A Novel Pilot Study Using Spatial Frequency Domain Imaging to Assess Oxygenation of Perforator Flaps During Reconstructive Breast Surgery

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Introduction: Although various methods exist for monitoring flaps during reconstructive surgery, surgeons primarily rely on assessment of clinical judgment. Early detection of vascular complications improves rate of flap salvage. Spatial frequency domain imaging (SFDI) is a promising new technology that provides oxygenation images over a large field of view. The goal of this clinical pilot study is to use SFDI in perforator flap breast reconstruction.

Methods: Three women undergoing unilateral breast reconstruction after mastectomy were enrolled for our study. The SFDI system was deployed in the operating room, and images acquired over the course of the operation. Time points included images of each hemiabdominal skin flap before elevation, the selected flap after perforator dissection, and after microsurgical transfer.

Results: Spatial frequency domain imaging was able to measure tissue oxygen-hemoglobin concentration (ctO2Hb), tissue deoxygenated hemoglobin concentration, and tissue oxygen saturation (stO2). Images were created for each metric to monitor flap status and the results quantified throughout the various time points of the procedure. For 2 of 3 patients, the chosen flap had a higher ctO2Hb and stO2. For 1 patient, the chosen flap had lower ctO2Hb and stO2. There were no perfusion deficits observed based on SFDI and clinical follow-up.

Conclusions: The results of our initial human pilot study suggest that SFDI has the potential to provide intraoperative oxygenation images in real-time during surgery. With the use of this technology, surgeons can obtain tissue oxygenation and hemoglobin concentration maps to assist in intraoperative planning; this can potentially prevent complications and improve clinical outcome.

Key Words: perforator flap, breast reconstruction, microsurgery, perfusion monitoring, near-infrared imaging, spatial frequency domain imaging

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A novel, autologous flaps are well-established options for breast reconstruction after mastectomy.1-5 Commonly used options include the pedicled transverse rectus abdominis myocutaneous, free transverse rectus abdominis myocutaneous, and deep inferior epigastric perforator (DIEP) flaps.6-11 The dissection of these flaps requires adequate perfusion to ensure success; in addition, the advent of microsurgical reconstruction is further associated with potential risks and complications of microsurgery.

Technological advances are necessary to improve flap perfusion and minimize donor-site morbidity. In particular, vessel thrombosis, partial and total flap loss, and fat necrosis are associated with poor outcomes.6,12 As early recognition has been shown to lead to improved flap salvage and success,12 it is vital that vascular compromise is detected effectively to expedite surgical and therapeutic intervention for free flap salvage. Currently, the standard of care to monitor perfusion is the clinical observation of skin warmth, color, capillary refill, and dermal bleeding. This subjective approach leads to a wide degree of variability in reported rates of flap loss.13 Methods to identify early flap failure intraoperatively include thermal, flow, perfusion, and oxygen monitoring.14 Each method has limitations precluding widespread use. We provide here a review of each method, purposely excluding computed tomographic angiography and magnetic resonance angiography because they are not widely used intraoperatively.

- Thermal monitoring using noncontact infrared imaging has been previously evaluated, particularly in the form of dynamic infrared thermography. However, this technique requires cooling the surface of the flap (cold challenge), which can prove challenging to implement routinely into clinical practice.15,16
- Flow monitoring by Doppler effect (performed either using ultrasound waves or optically)17-20 and by laser speckle imaging21-22 has been used to assess blood flow in microsurgically anastomosed arteries and veins. However, such methods are typically noisy, can require vessels to be exposed, can require contact, suffer from surrounding vessel interference, and provide limited information about presence or absence of flow.
- Perfusion assessment, or angiography, can be performed intraoperatively using optical fluorescence imaging.23,24 This method attempts to monitor flaps by characterizing perfusion at the skin or superficial capillary level.25,26 Although this method can reliably visualize perforator arteries in flaps, it remains by nature superficial and requires the injection of an exogenous contrast agent.
- Tissue oximetry probes have been introduced for flap monitoring, as they are easy to use and are relatively inexpensive.27-30 The main advantage of this approach is direct measurement of tissue oxygenation, a surrogate for perfusion and indicator of tissue status. Such methods are intuitive to most clinicians as the technology is similar to pulse oximetry, which was introduced in the 1970s.31-33 These methods rely on light absorption from tissue endogenous
Spatial frequency domain imaging (SFDI) is a noncontact optical imaging method that allows for fast acquisition (<1 second) and accurate interrogation of tissue optical properties over large fields of view (>100 cm²). This method, when performed at multiple near-infrared (NIR) wavelengths allows measurement of concentrations of tissue constituents (namely, oxyhemoglobin, deoxyhemoglobin, lipids, and water) using Beer’s law at depths typically less than 5 mm. Briefly, the SFDI system consists of a multispectral NIR light source and a projector to illuminate the field with invisible spatially modulated patterns, a camera to acquire images, and a computer to process and display the images to the surgeon. The main advantage of this approach over other continuous wave multispectral wide-field imaging methods is that absorption of tissues is separated from scattering through measurement and not through empirical assumptions.

