Remote Monitoring of Cardiac Implantable Electronic Devices

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ABSTRACT. Pacemakers, implantable cardioverter-defibrillators, and cardiac resynchronization devices use advanced diagnostic functions to continuously monitor the patient and the device itself. Follow-up is necessary after implantation to provide individualized treatment and to detect malfunctions. Routine office visits, however, do not provide access to these data on a timely basis. Transtelephonic pacemaker follow-up has been used as an adjunct to scheduled visits. Advanced remote monitoring of implanted devices is rapidly expanding due to technological innovation in this field and promising results of large clinical trials. It has been shown to decrease patient morbidity and mortality, and to help with the management of heart failure and atrial fibrillation. The technology is widely accepted by physicians and patients alike and is more cost-effective than conventional follow-up. Ongoing studies are evaluating whether remote follow-up may guide anticoagulation for atrial fibrillation, facilitate early hospital discharge after device implantation, or completely replace routine office visits. However, lack of standardization may slow down widespread utilization of recent advances. The outlook for this technology is excellent and remote monitoring will continue to improve patient-centered, cost-effective follow-up of implanted cardiac devices.

KEYWORDS. cardiac resynchronization therapy, follow-up, pacemaker, implantable cardioverter-defibrillator, remote monitoring.

Follow-up of implantable cardiac devices

Patients require follow-up after implantation of a pacemaker (PM), implantable cardioverter-defibrillator (ICD), or cardiac resynchronization therapy device (CRT) to assess and adjust diagnostic, pacing, and antitachycardia functions. The frequency of follow-up is determined by multiple factors, including patient comorbidities, geographic accessibility to medical care and type of the implanted device. Current guidelines recommend in person follow-up within 72 h after implantation of any device, with the next visit at 2–12 weeks. Thereafter, in person or remote follow-up is recommended every 3–12 months for pacemakers, and every 3–6 months for ICDs or CRT-defibrillators (CRT-D). Once every year, in person follow-up is required for any device, even if more frequent remote follow-up is performed. At signs of battery depletion, in person or remote follow-up every 1–3 months is needed.

In practice, office follow-up visits are frequently uneventful, revealing no relevant changes related to the implanted device or the patient’s condition. There were an estimated 1.6 million PM and 2.1 million ICD follow-up encounters in 2007 in the USA alone, causing a significant burden on the cardiovascular workforce and contributing to increasing health-care costs.

Remote monitoring systems

The importance of pacemaker follow-up was recognized soon after these devices were introduced half a century ago. The authors report no conflicts of interest for the published content. Manuscript received April 15, 2011, final version accepted May 23, 2011. Address correspondence to: Mark H. Schoenfeld, MD, Clinical Professor of Medicine, Yale University School of Medicine, Director, Cardiac Electrophysiology and Pacemaker Laboratory, Hospital of Saint Raphael, 330 Orchard St, Suite #210, New Haven, CT 06511. E-mail: mschoenfeld@srhs.org
ago, as early leads and generators were unreliable. After
decades of technological innovation, hardware-related
issues of the generators are now extremely rare;
however, lead-related complications are still not uncom-
mon (dislodgement, insulation failure, fracture etc.).
Introduction of sophisticated devices, such as ICDs and
CRTs, significantly increased the workload of a follow-
up session due to the large amount of diagnostic data
that needs to be interpreted and device settings that have
to be optimized for the patient. However, most office
follow-up visits are routine procedures that rarely
require extensive changes to the settings.

The first rudimentary remote pacemaker follow-up
system, transtelephonic monitoring was introduced
40 years ago. It allows the assessment of battery
condition with basic sensing and pacing data.3 This
communication method requires simultaneous active
participation from both the patient and medical staff;
however, it cannot provide data regarding advanced
functions, such as arrhythmia events or pacing statistics,
which are essential in the follow-up of advanced
devices.4 Most major device manufacturers have recently
developed and introduced remote PM/ICD/CRT mon-
toring systems.5,6 Although the basic concept is similar,
the hardware and software for each device company is
unique and the systems are only compatible within a
specific product line (Figure 1).

