Extracellular electrograms, recorded directly from the heart, are the hallmarks of invasive cardiac electrophysiology and provide information about the electric status of the underlying myocardium. These electrograms are generated by depolarization of cardiomyocytes that generates transmembrane currents in extracellular space and potential differences due to electric resistance of the extracellular medium. In healthy myocardium, the basic configuration of the extracellular electrogram is simple. Under pathological conditions, however, electrograms may consist of multiple components and long duration, which have been attributed to abnormal conduction and arrhythmogenicity. Although complex and fractionated electrograms are presently considered to be a real phenomenon, this was not the case some 20 years ago. Debates about the “fact or artifact” of complex and fractionated electrograms were common. At that time, electrograms consisting of multiple “high frequency” components with low amplitudes and long duration, termed “fractionated,” were recorded in patients with healed myocardial infarction during endocardial mapping. Several investigators presumed that fractionated electrograms were artifacts, resulting from movements between electrode and myocardium, filter characteristics of amplifiers, or represented far field effects. A classic example to support the “artifact theory” was the recording of continuous activity from the jello brain. In those days, much attention had been paid to fractionated electrograms in healed myocardial infarction to guide catheter ablation or antiarrhythmic surgery. Although artifacts indeed may cause complex electrograms, most of the complex and fractionated electrograms are a “fact” and caused by the peculiar behavior of activation fronts, due to structural and electric complexity of the underlying myocardium. This article delineates the origin of the unipolar extracellular electrogram, reviews the circumstances that cause fractionated and complex electrograms, and discusses the impact of the recording technique on detection and interpretation of multifaceted electrograms.

**Origin and Configuration of Unipolar Extracellular Electrograms**

Extracellular electrograms arise because of transmembrane currents that occur due to differences in the axial voltage gradient at the interface between activated and inactivated myocardial cells. This is schematically illustrated in Figure 1, which shows a myocardial bundle that is at rest at the right and activated at the left. Activation in the bundle moves from left to right. The action potential causes an axial voltage gradient between activated (+20 mV) and inactivated (−90 mV) sites resulting in axial current flow from the activated toward the inactivated site. Current amplitude is large at the position of the wave front where the voltage gradient is high (site B, 4 arrows). At the left (site A) and the right (site C), the voltage gradient is less steep and the resulting axial current is low (1 arrow). The sum of currents flowing toward a point must be equal to the sum of currents flowing away from that point. This implies that at the back of the activation front (site A), current must flow through the membrane into the cell (3 arrows at the membrane pointing inward), and, in front of the activation front (site C) current flows outward through the membrane (3 arrows at the membrane pointing outward). Thus, the activation front operates as a current dipole, which injects current into extracellular space at the front and retrieves current from extracellular space at the back. Current through extracellular space, which has electric resistance, generates an extracellular voltage difference. The corresponding extracellular potentials, which have been determined by Spach et al for a Purkinje fiber, are illustrated in the upper part of Figure 1. Extracellular potentials are positive in front of the activation front, zero at the activation front, and negative at the back of the activation front. The amplitude of the voltage increases closer to the current dipole. The extracellular potential field moves with the activation front and a recording electrode positioned at a site where the activation front is passing will therefore record an increasing potential when the front is approaching. The potential reaches a maximum when the activation front is very close to the recording site. Then, the potential rapidly decreases to zero and becomes negative with its minimum just as the wave front has past. Thereafter, the negative potential returns to zero as the wave front proceeds. Thus, the extracellular electrogram that is generated by a passing wave front is biphasic, a positive deflection followed by a negative one.
This is illustrated in Figure 2, which shows a myocardial bundle that is stimulated electrically at the left. Thus, activation starts at the left and moves to the right. At site B, the wave front is passing and a simple biphasic deflection is generated. At the left, where activation starts, there is no approaching wave front and the local electrogram only has a negative deflection. At the far right, site C, the activation front stops. The recording site just faces the area with positive potentials and consequently the electrogram only has a positive deflection. Thus, in principle, the unipolar electrogram is simple and comprises only a biphasic or monophasic complex. In addition, the configuration of the unipolar electrogram provides information about the activation front at the recording site: (1) an initial negative deflection at the site where activation starts, (2) a biphasic deflection at the site where activation passes, (3) a positive deflection at sites where activation stops, and (4) the downstroke of a unipolar electrogram, the intrinsic deflection, coincides with the upstroke of the action potential of the myocardial cells underneath the electrode. The initial negative deflection of the unipolar electrogram is helpful in detecting the site of origin of focal activation but may also point to a site of pseudo initial activation such as the exit site of an infarct related ventricular tachycardia (VT). Stevenson and Soejima propose the use of unipolar recordings especially for mapping of focal arrhythmia sources. Haissaguerre et al suggest the use of unipolar recordings to determine accessory pathways, whereas Ito et al recommend the unipolar recording technique for atrioventricular junction ablation, which can be achieved more efficiently and with fewer radiofrequency energy application when guided by unipolar recordings than by bipolar recordings alone.