Our group has previously validated a clinically compatible SFDI system for intraoperative oxygenation imaging during porcine skin flap, bowel, and liver vascular occlusion comparing extracted oxygenation maps with a clinically approved oxygenation probe (ViOptix, Fremont, Calif). This previous study presented the detailed design of the imaging system, its validation during preclinical experiments and its translation into the clinic. In the current study, we present the results obtained with our SFDI system during our 3 patient, first-in-human, pilot study of flap oxygenation imaging during reconstructive breast surgery.

**METHODS**

**SFDI Clinical System**

The SFDI system used in this study has been optimized for intraoperative use. It is based on the Fluorescence Assisted Resection and Exploration (FLARE) clinical imaging system on which a projector and a multispectral NIR source have been integrated allowing the performance of SFDI acquisitions over a 16 × 12-cm field of view at a 45-cm working distance. This SFDI system meets all requirements for mechanical and electrical safety as defined in IEC 60601. Briefly, 1 SFDI acquisition consists of projecting 6 wavelengths onto the surgical field (670, 730, 760, 808, 860, and 980 nm) with 3 sets of patterns. These wavelengths have been selected to permit rapid and reliable oxygenation imaging. One set of pattern permits profilometry measurement and is used to correct for the sample’s surface profile variations. Two sets of patterns are used to extract optical properties. The acquisition is performed simultaneously on 2 NIR cameras, allowing the acquisition of 2 wavelengths at a time. Images acquired in real time during surgery are then processed to extract maps of \( \text{ctO}_2 \text{Hb} \), tissue deoxyhemoglobin concentration (\( \text{ctHHb} \)), and tissue oxygen saturation (\( \text{stO}_2 \)). A schematic and an actual picture of the SFDI system are provided in Figure 1. Extensive details about the FLARE system and the SFDI clinical system can be found elsewhere.

**FIGURE 1.** SFDI system. A, Schematics of the system. The light from the NIR light source is coupled into the projector and patterns of NIR light projected onto the field. Wavelengths shown in green and red are collected by NIR cameras 1 and 2, respectively. B, Picture of the clinical imaging system composed of a cart containing the NIR light source, control electronics, computer, mast, and arm holding the adjustable imaging head.
First-in-Human Pilot Study

The clinical pilot study was approved by the institutional review board of the Beth Israel Deaconess Medical Center and was performed in accordance with the ethical standards of the Helsinki Declaration of 1975. The institutional review board deemed the SFDI system a “nonsignificant risk” device. Study subjects were women undergoing unilateral or bilateral mastectomy and reconstruction with a microsurgical DIEP flap. Informed consent was obtained before surgery. Patient demographics include age range from 41 to 65 years, with 2 patients undergoing immediate unilateral reconstruction and 1 delayed unilateral reconstruction. Flap weights ranged from 326 to 890 g and perforators were chosen based on clinical observation (see Table 1).

Trained personnel deployed the imaging system into the operating room. The system was draped in sterile fashion using a shield/drape combination that could be applied by a single person (scrub nurse). After draping, the imaging head entered the sterile field and was positioned at a fixed distance. Images were acquired to extract profile information and optical properties at several wavelengths, as described previously.

