Study Library
CardioSecur’s one-of-a-kind ECG Technology

The 12-lead electrocardiogram (ECG) is the worldwide gold standard for the detection of myocardial ischemia. 12 leads are usually derived from 10 electrodes. The EASI lead system is an alternative to the conventional method, resorting to vector electrocardiography and allowing a three-dimensional view of the heart. EASI was described by Dower et al. in the 1980’s and uses only 5 electrodes (4 recording + 1 grounding) to derive the standard 12 leads. The agreement between an EASI derived 12-lead ECG and a standard 12-lead ECG has been shown in many reviewed studies.

CardioSecur’s technology is based upon the EASI standard, additionally removing the grounding electrode to allow for the registration of a derived 12-lead ECG using only 4 electrodes. The agreement of the modified CardioSecur setting has been clinically proven.

More information from fewer electrodes

Using vector electrocardiography CardioSecur not only reduces the number of electrodes necessary to record an ECG, but it also expands the information attained. In addition to the standard 12 leads (I, II, III, aVL, aVR, aVF, V1-V6), CardioSecur records V7-V9 and VR3-VR9. By recording these additional leads, CardioSecur’s ECG is one step ahead as it remains compliant with recommendations from the European Society of Cardiology (ESC) to also record posterior leads when myocardial infarction is suspected. The value of including additional leads has been shown in many studies, and includes the potential to reduce unnecessary “rule-out MI” admissions.

Personalized ECG Analysis

CardioSecur offers the highest possible sensitivity for detecting myocardial ischemia by comparing the user’s ECG with a previously recorded reference ECG (also recommended by ESC). When first using the CardioSecur Active system, the user records a reference ECG—all subsequent measurements are compared to this reading. This personalized comparison allows for detection of ECG changes, even when preexisting ECG abnormalities are present (left bundle branch block, ST deviation, prior MI changes). The intra-individual ECG comparison also offers the opportunity to detect ECG changes that have pseudo-normalized, in particular those of the ST segment and the T-waves. The serial use of CardioSecur also allows for detection of “silent” myocardial ischemia in patients without symptoms.

The ECG Algorithms

CardioSecur’s algorithms are based upon recommendations of cardiovascular societies and standards. The sensitivity of the algorithms has been proven in an internal validation study including more than 120,000 ECGs based on a normatively standardized ECG database. The CardioSecur algorithms analyse the ECG for changes in heart rate, rhythm and circulation. Detected changes form the basis for the user’s personalized feedback. For example, guidelines for diagnosis of ST segment elevation acute myocardial infarction require the presence of at least 1 mm (0.1 mV) J-point elevation in at least 2 anatomically contiguous leads (with the exception of V2-V3 where the criteria are an elevation of >0.2 mV for men > 40, >0.25 mV for men < 40, and >0.15 mV for women). If changes that fit these criteria are present, the user receives feedback that immediate medical attention is necessary.
References

Internal Research


3. van Langenhove, Glenn and Bruno Schwagten. The Revealing timely ECG changes Decreases the likelihood of Undesireable Cardiac Events-Trial (REDUCE-Trail). Poster presented at ESC Congress 2014: Barcelona.

About the EASI Technology


**ESC Guidelines**


The Value of Additional ECG Leads


Klaus Bonaventura¹, Ernst Wellnhofer² and Eckart Fleck²

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Background: Electrocardiograms (ECGs) derived by the transformation of three bipolar quasi-orthogonal leads have, according to EASI, been introduced for many years for use in emergency situations and for the monitoring of patients during the acute phase of myocardial infarction (MI). Theoretically, a further reduction and simplification of the classic EASI setting of five electrodes may even improve acceptance of the derived 12-lead ECG in these critical situations, especially in the telemedical use and for monitoring of cardiovascular patients. The objective of the present study was to evaluate the comparability of the 12-lead ECG derived by a system that reduces the classic EASI setting from five to four electrodes with the standard 12-lead ECG in the detection of acute MI induced during percutaneous transluminal coronary angioplasty (PCI).

Methods: To determine whether a 12-lead ECG derived from a reduced EASI setting using only 4 electrodes would demonstrate typical ST-segment changes of ischemia during PCI (percutaneous coronary intervention) 24 patients with overall 148 episodes of balloon-induced myocardial ischemia were monitored with continuous 12-lead ST-segment monitoring during PCI. A derived 12-lead ECG was registrated by the 4-electrode system. 2 blinded cardiologists not involved in the intervention compared both ECGs for each patient.

Results: Of the 148 episodes of balloon inflation recorded with the derived ECG, 104 (70.3 %) were associated with typical and significant ischemic ST-segment changes during balloon inflation. The amplitudes of these ST deviations were similar to those observed during transient myocardial ischemia observed in clinical settings (median peak ST deviation, 234 microV). There was agreement regarding presence or absence of ischemia in 147 of 148 episodes, with both derived and standard electrocardiographic methods (>99 % agreement). With use of the standard ECG as the ‘gold standard’ for ischemia diagnosis, there were no false negatives (0 %) and only one false-positive (0.7 %) with the derived ECG. There was no significant difference between the two techniques by linearity tests (p>0.1). Bland–Altman analysis showed no significant bias. Moreover, both methods demonstrated 100% concordance with respect to localization of myocardial ischemia (anterior, inferior and lateral).

Conclusions: The new 4-electrode set 12-lead ECG is as an alternative to the standard 12-lead ECG with 10 electrodes in emergency situations and for cardiovascular monitoring.

DOI: 10.15420/ecr.2012.8.3.179
2. Comparative Study of the CardioSecur Pro ECG system with the EASI Philips M2601B.

Abstract presented: eCardiology Congress, 2016: Berlin

David Triebl, Peter Kenedi, Istvan Predas, Adam Szekely, Markus Skribek, Markus Riemenschneider
Personal MedSystems GmbH Frankfurt, Central Hospital of the Hungarian Defence Forces Budapest

**Aim of the Study:** A comparative study was conducted to validate the ECG measurements of CardioSecur. CardioSecur is a tablet-based ECG system using 4 electrodes to derive a 22-channel ECG (standard 12-leads + V7-V9 and VR1-VR9). The technology of CardioSecur is based on the calculation of 12 leads from 4 electrodes which are comparable to the standard 12 lead ECG.

