Circadian variation in witnessed out of hospital cardiac arrest

L H Soo, D Gray, T Young, J R Hampton

Abstract

Objectives—To examine the effect on circadian variation of out of hospital cardiac arrest according to the underlying aetiology and presenting rhythm of arrest, and to explore strategies that might help to improve survival outcome using circadian variation.

Design—Population based retrospective study.

Setting—County of Nottinghamshire with a total population of 993 914 and an area of 2183 km².

Subjects—Between 1 January 1991 and 3 December 1994, all witnessed cardiac arrests attended by the Nottinghamshire Ambulance Service, of which 1196 patients had a cardiac cause for their arrest (ICD, 9th revision, codes 390-414 and 420-429) and 339 had a non-cardiac cause.

Results—The circadian variation of the cardiac cases was not significantly different from that of non-cardiac cases (p = 0.587), even when adjusted for age, sex, or presenting rhythm of arrest. For cardiac cases, the circadian variation of those who presented with ventricular fibrillation was significantly different from those presenting with a rhythm other than ventricular fibrillation (p = 0.005), but was similar to the circadian variation of bystander cardiopulmonary resuscitation (p = 0.306) and survivors (p = 0.542). Ambulance response time was also found to have a circadian variation.

Conclusions—There is a common circadian variation of out of hospital cardiac arrest, irrespective of underlying aetiology, where the presenting rhythm is other than ventricular fibrillation. This is different from the circadian variation of cases of cardiac aetiology presenting with ventricular fibrillation. The circadian variation of ventricular fibrillation, and consequently survival, may be affected by the availability of bystander cardiopulmonary resuscitation and the speed of ambulance response.

(Heart 2000;84:370–376)

Keywords: out of hospital; cardiac arrest; circadian variation

A wide range of cardiovascular disease has been found to have circadian patterns, but the reason for this remains unclear. Circadian variation in hypertension, sudden death, ventricular tachyarrhythmias, myocardial infarction, unstable angina, acute cerebrovascular disease, stroke, subarachnoid haemorrhage, and ruptured aortic aneurysm seem to follow a circadian pattern. These events occur with greater frequency in the morning or early afternoon. In some of these studies, however, additional peaks have been reported in the late afternoon and early evening.

Cardiac arrest may have either a cardiac or a non-cardiac aetiology and may present with a variety of arrhythmias, including ventricular tachycardia, ventricular fibrillation, asystole, and electromechanical dissociation. Where prompt basic life support is available in the community, survival chances are greatest. With the exception of a few places in the USA, survival rates for those who reach hospital alive have been disappointing low. We wondered whether there was any evidence of circadian variation in cardiac arrest occurring in our health district. Knowledge of this might help us to develop strategies aimed at improving the appalling chances of survival from out of hospital cardiac arrest.

Methods

We conducted a retrospective study of all incidents of cardiac arrest in adults (defined as over 18 years of age) attended by the Nottinghamshire ambulance service from January 1991 to December 1994. Only witnessed events, where the patient was either observed or heard to collapse, were included. Unwitnessed cases were excluded because of the uncertainty of the timing of arrest.

Data were collated from the Nottinghamshire ambulance patient report forms, records of the ambulance dispatch and control unit, and all Nottinghamshire accident and emergency departments, coronary care units, and intensive care units. The presenting rhythm for each episode of cardiac arrest was obtained from the printout of the defibrillator unit; the patient report form was reviewed where the presenting rhythm was not recorded but an attempt at resuscitation was made. The coroners' records and inpatient case notes were also examined to identify all those sustaining a cardiac arrest from a “cardiac” cause (International classification of diseases, ninth revision (ICD-9), codes 390 to 414 and 420 to 429), usually determined from a necropsy examination but otherwise from death certificate records. Deaths attributed to a non-cardiac cause included drug overdose, suicide, drowning, hypoxia, exsanguination, cerebrovascular accident, subarachnoid haemorrhage, trauma,
ruptured aortic aneurysm, and pulmonary thromboembolism.