The implanted device incorporates a short-range
telemetry antenna, which communicates with the remote
transmitter using radiofrequency signals, on an inter-
mittent basis. This may either require active patient
participation, such as placing a telemetry wand over
the device, or may be completely automatic, when the
transmission occurs at given time intervals if the
transmitter is within reception range. Most companies
use a configurable schedule for transmission, with
an option for patient-triggered transmission. The two
extremes of transmission intervals can be classified as
“remote follow-up”, when transmission is performed
once every several months, similar to an office visit, and
“remote monitoring”, when transmission occurs daily,
allowing more rapid detection of an adverse event. Truly
continuous remote monitoring is not feasible with
current battery technology as the energy consumption
during the telemetry session is significantly higher than
during normal device operation. The relatively low data
transfer speed with the short time available for tran-
smission (up to a few minutes, to be convenient for the
patient) limits the amount of information that can be
communicated. Current remote monitoring technology
does not provide the complete data acquisition that can
be accomplished during an office visit. Intracardiac
electrogram (IEGM) is an essential part of diagnostic
information, but requires large amount of data to be
transmitted. However, even short segments of IEGM
can be of significant diagnostic help: the RIONI study
(Biotronik, Lake Oswego, OR) found that IEGMs with a
mean length of 4.4 ± 1.5 s were adequate to assess the

Figure 1: General set up of current remote monitoring systems. Features that are still not widely implemented or experimental
are displayed in italic. Remote programming is not available currently.
appropriateness of ICD therapy in 92% of cases and were helpful for arrhythmia classification in 93.4%. The small computing capacity of the implanted devices limits the use of advanced data processing methods (crunching, encryption). The communication may be intercepted and altered; however, no significant patient safety breaches have been reported so far.

The patient receives feedback regarding the success or failure of the transmission. Some systems may be used to gather and transmit data from different home monitoring devices (such as scales, blood pressure device, hemodynamic monitor). Future improvements may include integration with further diagnostic modalities or medication management systems (digital pillbox). The transmitter forwards data to the central databank using a phone line or cellular network. Mobile systems are preferred in active (working, travelling) patients. Transmitters using the cellular network may include roaming capability, allowing the patient to travel while being monitored.

The server of the databank performs triage of the incoming data and notifies the physician’s office via internet, fax or phone message. The clinician (usually a mid-level provider) evaluates the triaged data and may notify the physician or the patient, adjust the transmission schedule, or configure alerts. Server-side data analysis with automated response may decrease the workload of the follow-up team. As the computing capacity of the database mainframe exceeds that of the implanted device, more detailed analysis of diagnostic information may be possible to improve data triage or facilitate automatic responses.

Remote reprogramming of correctable malfunctions (such as oversensing issues) may spare an office visit or avoid an adverse event (Figure 2). Despite technical feasibility, current systems do not allow remote (nevertheless automatic) reprogramming. As most implanted devices already automatically adjust their settings (such as pacing threshold management), this would be a logical expansion of an already used feature, using the greater computing capacity of a central server.

Feasibility and patient safety with remote monitoring

Most clinical studies have addressed safety and feasibility issues, as remote monitoring is a relatively new technology (Table 1). Patient and clinician acceptance was found to be high in early trials. There are significant differences between different manufacturers’ systems, which makes generalization of data difficult.

The ALTITUDE registry (Boston Scientific, St. Paul, MN) compared outcomes between patients followed in the device clinic only and those who also regularly transmitted remote data, on average four times a month. For the 69,556 ICD and CRT-D patients receiving remote follow-up, 1- and 5-year survival rates were higher compared with the 116,222 patients who received device follow-up in device clinics only (mortality reduction 44% with ICD, 56% with CRT-D). The differences were not affected by patient characteristics such as age, gender, implanted device year or type, economic or educational status, although this was not a randomized study.

