinsed with deionized water. The vial was then centrifuged at 10,000 rpm for 5 minutes. The bacteria isolates were transferred back to the weighing dish for spectral acquisition. The rinsing process was repeated until fluorescence reduction was appreciable.

### 2.5 Data Analysis

For the effluent samples, all spectral preprocessing was performed in GRAMS/AI software (Thermo Fisher Scientific, Madison, WI). Raman spectra were truncated to 1800-400 cm⁻¹ and baseline corrected with a sixth degree polynomial. For effluent samples, spectral subtraction of blood was performed if spectral interference of blood was noted.

Hierarchical clustering of *Acinetobacter baumannii* isolates was performed in Unscrambler X 10.1 software (CAMO Software, Woodbridge Township, NJ). Prior to classification, spectra were transformed with a first derivative function (5th order, 13 points) and truncated to 930-1080 cm⁻¹.
3. RESULTS

3.1 Wound Effluent

The spectral profiles of wound effluent components are displayed in Figure 1 – plasma, whole blood, human mesenchymal stem cells, bacteria and immunoglobulin G, respectively.

![Figure 1. Spectral comparison of Raman spectra of wound effluent components – A) plasma, B) whole blood, C) cells, D) bacteria, and E) immunoglobulin G.](image)

Major bands exhibited in the spectra 1665 cm\(^{-1}\) (amide I), 1620 cm\(^{-1}\) (C=C), 1557 cm\(^{-1}\) (C-C ring stretching), 1450 cm\(^{-1}\) (CH\(_2\) scissoring), 1340 cm\(^{-1}\) (\(\phi\)CH\(_2\)), 1270 cm\(^{-1}\) (amide III), 1245 cm\(^{-1}\) (amide III), 1070-1080 cm\(^{-1}\) (C-O stretch), and 1004 cm\(^{-1}\) (C-C aromatic ring). [25, 33-40] In plasma and whole blood, bands at 1620 cm\(^{-1}\) and 1557 cm\(^{-1}\) are particularly prominent. In plasma and whole blood, these bands can be attributed to hemoglobin. Only the cell and bacteria spectra in Figures 1C and 1D exhibit a significant, broad band at 1070 cm\(^{-1}\); this band confirms the presence of nucleic acids. While there is a significant amount of overlap in some of the spectra, each demonstrates uniqueness when examined as a whole.

Some of these spectral features are clearly present in the spectrum of fresh wound effluent (Figure 2A), namely whole blood. After the Raman spectrum of whole blood is subtracted from the Raman spectrum of effluent, the resulting spectrum (Figure 2B) shares spectral features with both Acinetobacter baumannii (Figure 2C) and cells Figure 2D), but most closely resembles the Raman spectrum of the bacteria. Additionally, the Raman spectrum of effluent can be used to monitor the amount of cellular matter (human or bacteria) throughout the course of treatment for the wounded warriors. Figure 3 shows the Raman spectra of wound effluent collected from the same patient at the fifth, sixth, seventh, and eighth surgical debridements. Evidence of cellular matter is apparent in debridements five through seven (Figures 3A-C), as denoted by the presence of the 1450 cm\(^{-1}\), 1240 cm\(^{-1}\), and 1004 cm\(^{-1}\) bands,
Figure 2. Raman spectrum of wound effluent before (A) and after subtracting whole blood from the spectrum (B). The resulting spectrum is compared to Acinetobacter baumannii (C) and cells (D).

Figure 3. Raman spectra of wound effluent collected from the same wound after the fifth (A), sixth (B), seventh (C), and eighth (D) debridement.
but becomes drastically reduced by the eighth debridement (Figure 3D). Bacteria counts for these samples also
decrease with time (3.5 x 10^5 CFU/mL, 2.1 x 10^4 CFU/mL, and 2.0 x 10^3 CFU/mL for the fifth, sixth, and eighth
debridements respectively).

3.2 Acinetobacter baumannii Isolates

We also used Raman spectroscopy to profile thirty isolates of *Acinetobacter baumannii*. Hierarchical clustering was
used to delineate the spectral relationships between the different strains of *Acinetobacter baumannii* (Figure 4),
specifically Spearman’s rank hierarchical clustering with complete linkage over the wavelength range of 930-1080
\text{cm}^{-1}. Apa1 digestion (restriction enzyme digestion of DNA) and optical mapping (high-resolution restriction
maps from single, stained molecules of DNA) were also performed on the isolates (data not shown) and subjected to
hierarchical clustering.

![Hierarchical clustering of 30 Acinetobacter baumannii isolates into subgroups of strains. Boxes highlight strains that were misclassified according to optical mapping and/or Apa1 digest data.](image)

Clustering by optical mapping and Apa1 digestion were compared to the results of the Raman spectral clustering.
Genetically determined subgroup assignments are indicated by the numbers to the left of the brackets, while gray
boxes indicate the strains that did not classify correctly according to Apa1 digestion and optical mapping.
Hierarchical clustering of the Raman spectra correctly classified 77% of the thirty isolates examined. Performance
of the classification technique could potentially be improved with alternate spectral preprocessing and additional
spectral region optimization.
4. DISCUSSION

Current microbiological methods use culture from tissue homogenate or from wound fluid directly to determine the species of bacteria present as well as to quantify the bacteria present. At the time of wounding, or the inoculation event, 100% of wounds are contaminated. Not long after inoculation wounds become colonized, though not all microorganisms are harmful to the host. The signs of infection can be obvious in a healthy patient, but may be less conspicuous in a sick patient. Some combat wounded soldiers suffer from an exaggerated inflammatory response, similar to that observed in acutely ill states like sepsis.[32] This exposes some noncosomial bacteria like Acinetobacter baumannii with the perfect opportunity for infection. Previously, critical wound colonization has been correlated with the inflammatory cytokine and chemokine profiles of combat-wounded soldiers.[5] There has been some controversy over the culture methodology used for bacterial quantification, though some schools of thought believe that timing of the timing of sampling is more critical than the sampling itself. Current methodologies require 24-48 hours for results from microbial tests. Raman spectroscopy, a noninvasive and nondestructive technique, holds promise for the development of faster microbial testing.

