In this paper we analyse a low-cost commercial chest belt to be integrated into a lifestyle counselling system as a source of heart rate data. We compared data from a Schiller ECG Holter device, which serves as a reference to a CardioSport device. Due to missing data in the CardioSport device caused by loss of contact with the body, the creation of special algorithms was necessary for synchronization and data validation. The results show that when using our synchronization algorithms the average absolute percentage error between the two signals was 2% with correlation of more than 99%. Using a data validation algorithm, we were able to get on average more than 70% of the signal with an absolute percentage error of 3% and a high average correlation of 99%. The mean RR interval values and standard deviation of RR intervals are very close to those of the reference device using both the synchronization and data validation algorithms. When using the data validation algorithm, the reference measurements produced only slightly better results with regard to false detections of atrial defibrillation than the CardioSport device. In conclusion, we found that with a simple preprocessing algorithm, CardioSport as a low-cost device can be safely integrated into a lifestyle support system as a telemedical solution.

**Keywords:** telemedicine, lifestyle counselling, heart rate monitor

### Introduction

Low-cost telemedical sensors are often used in modern ambient assisted living (AAL) telemonitoring and self-management systems for providing inputs to medical intelligence algorithms [1]. Such systems extend the scope of traditional health care that is based purely on data measurement. However, the proper interpretation and reliability of the results depends on the reliability of the measured data and the sensor itself. Nevertheless, there are still surprisingly few reviews reported in the literature to date on the validation of the information content of such low-cost sensors compared to the clinically accepted reference device. An example of a device that was tested for validity is the SenseWear HR Armband [2]. In this study, they used the reference device simultaneously with the tested device as a way of validating data. However, most of the compared devices are expensive high-end devices, which present an obstacle for their wide use in telemedicine.

In this proof-of-concept paper, we analyse a simple commercial chest belt chosen to be integrated into the Lavinia lifestyle mirror system [3] as a source of heart rate (HR) data. In the Lavinia system, the HR signal of the patient will be used to (i) estimate the calories burnt by physical activity, (ii) calculate the heart rate variability (HRV) in order to detect periods of mental or emotional stress, and (iii) analyse arrhythmia patterns (Poincare plots) for atrial fibrillation detection. Our approach involves the comparison of the HRV and Poincare plots computed from the filtered chest belt signal, with those parameters computed from a reference Holter device.

### Methods

#### Measurements

Two devices were used simultaneously by a healthy volunteer over a 24 hour period. A Schiller MT-101/MT-200 Holter device was our reference device designed for clinical use. The chest belt was a CardioSport TP3 Heart Rate Transmitter device. Since this device does not have its own memory for storing data, we used a Nexus 7 tablet with Android version 4.4.2 to connect the device via the Bluetooth 4.0 protocol and store the measured data on the tablet. Although both devices were worn by volunteers for 24 hours, only 12 hours of the overall signal were used for analysis due to frequent detachments of the device from the body during nighttime. The measurements of 12 hours were repeated on 4 additional healthy male subjects.

#### Signal Analysis

The direct comparison of measured data was not possible due to the different designs of the reference and the telemedical devices. However, we wanted to
compare signals directly in terms of time and also to
develop a data validation algorithm for removing the
noisy parts of the CardioSport device measurements
reliably without using the reference data. The problem
was that the chest belt was not firmly attached to the
body and sudden movements of the device caused signal
loss. Therefore, we needed to create a software module
for synchronization and data validation before any
analysis. Data validation means removing obviously bad
data (artefacts) and keeping only “good” data segments
of sufficient length, because, as a rule of thumb, both
HRV and Poincare plot computations require data
chunks of at least 5 minutes. Even though the data
validation algorithm removed a considerable amount of
data from the original signal, we still had enough useful
data for analysis from the daytime.