**Methods:** To assess both the technical and the medical comparability of the systems, the setup was divided into two procedures, the first test covering the medical diagnostic accuracy of the two systems, the second test covering the technical comparability of the ECG signal, generated with the reduced lead system. In the first test, ECG measurements were taken from 41 individuals with both systems. A clinical diagnosis was made on both ECGs and the orientation of the P, R, S, and T waves were evaluated and compared. To assess the technical waveform of the two systems, ECGs were simulated with an ECG simulator to ensure identical electrical input on both systems. These ECGs were simulated at frequencies between 30 and 180bpm. Additionally, pathological ECG patterns were simulated and recorded with the systems. These waveforms were compared with respect to morphology and height of the electrical signal in the standard 12 leads of both systems.

**Results:** The clinical diagnosis for 41 measured patients was identical in the ECGs measured with both CardioSecur and Philips M2601B. This ensures equal clinical sensitivity and specificity in both devices. The additional 10 leads of CardioSecur were not part of the study as the Philips device does not offer this option. The second test, covering a technical analysis including waveforms and peak heights, revealed differences in the heights of the R and S wave. CardioSecur showed an absolute peak 10% higher than the Philips device. This can be explained by the use of different filter settings in the devices. The Philips M2601B clearly states that the recorded ECG may not be used for ST-segment evaluation. CardioSecur uses a filter setting compliant with the regulatory standards to allow for ST-segment evaluation. Consequently, a difference in absolute peak height can also appear. Morphologically, all patient ECGs showed identical orientation of the measured parameters. The simulated ECGs showed identical morphology for all measured settings.

**Conclusion:** This study has shown that the clinical information in the CardioSecur device is identical to the information of ECGs of the Philips M2601B device. A closer examination of the raw signal in peak heights arises from the different filter systems. Morphologically, the orientation of all recorded ECGs and R-wave progression were identical in all measured ECGs. Therefore, the diagnostic capabilities of the CardioSecur device can be seen as wholly comparable to those of the Philips ECG. The possible benefits of an additional 10 leads in the CardioSecur device will be the subject of future studies.
3. The Revealing Timely ECG Change Decreases the Likelihood of Undesirable Cardiac Events-Trial (REDUCE-Trial).

Presented: European Society of Cardiology Conference, 2014: Barcelona

van Langenhove¹, Schwagten B¹.

1. ZNA Middelheim, Antwerpen, Belgium.

**Background:** ECG technology is extremely useful in the diagnosis of a wide variety of cardiac diseases. Plenty of arrhythmic and ischemic conditions however are hard to diagnose and therefore treat because they don’t appear during the physician’s consultation. For cases with persistent or recurrent problems single-lead event recorders, holters or implantable devices have been developed to diagnose the underlying disease or symptoms. However due to their susceptibility to artefacts, the fact that they don’t provide 12-lead ECG information data and a necessary invasive procedure, they may not lead to satisfying results in all patients (1).

**Purpose:** To investigate whether the mobile 12-lead ECG CardioSecur™ (Personal MedSystems GmbH) is offering additional value in the management of patients with cardiac preconditions. CardioSecur™ is a mobile 12-lead ECG device based on the validated EASI-ECG-technology and allows for a 12-lead ECG using four electrodes only. Once the patient has recorded a reference ECG on the device he can perform control-readings. An algorithm that is based on clinical guidelines will detect ECG changes between those two ECGs and give the patient a recommendation to act. For example, in the case of minor ECG changes that patient will receive a yellow-warning and the information to make an appointment with his doctor. If major ECG changes occur, the patient will receive a red warning telling him to contact his physician immediately. If no ECG changes are detected, and also if minor changes are detected, the patient is informed that he should see a physician if symptoms pertain for longer than 20 minutes. ECG data can be transmitted to a database that can be accessed by the physician or medical institution.

**Methods:** This is a monocentric, single-armed non-randomized trial. Patients were asked to undertake measurements with the device once weekly and every time they were experiencing symptoms over a maximum period of three months. Subsequently patients were followed up for nine months. At inclusion and after follow-up patients were underwent a comprehensive diagnostic assessment consisting of a standard 12-lead ECG, echocardiogram and exercise-ECG. Inclusion criteria: 18 - 80 years of age plus CABG, PCI, AMI in the last 12 months, angina pectoris treated pharmacologically, significant rhythm disturbance for which they received either a pharmacological or electrophysiological intervention, or recurrent palpitations of unknown origin in the past. Ability to handle device, regular access to the internet, signed the informed consent form. The following outcomes were assessed: Is the device able to detect ECG-changes and give the patient a correct recommendation and clinical relevance. The local ethic’s committee at the ZNA Middelheim approved the study, that was conducted in line with the guidelines for GCP and the declaration of Helsinki.

**Results:** In total 51 patients were recruited between 11/2011 and 03/2012. Patient characteristics and main symptoms are shown in table 1.
Patients recorded in total 1.237 ECG-readings with 2.2% of the measurements being symptom-induced and the rest being undertaken during weekly measurements. In five patients (9.8%) the CardioSecur™ device showed its clinical relevance: It diagnosed a new or so far undiagnosed condition and led to a successful treatment. A full overview of the results of the readings, number of critical results, diagnosis and interventions performed is given in table 2.

<table>
<thead>
<tr>
<th>Patient Reference ID</th>
<th>Result</th>
<th>n of Readings</th>
<th>Diagnosis</th>
<th>Intervention</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>red</td>
<td>5</td>
<td>90% stenosis</td>
<td>PCI</td>
</tr>
<tr>
<td>2</td>
<td>red</td>
<td>4</td>
<td>Paroxysmal AF</td>
<td>PVI</td>
</tr>
<tr>
<td>3</td>
<td>yellow</td>
<td>46</td>
<td>Monofocal ventricular premature beats, with bi- and trigemina</td>
<td>Focal ablation</td>
</tr>
<tr>
<td>4</td>
<td>yellow</td>
<td>18</td>
<td>AV nodal re-entry tachycardia</td>
<td>Ablation</td>
</tr>
<tr>
<td>5</td>
<td>yellow</td>
<td>7</td>
<td>Paroxysmal AF</td>
<td>PVI</td>
</tr>
</tbody>
</table>

Table 2: Results of clinical value

Out of those five patients, one patient (ID 1) was suffering from a severe ischemia. Four patients had arrhythmias with two suffering from atrial fibrillation (ID 2 & 5), one from monofocal ventricular premature beats (ID 3), with bi- and trigemina and one from AV nodal re-entry tachycardia (ID 4). During the study period, no events were reported that the device should have been able to detect. Patients reported a high ease of use and 80% would recommend the device to a friend or family member.