In this preliminary study, Raman spectroscopic profiling of wound effluent during wound surgical debridements demonstrates a decrease in bands associated with cellular material, notably 1450 cm$^{-1}$, 1240 cm$^{-1}$, and 1004 cm$^{-1}$; these changes in the spectral profile of the wound effluent are possibly indicative of reduced bacterial load over the course of wound healing. In addition, Raman spectroscopy was able to correctly classify 77% of thirty Acinetobacter baumannii isolates into their respective strain subgroups. These results corroborate genetic mapping data performed on the same Acinetobacter baumannii isolates. Thus, it is conceivable that the Raman spectrum collected from the wound effluent itself could be compared to Raman spectra of bacterial isolates for identification. This kind of rapid assessment may eventually help to direct antibiotic therapy and prevent over- or under-treatment of bacterial infection.

We have demonstrated that Raman spectroscopy can be utilized to examine wound effluent, in addition to bacterial isolates. One advantage of Raman spectroscopy is that it can be employed in a non-invasive manner, such as a fiber probe-coupled system. It is possible to incorporate such a Raman spectroscopic system into the operating room. For microbial studies, however, a microscope coupled Raman system will allow for probing of smaller sample sizes, such as single colonies of bacteria.

5. CONCLUSIONS

This study demonstrates the potential of vibrational spectroscopy as a technique capable of affording an objective measurement regarding wound effluent colonization. Such a capability could allow for real-time point of care analysis of wounds, allowing subjective decisions to be supplanted by objective data. This is a critical need as constraints on medical education reduce clinical exposure and decision-making is moved from the subjective arena to personalized, data driven decisions. The use of such methodologies as presented herein, may allow for evaluation of wound bioburden in a shorter time frame than is currently possible. We need to expand the scope of our study to a larger patient population and a more diverse microbial to better delineate Raman spectroscopic trends to develop a classification model for wound infection.

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7. REFERENCES

Vibrational spectroscopy: a tool being developed for the noninvasive monitoring of wound healing

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Eric A. Elster
Vibrational spectroscopy: a tool being developed for the noninvasive monitoring of wound healing

Nicole J. Cranea and Eric A. Elstera,b,c

Abstract. Wound care and management accounted for over 1.8 million hospital discharges in 2009. The complex nature of wound physiology involves hundreds of overlapping processes that we have only begun to understand over the past three decades. The management of wounds remains a significant challenge for inexperienced clinicians. The ensuing inflammatory response ultimately dictates the pace of wound healing and tissue regeneration. Consequently, the eventual timing of wound closure or definitive coverage is often subjective. Some wounds fail to close, or dehisce, despite the use and application of novel wound-specific treatment modalities. An understanding of the molecular environment of acute and chronic wounds throughout the wound-healing process can provide valuable insight into the mechanisms associated with the patient’s outcome. Pathologic alterations of wounds are accompanied by fundamental changes in the molecular environment that can be analyzed by vibrational spectroscopy. Vibrational spectroscopy, specifically Raman and Fourier transform infrared spectroscopy, offers the capability to accurately detect and identify the various molecules that compose the extracellular matrix during wound healing in their native state. The identified changes might provide the objective markers of wound healing, which can then be integrated with clinical characteristics to guide the management of wounds. © 2012 Society of Photo-Optical Instrumentation Engineers (SPIE). [DOI: 10.1117/1.JBO.17.1.010902]

Keywords: wound healing; acute wounds; chronic wounds; combat wounds; Raman spectroscopy; Fourier transform infrared spectroscopy.

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1 Introduction

There is no healthcare specialty that is free from the morbidity and costs of wound development in a patient. In 2009, U.S. hospitals discharged over 1,300,000 patients with chronic wounds and more than 547,000 with traumatic wounds (classified as >10% body surface area burn or open wound).1 U.S. healthcare costs related to wound treatment are well over $20 billion yearly, and the impact of wound healing on these expenditures is extensive.2 In addition, if every surgical procedure is considered a case of an acute wound, the significance of wound healing is simply tremendous.

Although the wound-healing process of acute wounds such as surgical incisions is fairly well understood, the modified wound-healing process encountered in patients with chronic wounds and some traumatic acute wounds still requires elucidation. Normal healing of an acute wound is directed by a cascade of growth factors and cell signaling that allows the wounds to repair quickly. Chronic wounds and some traumatic acute wounds are much slower to heal and behave differently for several underlying reasons. There may be a pathologic process such as infection that prevents the wound from healing normally. Additionally, wound healing may be complicated by a prolonged inflammatory phase that inhibits normal levels of chemical mediators and cell recruitment. Finally, the patient’s general condition contributes to the rate of wound healing: malnutrition and comorbidities such as diabetes are associated with impaired wound healing.3 Improved objective assessment of wounds would be conducive to better treatment of them, which might result in faster healing times, decreased infection rates, and decreased local and systemic complications of injury. For instance, if visits to the operating room were reduced by one instance per patient for 140 patients at one hospital, the cost savings would be over $2 million. The eventual timing of wound closure is often subjective, and there exists a need for an objective evaluation of the molecular environment of wounds throughout the wound-healing process. The use of vibrational spectroscopy and imaging for increased diagnostic accuracy and better wound treatment can produce improved clinical outcomes and decreased patient morbidity, resulting in an earlier return to an improved quality of life.

