



# **An Application of LARY\_C: Discrete Wavelet Transform Applied to Heart Rate Variability Analysis. Assessment of the Autonomic Nervous System Behaviour in Control and Iron-Deficient Anemic Infants**

Claire Médigue, Julie Bestel, Sophie Renard, Jean Clairambault, Marcelo Garrido, Felipe Pizarro, Patricio Peirano

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***An application of LARY\_C: discrete wavelet transform applied to heart rate variability analysis. Assessment of the autonomic nervous system behaviour in control and iron-deficient anemic infants***

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THÈME 4



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de recherche***





# An application of LARY\_C: discrete wavelet transform applied to heart rate variability analysis. Assessment of the autonomic nervous system behaviour in control and iron-deficient anemic infants

Claire Médigue, Julie Bestel, Sophie Renard, Jean Clairambault, Marcelo Garrido, Felipe Pizarro and Patricio Peirano

Thème 4 — Simulation et optimisation  
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**Abstract:** Our aim was to assess the Autonomic Nervous System (ANS) behaviour in six months old iron-deficient anemic and control infants during sleep.

We used LARY\_C, a software package, developed in the synchronous, data flow, parallel language SIGNAL, primary dedicated to the ANS analysis, through the study of physiological signals and their correlations.

We applied the discrete wavelet transform to heart rate variability analysis, which reflects the SNA activity. The method provides suitable frequency resolution in low frequency domain, which is precisely of physiological interest.

Significant differences between iron-deficient and control infants are shown in both quiet and active sleep, suggesting an autonomic dysfunction or delay in iron-deficient anemic infants.

**Key-words:** time-frequency method, discrete wavelet transform, synchronous language, autonomic nervous system, physiological signals, heart rate variability, sleep, iron deficiency anemia

*(Résumé : tsvp)*

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## **Une application de LARY\_C: la transformée en ondelettes discrète appliquée à l'analyse des variabilités du rythme cardiaque pour l'étude du système nerveux autonome chez des enfants contrôles et anémiques par carence en fer**

**Résumé :** Notre but était d'étudier l'impact de la carence en fer sur le comportement du Système Nerveux Autonome (SNA) de bébés de six mois pendant le sommeil, en comparant deux groupes, l'un contrôle et l'autre atteint d'anémie par déficience martiale.

Nous avons utilisé LARY\_C, logiciel développé dans le langage synchrone flot de données et parallèle SIGNAL, essentiellement dédié à l'étude du SNA, à travers l'analyse des rythmes physiologiques et de leurs corrélations.

Nous avons appliqué la transformée en ondelettes discrète à l'analyse des variabilités du rythme cardiaque, reflet de l'activité du SNA. Cette méthode donne une bonne résolution fréquentielle dans le domaine des basses fréquences, qui sont précisément les fréquences d'intérêt physiologique.

Des différences significatives sont apparues entre les bébés carencés en fer et les bébés contrôles, pour chacun des stades de sommeil calme et agité; ceci suggère un dysfonctionnement ou un retard de développement du SNA chez les bébés anémiques par déficience en fer.

**Mots-clé :** méthodes temps-fréquence, transformée en ondelette discrète, langage synchrone, système nerveux autonome, signaux physiologiques, variabilité cardiaque, sommeil, anémie, carence en fer

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## Contents

<b>1</b>	<b>Introduction</b>	<b>5</b>
<b>2</b>	<b>The wavelet transform</b>	<b>7</b>
2.1	Recall: basics on the spectrogram . . . . .	7
2.2	Continuous wavelet transform . . . . .	8
2.3	Filtering interpretation . . . . .	8
2.4	Discrete transform . . . . .	8
2.5	Implementation . . . . .	9
<b>3</b>	<b>Clinical material and methods</b>	<b>9</b>
3.1	Subjects . . . . .	9
3.2	Recording . . . . .	9
3.2.1	General procedure . . . . .	9
3.2.2	Coding sleep states . . . . .	11
<b>4</b>	<b>Signal processing</b>	<b>11</b>
4.1	Environment . . . . .	11
4.1.1	The SIGNAL language . . . . .	11
4.1.2	LARY_C . . . . .	12
4.2	Description of the global processing . . . . .	12
4.2.1	ECG signal processing . . . . .	12
4.2.2	Respiratory signal processing . . . . .	15
4.2.3	DWT applied to HRV analysis . . . . .	17
4.3	Some practical aspects . . . . .	21
4.3.1	Synchronisations between resulting rhythms . . . . .	21
4.3.2	Visual criteria . . . . .	23
<b>5</b>	<b>Statistical processing</b>	<b>24</b>
5.1	Classical hypothesis tests . . . . .	24
5.2	Multivariate Data Analyses (MDA) . . . . .	24
<b>6</b>	<b>Clinical results</b>	<b>24</b>
6.1	Between-group differences in Quiet Sleep . . . . .	24
6.2	Between-group differences in Active Sleep . . . . .	27
6.3	Between-state differences in each group . . . . .	28

<b>7</b>	<b>Discussion</b>	<b>29</b>
7.1	About signal processing methods . . . . .	29
7.2	About Statistical methods . . . . .	31
7.3	About clinical results . . . . .	32
7.3.1	Influence of the sleep on the neurovegetative functions . . . .	33
7.3.2	Hypothesis on mechanisms underlying iron deficiency . . . .	34
<b>8</b>	<b>Conclusion</b>	<b>35</b>



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## 1 Introduction

This study is carried out on six months old infants, control and anemic, during a nap. We wanted to assess the influence of Iron Deficiency Anemia (IDA) on the Autonomic Nervous System (ANS) maturation according to sleep states.

This study, as part of an ECOS-CONICYT programme between France and Chili, is made in collaboration with INTA - University of Chili (department of neurophysiology on developpment).

The Iron Deficiency Anemia (IDA) continues to be the most common single nutrient deficiency in the world. About 20 to 25% of the infant population worldwide is affected [1]. Several recent studies provide strong evidence that IDA alters infant development and behaviour [7, 8]. Recent research on developmental patterns of Heart Rate Variability (HRV) provide measures that are thought to reflect central nervous system functioning earlier and more closely than tests of mental and motor development [2, 3, 4]. Indeed, the HRV reflect the control of the ANS, and the resulting HRV patterns reflect the continuous feedback between the central nervous system and peripheral autonomic receptors. The ANS is the part of the Nervous system involved in regulating all the vital functions, particularly the cardio-vascular and respiratory systems, to maintain the vital balance and to allow the organism to adapt to changes in its state. The ANS has two components (sympathetic and parasympathetic (vagal)) which have an antagonist effect on the heart; the balance between these two components varies in function of sleep and waking states; in quiet sleep, the parasympathetic tonus is dominant whereas in active sleep a lot of sympathetic stimulations of different origins arise.

The study of the Heart Rate (HR) and it's variabilities is already well known as a noninvasive mean of investigating the ANS [24]. The HRV being a function of sleep states, it is so necessary to separately study the heart rate during quiet sleep and active sleep. On the raw rhythms, only the global, resulting effects of the two components, of the ANS balance are seen. On the HR, for example, the global effect of a sympathetic activation is an acceleration, whereas the effect of a parasympathetic activation is a deceleration. It is possible to describe more precisely the participation of each composant of the ANS, through HR variabilities (HRV) (Fig. 12). Among the frequencies bands known to reflect the ANS activity, one can consider:

- An high frequency band (HF): "Heart rate fluctuations at respiratory frequencies are recognized as a noninvasive measure of cardiovagal activity and reflect more modulation of firing of baroreceptors and of cardiac and pulmo-

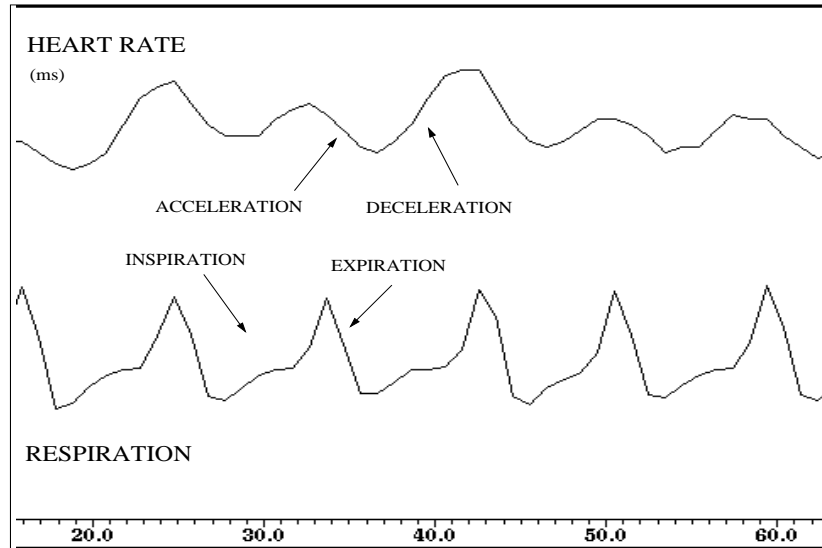


Figure 1: Respiratory Sinus Arrhythmia visualised on the heart rate and the raw respiratory signal, together resampled at 4Hz

nary volume receptors than an absolute measure of vagal tone” [6].