Timely intervention based on early detection of device- or patient-related issues may avoid more severe complications. Preliminary data from the ECOST trial (Biotronik) show that patients with the Medtronic Sprint Fidelis lead experienced lead fracture requiring lead replacement in 7.5% over an average of 22 month follow-up, all of these events were preceded by oversensing of noise artifacts or an abrupt rise in pacing impedance noticeable on remote transmissions (Figure 3). The PREFER trial (Medtronic, Mounds View, MN) compared traditional 2-month transtelephonic monitoring (with 6-month office visits) and remote transmission at 3-month intervals (with 12-month office visits). More events requiring action (new onset or sustained atrial tachyarrhythmia, ventricular arrhythmias, lead or battery issues) were detected in the remote group and the mean time to first diagnosis was shorter (incidence 45% versus 37.6%, time to detection 5.7 versus 7.7 months). Sixty-six percent of the events in the remote arm were identified remotely, compared with 2% in the control arm; conventional transtelephonic transmissions are less sensitive to pick up adverse events. Transtelephonic monitoring was found to be most useful to monitor battery longevity.

Automatic interpretation of interrogated data may reduce the workload of professionals involved in follow-up. A rule-based data filter was generally well accepted and found to be accurate in a multicenter study. Automatic clinician alert triggers used with remote follow-up reduce time needed to action: the CONNECT trial (Medtronic) randomized 1997 patients and followed them at 1, 3, 6, 9, 12, and 15 months (patients in the remote arm had only remote follow-up at months 3, 6, 9, and 12). The median time to clinical decision after a detected event was reduced from 22 days in the in-office arm to 4.6 days in the remote arm, and the mean length of stay per cardiovascular hospitalization visit decreased from 4.0 days in the in-office arm to 3.3 days. Only 55% of the triggered automatic alerts were transmitted successfully to the clinician due to configuration errors or patient location (the study used a land line-based system). Automated, event-based clinician alert was thus able to improve clinical outcomes even when the routine remote follow-up was not performed more frequently than the office follow-up. The ongoing REACT study (St. Jude Medical, St. Paul, MN) will address similar questions.

Increasing the time interval between office follow-up visits may be safe if adequate remote monitoring is performed. The TRUST study (Biotronik) compared remote monitoring and conventional follow-up at 3, 6, 9, 12, and 15 months after implantation. Office visits at 6, 9, and 12 months were performed in the remotely monitored group only if necessary based on remote data. As a result, 85.8% of these were withheld, in-office device evaluations were reduced by 45% without significantly affecting mortality (3.4% remote versus 4.5% in conventional) or adverse event rates (10.4%/year in both groups). At the mandatory 3-month follow-up
office visit, 6.6% of patients from both groups had findings requiring action, with similar distribution: reprogramming (78.4% versus 72.6%), antiarrhythmic medication change (21.9% versus 29.6%), or lead or generator revision (4.4% versus 3.2%). The frequency of unscheduled office visits was low, but higher in the remote group (0.78 versus 0.50/year). Median time to evaluation of arrhythmic events was less in the remote group (2 versus 36 days). More device-related adverse events were observed in the remote group (4.4% versus 1.4%); of these 47% were asymptomatic, but 32% required surgical correction.17

Data from the InSync ICD registry (Medtronic) show that the need to re-program the device during follow-up decreases significantly 6 months after implantation and there is no need to alter device settings even after shocks in 49% of cases.18 Although there are specific device- or arrhythmia-related issues, which are best addressed by a cardiac device specialist, most patients are followed by multiple health-care providers (general cardiologist, heart failure specialist, primary care etc.), which can make office visits redundant. The ongoing ANVITE trial (Biotronik) will compare adverse event rates and quality of life measures of 12-month office ICD follow-up intervals and advanced daily remote monitoring (threshold, impedance, IEGM), with 3-month office follow-up. The VIRTUE trial (Biotronik) will investigate the efficacy and safety of daily remote monitoring with a goal to completely replace routine device office visits over a 7-year follow-up period.