2 Wound Pathophysiology and the Process of Wound Healing

Several parameters are used to classify wounds: the layers of tissue involved, the origin and duration of the wound, and the type of wound closure used (i.e., surgical closure with sutures or formation of scar tissue). Origin and duration dictate whether a wound is classified as chronic or acute. Wounds resulting from trauma or surgery are acute wounds and generally proceed normally through the wound-healing process. An incision site in the abdomen, a third-degree burn, or a crushed limb...
is termed an “acute wound.” Wounds arising from chronic inflammation, repetitive insult, or vascular compromise that fail to heal normally or in a timely manner are called “chronic wounds.” Pressure ulcers and diabetic foot ulcers are examples of chronic wounds. Acute wounds generally begin with a single, abrupt insult and progress through the healing process in an orderly manner. Conversely, chronic wounds are usually caused by a pathologic process such as infection or poor circulation.

In general, the wound-healing process proceeds through regeneration and/or repair. “Wound regeneration” is the renewal of the damaged tissue with healthy tissue that is the same, whereas “wound repair” is the replacement of the damaged tissue by scar tissue. Wounds that are confined to the superficial layers of skin heal by regeneration, but wounds that penetrate deep into the subcutaneous layers are not able to regenerate and heal by scar formation. The overall sequence of events that precedes injury is thought to be similar for chronic and acute wounds whereby chronic wounds simply stall at one or more stages during the wound-healing process.4

The first step in wound healing is hemostasis, the vascular response that triggers platelet activation and aggregation, clot formation, and vasoconstriction. The second step in wound healing is inflammation—capillaries vasodilate, and neutrophils and macrophages migrate to the wound bed to debride the wound and secrete growth factors to promote angiogenesis and connective tissue synthesis (tissue inhibitors of matrix metalloproteases, matrix metalloproteases, transforming growth factor-α and transforming growth factor-β, interleukin-1, interleukin-6, interleukin-8, epidermal growth factor, and keratinocyte growth factor). The third step in wound healing is proliferation, a multistep process involving epithelialization (early formation of the new wound bed from fibroblasts), neangiogenesis (induction of new vasculature), and matrix and/or collagen deposition. The final step in wound healing is wound contraction and maturation and/or remodeling—the wound edges close, and a stronger, more orderly matrix forms scar tissue.3

Numerous factors that can affect the wound-healing process make an already complicated process even more difficult to accurately assess. These factors include age, stress, nutrition, tissue perfusion and oxygenation, infection, and other comorbidities, such as obesity, diabetes mellitus, immunosuppression, pulmonary disease, renal disease, and vascular disease. Unfortunately, in some cases, wound healing is complicated by dehiscence, in which “closed” wounds fall apart and reopen. The events leading up to wound dehiscence are not well understood but are suspected to result from an intensely exaggerated inflammatory response.4

Currently, wounds are evaluated on the basis of parameters such as location of injury, adequacy of perfusion, gross appearance of the wound, wound tensile strength, and the patient’s general condition. Although parameters such as the location of injury, the gross appearance of the wound, and the patient’s general condition are fairly obvious and can be reasonably assessed, parameters such as the adequacy of perfusion and tensile strength are not readily quantifiable during surgery. It has previously been demonstrated that there is a greater incidence of associated vascular injury in slowly healing wounds than in normally healing wounds.5 It is also well established that the tensile strength of the wound is dependent on collagen deposition.6 There exists a need for technologies that can be used to noninvasively and objectively assess these challenging parameters.

3 Raman and Fourier Transform Infrared Spectroscopy

Raman and Fourier transform infrared (FTIR) spectroscopy are types of vibrational spectroscopy that measure the vibrational frequencies of molecules as the molecules are excited by incident photons. Every molecule has a unique fingerprint of vibrational frequencies, which makes Raman and FTIR spectroscopy highly specific techniques for molecular identification. Both techniques can be employed noninvasively, making them ideal for biomedical applications. Raman spectroscopy and FTIR spectroscopy are sometimes referred to as “sister” techniques and provide complementary information about molecules, but they differ in several fundamental ways.

Raman spectroscopy arises from the inelastic scattering of ultraviolet, visible, or near-infrared light when a photon interacts with a molecule. Raman scattering is an inherently weak process, and, as such, samples are typically illuminated by laser light. Light scattered by the sample is diffracted into individual wavelengths by a spectrophotograph and collected by a detector such as a CCD or CMOS sensor.7 Raman systems can be coupled to a microscope and motorized stage for high-resolution imaging8–14 or to a fiberoptic probe for bulk in vivo sampling.15–20 Raman spectroscopy’s independence from a specific sample thickness and lack of spectral interference from water make it an ideal technique for biomedical applications. One disadvantage of Raman spectroscopy in the biomedical arena, however, is its inherently weak signal, which can be overwhelmed by sample fluorescence. Often this is overcome by excitation in the near-infrared region of the spectrum where biological molecules tend not to fluoresce. There are other advanced configurations and applications of Raman spectroscopy, but they lie outside the scope of this review.21–25

FTIR spectroscopy consists of the absorbance of frequencies of light by a molecule that contains the same vibrational frequencies within its molecular bonds. A beam of infrared light is passed through or reflected by a sample. Some light is absorbed by the sample’s vibrational frequencies, and the remaining light is transmitted to an interferometer and then collected by a detector, such as a mercury cadmium telluride photoconductive detector or an indium gallium arsenide photodiode detector.26 As with Raman spectroscopic systems, FTIR systems can be coupled to a microscope27–39 or a fiberoptic probe.40 FTIR spectroscopy is sensitive to the presence of water, however, and in vivo sampling can be challenging. One disadvantage of FTIR spectroscopy is that it requires that light be able to pass through the sample and thus is confined to use with thin samples, such as tissue sections on optically transparent windows.

