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Rapid Fluctuations in Atrial Fibrillatory Electrophysiology Detected during Controlled Respiration

Fredrik Holmqvist, Martin Stridh*, Johan EP Waktare†, Johan Brandt‡, Leif Sörnmo*,
Anders Roijer, Carl J Meurling

Department of Cardiology, Lund University Hospital, Lund, Sweden, *Department of Electroscience, Lund Institute of Technology, Lund, Sweden, †Cardiac Department, Blackpool Victoria Hospital, United Kingdom and ‡Department of Thoracic Surgery, Lund University Hospital, Lund, Sweden

Running Head: Controlled Respiration and AF Electrophysiology

Contact information:
Dr Fredrik Holmqvist
Dept of Cardiology
Lund University Hospital Tel: +46-(0) 46-17 35 18
SE-221 85 Lund Fax: +46-(0) 46-15 78 57
SWEDEN E-mail: Fredrik.Holmqvist@kard.lu.se
ABSTRACT

Introduction: Heart rate during sinus rhythm is modulated through the autonomic nervous system generating short-term oscillations. The high-frequency components in these oscillations are associated with respiration, causing sinus arrhythmia, mediated by the parasympathetic nervous system. The purpose of this study was to evaluate whether slow controlled respiration causes cyclic fluctuations in the frequency of the fibrillating atria.

Methods and Results: Eight patients (four female; median age 63 years, range 53 to 68) with chronic atrial fibrillation (AF) and 3rd degree AV block treated by permanent pacemaker were studied. Electrocardiogram was recorded during baseline rest, during 0.125 Hz frequency controlled respiration and finally during controlled respiration after full vagal blockade. Fibrillatory frequency was calculated using frequency analysis of the fibrillatory ECG (FAF-ECG) for overlapping 2.5 s segments and spectral analysis of the resulting frequency trend was performed to determine the spectrum of variations of fibrillatory frequency.

Normalized spectral power at respiration frequency increased significantly during controlled respiration from 1.4 (0.76 to 2.0) (median and range) at baseline to 2.7 (1.2 to 5.8) (p = 0.01). Following vagal blockade the power at respiration frequency decreased to 1.2 (0.23 to 2.8) (p = 0.01).

Conclusion: Controlled respiration causes cyclic fluctuations in the atrial fibrillatory frequency in patients with long duration AF. This phenomenon seems to be related to parasympathetic modulations of the atrial fibrillatory refractory period.

Keywords: Atrial fibrillation, Autonomic nervous system, Spectrum analysis, Non-invasive
**INTRODUCTION**

Cyclic fluctuations in heart rate during sinus rhythm vary in response to a range of physiological regulatory systems. These fluctuations have been utilized to study autonomic regulation of human cardiovascular physiology.\(^{(2, 12, 28)}\) The origin of variations in the sinus interval is not fully explained, but appears to be predominantly determined by interactions between the autonomic nervous system, hormonal efferent systems and cardiac pacemaker cells.\(^{(12, 28)}\)

Analysis of heart rate variability by spectral methods (frequency domain analysis) is widely used to study autonomic tone and variations thereof. For example, fluctuations in the so-called high-frequency band (HF, 0.15 to 0.4 Hz) correspond to normal respiratory frequencies and are thought to reflect parasympathetic tone.\(^{(26)}\) These fluctuations have been shown to be abolished by parasympathetic withdrawal.\(^{(1, 2, 28)}\)

Although variations in beat-to-beat intervals (i.e. RR intervals) are usually examined in frequency domain analysis studies, it is assumed that during sinus rhythm the variability of heart rate is determined by changes in autonomic input to the sino-atrial node (i.e. it is assumed that variations in the AV interval do not significantly influence total heart rate variation). Studies on heart rate variability in settings outside of sinus rhythm are sparse, but occasional studies during atrial tachycardia\(^{(15)}\) and atrial fibrillation (AF)\(^{(34, 35)}\) have been performed. These utilized analysis of RR interval data, which represents a potential significant weakness.\(^{(8)}\) During AF rhythmic fluctuations in RR intervals are potentially determined by both AV-nodal conduction and atrial input frequency.\(^{(23)}\) Thus, spectral analysis of RR intervals during AF cannot be assumed to reflect solely or predominantly atrial events as it may be primarily determined by AV nodal factors.