Conclusions: We showed that CardioSecur™ system is an important tool for diagnosing cardiovascular disease and adds value in the management of patients with cardiac diseases such as rhythm disturbances and ischemic episodes. Further research is needed to validate this first study in larger patient cohorts and assess the long-term effects of this 12-lead mobile ECG.

4. The ECGD: a derivation of the ECG from VCG leads.


Dower GE

No Abstract available.

PMID: 6736842
5. On deriving the electrocardiogram from vectorcardiographic leads.


Dower GE, Machado HB, Osborne JA

Abstract
The issue of whether a traditional or scientifically based system for applying electrodes to the body for routine electrocardiography may be resolved by deriving the 12-lead ECG from the Frank XYZ signals. The result, the ECGD, is sufficiently close to the ECG for serial comparisons to be valid. Reducing data acquisition to the XYZ signals alone has several technical advantages. These have been realized with the introduction of a computer system employing the ECGD at a large general hospital. Plotting the lead vectors of the ECGD on Aitoff’s projection of the sphere brings out important relationships between the leads, one to another, and to the spatial directions of the QRS and T vectors. Reversing the polarity of a VR enhances the sequential relationship between the limb leads; this is taken advantage of in an educational display generated by the computer.

PMID: 6993081
6. Deriving the 12-lead electrocardiogram from four (EASI) electrodes.

*Journal of Electrocardiology*, vol. 21, supplement, pp. 182-7.

Dower GE, Yakush A, Nazzal SB, Jutzy RV, Ruiz CE

**Abstract**

Computerized interpretation of the electrocardiogram has now advanced to computerization of the electrocardiograph, resulting in greatly increased versatility, including the capacity for adapting to a variety of lead systems rather than being tethered to the old Einthoven-Wilson-Goldberger (EWG) system. Many varieties of display beyond the 12-lead ECG are also available in software. To date, these new and interesting capabilities have scarcely been exploited. The EASI lead system uses the E, A, and I electrode positions of the Frank lead system, plus an electrode, S, positioned over the upper end of the sternum and, if necessary, ground (anywhere convenient). Its outputs form quasi-xyz signals, x'y'z', that can be approximately transformed into xyz signals by means of a matrix derived from the EASI lead vectors. The result forms a good basis for deriving the 12-lead ECG, using previously published coefficients for the Frank lead system. The match with the conventional ECG can then be improved by statistical means. The results are surprisingly good, and certainly of clinical value. Recent widespread interest in silent ischemia and its detection through Holter monitoring suggests an immediate application which has been rendered practical by the recent introduction of three-channel recorders. The EASI electrode positions give technically satisfactory Holter recordings. Very compact three-channel, multiplexed, radio telemetry equipment is now commercially available and provides another application for the EASI 12-lead ECG. (ABSTRACT TRUNCATED AT 250 WORDS)

PMID: 3216172
Electrocardiographic systems with reduced numbers of leads–synthesis of the 12-lead ECG.


Tomasic I, Trobec R

Abstract
Systems with reduced numbers of leads that can synthesize the 12-lead electrocardiogram (ECG) with an insignificant or a small loss of diagnostic information have been proposed. The advantage over standard 12-lead ECG systems is the smaller number of measurement sites (i.e., electrodes) and, consequently, fewer wires. In this paper, we review all the important systems with reduced numbers of leads together with the methodology for synthesizing the leads. The fundamental theoretical background necessary to understand the most important concepts related to the synthesis is included. The presented theoretical and experimental justifications for the synthesis show that it is not necessary to measure a large number of leads directly, because the standard 12-lead ECG and arbitrary additional leads can be synthesized. Various approaches to evaluating the synthesized 12-lead ECG are defined and explained, and a number of systems that synthesize 12-lead ECG are presented as they were introduced in the literature. We cover the developments and improvements from the 1940s to the present day. The systems are classified on the basis of the synthesis method used, the approach to the evaluation of the synthesized ECG (depending on the measurement sites used), and on the number and types of leads employed. Based on a detailed assessment of state-of-the-art systems, open problems and challenges are highlighted, while further developments of electrocardiographic systems are envisaged.

PMID: 23708809
A vector-based, 5-electrode, 12-lead monitoring ECG (EASI) is equivalent to conventional 12-lead ECG for diagnosis of acute coronary syndromes.


Wehr G, Peters RJ, Khalife K, Banning AP, Kuehlkamp V, Rickards AF, Sechtem U

**Aims:** The conventional 12-lead electrocardiogram (cECG) derived from 10 electrodes using a cardiograph is the gold standard for diagnosing myocardial ischemia. This study tested the hypothesis that a new 5-electrode 12-lead vector-based ECG (EASI; nPhilips Medical Systems, formerly Hewlett Packard Co, Boeblingen, Germany) patient monitoring system is equivalent to cECG in diagnosing acute coronary syndromes (ACSs).

**Methods:** Electrocardiograms (EASI and cECG) were obtained in 203 patients with chest pain on admission and 4 to 8 hours later. Both types of ECGs were graded as ST-elevation myocardial infarction if at least 1 of the 2 consecutive recordings showed ST elevation more than 0.2 mV, as ACS if one or both showed ST elevation less than 0.2 mV, T-wave inversion, or ST depression. Otherwise, the ECG was graded negative.