Both Raman spectroscopy and FTIR spectroscopy offer the capability to accurately detect and identify the various molecules that compose the extracellular matrix in their native state during wound healing. They are both imaging techniques in which the precise biochemical composition of biologic samples can be obtained by noninvasive and nondestructive means.41–44 Both have been proven to be effective in studying tissues at the molecular level using diverse clinical and diagnostic applications, including the analysis of cellular structure and the determination of tumor grade and type.42,45–48 Pathologic alterations of wounds are accompanied by fundamental changes in the molecular environment that can be analyzed by
vibrational spectroscopy. The identified changes might provide the objective markers of acute wound healing, which could then be integrated with clinical characteristics to guide the management of traumatic wounds. For instance, changes in collagen vibrational bands could be correlated with alterations in collagen deposition and reepithelialization of the wound bed.

Fig. 1 Infrared characterization (factor analysis conducted over the 1185 to 1475/cm region) of wounded and nonwounded areas six days after wounding is shown. (a) Optical image of an unstained section with the edge of the wounded area marked by a vertical dashed line. (b—g) The score images are shown for various components of the tissue. (b) f1 is the stratum corneum and part of the viable epidermis. (c) f2 is the suprabasal epidermis. (d) f3 is the basal epidermal layer. (e) f4 is the outer leading edge of the migrating epithelial tongue. (f and g) f5 and f6 are the collagen-rich areas, respectively. (h) The factor loadings of f1 to f4 are characteristic of keratin-rich areas. The factor loadings of f5 and f6 are characteristic of collagen-rich areas. Reprinted with permission from John Wiley and Sons [J. Cell. Mol. Med. 12(5B), 2145–2154 (2008)].

Fig. 2 Factor analysis of a confocal Raman dataset delineates skin regions near a wound edge 0.5 days after wounding. Data analysis was conducted over the 800 to 1140/cm region, yielding four factor loading images that map to anatomically distinct regions in the skin. (a) The spatial distribution of scores for f1 highlights the stratum corneum region of the skin, which is rich in keratin-filled corneocytes and lipids. (b) f2 shows high scores in the underlying epidermal region. (c) High scores for f3 reside near the dermal-epidermal boundary region. (d) The size, location, and spatial distribution of several smaller regions with high scores for f4 are identified as cell nuclei. (e) Factor loadings reveal several spectral features specific to the microanatomy of the epidermis in human skin. Reprinted with permission from John Wiley and Sons [J. Cell. Mol. Med. 12(5B), 2145–2154 (2008)].
4 Vibrational Spectroscopic Studies of Wound Healing

4.1 Wounds

The application of vibrational spectroscopy, such as Raman spectroscopy and FTIR spectroscopy, to study wound healing is a developing field of interest. Both ex vivo and in vivo models of wound healing have been explored in animals and humans, but all studies published to date have focused on acute wounds versus chronic wounds.

In all surgical cases, an acute wound is inflicted once a surgical incision is made. Thus, all surgical wounds are classified as acute wounds and are typically examples of the normal healing process. In early ex vivo studies by Wijelath and co-workers, FTIR attenuated total reflection (ATR) spectroscopy illustrated modified healing patterns in arterial grafts implanted into dogs. Standard histological analysis of the graft implants showed little or no activity in the first 10 days after implantation, but FTIR-ATR spectroscopy demonstrated changes within the fibrin layer of the graft that could be correlated to endothelialization of the wound.51,52 Gough et al. utilized synchrotron FTIR spectroscopy to study wound healing in humans have been performed on ex vivo biopsies of wounds. In 2008, Mendelsohn et al. utilized both FTIR and Raman spectroscopy to correlate spectroscopic changes with the reepithelialization of the wound bed of cutaneous incisional wounds.53 Spectroscopic results were compared directly with immunohistological images of serial tissue sections and gene array analysis data. FTIR images collected four days after wounding precisely depicted the keratin-rich migrating epithelial tongue from the collagen-rich wound bed with focal data analysis of the 1185/cm to 1475/cm spectral region (Fig. 1). Similar spectral features are exhibited by factors 1 to 4 (f1 to f4), but the factors are spatially distinct within the sample itself. These represent keratin-rich areas confirmed by immunohistochecmistry. Factors 5 and 6 are spectrally distinct from factors 1 to 4 and represent collagen-rich areas of the sample. Confocal Raman microspectroscopic images of tissue sections demonstrate the time dependence of elastin distribution in the wound up to six days after wounding (Fig. 2).54 By day 2, the elastin distribution (f1) and the distribution of a collagen factor (f3) were significantly decreased, whereas the distribution of a second collagen factor (f2) decreased. Their study clearly demonstrates the utility of vibrational spectroscopy and imaging to monitor component-specific changes in skin in an acute wound-healing model.