This study set out to explore whether in patients with atrial fibrillation, vagal induced changes in atrial electrophysiology can be detected during rhythm-controlled respiration. This is a time frame much shorter than previously examined for autonomic influences on refractoriness. We utilized time-frequency analysis\(^{(14, 16, 24, 27, 32)}\) which examines atrial electrophysiological
METHODS

Study population

Patients with permanent AF and complete heart block treated by permanent pacemaker were recruited. Exclusion criteria were pharmacological treatment with sympathomimetic or anti-cholinergic drugs, AF duration exceeding three years, or clinical conditions that could affect the autonomic nervous system, such as diabetes, hyperthyroidism, alcohol abuse, smoking or recent cardiovascular or cerebrovascular events. All patients gave written informed consent; the study was approved by the local Ethics Committee and complied with the Declaration of Helsinki.

Study Protocol and Data Acquisition

In line with recommended practice for autonomic studies, all studies were performed during the morning and patients were instructed to abstain from eating for four hours and from caffeine for 24 hours prior to commencing the study. Cardiovascular drugs, other than direct sympathomimetic or anti-cholinergic drugs, were not discontinued. On arrival the patients’ pacemakers were programmed to pace at 60 bpm in unipolar VOO mode for the duration of the study.

Following the attachment of ECG electrodes and other preparation, patients were rested supine for 15 minutes in a quiet, temperature-controlled room. Electrocardiogram acquisition was then begun, and five minutes of baseline data was acquired. Subsequently, controlled respiration with four seconds inspiratory and four seconds expiratory phase duration was begun using auditory guidance (i.e. eight second cycles, respiratory frequency = 0.125 Hz) and a second five-minute period of data for analysis was acquired once the patient was in a stable synchronized respiratory pattern. Finally full vagal blockade was induced by intravenous administration of atropine (0.02 mg/ kg). After a five minute interval, 0.125 Hz controlled respiration was again commenced and
a final five-minute period of ECG for analysis was acquired. Real time ECG was monitored throughout the study and blood pressure was manually measured every minute.

Data Analysis

(A) ECG Acquisition
A standard 12-lead ECG was acquired using a custom made optically isolated PC card (Siemens Elema AB, Solna, Sweden). The digital signal (1 kHz sampling rate, 16 bit analogue-to-digital conversion, 0.6 µV amplitude resolution) was transferred to a personal computer, where the data was written to a file for subsequent off-line processing. Electrocardiogram signal acquisition was continuous throughout the study, but the periods for subsequent analysis were noted at the time of recording.

(B) Pacing Spike and QRST Removal
The initial stage comprised identification and digital subtraction of the unipolar pacing spike. Subsequently QRST cancellation was performed using a spatiotemporal approach(33) which involves multiple templates, allowing an improved matching to each individual complex and leaving less ventricular residua than conventional processing.

In this study, only data from lead V1 was used for further analysis, although V2 and V3 data were used in the QRST cancellation process.

(C) Frequency Analysis to Detect Fluctuations at Respiratory Cycle Lengths
A new method for fibrillatory frequency (FF) trend estimation was used based on the logarithmic Fourier transform of overlapping (one every second) 2.56 second segments of ‘atrial fibrillatory ECG’. (32) An example of the resulting time-frequency distribution is shown in Figure 1d. The frequency estimation was performed by aligning the spectrum of each new signal segment to a template spectrum with known main peak position. The advantage with such a procedure is that the entire energy of the signal (both fundamental frequency and harmonic pattern representing the rate and waveform shape of the signal, respectively) is used to obtain detailed frequency
estimation. Estimation accuracy was ensured by requiring that both the peak magnitude and the ratio of the peak magnitude to the noise level of the spectrum exceeded certain thresholds. For each new spectrum, a template spectrum was formed by averaging frequency-aligned versions of previous spectra. All frequency estimates were used to provide a time series (see the frequency trend in Figure 1e) for a second spectral analysis using the fast Fourier transform (FFT) to detect fluctuations of the peak frequency value within the band 0.04-0.4 Hz see Figure 1f for an example of such modulation spectra. This range was chosen such that it corresponds to the range of the conventionally tested high and low frequency bands (HF, 0.15-0.4 Hz and LF, 0.04-0.15 Hz respectively) examined in heart rate variability studies, and contains the frequency of interest (0.125 Hz). The spectra were estimated from segments of five minute duration from each of the three intervals of interest for analysis, namely, the initial period of quiet rest, baseline (B), the period of controlled respiration (CR) and, finally, the period of controlled respiration post-atropine administration (PA).