Above 0.15 Hz, it reflects the parasympathetic tone alone; on the HR, it is called Respiratory Sinus Arrhythmia (RSA) and it is the part of variability related to the respiratory cycles; RSA is described as an increase in heart rate during inspiration and a decrease during expiration (Fig. 1). Within mechanisms involved, the blockade of the centrally generated inspiratory drive and the blockade of lung receptors during inspiration are responsible for a phasic vagal inhibition [17].

- A middle frequency band (MF): Between 0.03 and 0.15Hz, it is related to the cardio-vascular control of the ANS, through the arterial baroreceptors; changes in heart rate and in peripheral vasomotor regulation are involved. The two ANS components, sympathetic and parasympathetic, have an antagonist chronotrope effect on the heart rate whereas the vasomotor activity involves the sympathetic effect on vessel contraction; the baroreflex activity, called Mayer waves, concerns periods about 10 heart beats and gives a peak rather around 0.1Hz in adult. As baroreflex responses derive from interactions between multiple reflex inputs, the situation is very complex and the frequential activity could be increased toward 0.1Hz in case of dominant vagal influence ; conversely, it could be reduced toward 0.05Hz in case of dominant sympathetic influence, with a strong peripheral vasoconstriction [24].
- Low and very low frequency bands (LF and VLF): lower frequency bands reflect probably not only the autonomic control and are involved in mechanisms of thermoregulation, variations in plasma hormones, including catecholamines, angiotensine ...

We used the Discrete Wavelet Transform (DWT) to analyse these heart rate variabilities, as it is developed in the medical library of LARY\_C. Our aim is to rely on the high frequential resolution improved by the DWT to compare the HRV of anemic and control infants.

## 2 The wavelet transform

### 2.1 Recall: basics on the spectrogram

The spectrogram is a classical time-frequency analysis tool that has been widely performed on non-stationnary signals:

$$S_f(t, \nu) = \left| \int_{-\infty}^{+\infty} \hat{f}(n) \hat{h}(n - \nu) e^{2i\pi n t} dn \right|^2 \quad (1)$$

where  $\hat{f}(n), n \in \mathbb{R}$ , denotes the Fourier transform of  $f \in L^2(\mathbb{R})$ . As shown by eq.(1), it simply consists in scanning the frequency domain by shifting a unique lowpass filter. Assuming that  $h$  is an analysing window with equivalent time-support  $\Delta t$  and bandwidth  $B$ , it provides a tiling of the time-frequency plane with uniform cells of fixed dimensions  $\Delta t$  and  $\Delta f = B$ , the product  $\Delta f \Delta t$  being constant and lower bounded (according to Heisenberg-Gabor's incertitude principle). Now, let us focus on another analysis tool, based on the wavelet transform.

## 2.2 Continuous wavelet transform

Let  $\Psi(t)$  be a function with good time and frequency localization properties. It has to fulfill the admissibility condition  $\hat{\Psi}(0) = \int_{\mathbb{R}} \Psi(t) dt = 0$ , and it is then called the mother wavelet. We define the family of its dilated and time shifted versions

$$\left\{ \psi_{(a,b)}(t) = \frac{1}{\sqrt{a}} \Psi \left( \frac{t-b}{a} \right), b \in \mathbb{R}, a \in \mathbb{R}_+^* \right\}$$

One then defines the CWT of  $f$  as the expansion of  $f$  onto the analysing set  $\{\psi_{(a,b)}\}$ :

$$C_f(a, b) = \langle f, \psi_{(a,b)} \rangle = \int_{-\infty}^{+\infty} f(t) \overline{\psi_{(a,b)}}(t) dt \quad (2)$$

$(a, b)$  taking on values in the half plane  $\mathbb{R}_+^* \times \mathbb{R}$ .  $C_f(a, b)$  is a time-scale representation of  $f$ .

## 2.3 Filtering interpretation

When dealing with finite energy signals:  $f \in L^2(\mathbb{R})$ , we have:

$$\int |f(t)|^2 dt = \frac{1}{C_{\Psi}} \int \int \frac{|C_f(a, b)|^2}{a^2} da db$$

where  $C_{\Psi}$  is a coefficient of normalization dependent on  $\Psi$  (see [10]). Given the admissibility condition,  $\Psi$  is band-limited with bandwidth  $B$  and central frequency  $\nu_0$ ; then its dilated version  $\Psi(t/a)$  has a spectrum centred around  $\nu_0/a$  with bandwidth  $B/a$ . The convolutive structure entering the definition of CWT (eq.(2))1 results in a  $Q$ -constant filter bank analysis, where  $Q = \frac{B}{\nu_0}$ . Therefore, the CWT yields a frequency-dependent tiling of the time-frequency plane:  $\Delta f$  and  $\Delta t$  vary depending on  $a$  (one at the expense of the other) while maintaining the product  $\Delta f \Delta t$  constant; it thus provides good frequency resolution in low frequencies. However, the CWT is informationally redundant, since it is a transformation from  $L^2(\mathbb{R})$  to  $L^2(\mathbb{R}^2)$ .

## 2.4 Discrete transform

To reduce the redundancy of the CWT, a discrete version has been proposed. Loosely speaking, the DWT can be understood as a CWT sampled on a discrete grid of the  $(a, b)$  plane. A dyadic grid is commonly proposed:  $a = 2^j, b = 2^j k$  where  $j \in \mathbb{Z}^*, k \in \mathbb{Z}$ . This particular tiling corresponds to a critical sampling of the time-scale

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plane that ensures no loss of information and possible reconstruction. Moreover, I. Daubechies showed that it was possible to find compact-supported orthonormal bases yielding discrete decomposition, which guarantees no loss of energy through the whole expansion [11]. A similar frequency interpretation as for the CWT holds for the DWT: given a signal sampled at  $F_s$  Hz, the wavelet at scale  $j$  acts as a bandpass filter between  $F_s 2^{j-1}$  Hz and  $F_s 2^j$  Hz. Besides reduction of redundancy, DWT allows efficient and fast algorithms.

## 2.5 Implementation

A fast implementation was proposed by S. Mallat [10]. Its principle is depicted in figure 2.  $G$  and  $H$  are respectively highpass and lowpass filters that form a Quadrature Mirror Filters pair ([12]). Downsampling by a factor of 2 allows using the same filters at each resolution level, and therefore leads to an iterative structure: the so-called pyramidal algorithm. Convolutions were computed by Radix FFT multiplications followed by IFFT. The wavelets used are orthonormal compact-supported Daubechies wavelets. At each scale  $j$ , the quantities displayed are normalized  $|C_f(2^j, 2^j k)|^2$ , and will now be referred to as Discrete Wavelet Coefficients (DWC).

## 3 Clinical material and methods

### 3.1 Subjects

Fifteen control and fifteen iron deficient anemic infants, 6 to 7 months old were studied. The healthy subjects are defined according in particular the following criteria: birth weight  $\geq 3.0$  Kg, singleton birth, no congenital or neonatal complications or anomalies, and no other nutrient deficiency. The diagnostic of iron-deficient anemia was based on biological criteria (as hemoglobin  $\leq 103g/l$ ) and subsequently on an increase in hemoglobin level of 10 g/l or more in response to iron therapy. The research protocol was approved by the Institutional Review Boards of the University of Michigan Medical Center, Ann Arbor, and of INTA, University of Chile, Santiago.

### 3.2 Recording

#### 3.2.1 General procedure

Standard polygraphic recordings were conducted in the sleep laboratory of the Developmental Neurophysiology Unit, INTA, University of Chile. They were performed

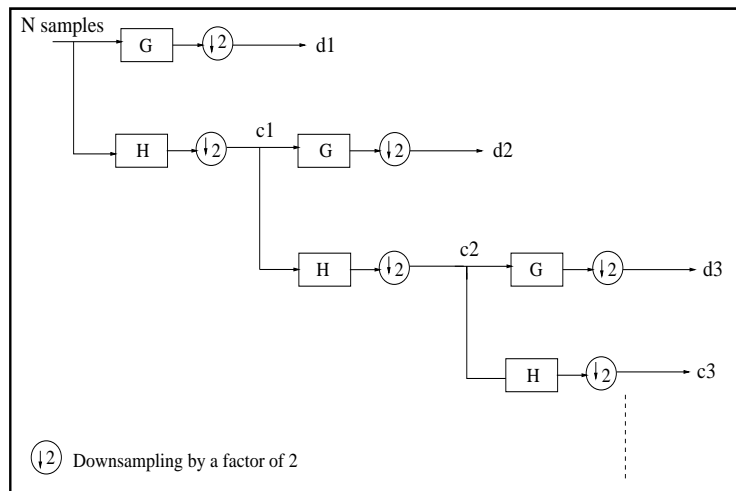


Figure 2: Pyramidal algorithm

during a spontaneous afternoon nap, in standardized conditions to limit the influences of environment, circadian rhythms and food intake on sleep-waking patterns and related physiological activities.