**Results:** Final diagnosis was identical in 177 patients (87%; 95% confidence interval (CI), 82%-91%; kappa=0.81; SE=0.035). ST-elevation myocardial infarction was correctly identified or excluded by EASI with a specificity of 94% (95% CI, 89%-97%) and a sensitivity of 93% (95% CI, 86%-97%; using cECG as the gold standard). Of 118 patients with enzyme elevations, an almost identical number (72 (61% by EASI) and 73 (62% by cECG) had ST elevations. Both techniques were equivalent in predicting subsequent enzyme elevation (identical, 108/143; 75% of ACS and ST-elevation myocardial infarction ECGs by ESI and cECG). Thus, both ECG methods had exactly the same specificity of 59% (85% CI, 48%-69%) and sensitivity of 91% (95% CI, 85%-96%) for detecting myocardial injury.

**Conclusion:** EASI is equivalent to cECG for the diagnosis of myocardial ischemia.

PMID: 16387045
9. The relative accuracies of ECG precordial lead waveforms derived from EASI leads and those acquired from paramedic applied standard leads.


**Abstract**

Accurate precordial electrode placement can be difficult in emergency situations leading either to loss of time or diminished accuracy. A possible solution is the quasi-orthogonal EASI lead system, with only five electrodes and easily defined landmarks to provide a derived 12-lead electrocardiogram (ECG). The purpose of this study was to test the hypothesis that precordial waveforms in EASI-derived ECGs have no greater deviation from those in gold standard ECGs, than do the precordial waveforms in paramedic acquired standard ECGs. Twenty paramedics applied the standard precordial electrodes employing the routine procedure. A certified ECG technician applied the 6 standard precordial electrodes in their correct gold standard positions, and the EASI electrodes. 12-lead ECGs were obtained from the paramedics’ standard leads, and derived from the EASI leads, for comparison with the gold standard ECG. In each precordial lead recording, 6 computer-measured QRS-T waveform parameters were considered. Differences between deltaEASI-gold standard versus deltaparamedic-gold standard were calculated for every waveform in every lead resulting in 720 comparisons. EASI and paramedic results were "equally accurate" in 47%, the paramedic was more accurate in 31%, and EASI was more accurate in the remaining 22%. The differences from gold standard recording of precordial waveforms in ECGs derived from the EASI leads and those acquired via paramedic-applied standard electrodes are similar. The results suggest that the EASI lead system may provide an alternative to the standard ECG precordial leads to facilitate data acquisition and possibly save valuable time in emergency situations.

PMID: 12942479
Abstract
The purpose of the study was to compare the EASI system with the standard 12-lead surface electrocardiogram (ECG) for the accuracy in detecting the main electrocardiographic parameters (J point, PR, QT, and QRS) commonly monitored in patients with acute coronary syndromes or heart failure. In this observational comparative study, 253 patients who were consecutively admitted to the coronary care unit with acute coronary syndrome or heart failure were evaluated. In all patients, two complete 12-lead ECGs were acquired simultaneously. A total of 6,072 electrocardiographic leads were compared (3,036 standard and 3,036 EASI). No significant differences were found between the investigate parameters of the two measurement methods, either in patients with acute coronary syndrome or in those with heart failure. This study confirmed the accuracy of the EASI system in monitoring the main ECG parameters in patients admitted to the coronary care unit with acute coronary syndrome or heart failure.

PMID: 28412843
11. Derived 12-lead electrocardiogram in the assessment of ST-segment deviation and cardiac rhythm.

*Journal of Electrocardiology*, vol. 39, issue 1, 2006, pp. 7-12.

Chantad D, Krittayaphong R, Komoltri C

**Background:** There are little data on the validation of 12-lead electrocardiogram (ECG) derived by the EASI lead system used for continuous monitoring in critical care settings.

**Objective:** The objectives of this study were to determine the accuracy of 12-lead ECG derived by the EASI lead system in the detection of ST-segment deviation and cardiac rhythm compared with the standard 12-lead ECG.

**Methods:** All patients admitted to the coronary care unit were studied. Kappa statistics was used to calculate the agreement between both ECG systems in the determination of cardiac rhythm and premature ventricular complex morphology. ST-segment analysis was performed in patients with acute coronary syndromes. Pearson correlation was used to correlate the ST-segment deviation between both techniques. The sensitivity and specificity of the determination of significant ST-segment deviation by the EASI lead system were calculated.

**Results:** There were a total of 282 patients enrolled in this study. There was a complete agreement in the interpretation of cardiac rhythm between the 2 methods (kappa = 1). Analysis of ST-segment deviation of 12-lead ECG also showed a significant correlation (correlation coefficient varied from 0.62 in lead I to 0.823 in lead aVF with a P value of <.001 in all leads) between the 2 methods with very high sensitivity and specificity in the detection of significant ST-segment elevation and depression.

**Conclusion:** The 12-lead ECG derived by the EASI lead system is an accurate and reliable information for the assessment of ST-segment deviation and cardiac rhythm in critically ill patients.

PMID: 16387043


**Abstract**

To compare the diagnostic yield of electrocardiograms (ECGs) recorded by 12 standard leads with that of 12-lead ECGs derived from 3 bipolar EASI leads, we analyzed pertinent ECG data for 290 normal subjects and 497 patients who had had a prior myocardial infarction (MI); the latter group comprised 36 patients with a non-Q MI, 282 patients with a Q-wave MI, and 179 patients with a history of ventricular tachycardia (VT). We first estimated statistically an optimal set of coefficients for deriving the 12 standard leads from EASI leads and assessed this transformation in terms of goodness of fit. To gauge the diagnostic information content of the recorded vs. derived 12-lead ECGs, we performed successively two-group diagnostic classification--based on the Cardiac Infarction Injury Score (CIIS)--separating each of the patient subgroups from the normal group; the classification was repeated for 200 sets of patients selected randomly (with replacement), and the results were plotted as mean receiver operating characteristics. We found that derived 12-lead ECGs correlated well with the recorded ones, and reproduced faithfully the diagnostic features needed for the CIIS. When the CIIS was determined from features of the recorded standard 12 leads, its mean diagnostic performance (assessed in terms of area under the receiver operating characteristics curve) was 0.9004 for detecting non-Q MIs, 0.9546 for Q-wave MIs, and 0.9919 for MIs complicated by a history of VT. When, instead, features of derived 12 leads were used to determine the CIIS, diagnostic performance remained virtually unchanged (at 0.8905, 0.9531, and 0.9906, respectively). We conclude that, in our population, EASI-derived 12-lead ECGs contain nearly the same diagnostic information as standard 12-lead ECGs.