**Bipolar Versus Unipolar Electrograms**

A bipolar electrogram is made by subtracting 2 unipolar electrograms recorded at sites that are usually close together (millimeter distance). In the clinical setting, a bipolar recording is often preferred over a unipolar one because the bipolar mode suppresses interference of the electric mains. This is due to the difference in the “field of view” of the unipolar and bipolar electrode. As shown before, an electrode will already pick up a signal generated by the wave front when it is at a distance. This implies that the electrode not only records activity underneath the electrode (the local event) but also at
signal analysis. In addition, multiple catheters are frequently applied during catheter mapping. Because the catheters are often in close proximity, they may touch each other and cause spiky artifacts. Interference of the mains often overlays the electrogram with a 50/60-Hz signal, frequently accompanied by higher harmonics. In sporadic cases, the interference fluctuates and may give the impression that intervals with fractionation arise in the electrogram. Another type of artifact, not related to the heart is the electromyographic interference. The deflections are real signals generated by the patient’s muscle. They often occur in intervals and such fractionation may be misleading, because it is not cardiac in origin. Such artifacts may especially arise when unipolar recordings are applied.

What Makes Electrograms Complex?

Artifacts

As already outlined in the introduction, movement artifacts can cause fractionated electrograms that may complicate signal analysis. In addition, multiple catheters are frequently applied during catheter mapping. Because the catheters are often in close proximity, they may touch each other and cause spiky artifacts. Interference of the mains often overlays the electrogram with a 50/60-Hz signal, frequently accompanied by higher harmonics. In sporadic cases, the interference fluctuates and may give the impression that intervals with fractionation arise in the electrogram. Another type of artifact, not related to the heart is the electromyographic interference. The deflections are real signals generated by the patient’s muscle. They often occur in intervals and such fractionation may be misleading, because it is not cardiac in origin. Such artifacts may especially arise when unipolar recordings are applied.

Filtering

To suppress interference from the mains, remote signals such as the ventricular complex in the left atrium, and baseline drift, electrograms are routinely filtered. This allows higher amplification without compromising the quality of the baseline. Filtering, however, may also artificially add deflections to the electrogram. In Figure 3, the bipolar electrogram from an ablation catheter with a 4-mm distal electrode and 1-mm proximal ring electrode (2-mm spacing) is bandpass-filtered (30 to 500 Hz), but once with the 50 Hz notch filter off and once with the notch filter on (Prucka System, General Electric). Such filtering clearly adds components to the electrogram that may be interpreted as fractionation and this may become a source of error in finding ablation targets if based on fractionated electrograms.

Remote Activation

Remote signals may affect the local electrogram and a delay between both may lead to an apparently fractionated signal. As shown before, the unipolar recording is more sensitive to remote activity than the bipolar one, but also the latter is not free from deflections caused by remote activations. Typically, deflections are caused by remote ventricular activation that may interfere with local atrial deflections in recordings from the atrium. Depending on the size and configuration of the atrium, the atrial deflection may appear as fractionated. However, usually the surface ECG is recorded simultaneously with atrial or ventricular signals, which allows detection of the ventricular component in the atrial recording. More complex and disturbing during atrial recordings may be deflections caused by remote activation in other parts of the atrium, as discussed in the next section.