Data were digitized at 250 Hz and simultaneously recorded on paper tracings and stored on optic disk.

Electrocardiogram (ECG) and respiratory signal were automatically analysed; concerning breathing, abdominal respiratory movements using a mercury strain gauge and nostril airflow using a thermistor were recorded.

Sleep states were visually determined. They were related with ECG and respiratory signals by the mean of a file including the time of the state change and its nature. Correspondence between paper tracings and digitized recordings was ensured by a time counter in LARY\_C: beside sleep state changes, it allowed also to locate special events, as rocking in the mother arms, which have to be discarded from the automatic analysis.

### 3.2.2 Coding sleep states

They were visually coded by the concordance of behavioral and electrophysiological criteria [8, 9], which are extracted from the electro-encephalographic (EEG), electro-oculographic (EOG) and electro-myographic (EMG) signals.

- Quiet sleep: eyes closed, no eyes movements, EEG patterns with predominant delta waves with or without spindles.
- Paradoxical state: eyes closed, rapid eyes movements, muscular atonia, predominant high frequency and low voltage EEG patterns and irregular respiration.
- Indeterminate sleep: mixed characteristics of the two previous states.

## 4 Signal processing

The approach given by LARY\_C depends on two points: the use of the synchronous language SIGNAL to analyse relationships between rhythms and the library of medical signal processing methods to construct these rhythms.

### 4.1 Environment

#### 4.1.1 The SIGNAL language

SIGNAL is a data flow, oriented synchronous language for describing relationships among signals at various sampling rates; signals are sequences of events with associated values. The conception of time in SIGNAL is a minimal one, serving only to describe an application in a determinist way (same computations always run in the same order); in particular, no duration is considered at this level. The discrete time associated with a signal is called the 'clock' of the signal. The relationships among events are:

- precedence: occurrences of a same signal are totally ordered;
- simultaneity: two occurrences of different signals may be simultaneous or not;
- functional: values associated with simultaneous events may be linked by a computation expression.

SIGNAL provides a small set of temporal operators, sufficient for expressing all kinds of synchronization relationships among the clocks of the signals.

The application may be seen as a directed graph; arcs represent signals and dependancies between inputs and outputs of the nodes; a node express temporal and functional constraints of the signals that it connects. The SIGNAL compiler verifies

the global coherence of this complex relational system, and builds the corresponding automaton for a target machine [13, 18, 19, 20, 21].

#### 4.1.2 LARY\_C

We elaborated a complete processing chain in LARY\_C, a software package for physiological rhythms analysis, previously described. [21, 27] ; LARY\_C is based on the SIGNAL industrial environnement of development , SILDEX <sup>1</sup>. LARY\_C is supported by UNIX and X Windows. The flow chart of the whole parallel processing for this application is shown on Fig.3; it is divided into the following steps:

**Cardio and respiratory rhythms detection** A parallel preprocessing of the raw signals gives the resulting rhythms:

- From the electro-cardiographic signal (ECG) to the heart rate (HR)

The first processing step is the extraction of the different rhythms: a QRS detection adaptative algorithm is applied on the ECG, then an equidistant resampling at 4 Hz [14] is applied on the heart rate (HR).

- From the respiratory signal, the instantaneous respiratory frequency is calculated by differentiation of the analytical signal.

**Heart rate processing** The Discrete Wavelet Transform, used as a time-frequency method, studies the heart rate variabilities. This analysis is done by 1024 points epochs.

**Cardio-respiratory parameters extraction** The delivered results are the HR (mean RR, DWC) and respiratory (frequency) variables averaged over 1024 points epochs (4.2 mn), in relation with the current sleep state.

## 4.2 Description of the global processing

### 4.2.1 ECG signal processing

From the ECG signal, we extract the heart rate. The different steps of the whole heart rate processing are represented in the Fig. 6, as they are implemented in LARY\_C.

The raw ECG is prefiltered with a Finite Impulse Response filter (FIR), in order to

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<sup>1</sup>TNI, Technopôle Brest-Iroise, ZI du Vernis, 29608 Brest Cedex (France)



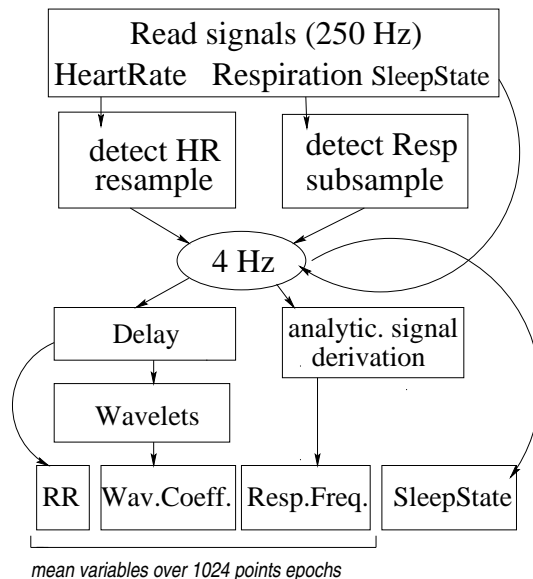


Figure 3: Flow chart of the parallel signal processing

derivate it and to reinforce the QRS complexes (Fig. 5). The detection is based on the correlation between the filtered ECG and the filter response whose shape is a derivated QRS: each time a detection occurs, the response is modified by the current QRS derivated complex on the ECG (eq. 3); the filter response is initialized with a derivated QRS shape and the forgetting factor  $a$  is fixed to 0.1 [27, 19].

$$\text{response}[i] = a * \text{response}[i - 1] + (1 - a) * QRS'[i] \quad (3)$$

An adaptative threshold detects the R peaks and returns the RR signal or heart rate; this RR signal is the serie of the distances between two R peaks, e.g. the number of ECG samples between two R peaks; this distance is usually translated in ms. This heart rate is in fact an unregularly sampled signal. An equidistant resampling [14] at 4Hz is applied (Fig. 6). Then an artefact detection is used, based on a clinical criterium of beat-to-beat variability: a RR can't be different from the previous one by more than 25%. An artefact so detected is replaced by the average of the  $n$  last valid RR. This method parameters have to be adjusted to each recording to make a compromise between the respect of the variability and the necessary detection of

the artefacts (Fig. 7).

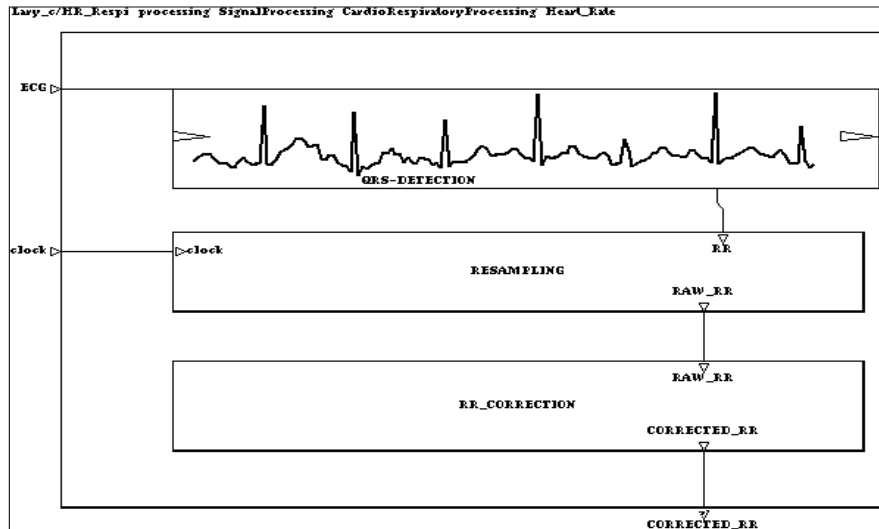


Figure 4: The modules of LARY\_C for Heart Rate processing, in the SILDEX graphical environment

*The 3 steps are the following ones:*

- *the QRS detection from the ECG to calculate the RR intervals or Heart Rate*
- *the Heart Rate resampling at 4Hz*
- *the detection and correction of the artefacts*

