Optical Imaging of Voltage and Calcium in Cardiac Cells & Tissues

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Abstract: Cardiac optical mapping has proven to be a powerful technology for studying cardiovascular function and disease. The development and scientific impact of this methodology are well-documented. Because of its relevance in cardiac research, this imaging technology advances at a rapid pace. Here, we review technological and scientific developments during the past several years and look toward the future. First, we explore key components of a modern optical mapping set-up, focusing on: (1) new camera technologies; (2) powerful light-emitting-diodes (from ultraviolet to red) for illumination; (3) improved optical filter technology; (4) new synthetic and optogenetic fluorescent probes; (5) optical mapping with motion and contraction; (6) new multiparametric optical mapping techniques; and (7) photon scattering effects in thick tissue preparations. We then look at recent optical mapping studies in single cells, cardiomyocyte monolayers, atria, and whole hearts. Finally, we briefly look into the possible future roles of optical mapping in the development of regenerative cardiac research, cardiac cell therapies, and molecular genetic advances.

Key Words: arrhythmia ■ fluorescence ■ multiparametric ■ optical mapping ■ optogenetics

The study of the electrophysiological properties of the healthy and diseased heart and the coupling between excitation and contraction in its component cells and tissues provide important mechanistic insights into how life-threatening cardiac arrhythmias can occur.1 This information may be used to aid in the generation of computer models to make predictions about arrhythmogenesis and ultimately for design of rational antiarrhythmic therapies.2 Traditionally, surface electrodes have been, and continue to be, used to measure extracellular cardiac electric potentials. However, this surface contact mapping suffers from low spatial resolution, low depth of field, far-field effects, and interference from electrical stimulation electrodes. Optical mapping using fluorescent probes to look at physiological parameters offers a higher resolution and less invasive method to study the electrical activity of the heart and cardiac cells. In addition, fluorescent methods have revolutionized the analysis of the molecular dynamics underlying the coupling between cardiac excitation and contraction.

Fluorescence techniques are widely used in cardiac research. Immunocytochemistry using immunolabeled fluorescent probes provides subcellular molecular structure data, whereas parameter-sensitive probes (ie, ion selective or voltage) loaded into live cells and tissue provide measurement of physiological function.3,4 Hence, the use of fluorescence has led to detailed mechanistic insight into the molecular and cellular processes that are vital to cardiac function. For example, the use of calcium-sensitive fluorescent probes led to discovery of the precise cellular and molecular mechanisms of cardiomyocyte excitation–contraction coupling5 and how this vital coupling is altered in...
pathophysiological states such as heart failure.6,7 The heart works as a functional syncytium of many electrically and mechanically connected cardiomyocytes, so it is imperative to study physiology of the whole heart or multicellular tissue preparations to provide a conceptual framework for single cell data.

Fluorescence imaging using voltage-sensitive dyes was first used by Salzberg et al8 in 1973 to record changes in axonal impulse propagation. Since then, fluorescent dyes have been used to study a wide array of electrically excitable organs and tissues. In cardiac tissues, simultaneous potentiometric dye fluorescence recordings from multiple sites have been made by arrays of photodiodes, photomultiplier tubes, laser scanning systems, charge-coupled device (CCD) cameras, and their more recent derivatives.9–11 Furthermore, whole heart optical mapping techniques have been developed and utilized to study electrophysiological mechanisms of cardiac arrhythmias.12,13 Since the early 1990s, optical mapping with CCD cameras has demonstrated to be an extremely useful tool in the analysis of the complex patterns of electrical wave propagation in both atrial and ventricular arrhythmias in isolated animal hearts.14

Now, optical mapping makes it possible to measure action potential and calcium wave propagation at high spatiotemporal resolution.15 The complex dynamic coupling between these two electrophysiological parameters is critical for proper cardiac function, thus underscoring the vital need to monitor multiple parameters simultaneously. New techniques are being developed to extend multiparametric imaging to include fluorescent monitoring of the metabolic state of the heart together with electrophysiological monitoring.16 Fluorescent reporters have been developed more recently for magnesium, sodium, potassium, pH, nitric oxide, redox state, and oxygen content. Even subcellular organelle electrophysiology now can be probed optically.17,18 Simultaneous multiparametric imaging of membrane voltage and pH for, example, will provide new insights into cellular and molecular mechanisms of arrhythmias in ischemia. The general goal of cardiac optical mapping is to provide a better understanding of cardiac electrophysiological function during health and disease. Cardiac optical mapping techniques have been reviewed extensively.11,19 Here, we discuss some of the new optically based developments for cardiac mapping since that time and present emerging technologies that will prove pivotal in the advancement of the cardiac optical mapping field.

### Optical Mapping Instrumentation and Fluorescent Probes

**Photodetectors for Optical Mapping**

Optical mapping experiments require a light source to illuminate cardiac preparations treated with a fluorescent reporter and a detector to sense the changes in fluorescence that represent physiological changes in transmembrane potential (V_m) or intracellular ion concentrations (e.g., Ca^{2+}). An optical mapping set-up is displayed schematically in Figure 1. The optical detector is perhaps the most essential component of the set-up.10 In this section, we review the various options for high-resolution optical mapping detectors (cameras) and focus on technologies that have emerged and are being utilized.