The standard imaging protocol included 4 sets of SFDI measurements. The first 2 SFDI measurements were taken before flap elevation by positioning the system 45 cm over each hemiabdomen. On the basis of standard practice, the surgeon chose to dissect 1 set of vessels (from the right or left side of the abdomen). A third SFDI measurement was taken after dissection of the vessels through the intramuscular course and isolation of the selected perforator vessels and vascular pedicle. The flap was then transferred to the chest and a microsurgical anastomosis was performed attaching the deep inferior epigastric artery and vein to the internal mammary vessels. A final measurement was performed using SFDI at this time point. Results from the SFDI measurements were not accessible to the operating surgeons in this feasibility study, thereby not altering the standard of care.

Once discharged from the hospital, patients were seen during the routine course of follow-up. If no emergent complications occurred, patients were seen at 1 week after discharge, then at 6 weeks, 3 months, and 6 months after surgery. The reconstructed breast and surgical sites were evaluated at the time of the follow-up with emphasis on perfusion issues such as fat necrosis and partial flap loss.

RESULTS

SFDI Oxygenation Imaging Results

Four SFDI measurements were taken during the DIEP flap procedure. These measurements were taken at crucial time points, including each abdominal hemiflap (discarded and chosen abdominal flap), images of the chosen flap after elevation (postflap elevation), and after microsurgical attachment (postflap transplant). Extracted concentration maps of tissue oxyhemoglobin (ctO2Hb),

<table>
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<th>Patient No.</th>
<th>Age, y</th>
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<th>Flap Weight, g</th>
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<td>41</td>
<td>Delayed</td>
<td>2 lateral and 1 medial perforators</td>
<td>890</td>
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<tr>
<td>2</td>
<td>55</td>
<td>Immediate</td>
<td>1 lateral row perforator</td>
<td>326</td>
</tr>
<tr>
<td>3</td>
<td>65</td>
<td>Immediate</td>
<td>2 medial and 1 lateral perforators</td>
<td>524</td>
</tr>
</tbody>
</table>

FIGURE 2. SFDI oxygenation imaging results—Patient 1. Columns, from left to right, include abdominal skin flaps after preparation (discarded and chosen), skin flap after elevation, and skin flap after attachment (transplantation). The first row presents the concentration of tissue oxyhemoglobin (ctO2Hb), the second row the concentration of tissue deoxyhemoglobin (ctHHb), and the third row the tissue oxygen saturation images (stO2).
tissue deoxyhemoglobin (ctHHb), and tissue oxygen saturation (stO2) are shown for each patient in Figures 2, 3, and 4 (top, middle, and bottom row, respectively).

Patient 1
Abdominal Flaps
The SFDI maps showed that both flaps had good oxyhemoglobin and oxygen saturation levels initially. Both abdominal flaps showed good tissue oxygen saturation levels (76% and 74%, respectively) in a region of interest chosen in the middle of the flap (dashed black square in Fig. 2). In this case, the right flap was chosen for harvest and transplantation. The chosen flap had a higher ctO2Hb level compared to the discarded flap (157 and 111 μM, respectively) and a higher ctHHb level (49 and 38 μM, respectively).

Postelevation
As the vascular pedicle was isolated and the chosen flap elevated, tissue oxygen saturation decreased from 76% to 74% in our chosen flap, with a 20% decrease in ctO2Hb (126 μM) and 12% decrease in ctHHb (43 μM).

Posttransplantation
Once the flap was transplanted, measurements showed tissue oxygen saturation decreasing from 76% to 69%, with a 26% decrease in ctO2Hb (100 μM) and a slight increase in ctHHb (45 μM).

Patient 2
Abdominal Flaps
The SFDI maps showed both flaps with good oxyhemoglobin and oxygen saturation levels initially. Both abdominal flaps had identical tissue oxygen saturation levels at 71% in a region of interest chosen in the middle of the flap (dashed black square in Fig. 3). Here the left flap was chosen for harvest and transplantation. The chosen flap had a higher ctO2Hb level compared to the discarded flap (123 and 104 μM, respectively) and a higher ctHHb level (51 and 41 μM, respectively).

Postelevation
As the vascular pedicle was isolated and the chosen flap elevated, tissue oxygen saturation increased from 71% to 73% in our chosen flap, with a 31% decrease in ctO2Hb (85 μM) and 39% decrease in ctHHb (31 μM).