Our group has used Raman spectroscopic mapping to monitor changes within the wound bed. Tissue biopsies were collected from Operation Iraqi Freedom and Operation Enduring Freedom combat-wounded soldiers at each surgical debridement during the wound-healing process.55 Spectral maps revealed differences in the amide I/CH2 scissoring band area ratios that correlated with wound outcome (Fig. 3), i.e., normal healing or impaired healing. Raman spectroscopic results were and increased cellularity56 as well as conformational changes within the proteins themselves.57

To date, published applications of vibrational spectroscopy to study wound healing in humans have been performed on ex vivo biopsies of wounds. In 2008, Mendelsohn et al. utilized both FTIR and Raman spectroscopy to correlate spectroscopic changes with the reepithelialization of the wound bed of cutaneous incisional wounds.53 Spectroscopic results were compared directly with immunohistological images of serial tissue sections and gene array analysis data. FTIR images collected four days after wounding precisely depicted the keratin-rich migrating epithelial tongue from the collagen-rich wound bed with focal data analysis of the 1185/cm to 1475/cm spectral region (Fig. 1). Similar spectral features are exhibited by factors 1 to 4 (f1 to f4), but the factors are spatially distinct within the sample itself. These represent keratin-rich areas confirmed by immunohistochecmistry. Factors 5 and 6 are spectrally distinct from factors 1 to 4 and represent collagen-rich areas of the sample. Confocal Raman microspectroscopic images of tissue sections demonstrate the time dependence of elastin distribution in the wound up to six days after wounding (Fig. 2).54 By day 2, the elastin distribution (f1) and the distribution of a collagen factor (f3) were significantly decreased, whereas the distribution of a second collagen factor (f2) decreased. Their study clearly demonstrates the utility of vibrational spectroscopy and imaging to monitor component-specific changes in skin in an acute wound-healing model.

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Fig. 3 Photographs are shown for a patient with a normal healing wound (a) and one whose wound healing was impaired (b). (c) This graph shows the percentage difference of the 1665-to-1445/cm band area ratios calculated from the first and last debridement 1665-to-1445/cm band area ratios for wounds classified as healing normally (black bars) and those wounds in which healing was classified as impaired (white bars).
corroborated with collagen gene expression profiles. In impaired healing wounds, a decrease in collagen-like bands was confirmed by decreased expression of the COL1A1 and COL3A1 genes (for type I and type III collagens, respectively). In addition to monitoring the wound bed itself, FTIR and Raman spectroscopy were utilized to monitor complications of wound healing, such as infection, the formation of biofilm from subsequent infection, and heterotopic ossification (HO), to which acute and chronic wounds are susceptible.

4.2 Infection

For acute wounds such as surgical incisions, infection is the most prevalent postsurgical complication. Chronic wounds provide a bed of growth for pathogens—they are warm, deep, and sometimes full of necrotic tissue. Chronic wounds are more often infected than acute wounds, but acute combat wounds present a subset of acute wounds with a high infection rate. Identifying the pathogens responsible for wound bioburden is especially important because the prevalence of multidrug-resistant bacteria is increasing, necessitating treatment with appropriate antimicrobial agents. Because of the specificity of Raman and FTIR spectroscopy, they can also be used to evaluate the bioburden of wounds. There have been numerous FTIR and Raman spectroscopic studies of microorganisms, many of which have been focused on rapid identification of the microorganisms.

Differences in the Raman spectral profile of three bacterial species as well as three bacterial strains are evident in Fig. 4 (unpublished data). Both *Klebsiella pneumoniae* and *Acinetobacter baumannii* are Gram-positive bacteria, whereas methicillin-resistant *Staphylococcus aureus* is a Gram-negative bacterium. Differences in the Raman spectral profile, however, are due not strictly to peptidoglycan content but to other structural differences in the proteins as well. Inherent chemical differences in different bacterial species and strains, as demonstrated in Fig. 4, make possible the high specificity of Raman spectroscopy. When the Raman spectra of wound effluent collected from two patients colonized with different bacteria are compared (Fig. 4), the spectral profiles show differences in amino acid content and alterations in glycosidic linkages.

4.3 Heterotopic Ossification

Another complication of wound healing, “heterotopic ossification,” is defined as the pathological formation of bone in soft tissue. HO formation has been observed following orthopedic surgery (total hip arthroplasty as well as acetabular and elbow fracture surgery), burn injury, traumatic brain injury, and spinal cord injury. HO formation is not commonly observed in civilian traumatic wounds without the presence of head injury or spinal injury and develops in only 20% and 11% of these patients, respectively. During the current military conflicts in Iraq and Afghanistan, HO has been a frequent and common clinical problem in soldiers with traumatic combat wounds. Currently, operative excision is the only treatment for mature, symptomatic HO. Identifying tissue that will develop into HO is not trivial, however, and can only be confirmed once mineralized tissue is evidenced on a radiograph. Tissue mineralization could easily be monitored with Raman spectroscopy. Information could be gained that would reveal the quality of the bone being formed during HO. For example, is the bone “normal” but developing in soft tissue, or is the bone “pathological,” developing by a different mineralization mechanism altogether?

While Raman and FTIR spectroscopy have been used extensively to study the process of biomineralization, they have not previously been used to provide insight into the pathological process of HO. We have collected Raman spectra of uninjured muscle, injured muscle, and “pre-HO” tissue (defined as palpably firm or “woody” tissue without roentgenographic evidence of HO) found within high-energy penetrating wounds (Fig. 5). When we compared uninjured to injured muscle, we found an apparent decrease in the 1340 and 1320/cm vibrational bands in the injured muscle as well as an increase in the 1266/cm vibrational band. This suggests collagen-specific alterations within the tissue as a result of traumatic injury. In one case, a patient exhibited “pre-HO” muscle during a debridement procedure.
Upon Raman spectroscopic examination, it was clear that the tissue was indeed mineralized, even in “soft” tissue areas. Mineral vibrational bands at 1,070, 960, and 591/cm, typical of a carbonated apatite, were prominent in the spectrum. These vibrational bands are attributed to the phosphate and carbonate stretching modes of bone. Thus, Raman spectroscopy can potentially be utilized to identify areas of tissue affected by early HO as well as areas of tissue that may be predisposed to HO formation.