STATISTICAL METHODS
Two techniques were used to establish circadian variation over a 24 hour period from midnight to midnight. It was first necessary to establish whether any circadian variation existed. All cardiac arrest events were recorded against a time scale. The frequency–time curve was scrutinised using spectral analysis, a technique that describes the data in cosinusoidal terms and tests for the presence of higher frequency components. Harmonic regression analysis was then used to determine the number of significant cosinusoidal components needed to adequately fit the data. The cosine wave variation over a 24 hour period was measured by the shift in nadir from 12 am (midnight), known as the phase, and by the maximum displacement during one cycle, defined as the amplitude. The degree of phase shift and the difference in amplitude from a flat line (representing no circadian variation) determine the significance of the component. We fitted models of the form:

\[ x_t = \mu_t + RC_\cos(2\pi t/P + \phi) + \epsilon_t, \]

where \( \mu_t \) is an added constant, \( R \) is the amplitude of the cosine wave, \( P \) is the period effect (for example, 24 h, 12 h, 8 h, 6 h), \( \phi \) is the phase shift, and \( \epsilon_t \) is the residual (the value needed to make the equality exact). A stepwise regression analysis was used to determine whether each cosinusoidal component was significantly different from zero.

The \( \chi^2 \) test was used to examine discrete variables in a tabulated time series of six 4 hourly intervals to determine whether there was any variation within a group or between groups. The Kruskal-Wallis test was used to make comparisons within a group of continuous variables in a tabulated time series of six 4 hourly intervals. A probability value of \( p < 0.05 \) was considered significant in all models. The S-Plus statistical package was used for all analyses.

Results
Nottinghamshire ambulance service has 12 ambulance stations that serve a population of one million. Each ambulance station normally provides one accident and emergency vehicle with a crew of two, usually a paramedic and a medical technician, around the clock. Both are trained to defibrillate, although a paramedic has completed additional training and skills. The shift duty for the crew of this vehicle is from 0600 to 1400, 1400 to 2200, and 2200 to 0600 hours. Most ambulance stations provide a second accident and emergency vehicle that is operational from 0800 to 1600 hours. A few larger ambulance stations serving urban city areas provide a third vehicle operating between 1000 and 1800 hours. In exceptional circumstances, a fourth vehicle can be mobilised.

STUDY POPULATION
From January 1991 to December 1994, the Nottinghamshire ambulance service attended 2362 out of hospital cardiac arrest cases; we excluded 748 cases, as a bystander did not witness the event. The mean (SD) age of this group was 63.2 (18.9), 502 (67.1%) were male, 246 (32.9%) were female, and 599 (80.1%) arrested at home. A further 79 had incomplete records; of these, only seven were admitted to a hospital ward and all died while in hospital. The remaining 1535 cardiac arrest events form the basis of this study.

The mean (SD) age of these 1535 cases was 65.6 (14.5) years, with 917 (59.7%) of patients being aged less than 70 years; 1126 (73.4%) were male, 409 (26.6%) were female, and 1046 (68.1%) arrested at home.
AETIOLOGY OF THE CARDIAC ARREST

Resuscitation was attempted by the ambulance crew in 1440 cardiac arrests (93.8%). In the remaining 95 cases (6.2%), death was attributed to a cardiac cause in 54 (3.5%); 43 (79.6%) after necropsy.

In 1142 cases (74.4%) where the ambulance crew attempted resuscitation, the arrest was considered to have a cardiac aetiology, falling to 827 (72.4%) after necropsy. The distribution of 339 arrests of non-cardiac aetiology (234 (69%) after necropsy), including those where there was no resuscitation attempt by the ambulance crew, is shown in fig 1.

CIRCADIAN VARIATION IN CARDIAC ARREST FROM ALL CAUSES

Figure 2 shows the number of witnessed arrests from all causes in the community over a 24 hour period and the circadian variation: 24 h, 12 h, and 6 h circadian effects were observed (p < 0.001). The number of witnessed arrests in the community was lowest between midnight and 0600; there was a sharp increase in the number of arrests occurring after this time, with peaks observed at approximately midday, 1600, and 2200 hours ($R^2 = 94.3\%$, $F$ value = 46.5, $p < 0.001$).

CIRCADIAN VARIATION OF CARDIAC ARREST OF CARDIAC AND NON-CARDIAC AETIOLOGY

Figure 3 compares the circadian variation in cardiac arrest of cardiac and non-cardiac aetiology. The fitted curve for cardiac cases shows peaks occurring at 1100, 1600, and 2200 hours. These are similar to the distribution of cardiac arrest from all causes, probably reflecting the number of cases of cardiac aetiology. Even so, a greater number of cardiac cases occurred later in the day at 1600 rather than at 1100 hours ($R^2 = 94.0\%$, $F = 44.5$, $p < 0.001$).