The propagating electrical impulse that triggers every heart beat is rapid and short-lived. Because of the low signal strength (ie, change in fluorescence over the background fluorescence [ΔF/F]) of current voltage-sensitive fluorescent dyes and the high speed of electrical wave propagation, high-speed low-noise photodetectors are required. There are currently only a limited number of useful detectors that offer high spatiotemporal resolution for cardiac mapping. The available technologies include photodiode arrays (PDAs), photomultiplier tubes, CCD cameras, and complementary metal oxide semiconductor cameras. Photomultiplier tubes offer high temporal resolution acquisition of fluorescence signals, but little to no spatial information. PDA detectors and CCD cameras have predominantly been used for cardiac imaging applications because of their high spatial and temporal resolutions.20–22

For optimal optical mapping of the heart and other cardiac preparations, CCD/complementary metal oxide semiconductor technologies are proving to be far superior to PDAs in terms of spatiotemporal resolution because of the massive number of pixels and fast data-streaming rates to a computer.
However, CCD sensors offer a slower full-frame rate than PDAs (because of the much larger number of pixels), but this can be overcome by increased pixel binning. Pixel binning, however, decreases spatial resolution, and thus it is imperative to utilize the optimal resolution relative to the preparation being used or the question being asked. For very low light level signals such as those from cell culture systems using monolayers or strands, newer CCD technology has been developed that provides increased sensitivity with low noise. The two CCD derivatives are electron-multiplying CCDs (EMCCDs) and intensified camera systems (intensified CCD/complementary metal oxide semiconductor).10

The most recently developed concept for increasing sensitivity of CCDs is on-chip electron multiplication, referred to as EMCCD technology.23 EMCCDs utilize impact ionization (an avalanche process) to produce secondary electrons before conversion to a voltage signal.24 An on-board fully chip-incorporated “gain register” provides electron multiplication in a serial process that involves the application of high electrical fields. Major benefits of EMCCDs over older CCD technology include lower readout noise at higher acquisition rates, lower multiplicative noise, and improved quantum efficiency compared to intensifiers. New EMCCDs show promise as high-resolution cameras, especially for cell monolayers that require high sensitivity and temporal resolution,25 and recently they have been suggested to provide the highest potential for fast imaging at very low light levels, as encountered in mapping of voltage and calcium waves in cultured cells.26 The latest thermoelectrically cooled EMCCD cameras permit user-specified EM gain, with minimal decrease in the signal-to-noise ratio, for low light level applications. For the whole heart, which emits high fluorescence, large pixel well depths and dynamic ranges are critical. Modern EMCCD cameras (for example, Evolve cameras from Photometrics and SciMOS cameras from Fairchild Imaging) with these features make for a camera that is versatile enough to be used for either whole heart optical mapping where signal intensity is strong or cell culture systems where fluorescence signal intensity is weak and dynamic fluorescence is fast (ie, action potential propagation). As photodetector and CCD camera technologies advance, so will the cardiac optical mapping field and our understanding and treatment of cardiac arrhythmias.

Illumination
Proper illumination is pivotal to successful optical mapping experiments. Whether the preparation is a Langendorff-perfused heart or a monolayer of cells, stable and even illumination during recording of fluorescence changes is important for high signal-to-noise recordings of physiological parameters. Multiparameter cardiac imaging often relies on switching between distinct wavelengths of light, thus making the choice of illuminating sources essential for these technically challenging experiments.

Traditional lighting methods have relied on xenon, mercury, or halogen lamps for illumination and excitation of fluorescent reporters.11 Illumination duration has been controlled by shuttering devices and moving parts, which can be impractical when stable and fast multicolor imaging is required (ie, multiparametric imaging). Stable illumination requires a highly regulated voltage supply and feedback control system, and multicolor illumination requires filter wheels, galvanometers, and shutters to switch between excitation wavelengths. Arc lamps are unstable because of plasma oscillations and thermal runway.27 Newer illumination technology, such as light-emitting diodes (LEDs), offers a more stable and easy-to-use alternative to traditional lighting sources for standard cardiac optical mapping.11,27,28

LED illumination is cost-effective, energy-efficient, portable, and flexible.10 The main concern with LED use is adequate cooling and sufficient output power. New high-power LEDs from ultraviolet to red wavelengths are now available with sufficient power and stability for more light-demanding preparations like the whole heart. Computer control of multiple LED lights allows for complex and rapid wavelength switching, thus enabling multiparametric and ratiometric imaging for quantitative fluorescence measurements.28,29 To date, the majority of optical mapping studies have utilized single illumination sources to study a single physiological parameter, typically using a potentiometric dye.

LED illumination, which easily enables multiparametric and ratiometric imaging because of the ability to rapidly change between excitation sources in real-time and during the experimental procedure, will increase the rate and breadth of knowledge acquisition in the cardiac optical mapping field.

Potentiometric Dyes
Several decades ago, molecular probes were discovered that embedded into the plasma membrane of neurons and cardiac cells and exhibited interactions with the electric field. The resulting molecular rearrangement of the probe within the membrane led to a change in fluorescence or absorption that reported alterations of membrane potential.30,31 Potentiometric probes enable high spatial resolution measurements of membrane potential in subcellular compartments (eg, mitochondria),32 single cells, monolayers of cardiac cells,33,34 and in the whole heart.35 Fluorescence of these dye molecules changes linearly with membrane potentials in the physiological range36 and modern probes can yield >10% fractional changes in fluorescence. Figure 2 illustrates the tight correlation between fluorescence changes induced by transmembrane changes in voltage and sharp microelectrode recordings in various anatomic regions of the heart.