Infarcted myocardium also is notorious for remote complexes caused by activation in healthy surrounding myocardium. In this case, combined unipolar and bipolar recordings can be very helpful as the unipolar electrogram puts the large remote component in the foreground, whereas the bipolar electrogram enhances the local component (Figure 4).

Adjacent Structures

Many structures in the heart that are activated at different times are anatomically located close to each other. Thus, local
activation recorded at one structure can be disturbed by (remote) activation generated by the other structure. This is important to realize, because it may affect decisions with regard to interventions. For instance, the right superior pulmonary vein (RSPV) is close to the superior caval vein (SCV) (Figure 5, left panel).18 Similarly, as discussed before, the left superior pulmonary vein and sometimes also the left inferior pulmonary vein are close to the left atrial appendage. Electric isolation of the pulmonary veins (PVs) by means of catheter ablation is a standard procedure to treat AF. Success of isolation is tested by recording activity from the PVs during sinus rhythm or atrial stimulation. If electric isolation is complete, no activity evoked at the atrium will be recorded at the PV. However, activity in adjacent structures might suggest that isolation is incomplete. The right panel of Figure 5 shows a circular mapping catheter in the RSPV and a conventional one in the SCV (in front of the left atrial appendage). Tracings in Figure 6 are obtained with these catheters during sinus rhythm and show that activity in the SCV (MAP 1,2) gives rise to remote deflections in the RSPV. Note that compared with the local RSPV activity (bold arrows) remote SCV deflections in the RSPV (open arrows) occur 2 to 1 and coincide with the SCV deflection, supporting their SCV origin. Such remote deflections in the RSPV (or other PVs) might erroneously be interpreted as failed isolation of the RSPV. During AF, the deflections might be interpreted as highly fractionated RSPV signals.

Anisotropy
Cardiac tissue is anatomically and electrically anisotropic, which is caused by the cell morphology in conjunction with the electric coupling between the cells, mediated by gap...
junction proteins. This results in faster propagation of activation parallel to the fiber direction as compared with propagation perpendicular to the fibers. The configuration of the unipolar extracellular electrograms recorded at sites where the activation front runs parallel to the fiber direction follows the simple rule of the biphasic deflection at the site where activation passes. However, Spach et al.\(^\text{19}\) have shown that electrograms recorded at sites where activation proceeds perpendicular to the fiber direction deviate from this simple concept and are more complex. At these sites, where activation passes, the associated biphasic deflection is preceded by a small negative one. This deflection arises because of a remote effect of the distant but large wave front that runs parallel to the fibers.\(^\text{6,19}\)

**Overlaying Structures**

In several parts of the heart, myocardial structures that often have different electrophysiological characteristics, overlay. These structures may be totally or partly isolated from each other by collagen and arise in various parts of the heart. An obvious example is presented by the bundle branches and Purkinje fibers, which are isolated from surrounding myocardial tissue by a sheath of collagen. Electric recordings from myocardium near these structures will result in electrograms that reflect both myocardial and Purkinje activity.\(^\text{20,21}\) These deflections can often be distinguished, because of differences in their characteristics (sharpness and amplitude). The AV junctional area is much more complex and separation of the different tissue types (atrial cells, transitional cells, and compact nodal cells) is not complete but comprises an intermingling of myocardial cells and fibrosis.\(^\text{22–24}\) At the atrial level, the overlaying structure of left atrial myocardium and excitable coronary sinus sleeve is of interest and may result in complex electrograms.\(^\text{25–27}\) Myocardial tissue from atrium and coronary sinus are isolated at most sites, but connections may be present at some sites. Therefore, recordings from the inside of the coronary sinus often show multiple deflections because of activation in the 2 different structures.

**Alterations in Conduction Velocity**

There are several causes for changes in conduction velocity that might affect the complexity of electrograms.

**Wave Front Curvature**

Electric barriers in the heart, being either anatomically or functionally determined, will force activation fronts to curve around the pivot points.\(^\text{28,29}\) At these sites, conduction slowing arises because of source-sink mismatches.\(^\text{30}\) As delay arises at the pivot point, the positive deflection caused by the approaching wave front and the negative deflection caused by the receding front are separated, which results in 2 deflections instead of 1 at that location.