Figure 2: Remotely transmitted intracardiac electrogram (IEGM) from a patient with a cardiac resynchronization therapy defibrillator (CRT-D) and chronic atrial fibrillation (AF). Biventricular stimulation is suboptimal: the device is in VVIR pacing mode and pacing at a rate of 73/min in both ventricles until spontaneous atrioventricular (AV) conduction improves and inhibits ventricular pacing (beats 7–9). Pacing statistics may underestimate the amount of effective biventricular stimulation as beats 11, 13, and 14 are more consistent with spontaneous ventricular activation despite being classified as biventricular paced (beat 11) or ventricular sense response beats (13 and 14). Simple adjustment of pacing settings may resolve this issue. BV, biventricular pace; VS, ventricular sensed event.
Table 1: Clinical trials assessing general safety and efficacy of remote monitoring systems. Ongoing trials were accessed on www.clinicaltrials.gov

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<td>ALTITUDE\textsuperscript{11}</td>
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<td>ANVITE</td>
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<td>Biotronik</td>
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<tr>
<td>CONNECT\textsuperscript{15}</td>
<td>CRT-D, ICD</td>
<td>Medtronic</td>
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<td>ICD</td>
<td>Biotronik</td>
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<td>PM</td>
<td>Biotronik</td>
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<td>Remote monitoring significantly decreased adverse event rate (9.2% versus 13.3%) and hospital stay was shorter by 34%</td>
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<td>PREFER\textsuperscript{13}</td>
<td>PM</td>
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<tr>
<td>QUANTUM</td>
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<td>Patient acceptance, quality of life: early (post implant) versus late (9 months later) initiation of home monitoring</td>
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<td>RAPID-RF\textsuperscript{21}</td>
<td>CRT-D</td>
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<td>REACT</td>
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<tr>
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<td>Biotronik</td>
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<td>Safety and efficacy of home monitoring with as-needed office visits versus conventional follow-up</td>
<td>Home monitoring reduced in-office device evaluations by 45% without affecting morbidity, detected generator and lead issues more frequently and earlier</td>
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<tr>
<td>VIRTUE</td>
<td>PM</td>
<td>Biotronik</td>
<td>123, RCT</td>
<td>Follow-up workload and safety: home monitoring with 12-month remote follow-up versus 12-month office visits</td>
<td>Ongoing, results in 2015</td>
</tr>
</tbody>
</table>

BP, blood pressure; CRT, cardiac resynchronization therapy (device); CRT-D, cardiac resynchronization therapy defibrillator; ICD, implantable cardioverter-defibrillator; PM, pacemaker; RCT, randomized controlled trial.
The incidence of adverse events is higher in the immediate post-implantation period and patients are routinely observed in-hospital after the procedure. Remote monitoring may decrease the hospital stay after pacemaker implantation or generator replacement. The OEDIPE trial (Biotronik) compared patients discharged from the hospital 24 h after a first PM implant or 4–6 h after replacement, followed for 4 weeks with daily remote monitoring, with a control group followed for 4 weeks according to usual practices. The remote group had a 4.1% absolute risk reduction in experiencing at least one treatment-related major adverse event (such as lead issues, death, myocardial infarction, or heart failure (HF) exacerbation). By study design, the mean hospitalization duration also became shorter by 34%. The QUANTUM trial (Biotronik) will compare the effects of immediate or delayed (9-month) initiation of automatic daily remote monitoring after ICD implantation on quality of life, patient perceptions, and frequency of patient–physician contacts.

Combined remote monitoring systems may provide more clinically relevant information but may require more patient interaction. The TRIAGE CRT feasibility study (Biotronik) explored the use of remote device monitoring with weight and blood pressure external telemonitoring in 66 patients with CRT-D. Patient compliance (days transmitted) was 83% for device remote monitoring and 71% for weight and blood pressure measurements over a 6-month follow-up period. The ongoing RAPID-RF (Boston Scientific) multicenter registry will assess a similar combined system.

Cost-efficacy

Comparison of cost-efficacy data is not straightforward given the significant differences in remote monitoring systems and health-care settings. The REFORM trial (Biotronik) compared ICD remote monitoring against conventional follow-up in 115 MADIT II-like patients (27-month follow-up, 3-month remote follow-up with office visits at 3, 15, and 27 months versus 3-month office visits). There was no difference in hospitalizations and mortality, whereas the difference in potential cost-savings was significant: 0.81 physician hours and 712 Euro costs saved each year per patient with remote monitoring. Office visits were reduced by 63.2% in the remote group.