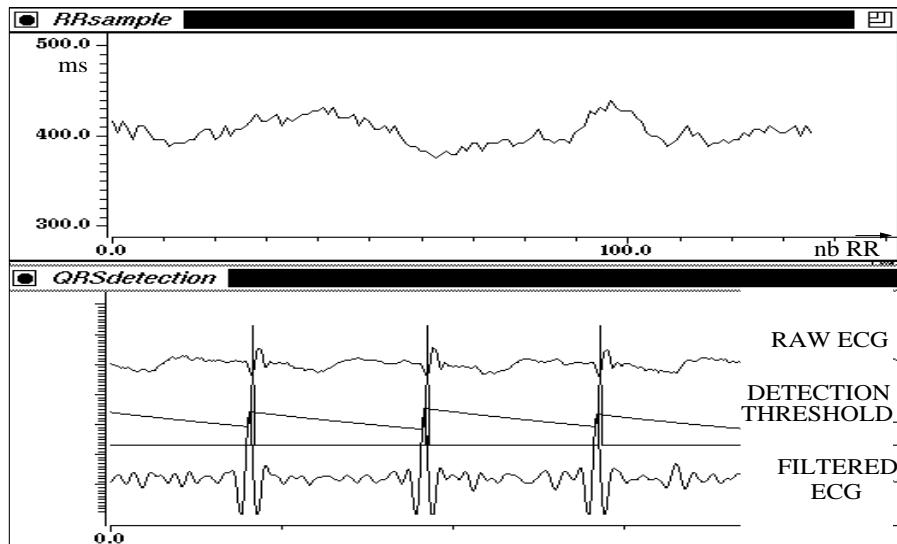


Figure 5: Illustration of the first step of the HR processing: the QRS detection. On the QRS detection screen (bottom), one can see the filtered ECG with the adaptive threshold; the vertical lines cutting the QRS complexes represent the events of detection; the distance between two detections (in number of ECG samples) is converted in ms and reported on the y axis of the RR signal screen (top).

#### 4.2.2 Respiratory signal processing

The instantaneous respiratory frequency is calculated by differentiation of the analytical signal (Fig. 8); this signal is obtained by a FIR-filter which computes the imaginary part (dephasing of 90 degrees); the domain where the filter is applicable is  $[.01-.49]$  normalised frequency. The frequency is the derivative of the arctangent of the analytical signal (Fig. 9). An illustration of the resulting signals is done in Fig. 10. We didn't take into account the amplitude of the respiratory (module

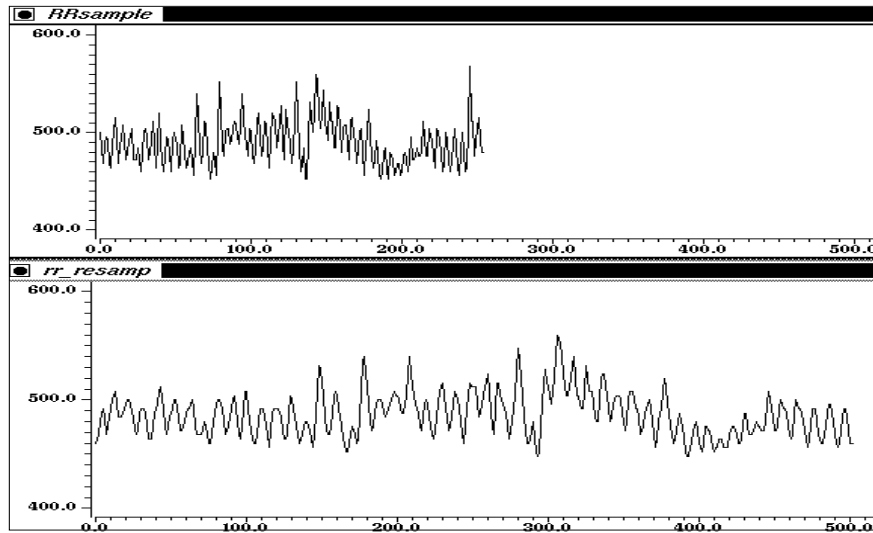


Figure 6: Illustration of the second step of the HR processing: resampling at 4Hz. The HR is expressed in ms; on the top, the raw RR signal, approximately around 2Hz (the mean RR is 500ms); on the bottom, the resampled RR signal; the x axis are expressed in number of RR samples.

of the analytical signal) because of the incertitude about the gains of the different recordings.

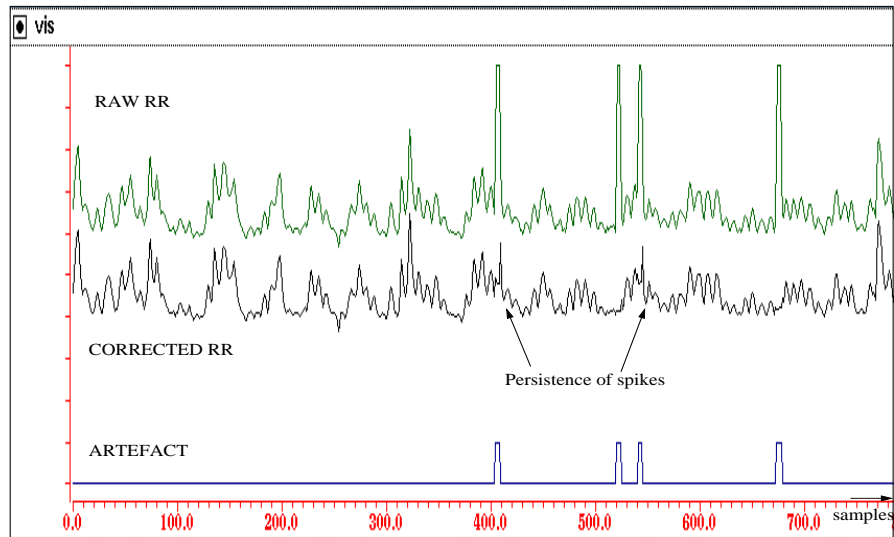


Figure 7: Illustration of the third step of the HR processing: detection and correction of the artefacts

*This screen shows the effects due to the compromise between the respect of the variability and the necessary detection of the artefacts; in this case, the detection threshold was fixed at a high rate not to smooth the real variability; but it induced the persistence of spikes.*

#### 4.2.3 DWT applied to HRV analysis

**Relations between wavelet coefficients and classical frequency bands** Table 1, Fig.11 and Fig.12 show the correspondence between DWC and the periods of physiological interest; c2 and c3 include the RSA, c4 and c5 the cardiovascular control; in particular, c4 closely contains the Mayer waves, corresponding to 10 heart beat periods.

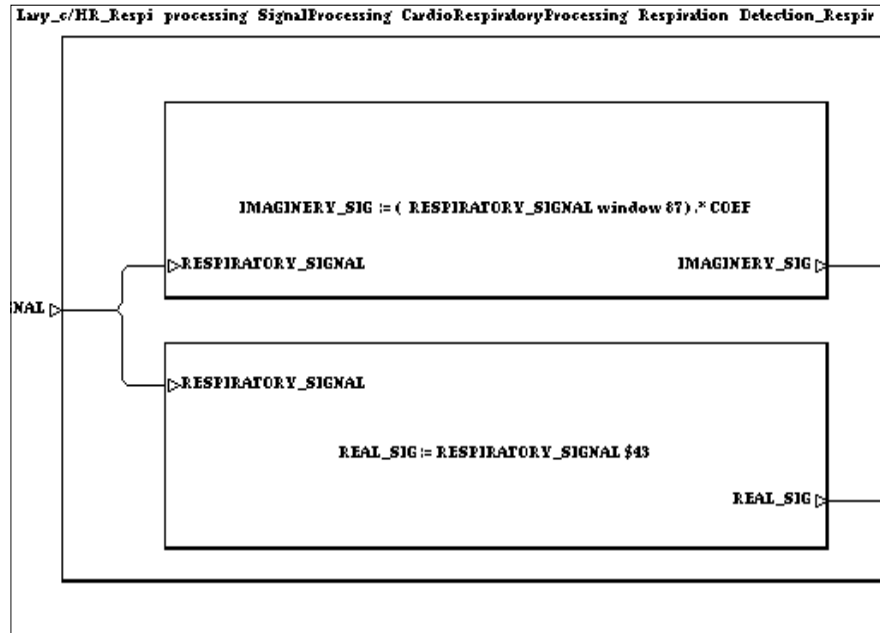


Figure 8: Respiratory signal processing: computation of the analytical signal in the SIGNAL language

*The imaginary part is computed by a FIR filter (dephasing of 90 degrees), with 87 coefficients; the delay induced by this filter is of 43 samples  $((n - 1)/2)$  and it can be compensated on the real part of the signal by the SIGNAL operator \$.*