There were relatively fewer arrests from non-cardiac causes, which may account for the smaller degree of variation. Even so, there were fewer events between midnight and 0600 hours, events increased during the day, and the number of arrests was lowest between midnight and 0600 hours ($R^2 = 66.0\%$, $F = 9.2$, $p < 0.001$) but the peak occurred later in the day at 1600 rather than at 1100 hours ($R^2 = 94.5\%$, $F = 41.5$, $p < 0.001$).

The 24 hour distribution of cardiac and non-cardiac cases of cardiac arrest was divided into six 4 hourly intervals (table 1) and no difference was found between the distributions of the two groups.

EFFECT OF AGE AND SEX

Age and sex did not appear to have any impact on the distribution of cardiac and non-cardiac cases (table 1).

PRESENTING RHYTHM AND OUTCOME

Six hundred and forty three patients (41.9%) were in ventricular fibrillation when the ambulance crew arrived. One hundred and sixty eight (10.9%) died at home and so were not transported to hospital, while 767 (50.0%) were taken by ambulance to an accident and treatment unit. Table 1 shows the number of witnessed arrests from all causes in the community over a 24 hour period and the circadian variation: 24 h, 12 h, and 6 h circadian effects were observed (p < 0.001). The number of witnessed arrests in the community was lowest between midnight and 0600; there was a sharp increase in the number of arrests occurring after this time, with peaks observed at approximately midday, 1600, and 2200 hours ($R^2 = 94.3\%$, $F$ value = 46.5, $p < 0.001$).

CIRCADIAN VARIATION OF CARDIAC ARREST FROM ALL CAUSES

Table 1 Effects of sex, age, and rhythm on the distribution of cardiac arrest cases of cardiac and non-cardiac aetiology by 4 hourly intervals

<table>
<thead>
<tr>
<th>Time intervals (24 h clock)</th>
<th>0000 to 0359</th>
<th>0400 to 0759</th>
<th>0800 to 1159</th>
<th>1200 to 1559</th>
<th>1600 to 1959</th>
<th>2000 to 2359</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>All cases (cardiac cases (%) / non-cardiac cases (%))</td>
<td>99 (8.3) / 28 (8.3)</td>
<td>93 (7.8) / 35 (10.3)</td>
<td>264 (22.1) / 72 (21.2)</td>
<td>285 (23.8) / 69 (20.4)</td>
<td>247 (20.7) / 73 (21.5)</td>
<td>208 (17.4) / 62 (18.2)</td>
<td>0.587</td>
</tr>
<tr>
<td>Male (cardiac cases (%) / non-cardiac cases (%))</td>
<td>80 (8.8) / 16 (7.5)</td>
<td>75 (8.2) / 22 (10.4)</td>
<td>189 (20.7) / 42 (19.8)</td>
<td>221 (24.2) / 45 (21.2)</td>
<td>191 (20.9) / 47 (22.2)</td>
<td>158 (17.3) / 40 (18.9)</td>
<td>0.805</td>
</tr>
<tr>
<td>Female (cardiac cases (%) / non-cardiac cases (%))</td>
<td>19 (6.7) / 12 (9.4)</td>
<td>18 (6.4) / 13 (10.2)</td>
<td>75 (26.6) / 30 (23.6)</td>
<td>64 (22.7) / 24 (18.9)</td>
<td>56 (19.9) / 26 (20.5)</td>
<td>50 (17.7) / 31 (22.3)</td>
<td>0.627</td>
</tr>
<tr>
<td>Aged &lt; 70 years (cardiac cases (%) / non-cardiac cases (%))</td>
<td>71 (9.9) / 17 (8.6)</td>
<td>63 (8.6) / 22 (11.1)</td>
<td>144 (20.0) / 47 (23.7)</td>
<td>150 (20.9) / 30 (15.2)</td>
<td>145 (20.2) / 40 (20.2)</td>
<td>146 (20.3) / 42 (21.2)</td>
<td>0.433</td>
</tr>
<tr>
<td>Aged &gt; 70 years (cardiac cases (%) / non-cardiac cases (%))</td>
<td>28 (5.9) / 11 (7.8)</td>
<td>30 (6.3) / 13 (9.2)</td>
<td>120 (25.2) / 25 (17.7)</td>
<td>135 (28.3) / 39 (27.7)</td>
<td>102 (21.4) / 33 (23.4)</td>
<td>62 (13.0) / 20 (14.2)</td>
<td>0.431</td>
</tr>
<tr>
<td>Presenting rhythm: other than VF (cardiac cases (%) / non-cardiac cases (%))</td>
<td>54 (10.1) / 16 (9.6)</td>
<td>53 (9.9) / 29 (10.7)</td>
<td>111 (20.8) / 56 (20.7)</td>
<td>103 (19.3) / 46 (17.0)</td>
<td>117 (21.9) / 62 (23.0)</td>
<td>46 (18.0) / 51 (18.9)</td>
<td>0.976</td>
</tr>
<tr>
<td>Not survived to hospital admission (cardiac cases (%) / non-cardiac cases (%))</td>
<td>82 (8.2) / 27 (8.7)</td>
<td>75 (7.5) / 34 (11.0)</td>
<td>212 (21.2) / 68 (21.9)</td>
<td>237 (23.7) / 61 (19.7)</td>
<td>213 (21.3) / 63 (20.3)</td>
<td>183 (18.3) / 57 (18.4)</td>
<td>0.375</td>
</tr>
</tbody>
</table>