A secondary objective of the CONNECT trial (Medtronic) was to compare cardiovascular health-care utilization between patients in the remote and in-office arms. As the mean length of stay of cardiovascular hospitalization was shorter in the remote group, the mean savings were estimated at $1,793 per hospitalization ($8,114 remote versus $9,822 in-office).

The ongoing EuroECO (Biotronik), EVOLVO (Medtronic) and TARIFF (St. Jude Medical) studies will also compare cost-efficacy and health-care utilization endpoints.

Management of atrial fibrillation

Atrial tachyarrhythmias are frequent causes of morbidity in PM and ICD recipients. A multivariate analysis of the MOST trial data demonstrated that the atrial high rate events recorded by pacemaker diagnostics are independent predictors of total mortality or non-fatal stroke.
(hazard ratio (HR) = 2.79) and chronic atrial fibrillation (HR = 5.93) in patients with sinus node dysfunction.24 Remote monitoring provides access to diagnostic device data on a timely basis, which may guide early intervention (Figure 4, Table 2). The TRENDS observational study (Medtronic) showed that thromboembolic risk was proportional to the atrial tachycardia/atrial fibrillation (AT/AF) burden, defined as the longest total AT/AF duration on any given day during a 30-day period. Two-thousand, four-hundred and eighty-six patients with PMs or ICDs and CHADS2 score > 1 were followed: at a mean follow-up of 1.4 years, annualized thromboembolic risk was 1.1% for zero-, 1.1% for low- (<5.5 h), and 2.4% for high- (>5.5 h) burden subsets.25

The IMPACT trial (Biotronik) investigates whether therapeutic action based on atrial high rate episode information can impact stroke incidence: initiation and termination of anticoagulation with coumadin is managed by a predefined plan in case an episode longer than 48 h is detected, potentially reducing the risk of stroke and systemic embolism, while avoiding the bleeding risks of prolonged anticoagulation. Patients with dual chamber ICD or CRT-D, CHADS2 score ≥1, without history of permanent AF or stroke and with no contraindications for anticoagulation are enrolled and undergo continuous daily remote surveillance. The IMPACT trial will assess the benefits of early optimization of CRT via daily remote monitoring, in addition to investigating the role of remote monitoring of atrial tachyarrhythmias in subjects with CRT-D and paroxysmal or persistent AF. The ongoing CASTLE-AF (Biotronik) uses daily remote monitoring in ICD patients with HF to compare the effect of radiofrequency catheter ablation on mortality and morbidity with that of conventional treatment.

Pacing may be attempted to prevent an AF episode if a predisposing condition is identified. Several atrial pacing algorithms were developed to suppress AF episodes by pacing at a rate that is slightly above the intrinsic rate.
Heart failure

Early detection of hemodynamic changes or fluid overload may help to guide interventions to avoid hospitalizations due to HF exacerbation. Remote monitoring of symptoms, weight, and blood pressure was cost-saving in a meta-analysis of 21 European multidisciplinary HF management trials (5,715 patients, 300–1,000 Euros saved per patient), decreased the number of hospitalizations by 30%, and provided a gain of 0.06 quality-adjusted life-years per patient.** Dedicated implantable HF devices provide additional data with continuous pressure and/or volume status monitoring. The CHAMPION trial (CardioMEMS, Atlanta, GA) showed that wireless monitoring of an implantable pulmonary artery pressure monitor in patients with NYHA class III HF decreased the rate of HF-related hospitalizations.