PMID: 11265716
13. Comparison of a vectorcardiographically derived 12-lead electrocardiogram with the conventional electrocardiogram during wide QRS complex tachycardia, and its potential application for continuous bedside monitoring.


Drew BJ, Scheinman MM, Evans GT Jr.

**Abstract**

Previous investigators published conflicting reports comparing a vectorcardiographically derived electrocardiogram (ECGD) with the conventional 12-lead one (ECG). Prior comparisons were obtained in adults during sinus rhythm, but never in patients with wide QRS complex tachycardia. The ECGD was evaluated during baseline rhythms in patients with varying cardiac diagnoses, and the diagnostic accuracy of the 2 methods was compared during 64 episodes of wide QRS complex tachycardia in 49 patients during cardiac electrophysiologic study. All leads of the 12-lead ECGD closely resembled the conventional ECG in baseline and tachycardia tracings, except leads V3 and V4. QRS voltages were less in the ECGD, resulting in an inability to detect left ventricular hypertrophy in one third of patients with that diagnosis. There was excellent agreement between the ECGD and ECG in diagnosing prior myocardial infarction (92%), ventricular preexcitation patterns (100%), bundle branch and fascicular blocks (100%), and axis deviation. The ECGD was equally as valuable as the ECG in the diagnosis of wide QRS complex tachycardia. There was perfect agreement between the 2 lead systems in application of the morphologic criteria differentiating supraventricular tachycardia with aberration from ventricular tachycardia in leads V1, V2 and V6, and for criteria requiring axis determination and measurement of RS intervals in the precordial leads. The ECGD tracings contained less muscle artifact during body movements (e.g., after direct-current defibrillation). In conclusion, the ECGD's close correlation with the ECG, and its technical superiority and simple 5 torso-positioned electrode configuration make it worth pursuing as an option for continuous bedside monitoring.

PMID: 1536110
14. Comparison of the standard ECG with the EASIcardiogram for ischemia detection during exercise monitoring.


Feldman CL, MacCallum G, Hartley LH

**Abstract**

The methodology for constructing the 12 lead ECG from Dower's EASI lead system-5 electrodes, all located at easy landmarks over bony areas of the thorax-has been recently updated with newly optimized coefficients. To test the ability of the updated EASIcardiogram to detect ischemia, 54 patients undergoing symptom limited, Bruce protocol exercise testing were studied with simultaneous standard 10 electrodes and EASI 5 electrode lead systems. Concordance between the two systems was 83%. In 34 patients with recent coronary angiograms sensitivity and specificity of the EASIcardiogram for detection of coronary disease were at least as good as that of the standard ECG. It is concluded that ST segment depression detected by the EASIcardiogram is very similar to that which is detected by the standard ECG and that the EASIcardiogram appears to have sensitivity and specificity at least equal to that of the standard ECG for detection of myocardial ischemia.

DOI: 10.1109/CIC.1997.647903
A Comparison between EASI System 12-lead ECGs and standard 12-lead ECGs for improved clinical nursing practice.


Lancia L, Pisegna Cerone M, Vittorini P, Romano S, Penco M

**Aims and Objectives:** This study was carried out to verify the accuracy of 12-Lead ECG, obtained through a continuous ECG monitoring system with five cables positioned in EASI mode, to identify basic ECG alterations.

**Background:** This study concerns continuous ECG monitoring systems in Coronary Care Units. Continuous ECG monitoring is an important device for nursing surveillance and is useful in decreasing adverse events.

**Design and Method:** Thirteen patients admitted consecutively to the Coronary Care Unit for Acute Myocardial Infarction underwent daily and simultaneous recording of a 12-lead ECG using both procedures: EASI ECG and STANDARD ECG. A sample of 1,164 ECG leads acquired in EASI mode was compared with a sample of as many ECG leads acquired using the standard procedure with a traditional cardiograph.

**Results and Conclusions:** In the Coronary Care Unit, Continuous ECG monitoring with five cables positioned in EASI mode is a valid alternative to the standard 12-lead ECG for cardiac rhythm abnormalities detection and for acute myocardial ischemia and old myocardial infarction assessment. Therefore, the EASI system might be advantageous for long-term patient monitoring.

**Relevance to clinical practice:** The EASI system represents a valid device for the nursing surveillance of patients who need continuous ECG monitoring, improves clinical nursing practice in Coronary Care Units, supports the reduction of adverse events such as cardiac arrest and reduces the hospital costs.

PMID: 18205693
Abstract
This study was performed to compare a derived 12-lead electrocardiogram (ECG) using a simple 5-electrode lead configuration (EASI 12-lead) with the standard ECG for multiple cardiac diagnoses. Accurate diagnosis of arrhythmias and ischemia often require analysis of multiple (ideally, 12) ECG leads; however, continuous 12-lead monitoring is impractical in hospital settings. EASI and standard ECGs were compared in 540 patients, 426 of whom also had continuous 12-lead ST segment monitoring with both lead methods. Independent standards relative to a correct diagnosis were used whenever possible, for example, echocardiographic data for chamber enlargement-hypertrophy, and troponin levels for acute infarction. Percent agreement between the 2 methods were: cardiac rhythm, 100%; chamber enlargement-hypertrophy, 84%-99%; right and left bundle branch block, 95% and 97%, respectively; left anterior and posterior fascicular block, 97% and 99%, respectively; prior anterior and inferior infarction, 95% and 92%, respectively. There was very little variation between the 2 lead methods in cardiac interval measurements; however, there was more variation in P, QRS, and T-wave axes. Of the 426 patients with ST monitoring, 138 patients had a total of 238 ST events (26, acute infarction; 62, angioplasty-induced ischemia; 150, spontaneous transient ischemia). There was 100% agreement between the 2 methods for acute infarction, 95% agreement for angioplasty-induced ischemia, and 89% agreement for transient ischemia. EASI and standard 12-lead ECGs are comparable for multiple cardiac diagnoses; however, serial ECG changes (eg, T-wave changes) should be assessed using one consistent 12-lead method.