5 Conclusions

The potential of vibrational spectroscopy to provide detailed information, noninvasively, about molecular and even structural changes within the components of the wound bed itself enable a more thorough understanding of the wound-healing process. Vibrational spectroscopic modalities such as Raman and FTIR spectroscopy can provide an objective means of evaluation by monitoring key components of wound bed reepithelialization, such as keratin, elastin, and collagen; by identifying and quantifying bacterial load; and by detecting HO. These techniques have the potential to offer improved objective assessment of combat wounds, resulting in faster healing times, decreased infection rates, and decreased local and systemic complications of injury. This, in turn, will produce improved clinical outcomes, decreased patient morbidity, and reduced medical costs.

Acknowledgments

The views expressed in this paper are those of the authors and do not reflect the official policy of the Department of the Army, the Department of the Navy, the Department of Defense, or the U.S. government. We are military service members (or employees of the U.S. government). This work was prepared as part of our official duties. Title 17 U.S.C. § 105 states, “Copyright protection under this title is not available for any work of the United States Government.” Title 17 U.S.C. § 101 defines a U.S. government work as a work prepared by a military service member or employee of the U.S. government as part of that person’s official duties.

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We certify that all individuals who qualify as authors have been listed; that each has participated in the conception and design of this work, the analysis of data (when applicable), the writing of the document, and the approval of the submission of this version; that the document represents valid work; that if we used information derived from another source, we obtained all necessary approvals to use it and made appropriate acknowledgments thereof in the document; and that each author takes public responsibility for it.

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Predicting Wound Outcome from Raman Spectroscopic Data: Univariate versus Multivariate Techniques

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Acute Combat Wounds
The management of modern traumatic war wounds remains a significant challenge for clinicians.

- Extensive osseous and soft-tissue damage caused by blasts and high-energy projectiles.

The ensuing inflammatory response ultimately dictates the pace of wound healing and tissue regeneration.

The timing of wound closure or definitive coverage is subjectively based.

Despite the use and application of novel wound-specific treatment modalities, some wounds fail to close, or dehisce.
Surgical debridements are performed every 2-3 days.

- remove devitalized tissue
- decrease bacterial load

Negative pressure wound therapy (NPWT) is applied between debridements. NPWT promotes wound closure.

Wound assessment involves:

- patient’s general condition
- injury location
- adequacy of perfusion
- gross appearance of the wound
Monitor wound healing *in vivo*, i.e. monitor wound healing during surgical debridements.

- Is it the best time to close the wound?
- Is the wound developing HO?
- Is the wound infected? With what?

Develop an objective and predictive model for wound healing.
The Toolbox
Our Toolbox

- Real-time PCR
- Multiplex Protein Assay
- Raman Spectroscopy
- FTIR Imaging
- Thermography
- Visible Reflectance Imaging
- Bayesian Belief Network modeling
Wound is surgically cleaned

- Collect 1 cm³ tissue biopsy from center of wound bed
- RT-PCR
- Raman spectroscopy
- Micro culture

NPWT is applied

- Serum is collected
- Protein assay
- Effluent is collected
- Effluent assay
Preliminary Study – Raman Mapping Results

- Previous study demonstrated the potential of Raman spectroscopic analysis of wound biopsies for classification of wounds as normal or impaired healing.


- Impaired healing wounds demonstrate a significant decrease in the 1665/1445 cm\(^{-1}\) band area ratio compared to normal healing wounds - these results were corroborated with collagen gene expression in the same samples.

*Could this type of analysis be extended to intact wound biopsies and ultimately obviate the need for excisional wound biopsies?*
Chronicling Wound Healing with Raman Spectroscopy
Approximately 1 cm² tissue biopsy is excised from the center of the wound bed.

Tissue is fixed in 10% neutral buffered formalin for storage.

Prior to spectral acquisition, samples are rinsed in 0.9% NaCl saline solution.

Examine multiple spots across the tissue.

14 accumulations, 5s spectrum
7 accumulations, 10s spectrum
Monitoring a Wound Over Time

**FIRST DEBRIDEMENT**

- Normal
- Dehisced

**FINAL DEBRIDEMENT**

- Normal
- Dehisced

- Spectral differences between biopsies collected from normal healing wounds (n=12) and dehisced wounds (n=13) are not readily apparent.
Peakfitting for Spectral Deconvolution

<table>
<thead>
<tr>
<th>Raman Shift (cm$^{-1}$)</th>
<th>Vibrational Band Assignment</th>
<th>Component</th>
</tr>
</thead>
<tbody>
<tr>
<td>860</td>
<td>$\nu$(C-C)</td>
<td>nucleic acids</td>
</tr>
<tr>
<td>920, 940</td>
<td>$\nu$(C-N), $\nu$(C-C)</td>
<td>nucleic acids, keratin</td>
</tr>
<tr>
<td>1004</td>
<td>$\nu$(C-C) ring</td>
<td>phenylalanine</td>
</tr>
<tr>
<td>1040</td>
<td>$\nu$(C-C) skeletal</td>
<td>nucleic acids, protein</td>
</tr>
<tr>
<td>1125</td>
<td>$\nu$(C-C), $\nu$(C-N)</td>
<td>nucleic acids, protein</td>
</tr>
<tr>
<td>1250</td>
<td>$\nu$ (C-N) and $\delta$(N-H); Amide III</td>
<td>protein</td>
</tr>
<tr>
<td>1320</td>
<td>$\delta$(CH$_2$) twisting</td>
<td>nucleic acids, protein</td>
</tr>
<tr>
<td>1445</td>
<td>$\delta$(CH$_3$) and $\delta$(CH$_2$) scissoring</td>
<td>protein</td>
</tr>
<tr>
<td>1555</td>
<td>$\nu$(C=O); Amide †</td>
<td>aromatic amino acids, heme</td>
</tr>
<tr>
<td>1665</td>
<td></td>
<td>protein</td>
</tr>
</tbody>
</table>
Univariate Data Set