Figure 4: Remote transmission of long-term implantable cardioverter-defibrillator (ICD) follow-up data. A transient decrease in the amount of atrial pacing, with increased ventricular pacing and persistent decrease in daily patient activity may be observed, which is followed by worsening intrathoracic fluid accumulation. These events preceded the onset of persistent atrial fibrillation by 1 month.
hospitalization by 36% over a 15-month follow-up period. Rate of system-related complications was only 1.4%.\textsuperscript{31} Integration of similar HF monitoring systems, either implanted or external, may provide additional benefits for patients with implanted arrhythmia devices. Patients who transmitted weight and blood pressure data weekly in the subgroup of CRT-D recipients in the ALTITUDE registry had the lowest mortality.\textsuperscript{11} The results of the randomized COMPASS-HF trial (Medtronic) were less convincing: 8% of 274 NYHA class III–IV HF patients with implantable hemodynamic monitors encountered system-related complications, and the rate of HF-related events did not decrease significantly in the remotely monitored group (21%). However, rate of HF hospitalizations decreased by 36%.\textsuperscript{32} The REDUCE-HF trial (Medtronic) will evaluate the safety and effectiveness of an implantable hemodynamic monitor (IHM), and its combination with ICD\textsuperscript{33} (Table 3).

Implanted devices may provide data on intrathoracic impedance, which correlates with fluid status and can thus give early warning of fluid overload. Retrospective analysis of 123 patients with CRT or ICD in the HOMECare Phase 0 registry (Biotronik) showed an increase in mean heart rate at rest and daily mean heart rate within 7 days preceding HF hospitalization in 70% of such events. Similarly, a decrease in the percentage of CRT was observed in 43% and a reduction in the patients’ daily activity in 30%.\textsuperscript{34} The HomeCare II and IN-TIME studies (Biotronik) will further investigate the role of daily intrathoracic impedance measurement and remote monitoring in patients with CRT-D or ICD in order to develop algorithms for device-based early detection and warning of HF deterioration accompanied by pulmonary congestion.\textsuperscript{35} The MORE-CARE and OPTILINK-HF studies (Medtronic) will analyze if remote monitoring of ICD and CRT-D devices with intrathoracic impedance monitoring and automated physician alarm trigger in case of fluid accumulation will affect the mean time between event onset time and clinical decision, mortality, morbidity, and total health-care system utilization.\textsuperscript{36}

CRT requires effective biventricular stimulation with optimal timing (Figure 5). Electrogram-based algorithms have been developed for atrioventricular and ventriculoventricular delay optimization and are currently being investigated in large clinical trials (SMART-AV, Boston Scientific and QuickOpt, St. Jude Medical).\textsuperscript{37,38} Monitoring the stroke volume via intracardiac impedance measurement with conventional pacemaker electrodes may also help optimization of the timing intervals; however, this method is still experimental.\textsuperscript{39}

Prevention of ventricular arrhythmia recurrence

ICD discharges adversely affect quality of life.\textsuperscript{40} Shocks are frequent even in the primary prevention population: 33.2% of patients receive at least one shock over a median of 45.5 months. Both appropriate and inappropriate shocks are associated with a significant increase in the subsequent risk of death (HR = 5.68 and 1.98, consecutively), mostly caused by progressive HF.\textsuperscript{41} The ALTITUDE registry data showed less, but still significant risk: appropriate and inappropriate shocks for ICD had a HR of 2.62. Appropriate shocks for CRT recipients had a HR of 2.09, whereas non-tachyarrhythmia reasons (noise and oversensing) were not associated with significantly increased risk of death in this group, although the sample size was small.\textsuperscript{11} As most adjuvant treatments (antiarrhythmic therapy, ablation, arrhythmia prevention pacing algorithms) may have significant side effects or high cost, careful risk-assessment for ventricular tachycardia/fibrillation (VT/VF) recurrence is essential. Early detection of high-risk conditions, such as increasing frequency and complexity of premature ventricular contractions (PVCs), short–long–short (SLS) PVC patterns, and increasing baseline heart rate, may help to guide therapy. Unfortunately, only a minority of VF episodes had such specific initiation patterns in the PainFree Rx II and EnTrust trials: 30% of VT/VF episodes were preceded by SLS cycles, part of these were VF events permitted or facilitated by pacing,\textsuperscript{32} VT, which occurs more commonly and may be more responsive to ATP, was initiated by a late PVC in 85%, early PVC in 13%, and SLS sequence in 2%, the pattern of initiation was similar for both monomorphic and polymorphic VTs in a study.\textsuperscript{43} The initiating sequences in non-ischemic cardiomyopathy are less well known. In survivors of idiopathic VF, the initiating PVCs seem to mostly have short coupling intervals, without pause dependency.\textsuperscript{44}