Potentiometric dyes are classified into fast-response and slow-response dyes based on their response times and mechanism of voltage sensitivity;37 fast-response dyes report voltage changes on the order of microseconds.38 The most typical used are the styryl dyes developed by Loew et al,39 of which the most commonly used are the aminonaphthylethenylpyridinium (ANEPP) dyes, di-4-ANEPPS and di-8-ANEPPS. Both dyes yield a uniform 10% change in fluorescence per 100-mV change in membrane potential, but di-4-ANEPPS rapidly internalizes in cells, thus making it useful only for short-term experiments. However, di-8-ANEPPS is better retained in the outer leaflet (longer membrane anchor) of the plasma membrane, which also contributes to its reported lower phototoxicity (related to higher photostability) to cells.40,41 Although di-4-ANEPPS is more commonly used for whole heart optical mapping, di-8-ANEPPS...
is also used for that purpose in some laboratories. Both ANEP dyes respond to increases of membrane potential (hyperpolarization), with a decrease in fluorescence emission when excited at their optimal wavelengths, which depends on the synthetic/biological preparation where the dyes are loaded (ie, the dyes’ properties are sensitive to the local environment). According to Loew et al, the ΔF/F ranges between 7% and 10%, depending on the preparation. Ratiometric cardiac action potential recording using di-4-ANEPPS and pulsed LED excitation recently has been described. Styryl voltage-sensitive dyes have been widely used and successfully used in cardiac cells and tissues. However, their utility has been somewhat limited because their excitation wavelengths are restricted to the short wavelength (blue to green) range. Longer excitation/emission wavelength probes (near infrared) can minimize interference from endogenous chromophores and improve recording from deeper tissue layers. Voltage dyes in multicellular preparations have been used extensively with much success because tissue is robust and provides fluorescence past the lens focus, providing a large ΔF/F. Single-cell and monolayer imaging, however, suffer from dye phototoxicity and low ΔF/F. New voltage-sensitive dyes that are less cytotoxic and provide larger ΔF/F are required to extend optical mapping to the single cell level. New longer wavelength voltage-sensitive dyes are red-shifted, which can be advantageous for combining with other dyes (eg, Ca²⁺), seeing through blood, and having bigger ΔF/F signals. Recently generated long wavelength voltage-sensitive dyes are designed to record cardiac action potentials (APs) from deeper layers in the heart. To do this, the emission spectrum of styryl voltage-sensitive dye was red-shifted by incorporating a thienyl group in the polymethine bridge to lengthen and retain the rigidity of the chromophore. Seven distinct dyes, called Pittsburgh I to IV and VI to VIII (Pittsburgh I–VIII) were synthesized in the Salama laboratory and characterized with respect to their spectral properties in organic solvents and heart muscles. Pittsburgh voltage-sensitive dyes exhibited two absorption, two excitation, and two voltage-sensitive emission peaks, each with very large Stokes shifts (>100 nm). As reported, hearts (rabbit, guinea pig, and Rana pipiens) were effectively stained by injecting a bolus (10–50 µL) of stock solution of voltage-sensitive dye (2–5 mmol/L) dissolved in dimethylsulfoxide plus low-molecular-weight Pluronic (16% of L64). Other preparations were better stained with a bolus of voltage-sensitive dye (2–5 mmol/L) Tyrode solution at pH 6.0. APs measured with a fast CCD camera showed that Pittsburgh I exhibited an increase in fractional fluorescence (ΔF/F=17.5%) per action potential at 720 nm (emission), with 550 nm excitation and ΔF/F of -6% per AP at 830 nm excitation. The long wavelengths, large Stokes shifts, high fluorescence change on membrane depolarization, and low baseline fluorescence make Pittsburgh dyes a valuable tool in optical mapping and for simultaneous mapping of APs and intracellular Ca²⁺. The spectra for potentiometric dyes have been tested almost exclusively in mammalian systems. Because optical mapping techniques are extended to studies using genetically malleable zebrafish and drosophila hearts, it will be critical to determine the spectra of these dyes in nonmammalian systems.

The Loew laboratory recently reported developing new potentiometric styryl dyes with red excitation wavelengths and near-infrared emission. Three dyes for cardiac optical mapping were investigated in depth from several hundred dyes containing 47 variants of the styryl chromophores. They recorded absorbance and emission spectra in ethanol and multilamellar vesicles, as well as voltage-dependent spectral changes in a model lipid bilayer. Optical action potentials were recorded in tissues of rats, guinea pigs, and pigs, and compared with those of di-4-ANEPPS. The voltage sensitivities of these new potentiometric indicators were shown to be as high as those of the widely used ANEP series of probes. In addition, because of molecular engineering of the chromophore, the new dyes provide a wide range of dye loading and washout time constants. These dyes will enable a series of new experiments requiring the optical probing of thick or blood-perfused cardiac tissues, as well as deeper recordings in the myocardial tissue.

In 2007, the first characterization of two new infrared styryl dyes di-4-ANBDQPO and di-4-ANBDQBS optimized for blood-perfused tissue and intramural optical mapping was reported. These dyes had excitation/emission wavelengths shifted >100 nm toward the red. Voltage-dependent spectra were recorded in a model lipid bilayer and a comprehensive examination of the new dyes was conducted in multiple cardiac preparations, including Langendorff-perfused pig hearts using Tyrode solution and a blood–saline mix, and in coronary-perfused pig right ventricular wall preparations. In cardiac tissues, the optimal excitation (650 nm) was >70 nm beyond the absorption maximum of hemoglobin, pointing to the potential utility to optically map through blood. Signal decay half-life attributable to dye internalization was 80 to 210 minutes, which is five-times to seven-times slower than for di-4-ANEPPS. In transillumination mode, ΔF/F was as high as 20%. In blood-perfused tissues, ΔF/F reached 5.5% (1.8-times higher than for di-4-ANEPPS). In effect, these dyes provide both high-voltage sensitivity and five-times to seven-times slower internalization rate compared to conventional dyes. They are optimized for deeper tissue probing and
optical mapping of blood-perfused tissue and also can be used for conventional applications. In the future, the continued development of new red-shifted voltage-sensitive dyes will pave the way for endoscopic mapping in vivo using cardiac catheterization, which may one day become a useful clinical tool to optically map and precisely diagnose arrhythmias and perhaps guide cardiac ablation procedures.