Figure 1 illustrates the process from ECG via ‘residual ECG’ to final spectral analyses.

**Post-Processing Analyses and Statistics**

From each power spectrum was calculated: total power (0.04-0.4 Hz); low frequency power (0.04-0.15 Hz); high frequency power (0.15-0.4 Hz); and power at the frequency of controlled respiration (0.125 Hz).

All values are expressed as median and range. Wilcoxon matched pairs test was used for comparison between paired samples. Spearman ranked correlation coefficient was used elsewhere. A p < 0.05 was considered statistically significant. All statistical analyses were performed using STATISTICA for Windows version 6.1 (StatSoft, Inc., Tulsa, OK, USA).

**RESULTS**

**Study Population**
Eight patients (four female, median age 63 years, range 53 to 68) were included. Four patients were taking RAAS-inhibitors, one patient each was taking a β-blocker and a calcium channel antagonist, two patients were not using any heart active drug. Echocardiography demonstrated a median left atrial diameter of 42 mm (32 to 53) and a median ejection fraction of 55% (35 to 55). The median AF duration was 17 months (range five to 32 months, defined as the elapsed time since last documented SR) and the patients had undergone a median of nine prior cardioversions (range 4 to 26) (Table 1).

**Data Availability**

All patients were able to satisfactorily perform the rhythm-controlled respiration and all provided recordings were suitable for analysis.

**Population Observations**

The median FF of patients at baseline was 6.9 Hz (6.1 to 8.2 Hz), with controlled respiration 7.0 Hz (6.0 to 8.2 Hz, p=NS) and following atropine injection 6.8 Hz (6.1 to 8.1 Hz, p=NS). Power at the frequency of respiration (P0.125) rose significantly during controlled respiration, from 0.15 ms² (0.06 to 0.25) at baseline, to 0.36 ms² (0.11 to 0.58) (p = 0.01). Following vagal blockade the P0.125 decreased to 0.13 ms² (0.02 to 0.42) (p = 0.02), which was not significantly different compared to baseline.

Inspection of full spectra demonstrated that there was considerable but variable leakage of low frequency noise into the region of interest. To correct for the significant variability in total spectral power, the data were modeled by normalizing for this value. The resultant trends were similar (Table 2). The ratio of P0.125 to 'high frequency power' (0.15 – 0.4 Hz) (P0.125/ PHF) was found to best reflect actual shifts in power and also corrected for the absolute pollution of the spectrum. P0.125/ PHF rose significantly during controlled respiration from 1.4 (0.76 to 2.0) at baseline to 2.7 (1.2 to 5.8) (p = 0.01) and fell to 1.2 (0.23 to 2.8) (p = 0.01) following atropine administration (figure 2). There was no significant difference between the baseline and post-atropine P0.125/ PHF (p = 0.40). There were no significant changes in other measured parameters.
**Observations in Individual Patients**

The response to controlled respiration was heterogeneous. Although all patients increased the P_0.125/ P_HF during controlled respiration followed by a decrease after vagal blockade, the amount of response was variable (table 3). A spectrum from a single patient is illustrated in figure 3. In this patient, and in some but not all of the other patients, there appears to be significant low frequency spectral power exhibited in the low frequency band. Some of this energy might represent artifacts, but at present quantification of absolute power in both the low and high frequency band, as can be done during sinus rhythm, is not validated. No differences in patient characteristics in terms of age, AF-duration, LA-diameter, previous cardioversions, heart-drugs and other diseases could be found between marked responders and the less responsive patients.