The Synchronization Algorithm

Our simple algorithm for signal synchronization uses a
sliding window that passes from the beginning of the
chest belt signal to the end and calculates the absolute
error between the two signals. When sliding finishes,
the location of the sliding window with the minimum
absolute error is considered as the point where the two
signals should be synchronized. This applies only if the
correlation of the data in the sliding window and the
same amount of data from the reference device are
higher than a minimum set by the user. If these
conditions are met, the algorithm copies data from the
sliding window into a newly generated third signal,
which represents the chest belt signal fully
synchronized with the reference signal. If conditions are not met, the
third signal is filled with zeros. Finally, the algorithm
extracts all the highly correlated segments from the third
signal ignoring zero values. Also, a file with all the
merged segments is generated for general analysis. The
algorithm uses the following 5 main parameters that can
be set up by the user:

1. window size: amount of data copied from the signal
   into the sliding window (default: 200),
2. window shift step: the number of samples by which
   we shift the sliding window in each iteration
   (default: 50),
3. absolute error window: amount of data used for
   calculating the minimum absolute error (default: 200),
4. maximum error distance: the number of samples by
   which we shift the absolute error window in order to
   find the minimum absolute error (default: 1000),
5. minimum correlation: minimum correlation,
   expressed as a percentage, required for the two
   signals to consider data in the chest belt signal as
   accurate (default: 97%).

Each parameter’s default value was determined
empirically. After running the synchronization process,
we obtained segments of highly correlated data. Fig.1
shows the distribution of the lengths of signal segments.
We can see that most segments are 3 to 18 minutes long.
The longest highly correlated segment with the
reference data is 110 minutes long. The default
parameter settings minimize the number of overly short
(< 5 min) segments. Most of the bad segments (Fig.2)
are shorter than one minute, and only one bad segment
was 60 minutes long.

Data Validation Algorithm

Another type of algorithm was used in the real
telemedical scenario for finding good parts of the signal
without relying on reference data. This implies finding
gaps and abnormal values and omitting them. First, we
compared the timestamp of each data point with the
timestamp of the previous one. If the difference between
the timestamps was longer than 3 seconds, we marked
this as a ‘gap’. The 3-second gap detection was enabled
by the chest belt’s buffering system that can tolerate
short detachments of the device from the body. In the
second step we identified abnormal values in the signal
that were treated as gaps. The abnormal values are
identified by observing the mean value of 20
neighbouring data points (10 before and 10 after a given
point). If the mean value differs from the value of the
current sample by more than 300 units, we consider it invalid and mark it as a gap/error in the signal. Finally, we extract the good segments from the signal with a length of more than 5 minutes.

We implemented the above algorithm in a simple software tool (Fig. 3). On the left-hand side we can load the two signals and set the parameter values as well as the amount of data to be analysed. The graph shows two signals after the synchronization process was completed. The user can examine signals by clicking the Previous and Next buttons. General statistics are shown in the middle part of the screen, while in the lower part, we can see the histogram, and save the histogram and results as a file. Two tabs in the top left-hand corner allow the user to switch between synchronization and data validation algorithms.

**Statistical Analysis**

Time and frequency domain analyses, correlation comparisons, mean absolute percentage errors, and the slopes of scatter plot diagrams were compared between two measurements for HRV analysis. The specificities of a self-developed atrial fibrillation detector algorithm were compared for atrial fibrillation analysis. The latter algorithm is based on the k-means clustering of Poincaré plots (consisting of RR intervals).

The time and frequency domain analyses for HRV were performed using Kubios HRV analysis software, while the rest of the analysis for HRV and atrial fibrillation was performed in Microsoft Excel. Atrial fibrillation detection was done using the MATLAB environment and the results were saved as Microsoft Excel workbooks.

**Results and Analysis**

**Heart Rate Variability**

After the synchronization process, we got strongly correlated (greater than 97%) synchronized data segments of various durations. Table 1 summarizes the duration of signals analysed.

<table>
<thead>
<tr>
<th>Subject</th>
<th>Duration (h:m:s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1</td>
<td>10:53:28</td>
</tr>
<tr>
<td>#2</td>
<td>8:45:40</td>
</tr>
<tr>
<td>#3</td>
<td>10:30:17</td>
</tr>
<tr>
<td>#4</td>
<td>7:46:56</td>
</tr>
</tbody>
</table>

Table 2 shows results in the time domain for Schiller and CardioSport devices after using the algorithm for the synchronization of signals. Time domain analysis shows similar values for mean RR values and standard deviation (STD RR in Eq. (1)). The average Mean RR values for the Schiller and CardioSport devices are 851 and 871 respectively. The average STD RR for the Schiller device is 108 and 110 for the CardioSport device.
The frequency domain analysis for the synchronization process is presented in Table 3. The absolute power was compared for very low frequencies (VLF: 0.00-0.04 Hz), low frequencies (LF: 0.04-0.15 Hz), high frequencies (HF: 0.15-0.4 Hz) and ratios between low frequencies and high frequencies (LF/HF). Results show no significant difference between Schiller and CardioSport device values. The average mean absolute percentage error (MAPE) between two signals is 2% with a high average correlation of close to 100%.