**Calcium-Sensitive Dyes**

Calcium cycling in cardiomyocytes is a vital component of cardiac excitation–contraction coupling.48,49 Cardiac excitation–contraction coupling is crucial for proper heart function, and the ubiquitous second messenger, \( \text{Ca}^{2+} \), is central to this elegant coupling.50 The action potential causes \( \text{Ca}^{2+} \) influx through activation of L-type voltage-gated \( \text{Ca}^{2+} \) channels. This \( \text{Ca}^{2+} \) triggers release of \( \text{Ca}^{2+} \) from intracellular stores of the sarcoplasmic reticulum that activates contraction. \( \text{Ca}^{2+} \) release from the sarcoplasmic reticulum is mediated by \( \text{Ca}^{2+} \) release channels (ryanodine receptors) that are activated by localized subsarcolemmal \( \text{Ca}^{2+} \) entry into the cell via L-type \( \text{Ca}^{2+} \) channels and this process is commonly referred to as \( \text{Ca}^{2+} \) induced \( \text{Ca}^{2+} \) release.5 In pathological conditions such as heart failure, dysregulation of cellular \( \text{Ca}^{2+} \) homeostasis may activate \( \text{Ca}^{2+} \)-dependent currents that can influence action potential duration and trigger spontaneous membrane depolarizations.51,52 In fact, mishandling of intracellular \( \text{Ca}^{2+} \) in cardiomyocytes contributes to contractile dysfunction and arrhythmogenesis in failing hearts.53,54 Therefore, simultaneous measurement of action potential and \( \text{Ca}^{2+} \) wave propagation is essential to provide mechanistic insight into acquired arrhythmias associated with heart failure and inherited \( \text{Ca}^{2+} \)-mediated arrhythmias such as catecholaminergic polymorphic ventricular tachycardia.55–57

To minimize perturbation of the \([\text{Ca}^{2+}]_i\) dynamics in cardiac cells and tissue, the choice of \( \text{Ca}^{2+} \) dye is critical for acquiring accurate measurements of the amplitude and time course of \([\text{Ca}^{2+}]_i\), transients. For cardiomyocytes and tissue, which show large and rapid changes in \([\text{Ca}^{2+}]_i\), a low-affinity and rapidly responding dye is necessary.58 Other widely used \( \text{Ca}^{2+} \) dyes, such as Fluo-4, Fluo-3, and Fura-2,59 have a relatively high affinity for \( \text{Ca}^{2+} \). This can artificially prolong the \( \text{Ca}^{2+} \) transient and confound interpretation (ie, the dye acts as a chelator and clings on to \( \text{Ca}^{2+} \) ions for too long). Low-affinity calcium dyes provide more accurate measurement of calcium dynamics.60 The most ideal \( \text{Ca}^{2+} \) indicator molecule would combine the option of ratiometry for amplitude quantification with low \( \text{Ca}^{2+} \) affinity, such as the newly developed Fura-4F dye.61 Ratiometric optical mapping has been technically challenging using traditional light sources that require moving parts for filter switching between excitation lights. Recently, this technological challenge has been overcome by the use of electronically controlled LED illumination, thus enabling quantitative assessment of calcium wave amplitudes and dynamics in whole hearts.28 Small molecule dyes are useful because of their high signal-to-noise ratio; there is a wide range of indicators with various excitation/emission spectra and affinities for \( \text{Ca}^{2+} \). Any untoward effects of small-molecule calcium dyes are easily overcome because of the ability to control the concentration of dye that enters cardiac cells. Thus, small-molecule calcium dyes are most commonly used for optical mapping experiments and this will likely continue into the future.

Genetically encoded \( \text{Ca}^{2+} \) indicator proteins represent a new generation of calcium-sensing molecules. Genetically encoded \( \text{Ci}^{2+} \) indicator proteins offer nominal advantages over small-molecule indicators such as Fura-2 and Fluo-4, which include cell-specific calcium mapping and the possibility for chronic imaging over days and weeks.62 A well-known limitation of genetically encoded \( \text{Ca}^{2+} \) indicator proteins, however, is their slow response time because of the slow on-and-off kinetics of calcium binding. This attribute makes genetically encoded \( \text{Ca}^{2+} \) indicator proteins less suitable for cardiac optical mapping, but development of genetically encoded proteins with faster response times will provide powerful new tools for cell-specific \( \text{Ca}^{2+} \) imaging within the whole heart.

**Optical Mapping Limitations and Solutions**

**Motion Artifacts**

One limitation of optical mapping of the heart is imaging artifact caused by cardiac contraction. Motion artifacts obscure the optical recordings of voltage or calcium transients and thus prevent accurate measurement of parameter dynamics. Mechanical, pharmacological, and imaging methods have been developed to overcome this limitation.19 Mechanical restriction of heart motion without physiological implication reportedly works for small rodent hearts.20 However, the heart is well-known to be sensitive to mechanical distortion,63 thus limiting the utility of this approach. Pharmacological approaches have included the use of calcium channel blockers64 or myosin inhibitors such as 2,3-butanedione monoxime and blebbistatin.65,66 The 2,3-butanedione monoxime is non-specific; it also blocks function of membrane ion pumps, including SERCA2, thus having significant effects on cardiac electrophysiology and limiting its usefulness. Blebbistatin is a specific myosin II inhibitor that reportedly prevents contraction but does not have any noticeable effects on cardiac electrophysiology,67 thus making it a more useful tool for cardiac optical mapping. However, it is important to consider that inhibition of the primary consumer of intracellular ATP (myosin) may confound experiments studying mechanisms of arrhythmias that can be influenced by the intracellular metabolic state, and any mechano-electric feedback is also eliminated that may play a critical role in arrhythmias.68,69 Additionally, the contribution of the myofilaments to cardiac arrhythmias is beginning to be realized, and eliminating their function in optical mapping experiments may have profound effects on the study of arrhythmia mechanisms.68,69 Despite their inherent limitations, motion blockers/uncouplers have helped optical mapping studies tremendously. Use of ratiometric imaging techniques20 and postacquisition motion tracking70 may permit the mapping of contracting tissue.