**DISCUSSION**

**Comparison of dynamic changes in the electrophysiologic properties of the fibrillating atria to other scenarios**

It is well known that fluctuations in heart rate during sinus rhythm are mainly determined by autonomic effects on the sino-atrial node. Specifically increased sympathetic tone increases and parasympathetic tone decreases the depolarization frequency of the specialized atrial pacemaker cells by respectively increasing and decreasing the rate of the spontaneous diastolic depolarization.(2, 6) However, the situation in the fibrillating atria is fundamentally different. The cycle length is the sum of the atrial myocyte effective refractory period, any possible excitable gap that may exist during AF(3, 19) and the action potential up-stroke time, which is related to the conduction velocity. Fibrillatory frequency is the inverse of the fibrillatory cycle length. Thus, rhythmic fluctuations in atrial FF may result from dynamic changes in the refractory period, as well as changes in the conduction velocity or of the excitable gap. The derived parameter of the FAF-ECG method, dominating atrial cycle length (DACL, inverse
of FF), from lead V1 has been shown to closely correlate to an invasively measured spatial mean of right atrial free wall cycle length.(14) This in turn has been shown to reflect atrial refractoriness.(7, 18, 36) In this study FF is used as an index of atrial refractoriness. While enhanced parasympathetic tone prolongs the ventricular refractory period(17) it has the opposite affect in the atria during sinus rhythm.(17, 21, 31) The effect on atrial myocardium during AF is less studied but the evidence that exists, suggest that acetylcholine and enhanced vagal tone shortens the effective refractory period(16, 34) as it does during sinus rhythm. Consequently, enhanced vagal tone would lead to a higher fibrillation frequency, this could however not be verified in this study. The response of vagal discharge is believed to be inhomogeneous, as the right atrium has more extensive parasympathetic innervation than the left atrium.(22, 37) This is consistent with the response of acetylcholine administration or vagal stimulation, demonstrating a more marked shortening of the refractory period in the right atria than in the left atria.(37), (31) Experimental studies are limited but those available have not shown an effect of acetylcholine on conduction velocity.(20, 31)

**Mechanism of Shift in Spectral Power in Response to Slow Controlled Respiration and Attenuation by Atropine**

Physiologically the atrial refractory period adapts automatically to changes in heart rate (i.e. depolarization rate). These changes occur immediately.(11) The autonomic nervous system and cardioactive drugs may also alter the electrophysiological properties of the atria.(24, 31) It has been demonstrated in previous experimental studies that cholinergic stimulation leads to a response within milliseconds, in contrast to the marked delay of response in the sympathetic modulation.(30) Moreover, it is well known that the length of the monophasic action potential can show rapid changes (i.e. beat to beat changes) even during high rate atrial depolarizations.(29) Thus, we conclude that it is theoretically possible for the atrial FF (i.e. the atrial refractory period) to be modulated at frequencies corresponding to respiration.
The shifts in spectral power after atropine administration occurred without any changes in respiration pattern, ventricular rate, blood pressure and body posture or body movements. Finally, although parasympathetic modulation was specifically examined in this study, the human autonomic nervous system is in practice a complex interaction. Simultaneous afferent inputs occur from both sympathetic and parasympathetic limbs and are influenced by the central nervous system modulation. However the methodology employed has demonstrated a detectable role of parasympathetic efferents during longstanding AF.

Atrial stretch(5) may also have been involved; cyclic change in left and right atrial pressure would have been accentuated by slow breathing. A direct effect of stretch on atrial electrophysiology was disproved as a mechanism for our observations by its abolition with atropine. This indicates that reflex arches involving cholinergic receptors are involved. It is likely that the efferent limb of these arches is parasympathetic, but an involvement of baroreceptors and sympathetic fibres in the afferent limb can not be excluded.

Preserved Dynamic Electrophysiological Changes in Chronic AF and Possible Mechanisms for Inter-Patient Variability

Two very interesting aspects of the present study are firstly the fact that dynamic changes in atrial fibrillatory rhythm were detectable among patients with long duration AF, and secondly the observed inter-patient variation. All patients had long duration of AF and thus by current understanding, electrical remodeling would be expected to be ‘complete’.(4, 13) The degree of spectral power shift to 0.125 Hz with controlled respiration was variable. Previous studies have shown that following pharmacological interventions (and autonomic modulation) the change in FF is more pronounced in those who have a lower initial value.(24, 25) This implies that the atrium is only susceptible to modulation if the FF is low. However there was no such trend seen in the present study.

The present study is too small and the population too heterogeneous to draw conclusions regarding the importance of the above factors, but the variable preservation of responsiveness to
autonomic stimuli is an important observation.

Methodological issues and Study limitations

Although all patients had long duration AF and complete heart block, the population was heterogeneous in ways which may be relevant. Complete heart block and AF are associated with, or directly caused by, a range of conditions such as congestive heart failure, other heart diseases, and with advanced age. All of these conditions are associated with attenuation of parasympathetic tone(10) and some of which were present in our patients. Conversely, half the patients in our study were taking RAAS inhibitors, and blockade of the renin-angiotensin system is known to increase the activity in the low-frequency band and to decrease the fluctuations at respiratory frequencies.(1, 2) One patient was taking a beta-blocker, which has been shown to shift the frequency distribution towards higher frequencies,(9) although the exact effects are complex the shift may be induced by augmentation of vagal tone.