All percentages are proportion of the total cases occurring over 24 hours.

VF, ventricular fibrillation.

---

**Table 1** Effects of sex, age, and rhythm on the distribution of cardiac arrest cases of cardiac and non-cardiac aetiology by 4 hourly intervals
Circadian variation in out of hospital cardiac arrest

For those presenting with ventricular fibrillation of cardiac aetiology, peaks in circadian variation occurred around 1100, 1500, and 2200 hours ($R^2 = 93.6\%$, $F = 41.2$, $p < 0.001$), with the highest peak being at 1500 hours (fig 4).

Harmonic analysis was not performed for non-cardiac cases presenting with ventricular fibrillation because the number of cases was small. The circadian variation for rhythms other than ventricular fibrillation was identical in both cardiac ($R^2 = 69.6\%$, $F = 10.8$, $p < 0.001$) and non-cardiac cases ($R^2 = 64.2\%$, $F = 8.5$, $p < 0.001$), with peaks occurring around 1130 to 1230 hours and around 2030 hours. This was confirmed when their distribution was compared by six 4 hourly intervals over a 24 hour period ($p = 0.967$) (table 1).

For cases of cardiac aetiology, we compared the distribution of ventricular fibrillation cases with other rhythms by dividing the 24 hour period into six 4 hourly intervals, as shown in table 2, and a significant difference was found between the two groups ($p = 0.005$). There was an expectedly low number of incidents of bystander resuscitation after midnight, which subsequently increased and reached a plateau after 0800 hours. The ambulance crew response times appear to make a small but significant difference, with slightly slower response times from midnight to 0359 hours and from 1600 to 1959 hours. The slower response times might be because there were fewer vehicles during these periods. The combination of infrequent bystander resuscitation and slower ambulance response times could account for the troughs seen in the circadian variation and presenting rhythm.

Table 2  The variation of cardiac cases over 24 hours of the ambulance crews response time intervals, bystander cardiopulmonary resuscitation, presenting rhythm, and non-survivors to hospital

<table>
<thead>
<tr>
<th>Time intervals (24 h clock)</th>
<th>0000 to 0359</th>
<th>0400 to 0759</th>
<th>0800 to 1159</th>
<th>1200 to 1559</th>
<th>1600 to 1959</th>
<th>2000 to 2359</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Response times by ambulance crew, median time (min) (interquartile range)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>VF (%)</td>
<td>7 (5 to 9)</td>
<td>6 (5 to 8)</td>
<td>6 (4 to 9)</td>
<td>7 (5 to 9)</td>
<td>6 (4 to 9)</td>
<td>7 (5 to 9)</td>
<td>0.022</td>
</tr>
<tr>
<td>Non-VF (%)</td>
<td>44 (44.4)</td>
<td>36 (38.7)</td>
<td>141 (53.4)</td>
<td>161 (56.5)</td>
<td>118 (47.8)</td>
<td>105 (50.5)</td>
<td>0.030</td>
</tr>
<tr>
<td>Unknown (%)</td>
<td>54 (54.5)</td>
<td>53 (57.0)</td>
<td>111 (42.0)</td>
<td>103 (36.1)</td>
<td>117 (47.4)</td>
<td>96 (46.2)</td>
<td>0.001</td>
</tr>
<tr>
<td>Bystander CPR (%)</td>
<td>0 (0)</td>
<td>4 (4.3)</td>
<td>12 (4.5)</td>
<td>21 (7.4)</td>
<td>12 (4.9)</td>
<td>7 (3.4)</td>
<td>NA</td>
</tr>
<tr>
<td>Survival to hospital admission (%)</td>
<td>17 (17.2)</td>
<td>18 (19.4)</td>
<td>52 (19.7)</td>
<td>48 (16.8)</td>
<td>34 (13.8)</td>
<td>25 (12.0)</td>
<td>0.023</td>
</tr>
<tr>
<td>Survival to hospital discharge (%)</td>
<td>10 (10.1)</td>
<td>8 (8.6)</td>
<td>21 (8.0)</td>
<td>18 (6.3)</td>
<td>16 (6.5)</td>
<td>15 (7.2)</td>
<td>0.823</td>
</tr>
</tbody>
</table>