**Photon Scattering Effects**

The majority of optical mapping studies to date have performed epicardial recording of transmembrane voltage signals. However, because of the thickness of the ventricular
Multiparametric Optical Mapping
Simultaneous Voltage and Calcium Optical Mapping

In 2004, multiparametric imaging was considered to be a field "...in its infancy." Today, this field has advanced extensively because of technological developments in LED illumination, optical filters, and fluorescent dyes. However, the bulk of cardiac optical mapping data to date has been generated by measuring only a single dye for a single parameter, typically monitoring either $V_m$ or intracellular $Ca^{2+}$. The complex interrelationship between these physiological parameters is critical for cardiac function and alterations of the normal dynamics of either can lead to cardiac arrhythmias. Membrane depolarization triggers intracellular $Ca^{2+}$ transients and changes of intracellular $Ca^{2+}$ can affect $V_m$ by modulating the function of multiple ionic currents such as L-type $Ca^{2+}$, chloride, and sodium–calcium exchanger currents. Therefore, to gain a precise understanding of the complex nature of the electric activity of every heart beat and to precisely identify mechanisms of arrhythmias, it is ideal to view both parameters simultaneously.

In 2000, Choi et al developed and reported a method to simultaneously map $V_m$ and $Ca^{2+}$ on the epicardium of perfused hearts by staining cells with a voltage-sensitive dye (RH-237) and loading also the cytosol with a fluorescent $Ca^{2+}$ indicator (rhod-2). The fluorescence characteristics of these two dyes are such that one illumination wavelength was used to excite each dye simultaneously; through the use of dichroic mirrors, the unique emission wavelength of light was directed to one of two PDAs. In these experiments, the image of the anterior region of the heart was focused on the array and each photodiode recorded activity from a 0.8-mm×0.8-mm area of epicardium, thus greatly limiting the area of view to a small portion of the heart. The use of multiple PDAs required a complicated six-step procedure to precisely focus and align the arrays, thus making this technology cumbersome and limiting its applicability to only the very engineering savvy cardiac researcher. An important feature of this optical apparatus is its extremely fast kinetic resolution (4000 frames/s; because of the small number of pixels) for each parameter to very accurately determine the temporal relationship between $V_m$ and $Ca^{2+}$. Though a tour de force at the time, this technology provided low spatial resolution (16×16 PDA) and was challenging to set-up and align.

Since that time, new detector configurations have been developed in an attempt to utilize higher-resolution CCD cameras and to simplify detector alignment. In 2004, Omichi et al used a two CCD camera system to increase spatial resolution of dual $V_m$-$Ca^{2+}$ mapping for the study of dynamics during ventricular fibrillation (VF). In this configuration the two cameras, with appropriate emission filters, were placed side-by-side and were manually aligned to view approximately the same area of the heart. Although the set-up provided higher spatial resolution using CCD detectors, the manual alignment of the multiple sensors introduced inaccuracies to the measurements. In 2009, Holcomb et al described a simplified method for detector alignment. The
advantage of this system over others is in the software camera calibration routine that eliminates the need for precise camera alignment for acquisition and processing of data, as is the case with complicated PDA alignment. Figure 1 shows an optical mapping set-up that uses a single camera (eliminating alignment needs) for multiparametric imaging (four excitation wavelengths), and Figure 4 shows simultaneous optical mapping of intracellular calcium and membrane voltage during sustained reentry in a neonatal rat ventricular myocyte monolayer, also using a single-camera system.

Some Recent Applications of Optical Mapping at Varying Levels of Integration

Optical mapping has contributed immensely over the past 20 years to the understanding of the dynamics of electric wave propagation in the most complex cardiac arrhythmias, including atrial fibrillation (AF) and VF; it also has advanced knowledge on the mechanisms of action of antiarrhythmic agents and the mechanisms of cardiac defibrillation, all of which will ultimately contribute to better care of patients. In the following sections, we briefly discuss some of the recent scientific highlights in this field of research.

Single Cell Optical Mapping

Single cell optical mapping offers the potential for higher throughput testing of treatment effects on the action potential than traditional patch electrode techniques. Additionally, voltage-sensitive dyes allow monitoring of cell excitation heterogeneities, providing spatial information not possible with electrode recordings. Windisch et al\(^8\) developed the first optical micromapping system in which single cell membrane potential was monitored optically using di-4-ANEPPS. However, single cells isolated from rodent hearts are only 5- to 10-μm-thick and in larger animals thickness may be just >10 μm, thus making light levels from potentiometric and calcium indicators challenging to detect. In addition, di-4-ANEPPS suffers from being cytotoxic to the extent that it has not been widely used for single cell electrophysiological mapping. More recently, Sharma et al\(^9\) used di-8-ANEPPS to study how cardiac cells respond to applied electric fields. Since then new technologies have emerged, including a new red-shifted voltage dye, di-4-ANBDQBS and EMCCD cameras have been used for optical AP recording in single cardiomyocytes.\(^9\) Importantly, as shown in Figure 5, loading di-4-ANBDQBS did not alter single cell APs recorded with a micropipette. The di-4-ANBDQBS yielded fluorescent signals with very high ΔF/F (19.2±4.1%) and signal-to-noise ratios (40±13.2), representing a major advancement in ANEP styryl dyes.\(^9\) Thus, the presented technique provides a unique opportunity for high-throughput noninvasive AP recording in isolated cardiomyocytes.