We excluded conditions and drugs which might have abolished autonomic modulations, but this compromise was made for logistical (patient recruitment) and ethical reasons (inadvisability of drug withdrawal in elderly patients with cardiac co-morbidity and without gain from participation in our study).

In this study we estimated changes in atrial FF using time-frequency analysis, which is an indirect assessment of atrial refractoriness. This method cannot discriminate between changes in atrial refractoriness and conduction velocity; the latter, however, is unlikely to change after atropine administration. The response of conduction velocity to controlled, slow respiration is uncertain, but since atropine attenuated the induced change (it is assumed that atropine does not affect the conduction velocity), it is likely that the observed change in atrial refractoriness is genuine. Time-frequency analysis from the V1 lead correlates to the fibrillatory cycle length of the right atrial free wall, and therefore direct deductions about the rest of the atria can not be made.

Equally, this study recruited patients with longstanding AF. Therefore inferences about shorter duration AF can not be made, although it would be expected that the dynamic responsiveness of
less re-modeled atria would be more pronounced.

The respiration was controlled by the use of auditory guidance but the definite respiration frequency was not objectively assessed during the study. Therefore it is possible that the full extent of the parasympathetic modulation of FF via respiration may be larger than found in the present study.

**Conclusions**

We found that slow controlled respiration causes cyclic alterations of AF rhythm at the respiratory frequency in patients with long lasting AF. The attenuation of this phenomenon by vagal blockade confirms that the observation is genuinely related to parasympathetic modulations of atrial electrophysiology. Our utilization of paced patients with complete heart block excludes any mechanism related to autonomic effects on the atrioventricular node and the cancellation of ventricular components of the ECG out-rules contributions of ventricular depolarization to the spectral analysis.
ACKNOWLEDGEMENTS

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GRANTS

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Effects of mild hyperkalemia on conduction velocity and effective refractory period in various parts of the


FIGURE LEGENDS

Figure 1 - Schematic illustration of the derivation of the modulation spectrum of the ECG, via QRST-subtraction and two subsequent fast Fourier transforms (FFT). The original ECG (a) is subsequently subjected to pacemaker spike removal (b) and QRST cancellation (c). By repeated measurements of the continuous recording a time-frequency distribution is obtained (d) from which the frequency trend is extracted (e). A final fast Fourier transform produces the modulation spectra of the different study phases (f).

Figure 2 - Response to controlled respiration, changes in P_{0.125}/P_{HF}. * p = 0.01, † p = NS. Wilcoxon matched pairs test.

Figure 3 - Example of modulation spectrum (patient a). The modulation spectra of the three different phases of the study are individually plotted. A clear peak at the frequency of respiration (0.125 Hz) is evident during the controlled respiration (CR) phase. It is not visible at baseline (B) and in spite of the same respiration pattern the peak is abolished following atropine administration (PA), suggesting a parasympathetic origin.
<table>
<thead>
<tr>
<th>Patient</th>
<th>Age</th>
<th>Sex</th>
<th>EF %</th>
<th>Left atrial diameter (mm)</th>
<th>AF Duration (months)</th>
<th>Number of cardioversions</th>
<th>Heart active drugs</th>
<th>Co-Morbidity</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>62</td>
<td>Female</td>
<td>55</td>
<td>41</td>
<td>7</td>
<td>10</td>
<td>diltiazem</td>
<td>none</td>
</tr>
<tr>
<td>b</td>
<td>64</td>
<td>Male</td>
<td>45</td>
<td>43</td>
<td>5</td>
<td>15</td>
<td>none</td>
<td>COPD</td>
</tr>
<tr>
<td>c</td>
<td>65</td>
<td>Female</td>
<td>35</td>
<td>37</td>
<td>9</td>
<td>4</td>
<td>metoprolol</td>
<td>HT, CHF</td>
</tr>
<tr>
<td>d</td>
<td>64</td>
<td>Female</td>
<td>55</td>
<td>32</td>
<td>30</td>
<td>4</td>
<td>none</td>
<td>none</td>
</tr>
<tr>
<td>e</td>
<td>54</td>
<td>Male</td>
<td>55</td>
<td>46</td>
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<td>7</td>
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<td>45</td>
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<td>26</td>
<td>spironolactone</td>
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<tr>
<td>h</td>
<td>53</td>
<td>Male</td>
<td>55</td>
<td>32</td>
<td>24</td>
<td>7</td>
<td>enalapril</td>
<td>HT</td>
</tr>
</tbody>
</table>