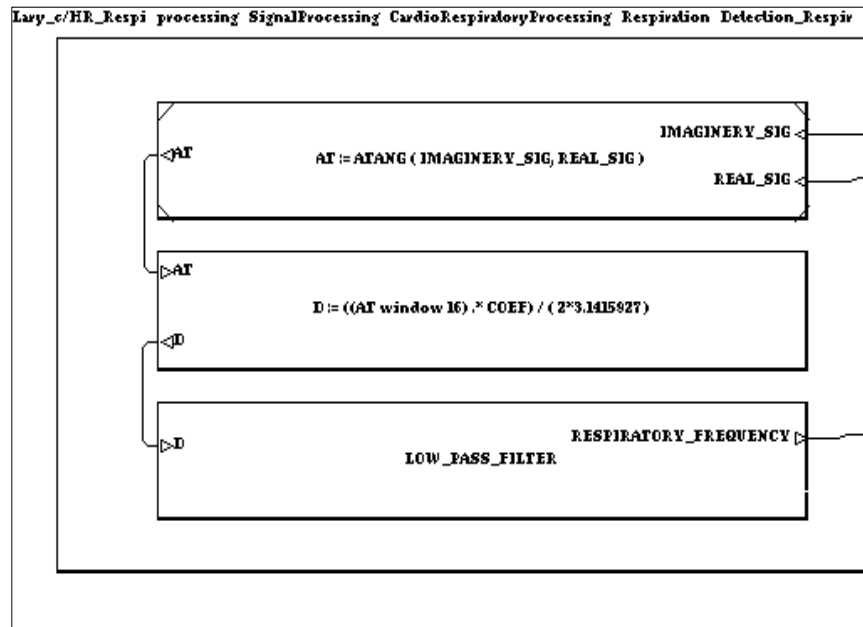


Figure 9: Respiratory signal processing: computation of the instantaneous respiratory frequency in the SIGNAL language

*The arc-tangent of the analytical signal is derived by a FIR, then low-pass filtered to produce the respiratory frequency.*

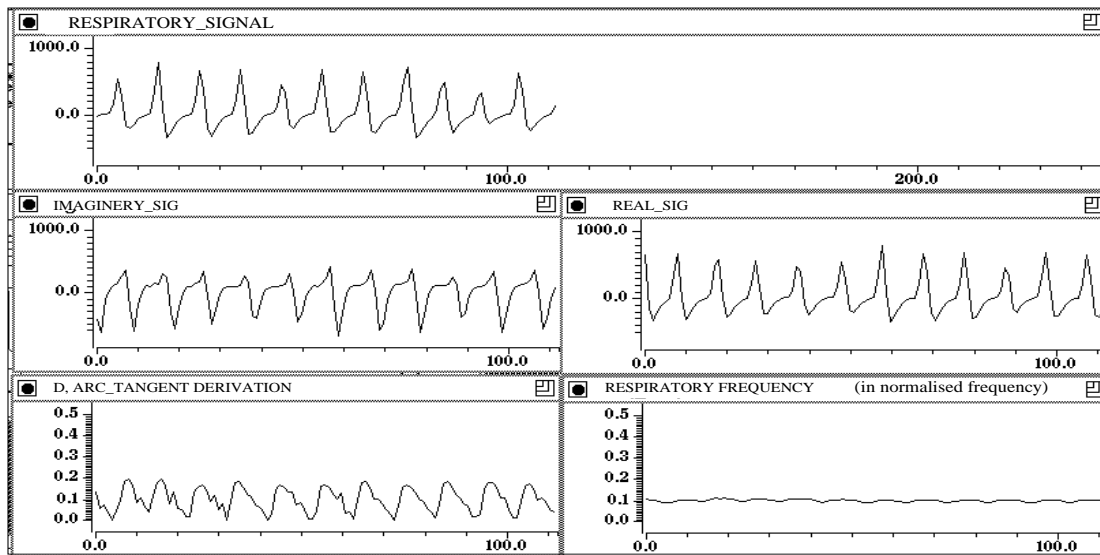


Figure 10: Respiratory signal processing: illustration of the different signals represented in the previous Fig.



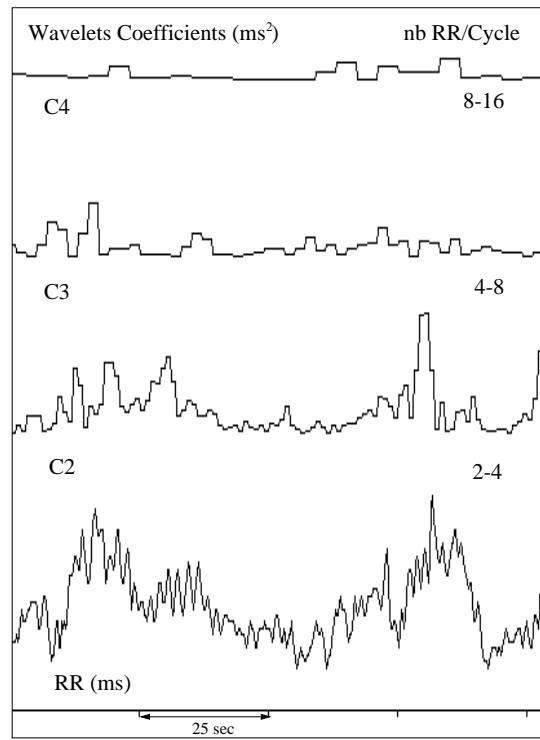


Figure 11: The first Discrete wavelet coefficients, corresponding to HF and MF

### 4.3 Some practical aspects

#### 4.3.1 Synchronisations between resulting rhythms

They were all resampled at 4Hz (Fig. 3) to be processed together and resynchronised because the different processing steps introduce delays; this was done by using filters with a linear phase, producing a constant time delay which can be exactly known. The Fig. 8 shows the details of implementation in the SIGNAL language. After resynchronisation, an averaging is done over the cardiac and respiratory variables; these variables are: heart rate, discrete wavelet coefficients, from C2 to C8, respiratory frequency. These means constitute the statistical units; they are done

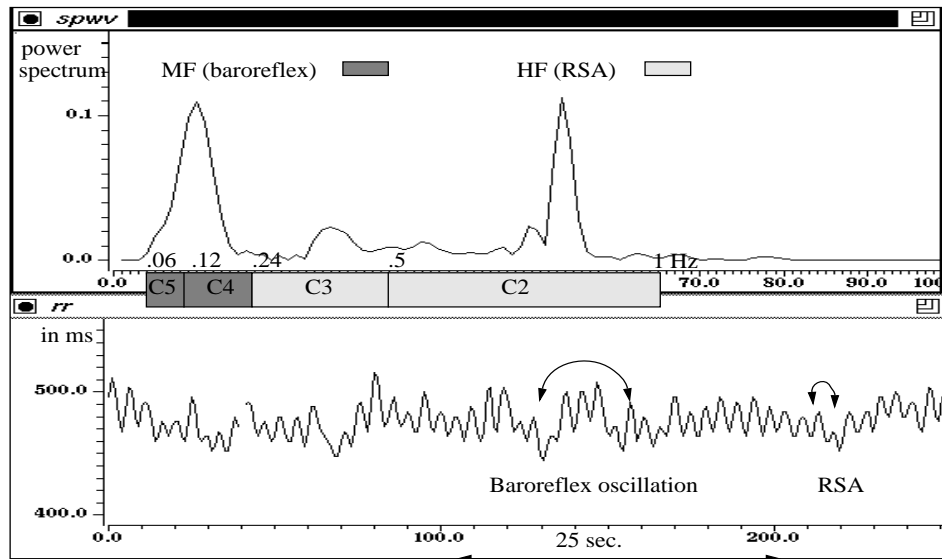


Figure 12: Correspondence between time and frequency representations of the heart rate

*A low-pass filter avoided the low frequencies to focus on the two frequency bands HF and MF; for this baby, the MF peak is centered on .16Hz, that corresponds to periods around 12 heart-beats; the HF peak is around .7Hz, that corresponds to periods around 3 heart-beats. The DWC are reported onto the spectrum to show the correspondence between the different representations.*

Table 1: Discrete wavelet coefficients and frequencies

DWC	N.Freq.	Freq. (Hz)	Period (nbRR)	Freq. Bands
c2	.125 .25	.5 .1	2 4	HF
c3	.062 .125	.24 .5	4 8	
c4	.031 .062	.12 .24	8 16	MF
c5	.015 .031	.06 .12	16 32	
c6	.007 .015	.028 .06	32 64	LF
c7	.003 .007	.012 .028	64 142	&
c8	.001 .003	.004 .012	142 285	VLF

N.Freq.:normalized frequency

each 1024 samples (about 4.2mn), which represent the window length for the wavelet processing; this size was chosen to get a good enough frequential resolution in low frequencies, according to the Heisenberg-Gabor's incertitude principle.