Optical mapping can be combined with other single cell methods such as stretching to elucidate the interdependence between the mechanical state of the myocardium and its electric activity.\(^9\) However, the information to date has been limited by the technical difficulties associated with stretching single myocytes and recording electrophysiological parameters optically, especially V_m. Recently, Nishimura et al\(^4\) combined the carbon fiber cell-stretching technique and ratiometric measurement of di-8-ANEPPS. A schematic and example of their system are depicted in Figure 6. They found that during systole, stretching caused depolarization that prolonged the action potential duration without affecting the peak amplitude. The effect, however, was only significant in the late phase of the AP. Stretching quiescent myocytes depolarized the membrane potential in amplitude and speed dependent ways and was suppressed by cytochalasin-D treatment, suggesting participation of the cytoskeleton in cardiac
mechanotransduction. Finally, ion replacement experiments revealed that although Na\(^+\) was the dominant charge carrier for large amplitude stretches, Ca\(^{2+}\) permeation was involved in small amplitude stretches, suggesting stretch amplitude ionic selectivity.

Monolayer Optical Mapping

The structural complexity of the heart has prompted the use of simplified cell culture systems to study excitation wave dynamics in cardiac tissues to precisely define cellular heterogeneity on impulse propagation.\(^{41,95,96}\) To understand cell network behavior, macroscopic imaging had to be used. However, cell culture systems present more challenges than the whole heart because fluorescence signals are depth-integrated.\(^{71}\) Whole heart fluorescence signals are recorded from myocardial tissue that can be millimeters in thickness, whereas cardiomyocyte culture systems (eg, neonatal rat ventricular myocyte monolayers) are only 5 to 10 \(\mu\)m in thickness.\(^{97}\) This is an order of magnitude thinner than the lower limit for the depth-of-field of useful macroscopic imaging lenses, thus making monolayer macro-mapping technically challenging. The first systematic application of optical mapping to record impulse propagation in strands of cardiomyocytes was in 1993 using PDAs.\(^{98}\) Bub et al\(^99\) then used a CCD-based system to map calcium in large areas of cardiac monolayers. For action potential measurements, styryl dyes (di-4-ANEPPS, di-8-ANEPPS, and RH-237) are most widely used for optical mapping in myocyte monolayers.\(^{100-102}\)

Multiparametric imaging of \(V_m\) and intracellular Ca\(^{2+}\) is also possible in monolayers (Figure 4). Dye pairs successfully used in simultaneous measurements of APs and intracellular calcium include di-2-ANEPEQ and calcium green,\(^{103}\) di-4-ANEPPS and indo-1,\(^{104}\) di-4-ANEPPS and fluo-4,\(^{105}\) RH-237 and rhod-2,\(^{87}\) and RH-237 and fluo-3/4.\(^{106,107}\) In some of these experiments, the dye spectra and their overlap were of primary interest; thus, measurements were not colocalized in space or time.\(^{105,107}\) Recently, small aggregates of human-induced pluripotent stem cell-derived cardiomyocyte (iPS-CMs) have been optically mapped to study \(V_m\) and intracellular Ca\(^{2+}\) (nonratiometrically) separately.\(^{108}\)

Optical Mapping of the Sinoatrial Node

The sinoatrial node (SAN) is the site of impulse initiation and represents a regionally heterogeneous structure\(^{109,110}\) in which thousands of pacemakers cells synchronize their activity by mutually entraining each other to generate each beat.\(^{111}\) Recently, the idea of mutual entrainment has been applied to also explain the mechanism of pacemaking at the level of the single SAN cell through the concept of beat-to-beat Ca\(^{2+}\) dependent regulation of rate and rhythm.\(^{112}\) Although controversial,\(^{113-115}\) the fact that membrane depolarization triggers intracellular Ca\(^{2+}\) transients and changes of intracellular Ca\(^{2+}\) can affect membrane potential by modulating L-type Ca\(^{2+}\), chloride, and sodium–calcium exchanger currents\(^{73-75}\) provides credence to the hypothesis that intracellular Ca\(^{2+}\) may be involved in the beat-to-beat regulation of SAN activity.\(^{116}\)

The cardiac impulse can originate from the SAN or from extranodal sites within the right atrium\(^{117}\) and discrete sinoatrial pathways have been hypothesized\(^{118,119}\) to explain the complex impulse conduction within the SAN and the observation that surface activation can occur from multiple sites at the same time. This hypothesis was not directly tested until 2009, when high-resolution optical mapping of the SAN was first performed. In this study, the canine SAN was optically mapped,\(^{120}\) and structural and functional evidence suggested discrete exit pathways that connect the SAN and atria. The canine SAN is histologically similar to the human SAN and
shares the three-dimensional structural complexity that smaller animal models lack. More recently, optical mapping was conducted in coronary-perfused preparations from nonfailing human hearts using di-4-ANBDQBS and blebbistatin to block contraction. It was shown that the human SAN is functionally insulated from the surrounding atrial myocardium except for exit pathways that serve as electric bridges between the nodal tissue and the atrial myocardium.

**Whole Heart Optical Mapping**

Understanding how electric waves spread through the heterogeneous anatomy of the heart is critical to understanding normal impulse propagation and cardiac arrhythmia mechanisms. Optical mapping of the whole heart has led to important discoveries about such phenomena as alternans, bidirectional tachycardia, atrial fibrillation (AF), and VF. We describe recent advances in whole heart optical mapping, including a review of advances in human heart mapping.

**Endoscopic Mapping of AF**

AF is the most common sustained arrhythmia seen by clinicians and is the most important cause of embolic stroke. Whereas high temporal resolution can be achieved by conventional catheter-based electric mapping, which is used in clinical electrophysiology laboratories as an aid to ablation, the limited number of intra-atrial electrodes that can be used simultaneously limits the spatial resolution and precludes any detailed tracking of electric waves during AF. During the past two decades, the ability to use optical mapping in the research laboratory has been instrumental in providing numerous insights into AF mechanisms. Many such insights have found their way to clinical practice, including the identification of sources that maintain fibrillatory activity. Importantly, the use of optical mapping has been accompanied with the development of software that permits combined time-domain and frequency-domain analyses of optical signals, which led to the discovery that AF can be driven by discrete sources of high-frequency periodic activity (rotors). The waves emanating from such rotors interact with either functional or anatomic obstacles in their path, resulting in the phenomenon of fibrillatory conduction. In addition, we now recognize important frequency gradients between the left and right atrium in AF. In most cases, the region with fastest activity is the posterior left atrium (PLA), which harbors the rotors that drive the overall arrhythmia. Of note, catheter-based AF ablation procedures in patients are most successful in terminating AF when targeting the PLA. In recent experimental studies, epicardial optical mapping on the PLA has been complemented with simultaneous endocardial mapping of the PLA using a dual-channel rigid borescope coupled to a CCD camera (Figure 7). To date, this has provided the best approach to visualize the patterns of activation in the most relevant region for AF maintenance and the demonstration that, at least in some episodes, the rotors that maintain AF correspond to three-dimensional scroll waves that span from the endocardium to epicardium.