Using the data validation algorithm, we extracted data points from the collected signals. The duration of the resulting signal is shown in Table 4. It is important to note that due to the noise on Schiller device recordings, we had to remove noisy parts from the original signal. Therefore, even though the signal was recorded continuously for 12 hours, overall duration is much less. Calculations show that in the worst scenario only 45% of the signal can be used for analysis using this data validation method, while in the best scenario this number reaches 95%. This leads to a conclusion that results are rather subject dependent.

The results of data analysis in the time domain after the removal of bad parts using the validation algorithm can be seen in Table 5. The mean RR intervals for Schiller and CardioSport devices are 851 and 871 and standard deviations are 104 and 106, respectively. The CardioSport device has slightly greater values, but these are practically identical.

The frequency domain analysis for the data validation process is presented in Table 6. The absolute power was compared for very low frequencies (VLF: 0.00-0.04 Hz), low frequencies (LF: 0.04-0.15 Hz), high frequencies (HF: 0.15-0.4 Hz) and ratios between low frequencies and high frequencies (LF/HF). As in the synchronization process, the results show no significant difference between the Schiller and CardioSport device values.

The minimum, maximum and average percentage errors on whole signals were calculated using 5 minute long sliding windows with one minute long shift steps (Table 7). Only one subject had a high maximum error value of 34%. By visual examination, it was determined that the cause of such a high error was the artefact of the Schiller device. In spite of that, the average error remained low (2%).
can be seen in our previous study [16].

otherwise to 

determine the 
plot 

We considered 30 RR intervals per iteration by analy-

We carried out the detection of atrial fibrillation (AFib) by analysing POINCARÉ plots consisting of 30 RR intervals. We considered 30 RR intervals per iteration and in each iteration after constructing the POINCARÉ plot we calculated the dispersion around the diagonal line and used k-means based cluster analysis to determine the number of the clusters. If the dispersion was too high (greater than 0.06) and the number of clusters was 1, or the number of clusters was more than 9; we assigned “AFib” to that series of RR intervals, otherwise to “Non-AFib”. The details of the algorithm can be seen in our previous study [16]. Since our data set did not contain real AFib cases, only specificity could be calculated with regard to the efficiency of detection. The evaluation of atrial fibrillation detection results for synchronized data validation can be seen in Tables 8 and 9.


Table 7: The minimum, maximum and average percentage errors

<table>
<thead>
<tr>
<th>Subject</th>
<th>Minimum error</th>
<th>Maximum error</th>
<th>Average error</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1</td>
<td>0.1%</td>
<td>3.5%</td>
<td>1.5%</td>
</tr>
<tr>
<td>#2</td>
<td>0.0%</td>
<td>7.7%</td>
<td>2.1%</td>
</tr>
<tr>
<td>#3</td>
<td>0.0%</td>
<td>33.9%</td>
<td>3.2%</td>
</tr>
<tr>
<td>#4</td>
<td>0.1%</td>
<td>6.7%</td>
<td>1.9%</td>
</tr>
<tr>
<td>#5</td>
<td>0.1%</td>
<td>5.1%</td>
<td>2.2%</td>
</tr>
<tr>
<td>Average</td>
<td>0.1%</td>
<td>13.4%</td>
<td>2.4%</td>
</tr>
</tbody>
</table>

Fig. 4 represents a typical relationship between CardioSport and Schiller devices. All gradient values are close to 1. The lowest slope value is 0.98 while the highest value is 1.02. The average mean absolute percentage error (MAPE) between two signals was 3% with a strong average correlation of 99%.

Atrial Fibrillation

We carried out the detection of atrial fibrillation (AFib) by analysing POINCARÉ plots consisting of 30 RR intervals. We considered 30 RR intervals per iteration and in each iteration after constructing the POINCARÉ plot we calculated the dispersion around the diagonal line and used k-means based cluster analysis to determine the number of the clusters. If the dispersion was too high (greater than 0.06) and the number of clusters was 1, or the number of clusters was more than 9; we assigned “AFib” to that series of RR intervals, otherwise to “Non-AFib”. The details of the algorithm can be seen in our previous study [16]. Since our data set did not contain real AFib cases, only specificity could be calculated with regard to the efficiency of detection. The evaluation of atrial fibrillation detection results for synchronized data validation can be seen in Tables 8 and 9.