Posttransplantation
Once the flap was transplanted, measurements showed relatively unchanged tissue oxygen saturation at 72%, with a 28% increase in ctO2Hb (109 μM) and a 32% increase in ctHHb (41 μM).

Patient 3
Abdominal Flaps
The SFDI maps showed both flaps with low oxyhemoglobin and oxygen saturation levels initially. The left flap showed tissue oxygen saturation at 68%, whereas the right flap showed tissue oxygen saturation at 55% in a region of interest chosen in the middle of the flap (dashed black square in Fig. 4). Here the right flap was chosen for harvest and transplantation. The chosen flap had a lower ctO2Hb level compared to the discarded flap (78 and 84 μM, respectively) and a higher ctHHb level (63 and 38 μM, respectively).

Postelevation
As the vascular pedicle was isolated and the right flap elevated, tissue oxygen saturation increased from 55% to 60%, with a 12% decrease in ctO2Hb (69 μM) and 27% decrease in ctHHb (46 μM).
Posttransplantation

Once the right flap was transplanted, measurements showed a decrease in tissue oxygen saturation from 60% to 54%, with a 17% decrease in ctO2Hb (57 \text{\textmu}M) and a slight increase in ctHHb (48 \text{\textmu}M).

Postoperative Follow-up

Our 3 patients were seen in clinic for follow-up at the time points described. Patients were examined for any flap perfusion issues and postoperative photographs were taken (Fig. 5). In all 3 patients, no vascular or perfusion complications were observed in follow-up. There were no areas of fat necrosis or partial flap loss identified postoperatively. All 3 flaps healed without any sequelae and the patients all went on to complete their reconstructive course.

Quantification of Results

Localized measurements from similar regions of interests (dashed black squares in Figs. 2–4) were quantified and their mean and standard deviation plotted (Fig. 6). Tissue oxyhemoglobin content, tissue deoxyhemoglobin content, and tissue oxygen saturation levels were evaluated for all 3 patients. Time points include comparison of chosen and discarded flap as well as the chosen flap post-elevation and post-transplantation.

Tissue Oxyhemoglobin (ctO2Hb)

All 3 patients had differences in ctO2Hb between the 2 hemiflaps. In patients 1 and 2, the side with higher ctO2Hb level was chosen for dissection and transfer (41% and 18% higher, respectively). In patient 3, the side with the lower ctO2Hb level was chosen (7% lower). After elevation and vessel dissection, the flaps on all 3 patients demonstrated a decrease in ctO2Hb (−20%, −31%, and −12%, respectively). After microsurgical anastomosis and flap transfer, ctO2Hb varied from patient to patient but all showed lower levels compared to baseline before vessel dissection (−36%, −11%, and −27%, respectively).

FIGURE 4. SFDI oxygenation imaging results—Patient 3. Columns, from left to right, include abdominal skin flaps after preparation (discarded and chosen), skin flap after elevation, and skin flap after attachment (transplantation). The first row presents the concentration of tissue oxyhemoglobin (ctO2Hb), the second row the concentration of tissue deoxyhemoglobin (ctHHb), and the third row the tissue oxygen saturation images (stO2).

FIGURE 5. Postoperative clinical images. All 3 women underwent unilateral delayed or immediate DIEP flap microsurgical reconstruction. Photographs shown were taken during postoperative follow-up at 6 months (patient 1), at 10 months (patient 2), and at 9 months (patient 3).
All 3 patients had differences in ctHHb between the 2 hemiflaps. The chosen flap in all 3 patients had a higher ctHHb level (+29%, +24%, and +66%, respectively). After flap elevation, the flaps on all 3 patients showed a decrease in ctHHb (−12%, −39%, and −27%, respectively). However, after flap transfer, there was a slight improvement in ctHHb level (+5%, +32%, and +4%, respectively), although not returning to baseline before vessel dissection.

In patient 1, the flap with higher stO2 was chosen (76% vs 74%); in patient 2, the 2 flaps had the same level (71%); and in patient 3, the flap with the lower level was chosen (55% vs 68%). After flap elevation and vessel dissection, stO2 remained consistent and was slightly lower in patient 1 (−3%), although increased in patients 2 and 3 (+3% and +9%, respectively). After flap transfer in patient 1, the flap showed a decrease in stO2 compared to baseline levels (−9%), with little change in patients 2 and 3 (+1% and −2%, respectively).