AF, atrial fibrillation; CHF, chronic heart failure; COPD, chronic obstructive pulmonary disease; EF, ejection fraction; HT, Hypertension
**TABLE 2. MEASURED PARAMETERS.**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Baseline</th>
<th>Controlled Respiration</th>
<th>Post Atropine</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>median</td>
<td>(range)</td>
<td>median</td>
</tr>
<tr>
<td>Mean arterial Blood pressure (mmHg)</td>
<td>90</td>
<td>(83 - 105)</td>
<td>92</td>
</tr>
<tr>
<td>Fibrillation frequency (Hz)</td>
<td>6.9</td>
<td>(6.1 - 8.2)</td>
<td>7.0</td>
</tr>
<tr>
<td>High frequency power (0.15-0.4 Hz band, ms²)</td>
<td>0.13</td>
<td>(0.03 - 0.21)</td>
<td>0.10</td>
</tr>
<tr>
<td>Low frequency power (0.04-0.15 Hz band, ms²)</td>
<td>0.24</td>
<td>(0.11 - 0.36)</td>
<td>0.20</td>
</tr>
<tr>
<td>[High freq.]/ [Low freq.]</td>
<td>0.49</td>
<td>(0.31 - 0.67)</td>
<td>0.49</td>
</tr>
<tr>
<td>Total power (0.04-0.4 Hz, ms²)</td>
<td>0.17</td>
<td>(0.06 - 0.25)</td>
<td>0.13</td>
</tr>
<tr>
<td>Power at 0.125 Hz (ms²)</td>
<td>0.15</td>
<td>(0.06 - 0.25)</td>
<td>0.36</td>
</tr>
<tr>
<td>Power at 0.125 Hz / High frequency power</td>
<td>1.4</td>
<td>(0.76 - 2.0)</td>
<td>2.7</td>
</tr>
</tbody>
</table>

* p = 0.01 compared to baseline, † p = 0.01 compared to controlled respiration, ‡ p = 0.02 compared to controlled respiration (Wilcoxon Matched Pairs Test)
Table 3. Changes in power at the frequency of controlled respiration.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Baseline</th>
<th>Controlled Respiration</th>
<th>Post Atropine</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$P_{0.125}/P_{HF}$</td>
<td>$P_{0.125}/P_{HF}$</td>
<td>$P_{0.125}/P_{HF}$</td>
</tr>
<tr>
<td>a</td>
<td>1,00</td>
<td>5,76</td>
<td>0,23</td>
</tr>
<tr>
<td>b</td>
<td>1,85</td>
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<tr>
<td>c</td>
<td>1,83</td>
<td>2,39</td>
<td>0,80</td>
</tr>
<tr>
<td>d</td>
<td>2,01</td>
<td>2,93</td>
<td>1,85</td>
</tr>
<tr>
<td>e</td>
<td>1,85</td>
<td>4,21</td>
<td>2,83</td>
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<tr>
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<td>0,95</td>
<td>1,23</td>
<td>1,03</td>
</tr>
<tr>
<td>g</td>
<td>1,05</td>
<td>3,63</td>
<td>1,33</td>
</tr>
<tr>
<td>h</td>
<td>0,76</td>
<td>2,42</td>
<td>0,27</td>
</tr>
</tbody>
</table>

All values represent the absolute values of power calculated at the frequency of respiration divided by the mean power value in the high frequency band ($P_{0.125}/P_{HF}$).
Original ECG (10 sec)

After pacemaker spike removal (10 sec)

After QRST cancellation (10 sec)

Time-frequency distribution (entire recording, 28 min)

Frequency trend (entire recording, 28 min)

Baseline  Controlled resp.  Post atropine

Modulation spectra
**FIGURE 2**

![Box plot showing median, 25%-75%, Min-Max for Baseline, Controlled Respiration, and Post Atropine.](image)

**FIGURE 3**

![Power spectra for B, CR, PA at 0.04 to 0.4 Hz.](image)