#### 4.3.2 Visual criteria

Table 2: Methods parameters

Parameters	Values
Sampling frequency on raw signals	250 Hz
Resampling frequency on resulting rhythms	4 Hz
Window length for means	1024 samples, 4.26 mn

Table 2 summarises the main parameters used in the signal processing chain. The respiratory frequency was detected in parallele with the DWC of the heart rate and averaged each 1024 samples; they are together displayed on the screen (Fig.13); this allowed us to follow the real respiratory frequency and to verify that it never slowed towards the Mayer waves [5]. For all the 30 recordings, the averaged respiratory frequency calculated from the raw respiratory signal is included between .35 and .7 Hz, that corresponds to C2 and C3. This visual inspection of the processing also allowed us to discard from the study the epochs with remaining artefacts,

except short spikes affecting only the first DWC as it is shown on Fig.14. These precautions and the length of the unit observation explain that in AS, where heart rate is very disturbed by artefacts, only 7 infants remain in each group compared to the (15 infants in each group).

## 5 Statistical processing

### 5.1 Classical hypothesis tests

They were performed on the variables averaged within one subject to assume independence of observations. We choose non parametric hypothesis tests, adapted for a few samples. Among them, the between-groups differences, control versus anemic, within the same sleep state were assumed by the unpaired Mann-Whitney'U test ( its parametric equivalent is the t-test unpaired). The between-state differences within the same group were assumed by the non parametric test of Wilcoxon which is a paired test, comparing differences within the same subject ( its parametric equivalent is the t-test paired).

### 5.2 Mutivariate Data Analyses (MDA)

We choose the most significant variables at the classical hypothesis tests to perform the following analysis: Principal Component Analysis (PCA), Linear Discriminant Analysis (LDA) and Factorial Discriminant Analysis (FDA). Their principle is based on the choice of the best linear combination of the chosen variables to describe a population and to explain the contribution of each of them in the between-groups discrimination.

For the Principal component analysis (PCA), as it is a purely descriptive method which makes no assumption on the statistical properties of the variables, all the 4 mn epochs were represented.

## 6 Clinical results

### 6.1 Between-group differences in Quiet Sleep

In QS (15 anemic v. 15 control), the addition of c2 and c3, corresponding to the global ASR, is significantly enhanced in control infants (Tab. 3, Fig 16). The DA, using the linear combination of HF(c2,c3) and LF(c7,c8) (c4,c5,c6 were not

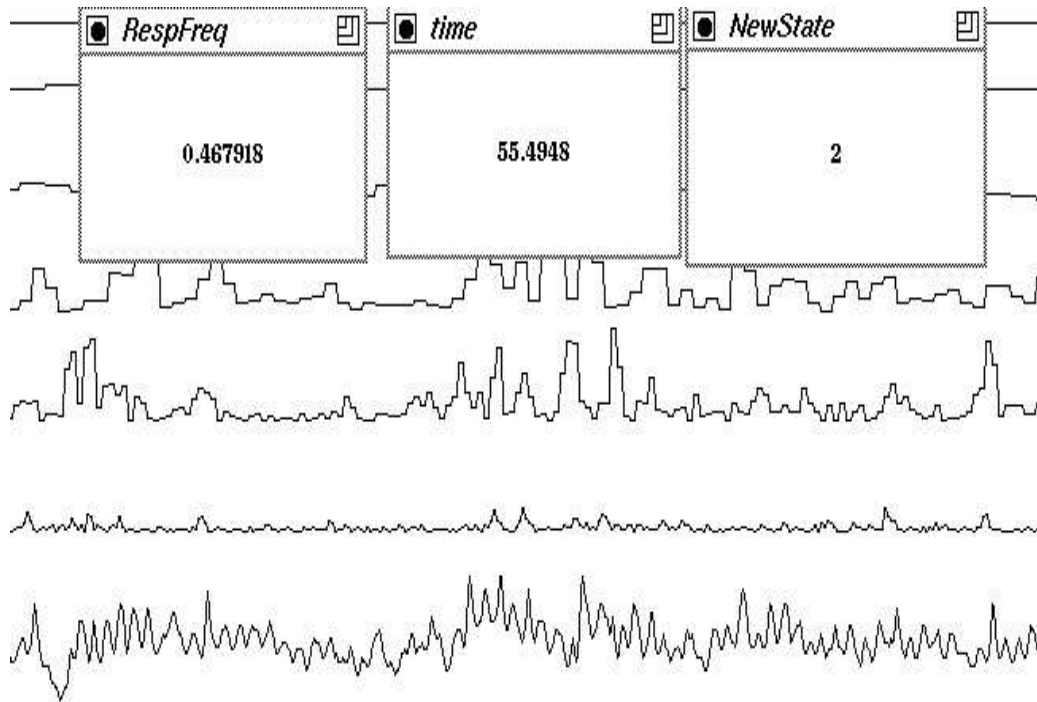


Figure 13: The Wavelet application on the Heart Rate during execution

*This view represents a part of the 1024 HR samples, unit of the wavelet processing; for each 1024 points epoch, the HR signal and its wavelet coefficients, the mean respiratory frequency, the current sleep state ( number 2 for QS) and the time (in mns) of the end of the epoch are displayed on the screen. So, it is possible to verify that C2 and C3 really contain the RSA.*

*From bottom to top, HR, C1 to C8; C1 was not of interest for this study.*

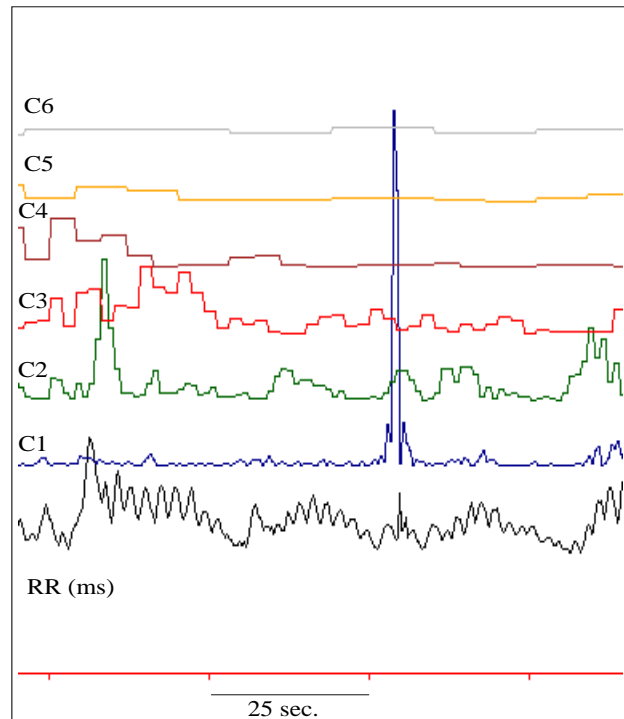


Figure 14: Effect of an HR artefact on the Discrete Wavelet coefficients  
*The spike on the HR is responsible for a very high frequency event which perturbs only the first coefficient; this one having no interest in physiological variabilities.*

relevant for discrimination) does'nt allow a very well discrimination (65% after cross-validation) but the correlation coefficient of the discriminant linear form (DLF, which provides the decision rule) shows the opposition between HF and LF (Tab. 4), as seen on the principal factor plane ( $> 80\%$  global variance) of the PCA. The first factor (x axis) represents the HF and LF powers with negative correlations whereas the second factor (y axis) represents the HF-LF opposition, with negative correlations for HF, positive for LF. Anemic and control are separated along these two axes.

Table 3: Between-group differences in Quiet Sleep

DWC ( $ms^2$ )	control		anemic		Mann-Whitney U test
	mean	StD	mean	StD	
c2+c3	77	68	50	65	$p \leq .03$ *
c2	33	28	22	21	$p \leq .07$
c3	44	46	28	46	$p \leq .04$ *

n:15 in each group; \*:  $p \leq .05$ 

Table 4: Correlation of DLF with DWC in Quiet Sleep

HF		LF	
c2	+.614	c7	-.143
c3	+.498	c8	-.404

## 6.2 Between-group differences in Active Sleep

In AS (7 anemic v. 7 control), c4, corresponding to baroreflex activity, is significantly enhanced in anemic infants (Tab. 5, Fig. 17). The DA using the MF(c4,c5) and a part of LF(c6) coefficients (the others are not relevant for discrimination), allows a discrimination of 78.6 %, with the strongest correlation of the DLF for c4 (-.93). No significant between-group differences were found in mean RR or in respiratory frequency, neither in AS nor in QS.