**Tissue Discontinuities**

A sudden change in conduction velocity may arise at tissue discontinuities. If the diameter of a myocardial bundle increases abruptly, current-to-load mismatch may occur at the connection site. The current that the thinner section delivers may not be sufficient to activate the thicker bundle instantaneously. This results in conduction delay and complex electrograms. The delay imposed by current-to-load mismatches is also reflected in the action potential at the discontinuity, which shows a step in the upstroke (Figure 7).\(^\text{31}\) Discontinuities in conduction velocity also arise at sites where tissue types with different excitability or coupling characteristics are connected. An extreme example is the interface between myocytes and fibroblasts. Fibroblasts couple electrically to myocardial cells and stretches of fibroblasts in between strands of myocytes are able to propagate the action potential, albeit at a large conduction delay.\(^\text{32}\) Impulse transmission along stretches of cardiac fibroblasts as long as 0.6 mm has been observed; conduction delay over this distance was 30 ms, corresponding to a apparent conduction velocity of 0.02 m/s. At present, there are no data available of the role of fibroblasts in situ. Data about conduction in surviving myocardial strands in infarcted myocardium makes it unlikely that fibroblasts play a major role in conduction slowing.\(^\text{33}\)

Evidence for fractionated electrograms caused by sudden changes in cell-to-cell coupling is supported by computer modeling. In a model for propagation, Lesh et al.\(^\text{34}\) showed that fractionated electrograms arose if a uniform wave front encounters a region of increased cellular coupling resistance.
Fibrillatory Conduction

Electrograms recorded at sites with structural and/or functional discontinuities may be close to normal during basic stimulation but become complex and fractionated after premature stimulation if sodium current availability is reduced. In the right atrium, the pectinate muscle network is an area with structural complexities where functional conduction block may occur. This was nicely illustrated in isolated, coronary-perfused sheep right atrium by Berenfeld et al.\cite{35} The investigators stimulated the Bachman bundle at different frequencies and recorded the electric activity of the crista terminalis and pectinate muscles optically. Up to a pacing rate of 6.3 Hz, no gross differences in the activation pattern occurred. At frequencies above 6.7 Hz, prominent zones of functional conduction block occurred, which were accompanied by increased complexity of the extracellular electrograms. As observed at tissue discontinuities, action potential upstrokes were complex and revealed multiple upstrokes.

Complex fractionated electrograms have been suggested as target areas for ablating AF.\cite{36} Intraoperative observations showed that these complex electrograms localized at areas of conduction block and pivot points for reentry.\cite{37} Recent studies show that wave front collision, functional conduction block, wave break, and wave fusion all may cause complex and fractionated electrograms in AF.\cite{38,39}

Asynchronous Conduction

In cardiac disease, structural and electric remodeling occur and may be accompanied by remodeling of the autonomic nervous system.\cite{39,40} Structural remodeling may involve changes in cell size, increased collagen deposition, and myocardial fiber disarray. Collagen deposition appears to be a major component in the remodeling process because the majority of cardiac diseases is accompanied by an increase in cardiac fibrosis.\cite{41,42} Although collagen constitutes the framework in which the cardiomyocytes are embedded to give the heart its mechanical rigidity, an increase in the amount of collagen will electrically separate the myocardial cells and prevent the formation of wide, coherent wave fronts. The electrophysiological and anatomic basis for fractionated electrograms recorded in regions where infarct healing caused separation of myocardial fibers was provided by Gardner in 1985.\cite{5} Despite normal transmembrane potentials, activation time was prolonged and inhomogeneous in areas where fractionated electrograms were recorded.

Increased collagen deposition often causes a complex network of intermingled collagen and myocardial fibers.\cite{33} Figure 8 shows the histology of an infarcted human papillary muscle at 3 different levels, 70 μm and 210 μm apart. The schematic drawing at the right shows that separated bundles at level A merge at level B but diverge again at level C. A schematic representation of the merging and diverging bundles is illustrated in Figure 9. When stimulating site A, activation can reach site B only by following the zig-zag route as shown (arrow). The recording electrode in the center of the preparation will not only record activity in the bundle underneath the electrode but also activity of wave fronts propagating in distant bundles. Because of asynchronous activation in the various bundles, multiple deflections will arise, which results in fractionated electrograms.