A reliable predictor of an imminent episode of VT/VF would have clinical utility if the device were capable of notifying the physician via the remote monitoring system or to provide preventive therapy. Such a predictor was tested in a test set of saved ICD IEGM records. With the analysis of heart rate and specific acceleration patterns in sinus rhythm in a 1.8-h time interval preceding a VT/VF event, sensitivity of 53–83% and specificity of 57–91% were achieved.\textsuperscript{45}

Intracardiac T wave alternans (TWA) is a marker of dynamic electrical instability of ventricular myocardium similar to ECG microvolt TWA. The concordance of these measurements was 87% in a study of 68 patients, the positive predictive value for VT/VF in 1 year was 14% for intracardiac TWA and 17% for surface MTWA, negative predictive value was 82% at 4 years for both.\textsuperscript{46} TWA may be a dynamic phenomenon: a study of 28 ICD patients showed that TWA increased before spontaneous VT/VF episodes, but not before supraventricular arrhythmias—this could be easily monitored continuously with the implanted device.\textsuperscript{47} Increasing incidence of non-sustained ventricular tachycardia (NSVT) may also be a sign of electrical instability. The length, but not the rate, of NSVTs on 24-h ambulatory ECG was a predictor of major arrhythmic events in non-ischemic patients in the MACAS study.\textsuperscript{48} Further studies are needed to investigate whether the results are applicable to the general ICD population or if NSVT suppression would change the risk.

The deleterious effects of right ventricular pacing are well known from previous studies, such as DAVID.\textsuperscript{49} Biventricular stimulation, on the contrary, may decrease the risk of VT/VF.\textsuperscript{50} Higher atrial pacing percentage was associated with an increased incidence of atrial
tachyarrhythmias and with respect to dual chamber pacing, a ratio of paced atrial beats $\geq 48\%$ combined with a ratio of paced ventricular beats $> 40\%$ was associated with an increased probability for VT in a prospective trial.\textsuperscript{51} Pacing interventions, such as application of a rate smoothing algorithm can be of importance to prevent episodes of torsade de pointes in long QT patients; however, data are mostly from small clinical studies.\textsuperscript{52} Rate smoothing has been used to prevent ventricular arrhythmia in the VAST study: with rate smoothing, 23\% of patients experienced a reduction in arrhythmias, 23\% saw an increase and the remaining 54\% had no difference and no individually useful predictors were found.\textsuperscript{53} Pacing algorithms may not be effective if the

### Table 3: Clinical trials assessing the efficacy of remote monitoring systems to manage heart failure. Ongoing trials were accessed on www.clinicaltrials.gov