PMID: 10688301
17. Comparison of standard and derived 12-lead electrocardiograms for diagnosis of coronary angioplasty-induced myocardial ischemia.


Drew BJ, Adams MG, Pelter MM, Wung SF, Caldwell MA

**Abstract**

To determine whether a derived 12-lead electrocardiogram (ECG) would demonstrate typical ST-segment changes of ischemia during percutaneous transluminal coronary angioplasty (PTCA), 207 patients were monitored with continuous 12-lead ST-segment monitoring during angioplasty. Additionally, to compare the derived and standard ECGs during known periods of ischemia with PTCA balloon inflation, 151 patients were recorded with both electrocardiographic methods during the procedure. Of the 207 patients recorded with the derived ECG, 171 (83%) had typical ischemic ST-segment changes during PTCA balloon inflation. The amplitudes of these ST deviations were similar to those observed during transient myocardial ischemia observed in clinical settings (median peak ST deviation, 225 microV). There was agreement regarding presence or absence of ischemia in 150 of the 151 patients recorded with both derived and standard electrocardiographic methods (> 99% agreement). With use of the standard ECG as the "gold standard" for ischemia diagnosis, there were no false-positive results and only 1 false-negative result with the derived ECG. Furthermore, there was nearly perfect agreement between the two 12-lead methods in terms of anterior versus inferior wall patterns of ischemia. Future studies are required to determine whether continuous monitoring with a derived ECG would improve diagnosis and lead to better patient outcomes.

PMID: 9068524
18. Comparability of 12-lead ECGs derived from EASI leads with standard 12-lead ECGs in the classification of acute myocardial ischemia and old myocardial infarction.


Rautaharju PM, Zhous SH, Hancock EW, Horácek BM, Feild DQ, Lindauer JM, Wagner GS, Pahlm O, Feldman CL

**Abstract**

We compared 12-lead electrocardiograms (ECGs) derived with an improved transformation matrix from EASI leads and standard 12-lead ECGs in the detection of acute myocardial ischemia and old infarction (MI). For the ischemia test, we used ECGs of 40 patients recorded prior to and at peak inflation during percutaneous transluminal coronary angioplasty, and for old MI we used test ECGs of 382 non-MI subjects and of 472 patients with prior MI documented by enzyme findings. Two experienced ECG readers served as separate, independent standards for lead-set comparisons, and the Philips ECG analysis program also classified the ECGs. The results showed no significant differences between the two lead sets in the detection of acute inflation-induced ischemia or of old MI according to coding by the electrocardiographers or the computer program. No significant differences were found between the electrocardiographers and the lead sets for acute ischemia. Classification differences between the electrocardiographers were larger than those between the lead sets for acute and old MI and were significant for the latter (P <.001). A more detailed comparison of the lead sets suggested a possible need for modified old-MI criteria and optimization of ST classification thresholds for acute ischemic injury, specific for the EASI 12-lead ECG. We conclude that the EASI-derived 12-lead ECG deserves serious consideration as an alternative to the standard 12-lead ECG in emergency situations and for monitoring in acute-care setting.

PMID: 12539097
Body position effects on the ECG: implication for ischemia monitoring.


Adams MG, Drew BJ

Abstract
Rotation of the heart in relation to surface electrocardiographic (ECG) electrodes when a patient turns to one side has been reported to cause ST-segment shifts, triggering false alarms with continuous ST-segment monitoring. We prospectively analyzed ST-segment and QRS complex changes in both standard and derived ECGs in 40 subjects (18 with heart disease and 22 healthy) in supine, right- and left-lying positions. Of the 40 subjects, 6 (4 cardiac, 2 healthy) developed positional ST deviations of 1 mm or more on the standard ECG. In the derived method, five of the same six subjects showed ST-segment deviation of which most occurred in the left-lying position. Positional ST changes were most frequent for males and for cardiac patients (33%). Changes in QRS complex morphology were common on the standard (28 of 40, 70%) and less frequent on the derived ECGs (17 of 40, 43%), occurring in both healthy and cardiac subjects. QRS axis changes occurred only on the standard ECG. It was concluded that (1) right and left side-lying positions frequently induce clinically significant ECG changes; (2) positional ST-segment deviation is less frequent than previously reported and is most likely to occur in males with cardiac disease; and (3) the derived method is less prone to positional QRS changes than the standard ECG.

PMID: 9375904
20. Value of a derived 12-lead ECG for detecting transient myocardial ischemia.

*Journal of Electrocardiology*, vol. 28, supplement, 1995, p. 211.

Drew BJ, Adams MG, Wung SF, Dower GE

No abstract available
21. Diagnosing ischemia from the bedside monitor.

*Prognostic Cardiovascular Nursing*, vol. 11, issue 1, 1996, pp. 45-6.

Drew BJ, Ide B

No abstract available
22. Derived 12-lead ECG. Comparison with the standard ECG during myocardial ischemia and its potential application for continuous ST-segment monitoring.


Drew BJ, Koops RR, Adams MG, Dower GE

No Abstract available
Abstract
To investigate the possibility of simplifying electrocardiogram (ECG) recording in children, we compared waveforms in conventional 12-lead ECGs to those derived from EASI leads in 221 children of various ages. The conventional 12-lead ECGs and the ECGs using EASI electrode positions were collected simultaneously. We developed and determined the value of age-specific transformation coefficients for use in deriving 12-lead ECGs from the signals recorded at the EASI sites. We compared the results of using age-specific coefficients to the results of using adult coefficients and studied the "goodness-of-fit" between the conventional and the derived 12-lead ECGs. The age-specific coefficients performed slightly better than the adult coefficients, and good agreement was usually attained between the conventional 12-lead ECG and the EASI-derived 12-lead ECG. Our conclusion is that EASI leads in children have the same high levels of "goodness-of-fit" to replicate conventional 12-lead ECG waveforms, as reported earlier in adults.

PMID: 12607193
24. Diagnostic conclusions from the EASI-derived 12-lead electrocardiogram as compared with the standard 12-lead electrocardiogram in children.