All percentages are proportion of total cases occurring within each 4 hour period. *Compared with distribution of VF rhythm of arrest cases. CPR, cardiopulmonary resuscitation; NA, not assessed; VF, ventricular fibrillation.
variation of ventricular fibrillation between midnight and 0800 hours and around 1900 hours.

Survival to both hospital admission ($R^2 = 62.8\%$, $F = 17.8$, $p < 0.001$) and to hospital discharge ($R^2 = 25.4\%$, $F = 25.2$, $p = 0.046$) for cases of cardiac aetiology were also found to show a circadian variation, with a single event peak occurring around 1300 hours (fig 5). However, the fitted harmonic curves had low $R^2$ value owing to the small number of events.

The distribution of those who were alive on reaching hospital and were subsequently discharged was examined concurrently with cases presenting with ventricular fibrillation rhythm (table 2); no difference between the circadian variation of the two groups was observed.

Discussion
The likelihood of a “sudden death” episode occurring and being witnessed is not constant throughout the day. We observed that there were three major time periods when a cardiac arrest was most likely to occur—an early peak in the morning around 1100 hours, a late peak in the evening around 2200 hours, and, for those whose arrest was from a cardiac disease, a third time period—around 1600 hours—also seemed to be critical.

There does not seem to be any consistent pattern to circadian variation in published reports. In Seattle and Berlin, the most important times for a cardiac arrest seemed to be around 1000 and 1800 hours. Ventricular fibrillation was most frequent around 2000 hours in the series by Peckova and colleagues, while Arntz and associates found that ventricular fibrillation occurred most often around 1000 hours, very similar to our own findings. Differences such as these probably arise for several reasons: methodological difficulties, not least in defining what is meant by “sudden death”; timing the onset of the acute event with any degree of accuracy; reliance on accurate data collection of ambulance staff rather than direct contact which increases the proportion of arrests designated as “witnessed ventricular fibrillation” ($p = 0.046$); accounting for events that are unwitnessed; excluding those who survive the acute event; and failing to establish from death certificate data or necropsy reports the underlying aetiology. We tried to address these potential problems by clearly defining sudden death, recording the response time to a call for out of hospital resuscitation by our ambulance service, which is continually monitored, comparing witnessed arrest of different aetiology to reduce the bias from unwitnessed events late at night, auditing the outcome of all patients in the study, and obtaining information on the cause of death, using necropsy data where available.

We are confident that our omission of unwitnessed events had little impact on our findings because the significance and pattern of the circadian variation remained unchanged even when all unwitnessed events are assumed to have occurred during late at night and early in the morning; and there is a low frequency (12.3%) of sudden cardiac death during sleep.

We did find that the periods when ventricular fibrillation were most likely to occur coincided with periods of faster response time by the ambulance crews, leading to early cardiac monitoring, as well as periods in which bystander resuscitation was more likely to be initiated, maintaining cardiac output until the ambulance crew arrived. These factors potentially affected the circadian variation of arrhythmia, particularly ventricular fibrillation, and could also account for the diverse findings reported so far.

We are unaware of any previous study that has examined circadian variation of out of hospital arrest of non-cardiac aetiology. This group of patients is not homogeneous—one third had some vascular disorder such as ruptured or dissected aortic aneurysms, subarachnoid haemorrhage, and cerebrovascular accidents. Such heterogeneity might be expected to lead to events being randomly distributed throughout the day and night, though others have reported circadian variation for some of the conditions in our series.

We were surprised to find that, with the exception of ventricular fibrillation, the circadian variation of the arrest rhythm in both cardiac and non-cardiac groups appeared to be identical. The triggers for a cardiac arrest in both groups are likely to be similar when the arrhythmia is something other than ventricular fibrillation, irrespective of the aetiology of arrest. Vigorous physical activity is one potential trigger, but this is disputed by others, including studies based on pathology and Holter monitoring. Changes in blood pressure, vascular tone, platelet activation, and hormonal concentrations of adrenaline (epinephrine), noradrenaline (norepinephrine), renin, and cortisol are examples of some physiological processes caused by a triggering activity.