The surgeon was blinded as to the results of the imaging system and clinical judgment was used during surgery to select the vessels and flap for harvest. For patients 1 and 2, SFDI confirms that the clinically chosen flaps for transplantation have higher oxygenation quantitatively with increased levels of oxyhemoglobin and oxygen saturation as compared with the discarded flap. However, for patient 3, SFDI suggests the discarded flap had better perfusion metrics, whereas the chosen flap was deemed clinically acceptable. Although these findings did not demonstrate clinical significance, knowledge of this information in patient 3 could have altered the course of surgery and may potentially improve outcomes in other cases.

The quantitative metrics identified with the SFDI system include tissue-level oxyhemoglobin, deoxyhemoglobin, and oxygen saturation. Importantly, the long-term postoperative clinical results from the transplanted flaps of our 3 patients correlated well with clinical results showing no areas of concern on our posttransplantation tissue oxygenation maps. Although large increases in deoxyhemoglobin are associated with compromised arterial perfusion surgical intervention. During elevation of the flap, SFDI would allow for assistance in the selection of flap used for harvest. During the dissection of the perforators and the isolation of the vascular pedicle, SFDI would allow identification of potential areas of diminished perfusion. After transplantation and microsurgical anastomosis, SFDI could allow for the immediate assessment of the flap status. The benefit to providing such measurements over large fields of view is the ability to identify areas of potential defect (hypoxia or ischemia) without prior indication for concern. Imaging at these time points may elicit immediate intervention whether it is the use of the contralateral flap, limiting the size of flap transferred, anticoagulation, or revision of the anastomosis.

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and large increases in oxyhemoglobin associated with compromised venous drainage, the thresholds for clinical perfusion deficits are still to be determined. In these 3 patients, although oxygen saturation was found to exhibit little variations, oxyhemoglobin and deoxyhemoglobin demonstrated larger variations throughout the course of dissection and transfer. In large animal studies from our laboratory, oxyhemoglobin and deoxyhemoglobin show distinct patterns of change that are more sensitive than oxygen saturation levels alone; these changes can potentially signal perfusion compromise at earlier time points.

Two distinct intraoperative patterns can be seen in our study (Fig. 6). In 1 pattern, seen in patients 1 and 2, higher oxyhemoglobin, deoxyhemoglobin, and oxygen saturation levels were seen in the chosen flap potentially demonstrating better overall perfusion. After elevation and microsurgical transfer, these levels were lower than baseline. The images obtained with SFDI correlated well with clinical assessment at the time of surgery in these patients with adequate tissue oxygen saturation levels (~70%), and high oxyhemoglobin concentration (150 μM), suggesting healthy flap status. In contrast, a different pattern is seen in patient 3, which depicts a flap with lower oxyhemoglobin and higher oxygen saturation levels after flap dissection. In this patient, the chosen flap showed some equilibration after flap dissection as demonstrated by an increase in oxygen saturation. After elevation and microsurgical flap transfer, the oxyhemoglobin and oxygen saturation were below baseline as previously mentioned. As free flaps undergo a period of hypoxia once harvested and reperfused, this transient period of decreased oxygenation in the flap was observed in both patterns. It would be interesting to understand how these metrics equilibrate over time and determine thresholds for inadequate perfusion. Finally, it is interesting to note that all patients had a similar deoxyhemoglobin concentration pattern over the procedure, with a decreased level after elevation and an increased level after transplantation.

The imaging system described also has the capability for simultaneous NIR fluorescence angiography and SFDI. This combination could allow for the identification of vascular structures, while also allowing for functional tissue perfusion mapping. Future work is underway to understand the relationship between flap outcome, fluorescence angiography, and SFDI.

CONCLUSIONS

We have designed and validated a system that is capable of wide-field tissue constituents imaging (oxyhemoglobin, deoxyhemoglobin, and oxygen saturation) based on SFDI and multispectral principles. Our SFDI system was translated to clinical use in a first-in-human pilot study, where skin flap oxygenation was imaged during reconstructive breast surgery intraoperatively. Our study demonstrates proof of concept and feasibility of using SFDI for tissue perfusion mapping.

REFERENCES