**Background:** Fewer electrodes on more easily located places would facilitate electrocardiogram (ECG) recording. To investigate the possibility of simplifying ECG recording in children, we compared the diagnostic conclusions when interpreting standard versus EASI-derived 12-lead ECGs. Our hypothesis was that the variation of the interpretation of standard versus EASI-derived 12-lead ECGs was not greater than the intrareader variation of the interpretation of standard ECGs.

**Methods:** The study included 221 children. The 2 lead systems were recorded simultaneously. Two experienced pediatric cardiologists interpreted the ECGs. First, the reader interpreted a set of 221 ECGs with randomly allocated standard and EASI-derived 12-lead ECGs. Next, the reader interpreted the complementary ECG set without having access to the first set. Finally, the reader reinterpreted the standard ECGs from 98 children.

**Results:** The variation of the interpretation of standard versus EASI-derived 12-lead ECGs was only slightly larger than the intrareader variation of the interpretation of standard ECGs.

**Conclusions:** For most of the electrocardiographic diagnoses, the conclusions from EASI-derived 12-lead ECGs were similar to those from standard ECGs. These findings support the suggestion that the EASI lead system is a potential alternative to the standard ECG in children.

PMID: 16644336
Abstract
The goals of electrocardiographic (ECG) monitoring in hospital settings have expanded from simple heart rate and basic rhythm determination to the diagnosis of complex arrhythmias, myocardial ischemia, and prolonged QT interval. Whereas computerized arrhythmia analysis is automatic in cardiac monitoring systems, computerized ST-segment ischemia analysis is available only in newer-generation monitors, and computerized QT-interval monitoring is currently unavailable. Even in hospitals with ST-monitoring capability, ischemia monitoring is vastly underutilized by healthcare professionals. Moreover, because no computerized analysis is available for QT monitoring, healthcare professionals must determine when it is appropriate to manually measure QT intervals (eg, when a patient is started on a potentially proarrhythmic drug). The purpose of the present review is to provide 'best practices' for hospital ECG monitoring. Randomized clinical trials in this area are almost nonexistent; therefore, expert opinions are based upon clinical experience and related research in the field of electrocardiography. This consensus document encompasses all areas of hospital cardiac monitoring in both children and adults. The emphasis is on information clinicians need to know to monitor patients safely and effectively. Recommendations are made with regard to indications, timeframes, and strategies to improve the diagnostic accuracy of cardiac arrhythmia, ischemia, and QT-interval monitoring. Currently available ECG lead systems are described, and recommendations related to staffing, training, and methods to improve quality are provided.


PMID: 28886621


PMID: 26320110
28. Third universal definition of myocardial infarction.

*J Am Coll Cardiol*, vol. 60, issue 16, pp. 1581-98.


PMID: 22958960
Study Objectives: To compare a new 22-lead ECG with the 12-lead ECG for diagnosis of acute myocardial infarction (AMI).

Design: Prospective study of all consenting patients presenting to the emergency department with chest pain.
Setting: Urban hospital ED.

Type of Participants: 163 patients admitted with a cardiac-related diagnosis and complete data sets of 22- and 12-lead ECG results and creatine kinase-MB analysis

Interventions: Patient care and existing protocols were unaltered, with the exception of including the new 22-lead ECG.

Measurements and main results: Forty-one of 163 patients had an AMI as defined by creatine kinase-MB analysis. The 22-lead ECG provided a statistically significant improvement in sensitivity (83%) for AMI diagnosis over the 12-lead ECG (51%) with specificities of 76% and 99%, respectively.

Conclusion: When combined with clinical judgment, the 22-lead ECG could provide a 97.6% sensitivity for AMI diagnosis while reducing unnecessary admissions for "rule-out MI" by 69%.

PMID: 1539875
Abstract
ST elevation myocardial infarction (STEMI) in the right ventricle (RV) associated with right coronary artery (RCA) occlusion is known to have high hospital mortality. The hypothesis tested in this study is: right precordial leads V4R and V5R help detect STEMI in the right ventricle. ECGs from 1,970 subjects were collected in Ruijin Hospital (n=1,342), Shanghai, China and Lund University Hospital, Lund (n=565), Sweden. All ECGs were recorded with additional leads on the right precordial location in V4R and V5R. Our results show that the subjects with middle to upper RCA occlusion often show ST elevation in leads V4R and V5R and ST depression in lateral leads I, aVL, V5-V6, and are often undetected as STEMI or AMI in the standard 12-lead ECG. We conclude that adding V4R and V5R to standard ECG recording in assessing patients presenting with acute coronary syndrome is an easy and convenient way to increases the sensitivity of STEMI detection.
Objectives: This study was done to determine whether electrocardiographic (ECG) isolated ST-segment elevation (ST) in posterior chest leads can establish the diagnosis of acute posterior infarction in patients with ischemic chest pain and to describe the clinical and echocardiographic characteristics of these patients.

Background: The absence of ST on the standard 12-lead ECG in many patients with acute posterior infarction hampers the early diagnosis of these infarcts and thus may result in inadequate triage and treatment. Although 4% of all acute myocardial infarction (AMI) patients reveal the presence of isolated ST in posterior chest leads, the significance of this finding has not yet been determined.

Methods: We studied 33 consecutive patients with ischemic chest pain suggestive of AMI without ST in the standard ECG who had isolated ST in posterior chest leads V7 through V9. All patients had echocardiographic imaging within 48 h of admission, and 20 patients underwent coronary angiography.

Results: Acute myocardial infarction was confirmed enzymatically in all patients and on discharge ECG pathologic Q-waves appeared in leads V7 through V9 in 75% of the patients. On echocardiography, posterior wall-motion abnormality was visible in 97% of the patients, and 69% had evidence of mitral regurgitation (MR), which was moderate or severe in one-third of the patients. Four patients (12%), all with significant MR, had heart failure, and one died from free-wall rupture. The circumflex coronary artery was the infarct related artery in all catheterized patients.

Conclusions: Isolated ST in leads V7 through V9 identify patients with acute posterior wall myocardial infarction. Early identification of those patients is important for adequate triage and treatment of patients with ischemic chest pain without ST on standard 12-lead ECG.