**Scroll Waves in the Human Ventricles**

VF, the most important cause of sudden cardiac death, has been defined as turbulent ventricular electric activity that precludes pumping of blood. The idea that three-dimensional scroll waves underlie the mechanism of turbulent electric wave propagation in human VF has been around for many years. A recent study combining optical mapping, mathematical modeling, and analysis of VF frequency in 11 species, from the mouse to the horse, indicates that the interbeat interval of VF scales as the fourth power of body mass. This suggests that there might be a strong similarity in the underlying mechanisms of VF in most, if not all, mammalian species, which may be of considerable fundamental and practical significance when analyzing the mechanisms of human VF. Recently, Nanthakumar et al provided the first direct demonstration of wavebreaks and rotor formation in severely diseased explanted human hearts. Wavefronts as large as the vertical length of the optical field were also found, which suggested a high degree of organization. New findings from simultaneous epicardial and endocardial multielectrode mapping, complemented by optical mapping, in patients with dilated cardiomyopathy suggested that during induced VF episodes, stable reentrant wavefronts occur in the endocardium and the epicardium. The same authors demonstrated a stable source in the endocardium, with a highly organized pattern in the local electrogram and a simultaneous and disorganized pattern in the epicardium (breakthroughs), consistent with the idea of three-dimensional scroll waves.

**Transmural Optical Mapping Using Optrodes**

One limitation of optical mapping is the lack of methodology for making three-dimensional measurements. Multiple intramural electric recordings can be obtained using plunge needle
electrode arrays. Optical mapping through the wall of the ventricles is technically challenging. Thin optical fibers offer a solution to this bioengineering problem. Optical fiber systems called “optrodes” have been developed for intramural measurements of V_m using voltage-sensitive dyes. In 2001, Hooks et al designed an optrode that consisted of a bundle of seven optical fibers enclosed in a glass pipette that was inserted into heart muscle. Laser excitation light was used and fluorescence emission (di-4-ANEPPS) was measured using PDAs. In 2003, Byars et al designed an improved optrode system that utilized a more stable light source (100-W mercury arc lamp) and the RH-237 dye, which has its absorption maximum close to very bright lines of the mercury lamp. The new optrode system also utilized better light-collecting optical fibers (ie, higher numeric aperture). Care must be taken because exposed fiber ends can form secondary sources of polarization when electric fields are applied. As a major innovation in design, the Fast laboratory ensures optimal light delivery into the tissue by polishing fiber ends at 45-deg and coating with reflective mirrors. This newest optrode design combines high signal quality (approximately 30% better signal-to-noise ratios) with increased durability for repeated use and reduced stimulus artifact. As optrode technology advances, so will our understanding of intramural impulse propagation in whole hearts, for which data are limited.

Human Heart Failure
Recent studies have utilized optical mapping techniques in an attempt to uncover important features of the electrophysiological substrate of failing human hearts. In 2010, Glukhov et al used di-4-ANEPPS to optically map action potentials in nonfailing and cardiomyopathic human left ventricular wedge preparations. The authors reported strong evidence of a transmural APD gradient in the human heart. Furthermore, it was concluded that heart failure results in the heterogeneous prolongation of APD, which reduces the normal transmural APD gradients. Immunostaining and fluorescent mapping of connexin43 expression revealed that downregulation of subepicardial connexin43 expression may underlie the electrophysiological remodeling evident in failing hearts. The first report of dual voltage and Ca^{2+} mapping of failing human hearts was published in 2011. Lou et al utilized multiparametric optical mapping to simultaneously map action potentials (RH-237) and calcium transients (rhod-2) in coronary-perfused left ventricular wedge preparations. Despite inherent limitations of the preparation, this study demonstrated transmural heterogeneity of excitation-contraction coupling and calcium handling in normal hearts and revealed the transmurally heterogenous remodeling of these properties in heart failure.

Panoramic Optical Mapping
During VF, activation waves are being conducted transmurally in three-dimension. One major limitation of traditional optical mapping is the two-dimensional single-side field of view. The anterior surface of the heart is typically mapped, which may complicate matters because the electrical wave that emanates as part of arrhythmic activity may originate, or be organized, outside this field of view. And although epicardial conduction velocities can be quantified, caution needs to be taken to interpret these values because of the curved surface of the heart. Apparent epicardial conduction velocity is an accurate representation of actual conduction velocity only when the wavefront propagates nearly parallel to the epicardial surface and field of view. Significantly more information regarding impulse propagation can be gleaned from panoramic imaging. Panoramic imaging involves the arrangement of detectors or mirrors around a preparation to allow acquisition of signal from the entire three-dimensional surface of the heart. Panoramic mapping data will enhance our ability to investigate the electrophysiology and nonlinear dynamics of the heart. Panoramic imaging provides an improved global perspective from which to investigate cardiac conduction in normal and diseased conditions. Panoramic systems have been described that provide one front view and two back mirror views of isolated hearts, thus extending single-camera optical imaging capabilities. Multiple PDAs have also been used to provide a panoramic view of impulse propagation in rabbit hearts.