Conclusion

Even though the CardioSport device may suffer from signal loss due to its design, we managed to determine that it can be safely used for telemedical purposes of measuring HRV and atrial fibrillation. We found only a few usable data segments that were less than 5 minutes long. With our algorithm that detects gaps and errors in

Table 8: Results from the synchronized data related to atrial fibrillation detection

<table>
<thead>
<tr>
<th>Subject</th>
<th>Number of iterations</th>
<th>Schiller MT-101/MT-200 system</th>
<th>CardioSport TP3 Heart Rate Transmitter</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1</td>
<td>331</td>
<td>26 8% 305</td>
<td>92% 31 9%</td>
</tr>
<tr>
<td>#2</td>
<td>1796</td>
<td>9 1% 1787</td>
<td>99% 3 &gt;1%</td>
</tr>
<tr>
<td>#3</td>
<td>1120</td>
<td>7 1% 1113</td>
<td>99% 5 1%</td>
</tr>
<tr>
<td>#4</td>
<td>1427</td>
<td>11 1% 1416</td>
<td>99% 16 1%</td>
</tr>
<tr>
<td>#5</td>
<td>964</td>
<td>46 5% 918</td>
<td>95% 45 5</td>
</tr>
<tr>
<td>Min</td>
<td>-</td>
<td>- 1% -</td>
<td>92% - &gt;1% -</td>
</tr>
<tr>
<td>Max</td>
<td>-</td>
<td>- 8% -</td>
<td>99% - 9%  -</td>
</tr>
<tr>
<td>Mean</td>
<td>-</td>
<td>- 3% -</td>
<td>97% - 3%  -</td>
</tr>
<tr>
<td>STD</td>
<td>-</td>
<td>- 3% -</td>
<td>3% - 4%  -</td>
</tr>
</tbody>
</table>

Table 9: Results from the data validation process related to atrial fibrillation detection

<table>
<thead>
<tr>
<th>Patient</th>
<th>Number of iterations</th>
<th>Schiller MT-101/MT-200 system</th>
<th>CardioSport TP3 Heart Rate Transmitter</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1</td>
<td>241</td>
<td>3 1% 238</td>
<td>99% 8 3%</td>
</tr>
<tr>
<td>#2</td>
<td>1879</td>
<td>29 2% 1850</td>
<td>98% 2 &gt;1%</td>
</tr>
<tr>
<td>#3</td>
<td>808</td>
<td>15 2% 793</td>
<td>98% 3 &gt;1%</td>
</tr>
<tr>
<td>#4</td>
<td>1296</td>
<td>10 1% 1286</td>
<td>99% 20 2%</td>
</tr>
<tr>
<td>#5</td>
<td>544</td>
<td>6 1% 538</td>
<td>99% 7 1%</td>
</tr>
<tr>
<td>Min</td>
<td>-</td>
<td>- 1% -</td>
<td>99% - &gt;1% -</td>
</tr>
<tr>
<td>Max</td>
<td>-</td>
<td>- 2% -</td>
<td>99% - 3%  -</td>
</tr>
<tr>
<td>Mean</td>
<td>-</td>
<td>- 1% -</td>
<td>99% - 1%  -</td>
</tr>
<tr>
<td>STD</td>
<td>-</td>
<td>- &gt;1% -</td>
<td>&gt;1% - 1%  -</td>
</tr>
</tbody>
</table>
the signal and removes them with an average effectiveness of more than 70%, which translates into having enough data to calculate HRV and atrial fibrillation from daytime measurements.

Regarding atrial fibrillation detection, we can conclude that by using the developed data validation algorithm the reference Schiller MT-101/MT-200 measurements produced only slightly better results with regard to false detections than the CardioSport TP3 Heart Rate Transmitter. In two cases the CardioSport measurements proved to be even better than Schiller records, which implies that some relatively simple heart rate recorders are equivalent to some Holter devices after signal processing using the data validation algorithm. We have to emphasize; however, that we have not performed any measurements on actual atrial fibrillating patients yet. Therefore, the investigation of the sensitivity of our atrial fibrillation detection algorithm under the presented circumstances could be the subject of further studies. In summary, the CardioSport as a low-cost device can easily be integrated into a lifestyle support system as a telemedical solution.

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REFERENCES