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<td>CHAMPION\textsuperscript{31}</td>
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<td>CardioMEMS</td>
<td>550, RCT</td>
<td>HF-related hospitalization, freedom from device and system-related complications: IHM with remote monitoring versus no monitoring after implant</td>
<td>30% reduction in HF hospitalizations at 6 months, 36% at mean 15-month follow-up with IHM+remote monitoring. Device or system-related complications only in 1.4%</td>
</tr>
<tr>
<td>COMPASS-HF\textsuperscript{32}</td>
<td>IHM</td>
<td>Medtronic</td>
<td>274, RCT</td>
<td>Freedom from system-related complications, freedom from pressure-sensor failure, reduction in the rate of HF-related events: patients with implantable hemodynamic monitors, treatment guided by remote monitoring versus no monitoring data</td>
<td>HF-related events did not differ significantly. Risk of HF-hospitalization was 36% lower in the IHM+remote monitoring group</td>
</tr>
<tr>
<td>HOMECARE Phase 0\textsuperscript{34}</td>
<td>CRT-P, CRT-D, ICD</td>
<td>Biotronik</td>
<td>123, registry</td>
<td>Daily home monitoring to monitor HF patients with ICD or CRT</td>
<td>Before HF hospitalization, 70% had increase in mean heart rate at rest and in mean heart rate over 24 h in the preceding 7 days, a decrease in the percentage of CRT was observed in 43%, reduction in the daily activity was in 30%</td>
</tr>
<tr>
<td>HomeCARE II</td>
<td>CRT-D, ICD</td>
<td>Biotronik</td>
<td>300, uncontrolled</td>
<td>Performance of intrathoracic impedance measurement with daily monitoring to predict HF exacerbations</td>
<td>Ongoing, results in 2011</td>
</tr>
<tr>
<td>IN-TIME\textsuperscript{35}</td>
<td>CRT-D, ICD</td>
<td>Biotronik</td>
<td>720, RCT</td>
<td>Mortality and hospitalization for HF, prediction of HF events: home monitoring versus conventional follow-up</td>
<td>Ongoing, results in 2012</td>
</tr>
<tr>
<td>MORE-CARE\textsuperscript{36}</td>
<td>CRT-D</td>
<td>Medtronic</td>
<td>1,721, RCT</td>
<td>Mean time between event onset time and clinical decision, mortality and morbidity: remote monitoring of device and intrathoracic impedance versus standard follow-up</td>
<td>Ongoing, results in 2014</td>
</tr>
<tr>
<td>Optilink-HF</td>
<td>CRT-D, ICD</td>
<td>Medtronic</td>
<td>1,000, RCT</td>
<td>Mortality or CV hospitalization: remote monitoring of device and intrathoracic impedance versus standard follow-up</td>
<td>Ongoing, results in 2013</td>
</tr>
<tr>
<td>REDUCE-HF\textsuperscript{33}</td>
<td>ICD w/PRM</td>
<td>Medtronic</td>
<td>1,300, RCT</td>
<td>Freedom from system-related complications, reduction of heart failure events: ICD+IHM with remote monitoring versus control</td>
<td>Ongoing, results in 2011</td>
</tr>
</tbody>
</table>

CRT, cardiac resynchronization therapy (device); HF, heart failure; ICD, implantable cardioverter-defibrillator; IHM, implantable hemodynamic monitor; PM, pacemaker; PRM, pressure monitoring; RCT, randomized controlled trial.

The Journal of Innovations in Cardiac Rhythm Management, July 2011
VT/VF is initiated by a single PVC without pause-dependency and the proarrhythmic potential must not be neglected.

**Conclusion**

Utilization of remote monitoring is rapidly expanding based on encouraging data from large clinical trials; however, lack of standardization is still a significant barrier to widespread acceptance of results. Completely automated mobile systems, instant transmission in case of an adverse event, live IEGM monitoring, remote programming, patient alert with geographic localization, and integration with other patient care/diagnostic modalities may further enhance the efficacy and can be incorporated into existing systems with little additional development. For implanted devices without remote monitoring capabilities, transtelephonic follow-up will remain a useful adjunct to office follow-up. The huge amount of data already obtained from clinical studies may be used develop risk-stratification models and test algorithms to predict future arrhythmia or HF events.

The outlook for this technology is excellent and further innovation will help to improve patient-centered, cost-effective follow-up of implanted cardiac devices.

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**Figure 5:** Intracardiac electrogram (IEGM) snapshot of a routine remote monitoring transmission. The patient has a cardiac resynchronization therapy defibrillator (CRT-D) and is in atrial flutter with complete atrioventricular block. The ventricular rate is biventricular paced in DDRI mode at 67/min. The morphology of the ventricular signal is not uniform despite regular pacing: beats 7 and 11 have a slurred onset. This is most consistent with intermittent loss of capture in one of the paced ventricles. However, this is unrecognized by the device as there are no ventricular sensed events, but may still be responsible for suboptimal CRT. There is occasional undersensing of atrial signals due to atrial blanking after ventricular pacing—this has no consequences in this patient. AS, atrial sensed event; AR, atrial event sensed in the refractory period; BV, biventricular pace.
Remote Monitoring of Cardiac Implantable Electronic Devices

References