25 WOUNDS

81 RAMAN SPECTRA

BAND AREA, BAND HEIGHT, BAND WIDTH

10 RAMAN FEATURES

2,430 DATA POINTS
Univariate Analysis of Band Areas

1665 cm\(^{-1}\) | 1555 cm\(^{-1}\) | 1455 cm\(^{-1}\) | 1665 cm\(^{-1}\)/1445 cm\(^{-1}\) | 1320 cm\(^{-1}\) | 1250 cm\(^{-1}\) | 1040 cm\(^{-1}\) | 1004 cm\(^{-1}\) | 1665 cm\(^{-1}\)/1004 cm\(^{-1}\) | 940 cm\(^{-1}\)

- **Healed**
- **Dehisced**

* indicates significant difference.
Predicting outcome based on univariate analysis of band areas would misclassify 31% of dehisced wounds and 33% of healed wounds.
Univariate Analysis of Band Areas

- We can also look for changes in the band area profiles of healed and dehisced wounds over time, i.e. first debridement versus final debridement.

For wounds that heal, there are no statistically significant difference between calculated band areas chronologically.

For wounds that dehisce, the band area at 1250 cm$^{-1}$ demonstrates a statistically significant difference between the first and final debridement ($p < 0.03$).
Multivariate Analysis: BBNs

• Bayesian Belief Networks fall under the umbrella of machine learning.

  Input = empirical data ➔ output = patterns or predictions that are believed to be responsible for generating the data

• Applies the Bayes probabilistic theory for analyzing data sets.

• Bayesian Belief Networks (BBNs) demonstrate probabilistic relationships – relationships in which knowledge of the value of a variable affects the belief about the likelihood of other variables taking certain values.

  ➢ Data mining
  ➢ Prediction
Bayesian Modeling Data Set: Raman Only

- 25 WOUNDS
- 81 Raman Spectra
- Band Area, Band Height, Band Width
- 10 Raman Features
- 2,430 Data Points
Bayesian Belief Network (BBN) machine learning can elucidate some of the relationships between band area changes.

- $1555 \text{ cm}^{-1} \to 1250 \text{ cm}^{-1} \to 1320 \text{ cm}^{-1} \& 1665 \text{ cm}^{-1} \to 1665/1445 \text{ cm}^{-1} \to 1004 \text{ cm}^{-1}$

- $1040 \text{ cm}^{-1} \to 1250 \text{ cm}^{-1} \to 1665/1004 \text{ cm}^{-1} \to 1004\text{ and 1455 cm}^{-1} \to 1665/1004 \text{ cm}^{-1}$
A naïve Bayes model for outcome as the target reveals that outcome is largely dependent on mean 1040 cm$^{-1}$ and mean 1455 cm$^{-1}$ band area ratios.

Note, this data set incorporates different collection parameters, operators and regions of the biopsy.
Here, we generated a naïve Bayes model based on a training data set of all debridements except final debridements and used it to predict the wound healing outcome for final debridements.

Unfortunately, the accuracy of the prediction model is only 68% for normal healing wounds and 64% for dehisced wounds.
Bayesian Modeling Data Set: Raman and Clinical Data

25 WOUNDS → 81 RAMAN SPECTRA

BAND AREA, BAND HEIGHT, BAND WIDTH

10 RAMAN FEATURES ← 8 CLINICAL PARAMETERS

3,078 DATA POINTS
BBN Analysis of Band Areas and Clinical Data: Data Mining

What are the relationships between clinical data and Raman spectroscopic data?
Here, we generated a naïve Bayes model based on a training data set of all debridements and used it to predict the wound healing outcome for final debridements.
Here, we generated a naïve Bayes model based on a training data set of all debridements except final debridements and used it to predict the wound healing outcome for final debridements.

We’ve improved the accuracy of the prediction model - **76%** for *normal healing* wounds and **76%** for *dehisced wounds*. 
Multivariate Analysis of Spectral Data Set

- 25 WOUNDS
- 81 RAMAN SPECTRA
- 525,366 DATA POINTS
- 6,486 RAMAN FEATURES
Support Vector Machine

Supervised classification method based on a non-probabailistic binary linear classifier.

- assumes that groups are separated by a wide boundaries
- places samples in the two classes as far apart as possible
- able to handle non-linear data
- can be susceptible to overfitting
- may perform better on data sets with more examples of training and fewer labels
Support Vector Machine

SUPPORT VECTOR MACHINE

Developed a subset of spectra for a training data set (N=61), randomly selected from the whole data set. Used the entire spectrum with no pre-processing (derivatization, smoothing, normalization).

<table>
<thead>
<tr>
<th>Actual</th>
<th>Predicted</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>Healed</td>
</tr>
<tr>
<td>Healed</td>
<td>36</td>
</tr>
<tr>
<td>Dehisced</td>
<td>0</td>
</tr>
</tbody>
</table>

For a test data set (N=20), **80%** of the wound spectra were classified correctly as dehisced.
Supervised classification method based on Bayes’ formula.

- assumes a normal distribution
- assumes that the variability between each group has the same structure
- uses PCA to reduce dimensionality
- places samples in the same class close to one another but samples not in the same class far apart
- may perform better than SVM in cases of many labels and little training data
Linear Discriminant Analysis

Developed a subset of spectra for a training data set (N=61), randomly selected from the whole data set. Used the entire spectrum with no pre-processing (derivatization, smoothing, normalization).