The chances of a Nottingham resident surviving an out of hospital cardiac arrest are generally very poor. Can we use knowledge about the circadian variation of out of hospital cardiac arrest to improve survival rates? The chances of survival are generally dismal when the presenting rhythm is asystole or electromechanical dissociation. This presumably reflects varying but catastrophic aetiology, so it is probably futile to expect to influence the outcome in this group. However, it might be more rewarding to identify trigger factors for this type of rhythm of arrest from future studies on circadian variation and to attempt modification of these factors as part of preventative measures.

The chances of surviving an out of hospital cardiac arrest are best when the arrhythmia is ventricular fibrillation. It is interesting that the circadian variation we observed among those who survived an out of hospital cardiac arrest matches the circadian pattern of ventricular fibrillation in Seattle, where the chances of surviving and being discharged alive are about
Circadian variation in out of hospital cardiac arrest

12.4%,\textsuperscript{19} although survival in Nottingham was only half that.

Our lower survival rate is most likely to reflect two factors. First, in very few cases (about one third of all those whose arrest is witnessed) is resuscitation started before the arrival of the ambulance crew. Providing training in basic life support for a large proportion of the community is one approach,\textsuperscript{15,\textsuperscript{34}} but it may be more cost–effective to focus attention on those with at-risk partners. Second, as we have previously reported,\textsuperscript{19} the response times of our ambulances were probably too long, at six minutes (median), to influence the outcome. Response times in Rochester\textsuperscript{26} and Seattle,\textsuperscript{41} 2.7 minutes and 3.6 minutes, have produced far superior survival rates. Knowing when cardiac arrest occurrence is most likely could facilitate the deployment of ambulance crew or suitably trained health professionals at peak times (which for ventricular fibrillation in Nottingham is from 1000 to 2200 hours). We might expect to improve the survival chances for 196 cases a year (66% of all ventricular fibrillation cases) presenting in ventricular fibrillation.

Strategies to improve survival chances following cardiac arrest from ventricular fibrillation continue to present a challenge. There are several alternatives. Effective preventive measures could be developed if we had a better understanding of the pathophysiological triggers to cardiac arrhythmia and an explanation for its predilection for certain times of the day. There are proposals to provide defibrillators in the community but these will benefit only those whose collapse occurs in a public place—about one quarter of our series—and in the presence of a reliably trained individual.\textsuperscript{30} When suitably trained and equipped ambulance personnel are at hand during an arrest, survival chances are greatly improved to about 40–57%.\textsuperscript{11,\textsuperscript{15,\textsuperscript{34}}}

While many episodes of ventricular fibrillation will continue to occur in the absence of trained personnel, ambulance controllers might benefit their local population by incorporating information on circadian variation into their computer programs advising on ambulance deployment.

Colour coded blood flow imaging in intravascular ultrasound

A 42 year old man was admitted to our hospital with sudden chest pain. The exercise ECG showed inferior ST segment depression suggesting ischaemia. Coronary angiography revealed a normal left coronary artery and severe proximal stenosis of the right coronary artery. The patient underwent direct percutaneous transluminal coronary angioplasty with coronary stent-graft implantation. The stent-graft is a combination of a metal stent with a membrane of polytetrafluorethylene. Angiography suggested good stent apposition without residual stenosis. With standard intravascular ultrasound (IVUS) full stent expansion was assessed (A, large arrow: stent-graft; small arrow: lumen), revealing a weak signal distal of the stent caused by partial absorption of the ultrasound signal from the specific stent graft material. In contrast to angiography and standard IVUS, colour coded blood flow imaging revealed incomplete apposition in the proximal entrance of the stent graft in more than 50% of the circumference documented by blood flow (red) in the two dimensional image (B, arrows) plane as well as in the sagittal reconstruction of the vessel (C, arrows).

Colour coded blood flow imaging is a recently introduced and commercially available technique encoding the rate of change of the backscatter echo of the blood cells into colour. Since the ultrasound beam is identical to the one used in standard IVUS, the colour information is displayed simultaneously with the standard IVUS information. Flow between the stent and the vessel wall was documented only with colour coding, representing incomplete attachment of the stent to the vessel wall (B and C, arrows). Colour coded blood flow imaging is a useful adjunct to standard greyscale IVUS in specific lesions.

DIETMAR H KOSCHYK
CHRISTIAN W HAMM
THOMAS MEINERTZ