Table 5: Between-group differences in Active Sleep

DWC ( $ms^2$ )	control		anemic		Mann-Whitney U test
	mean	StD	mean	StD	
c4	6.2	3.9	12	6.7	$p \leq .04$ *

n: 7 in each group

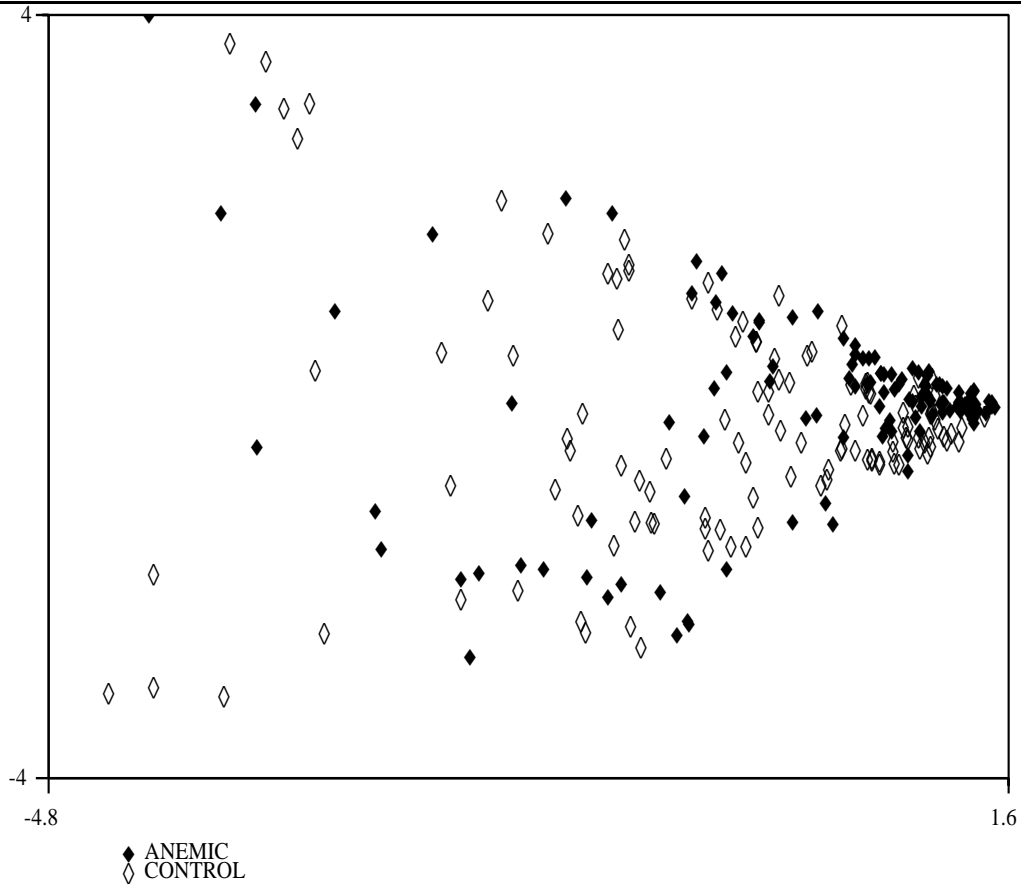


Figure 15: Principal component analysis in Quiet Sleep

Projection of all the 4mn HR epochs (125 control/125 anemic) onto the principal factor plane, according to the linear combination of HF ( $c_2, c_3$ ) and LF ( $c_7, c_8$ ) coefficients.

$$x = -.59 c_2 + -.64 c_3 + -.74 c_7 + -.71 c_8$$

$$y = -.67 c_2 + -.61 c_3 + .52 c_7 + .71 c_8$$

### 6.3 Between-state differences in each group

In the control group, QS and AS are significantly opposed by  $c_2$ , enhanced in QS and  $c_8$ , enhanced in AS (Tab 6). The DA, based on the HF( $c_2, c_3$ )/LF( $c_7, c_8$ ) coefficients



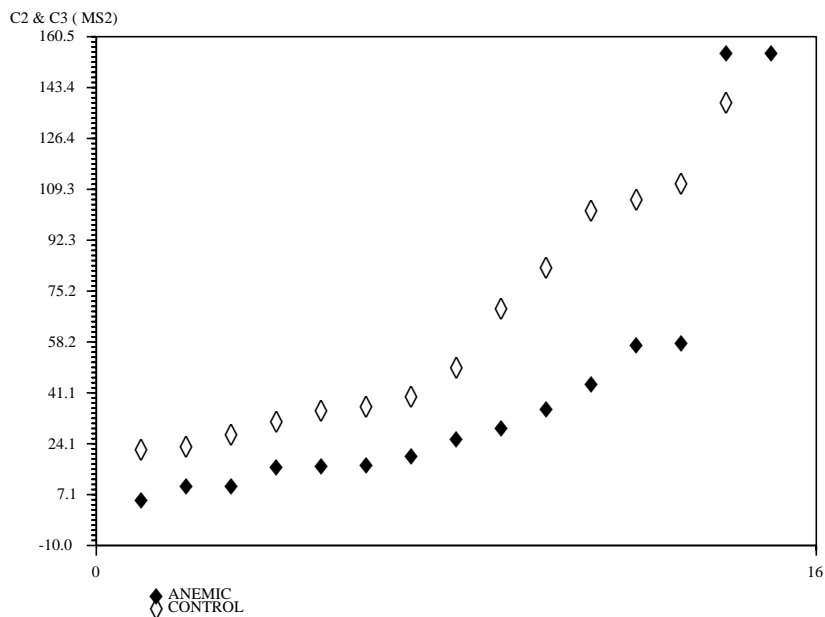


Figure 16: Between-group differences in Quiet Sleep: Discrete Wavelet Coefficients C2 and C3 corresponding to RSA

*The DWC of 15 anemic (black diamond) versus 15 control (white diamond) infants are plotted by growing order along the x axis; each point represents the mean value of the C2 and C3 coefficients for one baby.*

allows a discrimination of 82 %, and the correlation coefficient of the DLF with the DWC shows the HF/LF opposition (Tab. 8). The mean RR has a nearly significant enhancement in QS ( $p \leq .06$ , 495 v. 465ms). In the anemic group, neither significant differences (except almost for c4) nor conspicuous results in LDA appeared (Tab 6).

## 7 Discussion

### 7.1 About signal processing methods

Previous studies used other time and time-frequency methods: the Short Time Fourier Transform (STFT), applied to HRV of newborns [35, 34, 16]; the Smooth Pseudo

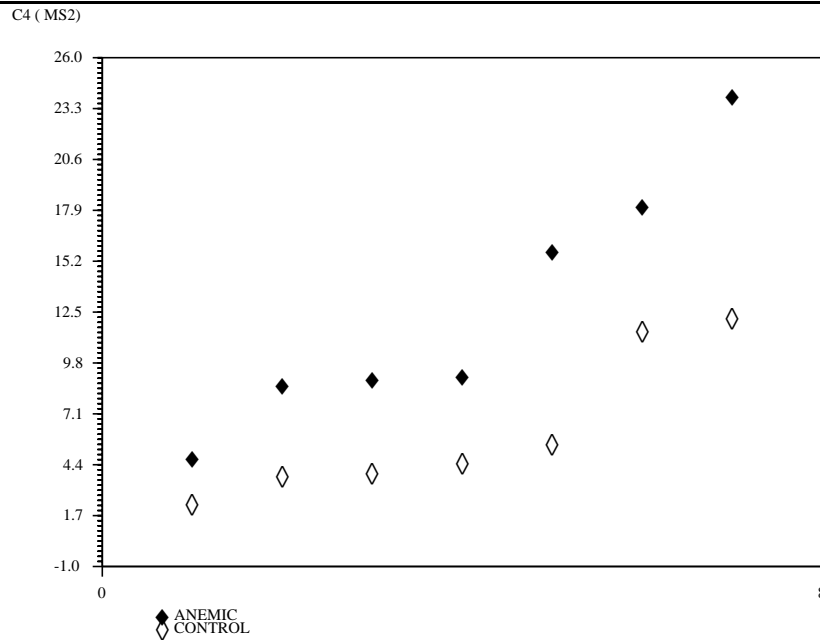


Figure 17: Between-group differences in Active Sleep: Discrete Wavelet Coefficient C4 corresponding to baroreflex activity  
*The DWC of 7 anemic (black diamond) versus 7 control (white diamond) infants are plotted by growing order along the x axis; each point represents the mean value of the C4 coefficients for one baby.*

Table 6: Between-sleep-states differences in control infants

DWC ( $ms^2$ )	AS		QS		Wilcoxon test
	mean	StD	mean	StD	
<b>c2</b>	<b>17</b>	<b>12</b>	<b>27</b>	<b>17</b>	$p \leq .02$ *
c4	6.2	3.9	7.0	4.0	NS
<b>c8</b>	<b>.31</b>	<b>.20</b>	<b>.14</b>	<b>.05</b>	$p \leq .02$ *

n=7; NS: non significant; \*:  $p \leq .05$

Wigner-Ville Transform (SPWVT) applied to HRV of transgenic mice and Periodic

Table 7: Between-sleep-states differences in anemic infants

DWC ( $ms^2$ )	AS		QS		Wilcoxon test
	mean	StD	mean	StD	
c2	25	20	30	28	NS
c4	12.6	6.7	7.8	6.2	$p \leq .06$
c8	.47	.48	.25	.23	NS

n=7; NS: non significant; \*:  $p \leq .05$

Table 8: Correlation of DLF with DWC in control infants

HF		LF	
c2	+.442	c7	-.271
c3	+.388	c8	-.680

Leg Movement Syndrome [31, 30, 33]; the cardio-respiratory Modified Complex De-modulation (MCD) [29, 28, 32, 33].