It is important to realize that a simple, biphasic electrogram could arise if activation proceeds parallel to the fiber orientation in the structure of Figure 9. This is the case if all the bundles at the left are activated at the same time. Indeed, there are still small wave fronts in the various bundles, but, because activation is synchronous now, the deflections they generate at the recording site occur simultaneously, resulting in a single deflection (electric signals at the same recording site

![Figure 8. Histological sections of a human papillary muscle stained for collagen with picrosirius red. Sections were taken perpendicular to the long axis of the papillary muscle. Sections are 70 and 210 μm apart. The right lower panel shows the distribution of collagen (red) and myocardium (yellow) for corresponding areas indicated by the squares in the sections. The upper area a shows multiple myocardial bundles separated from each other by collagen. A number of bundles in area a merges in area b of section B but appear to divide again at level C. These data show that the infarcted papillary muscle is characterized by merging and diverging myocardial bundles.](image)

![Figure 9. Schematic representation of infarcted myocardium consisting of merging and diverging myocardial bundles. Activation induced by stimulation at site A can reach site B only by following the indicated zig-zag path (arrow). The electrode in the middle will record activation in the various bundles. However, because activation in these bundles is asynchronous, the electrode will record multiple, separated deflections.](image)
Fractionated Electrograms

Role of Conduction Parameters in Fractionated Electrograms

Uniform Versus Nonuniform Conduction Slowing

Although slow conduction has been associated with fractionated electrograms, it depends on the mechanism by which conduction slowing is induced, whether fractionation indeed occurs. As illustrated before in experimental and modeling studies, discontinuous conduction will cause fractionation of the electrogram. In contrast, a homogeneous reduction of the conduction velocity may not result in split electrograms. Jacquemet et al.45 induced slow conduction in a 2D computer model for propagation of the electric impulse in 3 different ways: (1) by a homogeneous reduction of the fast inward sodium current, (2) by a uniform decrease in transverse coupling of myocardial cells, and (3) by introducing a set of sodium current. The authors suggested that these high-frequency fractionated electrograms correlated with the concentration of the drug. In addition, injection of acetylcholine into the anterior right ganglionated plexus too resulted in complex, fractionated electrograms in isolated sheep heart during AF. This model of AF showed regular, highly organized activity in the posterior left atrium. Fractionation of activity occurred at the border of the dominant frequency area where most distinct fractionated activity borders the area with most regular activity. Here, electrograms are most fractionated and the authors suggest that these high-frequency fractionated electrograms can be used to localize sources of AF at the posterior left atrium.

Autonomic Nervous System

Although several studies provide evidence for heterogeneous anatomy and areas of functional conduction block as drivers for complex fractionated atrial electrograms, Lin et al.49-50 suggested an autonomic basis for the formation of these atrial electrograms. In a dog model, the investigators showed that by local application of varying concentrations of acetylcholine during AF, the incidence of inducing local complex fractionated electrograms correlated with the concentration of the drug. In addition, injection of acetylcholine into the anterior right ganglionated plexus too resulted in complex fractionated electrograms. The complex electrograms oc-
curred distant from the site the drug was injected and were eliminated by ablation. The investigators postulated that the induction of complex fractionated electrograms by local application of acetylcholine excites the autonomic nerve terminals, which resulted in activation of distal ganglionated plexus. These plexus cause fractionation of electrograms by releasing neurotransmitters that modulate atrial wave front stability by causing wave breaks.