<table>
<thead>
<tr>
<th>Actual</th>
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<tbody>
<tr>
<td>Healed</td>
<td>Healed</td>
</tr>
<tr>
<td></td>
<td>Dehisced</td>
</tr>
<tr>
<td>Healed</td>
<td>33</td>
</tr>
<tr>
<td>Dehisced</td>
<td>3</td>
</tr>
</tbody>
</table>

For a test data set (N=20), 90% of the spectra were classified correctly as dehisced.
Which Model to Choose?

- The models shown require significant optimization.
- We need to ensure complete data sets for training (include different acquisition parameters, multiple acquisitions, different instrument operators – i.e. build robustness into our model).
- The final evaluation of the prediction model will require both the highest sensitivity and specificity.
- Preliminary results demonstrate promise for the use of Raman spectroscopy to predict wound outcome.
Ongoing Projects

• Currently, we are continuing to build our biopsy database for Bayesian Belief Network modeling and other multivariate analysis techniques.

• We have over 250 tissue biopsies from over 50 patients, most of which we have collected spectra in triplicate (over 4 years).

• Peak fitting needs to be performed over the entire spectrum to determine which vibrational bands provide optimal sensitivity and specificity. Some regions of the spectrum require a more in depth peak fitting.

• Other aspects of the spectra, such as the fluorescence background, may also provide indications of the status of the tissue overall.
Project Goals

• The final prediction model will have to incorporate over 1.4M data points.

• Validate spectroscopic wound healing prediction models in a military and civilian population.
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WRNMMC
*General Surgery, Orthopaedics and Rehabilitation*

http://www.med.navy.mil/sites/nmrc/Pages/ccc_regenmed.htm
Disclaimer

• The multidisciplinary care of these patients would not have been possible without the dedicated efforts of everyone at WRAMC and NNMC. Both civilian and military personnel have rendered skilled and compassionate care for these casualties. All of our efforts are dedicated to those who have been placed in harm’s way for the good of our nation.

• The views expressed in this manuscript are those of the authors and do not reflect the official policy of the Department of the Army, Department of the Navy, the Department of Defense or the United States Government.

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• This study was approved by the Walter Reed National Military Medical Center Institutional Review Board in compliance with all Federal regulations governing the protection of human subjects.
In Operation Iraqi Freedom (OIF) and Operation Enduring Freedom (OEF), many blast injuries result in limbs with severe soft-tissue, bone, and vascular injuries (“mangled extremities”). These wounds are potentially mortal and the drawback of current methods to provide temporary perfusion, and the inability to measure extremity blood perfusion. A Yorkshire swine model has been developed in which two types of extremity injuries are simulated—via spectroscopic imaging to evaluate the limb locally. Infrared imaging shows us that limb temperature decreases at Day 1 (D1), Day 3 (D3) and Day 7 (D7).

Heart rate, body temperature, resting respiration, end tidal CO₂, pH, Nitrites, Ketones, Blood, Tissue K+, Na+, Lactate, protein, CK, AST, K+, Ca2+, Na+, Histopathology, Unobilogen, Nitrites, Blood, Protein, Ketones, Chemistery Panel, BUN, CR, AST, K+, Na+, Lactate, pH, Creatine phosphokinase (CK) levels may be the most sensitive indicator of muscle injury and may also be predictive of development of renal failure.

Glantounis et al. reported increased levels of ALT (p<0.05). Increase in potassium levels in all control groups except 2hr tourniquet. The significant time points are 120 minutes to baseline).

Limb ischemia is limited by the combination of devastating injuries, the drawback of current methods to provide temporary perfusion, and the inability to measure extremity blood perfusion. A Yorkshire swine model has been developed in which two types of extremity injuries are simulated—via spectroscopic imaging to evaluate the limb locally. Infrared imaging shows us that limb temperature decreases at Day 1 (D1), Day 3 (D3) and Day 7 (D7). The animals were observed twice a day for seven days. Their behavior was rated on general clinical appearance, food and water consumption, and provoked behavior.

STUDY DESIGN

Female Yorkshire swine of 3-4 months of age and 40-60kg in weight were used for this study. Samples were collected every hour during ischemia, after 30 minutes of reperfusion and at Day 1 (D1), Day 3 (D3) and Day 7 (D7).

• Female Yorkshire swine of 3-4 months of age and 40-60kg in weight were used for this study.
• Samples were collected every hour during ischemia, after 30 minutes of reperfusion and at Day 1 (D1), Day 3 (D3) and Day 7 (D7).
• Heart rate, body temperature, resting respiration, end tidal CO₂ and oxygen saturation were monitored throughout the procedure.

The animals were observed twice a day for seven days. Their behavior was rated on general clinical appearance, food and water consumption, and provoked behavior.

RESULTS AND DISCUSSION

CONCLUSIONS

• The differences seen between all control groups can be attributed to the length and type of ischemia.
• Tourniquet groups resulted in a stronger pathophysiologial response compared to occlusion.
• It is possible that a longer period of ischemia and a modification on where the vessel is occluded is needed for the occlusion groups to show larger physiological response. This modification would help compare and evaluate the different types of vascular injury that can be sustained in combat and civilian scenarios, thus establishing a well developed large animal model for limb ischemia.
• Although variable, these results provide us with a baseline for understanding the effects of IRI, the modifications that are needed for developing a better model and help us indicate which parameters to take under consideration when evaluating limb viability.

REFERENCES

The authors thank the surgical residents Dr. Jason Radleski, Dr. Earl Lee and Dr. Joe Caruso for contributing to this protocol. We also thank the Department of Regenerative Medicine and the Department of Pathology at the Naval Medical Research Center and the Department of Surgery at the Uniformed Services University of Health Sciences.

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