Future Directions for Cardiac Optical Mapping
Bioengineering
The advent of iPS cell technology has generated much excitement in the cardiac research community, ushering in the era of regenerative and personalized medicine. The use of iPS cells for therapies obviates many of the boundaries imposed by using human embryonic stem cells. Though some researchers proceed into this field with caution, the excitement and possibilities of iPS cell technology have increased the urgency to improve bioengineering and optical mapping methodologies to enable the mechanistic study of human cardiac preparations manifesting disease phenotypes, drug testing in vitro, and novel cellular therapies. For investigation of inherited cardiomyopathy and arrhythmia mechanisms, human iPS-CMs offer unique advantages over commonly utilized transgenic animal model systems. First, animal model systems do not precisely recapitulate excitation-contraction-coupling mechanisms of human cardiac cells, thus making the use of iPSC-CMs more attractive. Second, the use of patient-specific iPSC-CMs permits direct study of specific mutations without the need for site-directed mutagenesis and complex transgenesis techniques that may result in unexpected compensatory mechanisms that mask the true cardiac arrhythmia phenotype. For example, in the case of catecholaminergic polymorphic ventricular tachycardia, caused by mutations of the ryanodine receptor, the production of patient-specific iPSC-CMs will enable isolation of the mutant channel, opening the door to detailed biophysical analysis of single channel function and response to pharmacological agents. Third, the generation of iPSC-CM monolayers will allow for the creation of patient-specific monolayers to study arrhythmia mechanisms and to determine the effect of potential therapeutics on impulse propagation. This offers a clear advantage over the commonly used neonatal rat ventricular myocyte monolayer system. Fourth, iPSC-CMs
provide a novel system for the study of cardiogenesis and may reveal important genetic and electrophysiological mechanisms of cardiac development in patients with congenital heart defects. Major advances, however, must be made in cardiac-directed differentiation techniques, especially for the maturation of the de novo myocytes, to significantly advance the field.

To date, most reports using iPSC-CMs to study electrophysiology have used single-cell patch-clamp techniques. However, the laborious nature of these techniques precludes efficient screening of electric wave propagation through a functional syncytium of cardiac tissue. Microelectrode arrays and optical mapping have been used on small human iPSC-CM aggregates, but observed conduction velocities have been reportedly very low (<2.5 cm/s). As innovative bioengineering approaches are developed to generate cardiac patches, hoping to repair infarcted hearts, optical mapping of these bioengineered constructs will be vital to assess function and electrical connectivity. Using a single-camera/LED illumination system, we recently imaged action potential (di-8-ANEPPS) and ratiometric calcium (fura-4F) transients and wave propagation in relatively large (diameter ≥1 cm) human iPSC-CM monolayers (unpublished work). The conduction velocity of wave propagation in these iPSC-CM monolayers was similar to that recorded in neonatal rat ventricular myocyte monolayers, approximately 22 cm/s. Figure 8 shows an example of a spontaneous activation spreading through the entire iPSC-CM monolayer. Thus, iPSC-CM monolayers offer an attractive in vitro human system for study of arrhythmia mechanisms and drug screening.

Optogenetics
Channel rhodopsins (ChRs) are protein channels that transport cations across the cell membrane in response to light stimulation. Four distinct channel rhodopsins have been discovered, channel rhodopsin-1 (ChR1) and channel rhodopsin-2 (ChR2) from Chlamydomonas reinhardtii and Volvox ChR1 (VChR1) and Volvox ChR2 (VChR2) from Volvox carteri. These channel rhodopsins function in nature as sensory photoreceptors that microalgae use for phototaxis (movement toward or away from light), optimizing light conditions for photosynthesis. In 2005, the light-sensitive gene was first delivered to cells using genetically engineered viruses, and pulses of light can now be used to precisely stimulate transfected excitable cells. This technique has gained widespread popularity in the neuroscience and cardiac fields and is now known as optogenetics.

The cardiac pacemaker initiates every cardiac contraction and in pathological conditions can be substituted by surgically implantable battery-operated devices. Recently, optogenetics has been used to optically control cardiac function in zebrafish hearts. Last year, Jia et al reported the first combination of optical excitation and optical imaging to capture light-triggered cardiac muscle contractions and high-resolution electrical waves in mammalian cardiac cells and tissue. More recently, Abilez et al demonstrated the potential of optogenetic control in human cardiac cells using a combined experimental/computational approach. It was demonstrated that ChR2 can be stably expressed in human pluripotent embryonic stem cells, which can then be differentiated into functional human cardiomyocytes using defined protocols. This was the first creation of ChR2-expressing human cardiomyocytes that could successfully be triggered using light. In computational work, they also showed the utility of optogenetics to generate propagated cardiac impulses. This technology may lead to a light-triggered biological pacemaker created from a patient’s own iPSC-CMs. However, this is a long way away and as optogenetic technology advances, it will be critical to utilize optical mapping techniques to provide detailed insight into the electrophysiological function of optogenetic manipulations.

Conclusion
Cardiac optical mapping has evolved to a powerful and essential technology for studying cardiovascular function and disease. Over the past two to three decades, the technological advances in cardiac optical mapping have provided new mechanistic insights into electromechanical function, arrhythmias, and disease mechanisms from the single cell to the whole organ. Its application continues to expand, even penetrating the fields of bioengineering and optogenetics. It was only in 2007 that submillimeter spatial resolution (0.65–0.85 mm) optical mapping demonstrated that VF in human hearts is associated with wave breaks and singularity point formation and is maintained by high-frequency rotors and fibrillatory conduction. With the development of new infrared voltage-sensitive dyes, optical mapping one day may be used to provide in vivo examination of AP propagation using sophisticated catheter-based mapping protocols. Last, but not least, optical mapping will also prove to be a pivotal tool in the development of regenerative cardiac research and therapies, including molecular genetic advances.
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