With SFT and SPWVT, the power spectrum is divided into several frequency bands, based on a physiological interpretation; the previous studies defined HF as periods from 2 to 8 heart-beats, MF from 10 to 25, and LF from 30 to 100. On the opposite, with the DWT used here, we try to give a physiological explanation to pre-fixed dyadic coefficients. So, some differences appeared (Tab.1), especially a very precise frequency band around Mayer Waves (8-16 heart-beats).

STFT doesn't allow a very good frequential definition, especially in low frequencies; SPWVT is more precise but interferences inherent in the method, compel us to avoid low frequencies and to limit the study to HF and MF. The MCD was used to study the HF only, in following the respiratory influence on the heart rate.

With DWC the frequential definition is well suited to low frequencies; with regard to this point, the window length of 1024 samples was chosen to get a good enough frequential resolution, according to the Heisenberg-Gabor's incertitude principle.

## 7.2 About Statistical methods

Even if great care is necessary to interpret the results, especially in active sleep, the same statistical methodology was already used for published studies [16, 34, 35];

they involved few individuals (8 full-term newborns, 8 intermediate, 8 premature). Concerning this study, 15 control versus 15 anemic infants in quiet sleep, 7 control versus 7 anemic infants in active sleep were the subject of the tests. The non-parametric tests are suitable for few samples and/or no Gaussian distribution; from six-seven individuals, the equivalence tables with parametric tests give reliable results. The discriminant analyses theoretically assume normality but are robust enough to tolerate violations of this normality assumption [36, 37].

We fixed the data window length at 1024 samples (about 4mn), which appeared to us as a good duration for an observation unit to take within subject variance into account. This variance effect is well shown by Principal Component Analysis, which uses all the epochs; it is made possible because the PCA is purely descriptive and makes no assumption on the statistical properties of the variables. On the opposite, the hypothesis tests were performed on the means over the epochs within one subject in a given sleep state to guarantee the independance of the observations.

### 7.3 About clinical results

We will try to explain the results of this study, particularly why the differences found between anemic and control infants don't involve the same neuro-vegetative variables in quiet and in active sleep.

In quiet sleep, anemic infants have a lower parasympathetic tone than controls, which confirms a previous study [32] and a not yet published work by P. Peirano. In active sleep, anemic infants have a greater sympathetic stimulation, in the range of the Mayer waves, than the control ones. This result is not yet confirmed (but not invalidated) by the previous study of P. Peirano. An explanation can be the very precise frequency definition around the Mayer Waves (from 8 to 16 heart-beats), given by the wavelet transform whereas the previous study used a larger frequency band (from 10 to 25 heart-beats). Nevertheless, the data are not exactly the same and the results remain to be confirmed on a larger population. In control infants only, results evidence the different levels of the autonomic balance according to sleep states: dominance of the parasympathic tone in QS versus the sympathetic one in AS.

To each sleep-waking state corresponds a particular physiologic state [38, 39]. A reciprocal interaction exists between the mechanisms which control the sleep-waking states and those which control the vegetative functions (“Carotide” comes from “to doze off” (karoun in greek), illustration of the hypnogenic effect of the baroreflex regulation).

### 7.3.1 Influence of the sleep on the neurovegetative functions

The QS is a state of very low metabolic consumption, with a maximal stability of the vegetative functions. The SA is a state under a “behavioral” control, motor perturbations (body or facial movements) modulate the respiratory and baroreflex control. In AS especially, two patterns, which can be evaluated on chin muscle [40] are considered; the tonic pattern is related to the muscle tone whereas the phasic one is related to the motor activity. In AS, muscle tone is often inhibited, giving way to a phasic activity.

**Sleep and respiratory control** During QS, the breathing is mainly under metabolic control, through chemoreceptors (central and peripheral), integrated in bulbo-protuberantial centers. Cortical connections are inactivated and few motor perturbations disturb the respiratory control during this state.

During AS, the breathing is mainly under control of a specific system, called “AS-state”, where tonic and phasic systems coexist. Some specific bulbo-protuberantial AS-neurons (non respiratory neurons) are supposed to change the activity of the spinal respiratory neurons, through tonic impulses. The phasic system modifies this background activity to adapt the respiratory control to the current movements. There is an inhibition of the hypothalamic control mechanisms and an exclusive influence of these brainstem neurons. This inhibition is thought to be responsible for a loss in the efficiency of the homeostatic regulations during AS [38].

These differences in the regulation mechanisms could explain why the differences in RSA are expressed in QS and not in AS: QS is the state where the bulbo-protuberantial interaction between the respiratory center and the vagus center can be the strongest, producing the RSA (vagal inhibition synchronised with firing of inspiratory motoneurons), whereas in AS these interactions are permanently modified. Moreover, the QS is supposed to be particularly vulnerable in the first year of life because of the transition from one to four stages [9], during this particular epoch of human ontogenesis.

**Sleep and cardiovascular control** During QS, there is a high cardiovascular stability, with a slower heart rate due to the negative chronotropic effect of a dominant parasympathetic tone and a subsequent diminution of the arterial blood pressure without change in peripheral resistances.

During AS, the cardiovascular regulation is more important, fluctuations of arterial blood pressure and heart rate often arise, according to phasic events. These fluctuations can be explained by the opposed effects of the tonic and phasic states:

during tonic AS, there is a global diminution of heart rate, arterial blood pressure and peripheral resistances, through a strong central sympathetic inhibition. On the opposite, during phasic AS, an increase in the vasoconstriction in skeleton muscles through an increase in sympathetic tone gives an increase in heart rate, arterial blood pressure and peripheral resistances.

Some experiments of sino-aortic denervation in the cat (Guazzi and Zancheti, cited in [38]) show the important role of the baroreflex in the cardiovascular control during AS: especially the phasic events give a very high instability of the arterial blood pressure, sometimes incompatible with a correct cerebral circulation. It seems that baroreflex are essential in cardiovascular regulation during AS and that the Nucleus Tractus Solitarius, first relay and integration center, is a very important structure for this homeostatic balance.

These differences in the regulation mechanism could explain why the differences in baroreflex activity between control and anemic infants are expressed in AS and not in QS: AS is the state where the baroreflex is the most evidently solicited, and so an abnormality can be better seen in AS than in QS.

One can make the assumption that anemic subjects show a delay in maturation of the cardiovascular control, with an instability of the baroreflex response, responsible for a higher activity in the range of the Mayer waves. However, as this higher activity could be only consecutive to an increase of the motor activity in the anemic group, we can't really conclude in favour of an abnormality but only in a higher solicitation of the baroreflex regulation.

### **7.3.2 Hypothesis on mechanisms underlying iron deficiency**

Iron is involved in lipid metabolism and is thought to have a role in production and maintenance of myelin, as it has been shown in the rat, for which a restriction in iron leads to poor myelination [41]. This defect, which alters neurotransmission, could affect parasympathetic fibers, entirely and late myelinated in postnatal period, and the sympathetic fibers myelinated on their pre-ganglion traject [42, 43]. It could explain both the effect on the RSA in QS and the effect on the baroreflex regulation in AS.

The results suggest a delayed or abnormal development of the ANS in anemic infants: smaller parasympathetic tone in QS, stronger baroreflex excitability in AS. Furthermore, the poor differentiation of heart rate variability parameters according to sleep states in the anemic group reinforces this hypothesis. A reduced SNA activity, especially of the vagal tone, could affect infants in an important way, in so far as ANS integrity is essential for a correct organization of physiological resources, for

an appropriate response to stress, for a good interaction of cardiovascular and respiratory control with sleep-waking states. This could contribute to poorer behavioral and developmental outcomes in anemic infants.

## 8 Conclusion

The use of the DWT is well suited for its good frequency resolution in low frequencies. Yet, it provides only a dyadic decomposition, which is inherent in the discretisation process; one might get better accuracy on the physiological frequency bands by performing continuous wavelet transform.

This study allows us to point out some physiological results which add further evidence to a delayed or abnormal ANS development in anemic infants. This suggests that iron is an essential micronutrient for the normal development of the autonomic control system.

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