PMID: 10483956
Background: A routine 12-lead electrocardiogram is commonly obtained to evaluate for possible acute myocardial infarction during the initial screening of patients with chest discomfort. Posterior myocardial infarction is commonly missed because it is not usually visible in the standard leads. In this study, we compared the sensitivity and specificity of posterior chest leads (V(7), V(8), and V(9)) and 12-lead electrocardiography in detecting posterior injury pattern during single-vessel percutaneous transluminal coronary angioplasty.

Methods and Results: Three posterior chest leads in addition to the routine 12-lead electrocardiogram were monitored simultaneously during single-vessel percutaneous transluminal coronary angioplasty of the right, circumflex, and left anterior descending coronary arteries in a total of 223 patients. Posterior injury patterns (95%) were detected mostly during circumflex coronary occlusion. Posterior leads were able to detect injury pattern in 49% (36 of 74) of patients, whereas the 12-lead electrocardiogram was able to detect only 32% (P <.04). When all 15 leads were used to detect all ST elevations, sensitivity increased to 57%, with a specificity of 98% for the circumflex coronary artery. If maximal ST depressions in leads V(2) to V(3) are considered to be from posterior myocardial injury, then the overall sensitivity is increased to 69%.

Conclusions: Posterior leads significantly increased the detection of posterior injury pattern compared with the standard 12-lead electrocardiogram. Using all 15 leads significantly further improved the detection of circumflex coronary-related injury pattern over the standard 12-lead electrocardiogram.
33. Improved detection of Coronary Artery Disease by Exercise Electrocardiography with the Use of Right Precordial Leads.


Michaelides AP, Psomadaki ZD, Dilaveris PE, Richter DJ, Andrikopoulos GK, Aggeli KD, Stefanadis CI, Toutouzas PK

**Background:** Exercise electrocardiography is an imperfect test for the detection of coronary artery disease. We attempted to improve the diagnostic accuracy of exercise testing as a noninvasive method for the detection of coronary artery disease by using a combination of the left and right precordial leads.

**Methods:** We studied 245 patients (218 men and 27 women) ranging from 32 to 74 years of age (mean [±SD], 52±8) who underwent treadmill exercise testing, thallium-201 scintigraphy, and coronary arteriography. During exercise testing, each patient had one electrocardiogram recorded with the standard 12 leads and 3 right precordial leads (V3R, V4R, and V5R), with the results for each set of leads recorded and analyzed separately.

**Results:** On the basis of coronary arteriography, 34 patients had normal coronary arteries, 85 had single-vessel disease, 84 had two-vessel disease, and 42 had three-vessel disease. The sensitivities of the standard 12-lead exercise electrocardiogram, exercise electrocardiography incorporating right precordial leads, and thallium-201 scintigraphy were 52 percent, 89 percent, and 87 percent, respectively, for the detection of single-vessel disease; 71 percent, 94 percent, and 96 percent for the detection of two-vessel disease; 83 percent, 95 percent, and 98 percent for the detection of three-vessel disease; and 66 percent, 92 percent, and 93 percent for the detection of any coronary artery disease. The specificities of the three methods for the detection of any coronary artery disease were 88 percent, 88 percent, and 82 percent, respectively.

**Conclusions:** Use of right precordial leads along with the standard six left precordial leads during exercise electrocardiography greatly improves the sensitivity of exercise testing for the diagnosis of coronary artery disease.

PMID: 9929523
Abstract
Criteria for reperfusion therapy in acute myocardial infarction require the presence of ST elevation in 2 contiguous leads. However, many patients with myocardial infarction do not show these changes on a routine 12-lead electrocardiogram and hence are denied reperfusion therapy. Posterior chest leads (V7 to V9) were recorded in 58 patients with clinically suspected myocardial infarction, but nondiagnostic routine electrocardiogram. ST elevation >0.1 mV or Q waves in ≥2 posterior chest leads were considered to be diagnostic of posterior myocardial infarction. Eighteen patients had these changes of posterior myocardial infarction. All 18 patients were confirmed to have myocardial infarction by creatine phosphokinase criteria or cardiac catheterization. Of the 17 patients who had cardiac catheterization, 16 had left circumflex as the culprit vessel. We conclude that posterior chest leads should be routinely recorded in patients with suspected myocardial infarction and nondiagnostic, routine electrocardiogram. This simple bedside technique may help proper treatment of some of these patients now classified as having unstable angina or non-Q-wave myocardial infarction.

PMID: 10072216
35. Value of posterior and right ventricular leads in comparison to the standard 12-lead electrocardiogram in evaluation of ST-segment elevation in suspected acute myocardial infarction.


**Abstract**

In this multicenter prospective trial, we studied posterior (V7 to V9) and right ventricular (V4R to V6R) leads to assess their accuracy compared with standard 12-lead electrocardiograms (ECGs) for the diagnosis of acute myocardial infarction (AMI). Patients aged >34 years with suspected AMI received posterior and right ventricular leads immediately after the initial 12-lead ECG. ST elevation of 0.1 mV in 2 leads was blindly determined and inter-rater reliability estimated. AMI was diagnosed by World Health Organization criteria. The diagnostic value of nonstandard leads was determined when 12-lead ST elevation was absent and present and multivariate stepwise regression analysis was also performed. Of 533 study patients, 64.7% (345 of 533) had AMI and 24.8% received thrombolytic therapy. Posterior and right ventricular leads increased sensitivity for AMI by 8.4% (p = 0.03) but decreased specificity by 7.0% (p = 0.06). The likelihood ratios of a positive test for 12, 12 + posterior, and 12 + right ventricular ECGs were 6.4, 5.6, and 4.5, respectively. Increased AMI rates (positive predictive values) were found when ST elevation was present on 6 nonstandard leads (69.1%), on 12 leads only (88.4%), and on both 6 and 12 leads (96.8%; p <0.001). Treatment rates with thrombolytic therapy increased in parallel with this electrocardiographic gradient. Logistic regression analysis showed that 4 leads were independently predictive of AMI (p <0.001): leads I, II, V3, V5R; V9 approached statistical significance (p = 0.055). The standard ECG is not optimal for detecting ST-segment elevation in AMI, but its accuracy is only modestly improved by the addition of posterior and right ventricular leads.

PMID: 9202344