**Distinguishing Local From Remote Activity**

Several techniques have been proposed to distinguish local from remote deflections in complex, fractionated electrograms.51–55 The question whether a deflection in a fractionated signal is local or remote is often difficult to answer, especially when recordings are made at only 1 position. Figure 11 shows tracings at 5 recording sites of an infarcted human papillary muscle. Histological investigation showed the presence of multiple, separated, parallel running myocardial bundles, as shown at the right. As outlined before, these bundles may connect at certain sites. Recording sites were 600 µm apart and positioned along a line perpendicular to the fiber direction. By comparing clustered deflections in the recordings, it is evident that the global propagation is from site e to site a. However, the actual trajectory of the activation from e to a is unclear. Presumably, the deflection marked by the black dot in tracing d is the local deflection. The arguments for this are the following: (1) Time-equivalent deflections with lower amplitude are present in the electrograms recorded at the other sites (circles); and (2) the amplitude of these deflections decreases with the distance from d (deflections along the left dotted vertical line).

The deflection in tracing d marked by x is generated by activation in the lower bundle. A time-equivalent deflection has its largest amplitude in tracing e (marked by the black dot in tracing e); the amplitude of the deflection fades away for electrograms recorded further away from site e. Thus, if multiple recordings are available, it is sometimes possible to distinguish local from remote components.

A problem arises if the recording area of the electrode covers multiple bundles. In that case, multiple deflections may be present that are all “local.” In the clinical situation, the mapping catheter is also used for ablation and its distal electrode is much larger than the ring electrode. With tissue contact, each electrode records but also more or less averages all signals generated by activation underneath its metal surface. Consequently, electrograms from the larger distal electrode may contain fewer high-frequency components than that from the smaller ring electrode. A bipolar electrogram from such electrode pair, both with good tissue contact, may then predominantly reflect events from the ring electrode, especially when the distal electrode is large (ie, 8 mm). Often, however, the proximal ring electrode is not in contact with myocardium. This (the larger distance to activated tissue), too, reduces the high-frequency components of the local electrogram and in that case it may mainly be the tip electrode that determines the bipolar electrogram morphology. Simultaneous recording of both the bipolar and the corresponding unipolar electrograms eliminates confusion about the origin of the various component of the bipolar electrogram. In addition, it reveals the direction of activation, which may be important in analysis and treatment of the arrhythmia.

**Conclusion**

The studies illustrated before show that the underlying mechanisms of fractioned and complex electrograms are diverse. They may reflect pure local effects but also may be caused by remote activity at the recording site where deflections caused by the local and distant activity merge. It is important to rule out this possibility if fractionated electrograms are used in the clinical setting as a guide for interventions. Discontinuous conduction and sudden changes in conduction velocity may play a role in structurally remodeled myocardium. They may point to areas with impaired conduction, but the involvement in arrhythmogenicity must be proven if such areas are used as target site for intervention. Homogeneous alterations in cell-to-cell coupling and excitability do not cause fractionation of electrograms, but electric remodeling in cardiac disease is often heterogeneous. In that case, irregular conduction may arise through areas with varying degrees of functional conduction block, resulting into fractionated wave fronts and fractionation of the electrograms. Complex and fractionated electrograms point to areas with abnormal propagation of the electric impulse, and it is conceivable that they are involved in arrhythmogenic processes and be an attractive target for treatment. In AF, fractionated and complex electrograms are currently being used as targets for ablation. Although there are data that suggest that fractionated electrograms might indeed be re-
lated to AF drivers, this outline illustrates that fractionation is a complex process the more so because cardiac disease is accompanied by structural, electric, and autonomic remodeling that all may affect fractionation of electrograms. Also, the role of fractionated electrograms as a guide to ablation of VTs is not yet clear. Weiner et al. observed that fractionated electrograms recorded during sinus rhythm were more numerous in the infarct border of patients with episodes of tachycardias and fractionated electrograms had longer duration. Sites with fractionated electrograms were supposed to delineate regions where reentry occurs. Kienzle et al., however, showed that fractionated electrograms were widespread in patients with infarct-related VT but also were found in regions outside the tachycardia origin. Recently, Haqqani et al. studied patients with and without sustained monomorphic tachycardia and healed myocardial infarction and showed that patients without VT had fewer fractionated, isolated, and very late potentials than patients with VT.

Thus, although fractionated electrograms refer to abnormal myocardial structure and conduction, a direct relation with tachyrhythmias remains unclear.

Disclosures

Dr. Wittkampf is a consultant of St. Jude Medical.

References

Fractionated and Complex Electrograms


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