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# Development of a Prototype for the Analysis of Multiple Responses of the Autonomic Nervous System

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#### Abstract

A modular hardware prototype is developed for the noninvasive acquisition, processing and transmission of biological signals to analyze the autonomic nervous system (ANS) in synchrony with the video recording of the pupil. The implementation includes 1) two noninvasive sensors, a pulse oximeter and an electrodermal activity sensor, 2) a module able to collect the information and send it to the PC via USB and 3) a graphic user interface (GUI) for visualization, synchronization and data saving. A series of experimental tests were performed to investigate the effect of different stimulations: light, dental occlusion, transcutaneous electrical nerve stimulation (TENS) and mental efforts. They indicate the reliability of the system and the importance of the joint detection of more signals for discriminating different states of the ANS. Specifically, heart rate, Galvanic response and pupil size were compared, showing some coherence in their oscillations and different discrimination capability in different conditions. Their joint detection is thus important for discriminating different states of the ANS.

*Keywords:* Pulse oximetry, Electrodermal Activity (EDA), Galvanic Skin Response (GSR), Pupillogram, Autonomous Nervous System (ANS), Transcutaneous Electrical Nerve Stimulation (TENS), Embedded systems.

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#### 1 1. Introduction

The autonomic nervous system (ANS) is the portion of the central nervous system that controls unconscious activities, such as visceral functions and homeostasis. It is divided into two main branches, the sympathetic and the parasympathetic, the first promoting the activation of a physiological response and the other inhibiting it. The ANS is profoundly affected by motions and somatosensory inputs and plays an important role in pain and stress modulation and perception.

Autonomic testing finds application in the clinical assessment of neurological disorders, particularly those affecting predominantly small nerve fibres [1]. Many studies have been devoted to the quantitative assessment of the ANS response, dating back to more than 3 decades [2][3][4]. However, most of the literature takes into account just one of the numerous physiological systems that are affected by the ANS in turn.

Very frequently the works that have dealt with ANS in different disorders 15 related to its dysfunction have focused on cardio-circulatory parameters [5]. 16 However, there are many other potential peripheral effects of ANS, that have 17 been largely overlooked. For example, a physiological system related to the 18 ANS can be investigated measuring skin conductance, reflecting the sweat-19 ing of the sweat glands [6]. Moreover, pupil is strongly affected by the ANS 20 [7]. The study of mydriasis and myosis (i.e., the dilation and contraction of 21 pupil) is usually done in a different context (vestibular system). However, 22 recent studies have indicated the possibility of characterizing the condition 23 of the ANS in healthy or pathological conditions, by analyzing the nonlinear 24 pupil oscillations [8][9][10][11][12][13]. In particular, the study of pupillary 25 dynamics was considered useful for evaluating the arousal state during men-26 tal effort due to cognitive tasks [14][15] and in relation to the involvement 27 of reward systems [16]. Furthermore, it has been suggested that the use 28 of different ANS parameters, including pupil size, may be useful for better 29 characterizing and quantifying the emotional component linked to the au-30 tonomous response [17]. Moreover, further evidence was provided that pupil 31 can be used to evaluate the state of emotional arousal as well as the generic 32 activation of the ANS [18]. 33

One of the limitations of the study of various ANS responses is the use of different instruments for the analysis of different signals. However, it was argued that pupillogram could provide useful information for the study of the arousal state and that it would receive a valid contribution from the association with signals already used for this purpose, such as skin conductance and electrocortical activity [19]. These observations would involve an important expansion of the combined and synchronous study of various parameters associated with pupillography to aspects not only of pathology, but also related to the emotional / affective state. Thus, important outcomes could be expected both for clinical patients and for any study involving the psychic assessment of the arousal state.

Only recently, more reactions of the ANS have been investigated simul-45 taneously [20][21]. The signals that have been often analyzed are the cardiac 46 pulse and the variations of skin conductance. In this study, we are inter-47 ested in the investigation of those signals jointly with the pupil response. 48 A modular hardware prototype is developed for the noninvasive acquisition, 40 processing and transmission of biological signals and is synchronized with 50 a commercial system for pupil investigation. Our present implementation 51 includes: 52

- two noninvasive sensors, a pulse oximeter and an electrodermal activity sensor (both often used in the study of the responses of ANS [22][23][24]);
- a module able to collect the information and send it to the PC via USB;
- a graphic user interface (GUI) for visualization and data saving.

Pulse oximetry and skin conductivity are acquired in synchrony with the video recording of the patient's dilation and constriction of the pupil (acquired by a commercial system [25]). The system was developed keeping low the production cost and energy consumption. It was tested in experiments from healthy subjects under different stimulations: light, dental occlusion, transcutaneous electrical nerve stimulation (TENS) and computational task.

# <sup>64</sup> 2. Design and Implementation

The recording system is shown in Figure 1. Two sensors, described below, are developed and are used together with a commercial system for pupil investigation [25].

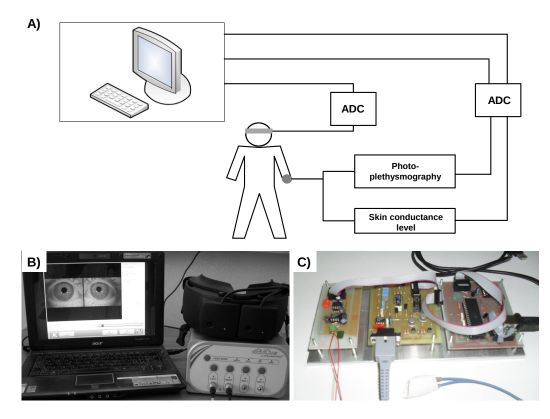


Figure 1: A) General scheme of the instrumentation used. B) Commercial system for pupil investigation [25]. C) Acquisition system recording photoplethysmogram and skin conductivity level.

## 68 2.1. Design of the sensors

#### 69 2.1.1. Pulse oximeter.

The first sensor measures the absorption of red and infrared lights that 70 pass through a patient's finger by light sensors. The light is generated with 71 2 LEDs that are controlled alternately. A photodiode receives both ambient 72 and modulated light from the LED and generates a current that is related 73 to the oxygen saturation and the cardiac frequency [26][27][28]. The pulse 74 oximetry signal (photoplethysmogram - PPG) has a frequency range between 75 1 and 10 Hz. The signal is affected by the line interference (50 Hz) and 76 the neon lamps that are usually used in offices and laboratories (giving an 77 interference at 100 Hz), which were removed by notch filters; moreover, a 78 high pass filter attenuated motion artifacts (notice that high performance 79 professional oximeters are stable to ambient light and motion artifacts [29], 80 but our low cost system had to rely on simple solutions). For the actual 81 implementation, the sampling frequency of the PPG is 220 Hz. The circuit 82 design is shown in Figure 2A and B. 83

#### <sup>84</sup> 2.1.2. Skin conductivity level sensor.

There are two skin resistance responses due to the electrodermal activity (EDA): the tonic and the phasic levels. The tonic level is the absolute level of resistance at a given moment in the absence of a measurable phasic response and is referred to as Skin Conductance Level (SCL). The phasic level or Skin Conductance Response (SCR) is superimposed on the tonic level and corresponds to the response to stimuli. The sum of the tonic and phasic response is the Galvanic Skin Response (GSR).

The skin conductance is obtained by measuring the current flow through 92 the skin in response to a constant applied voltage. The circuit design is shown 93 in Figure 2C. For bipolar recordings, 0.5 V is recommended [30]. For a reso-94 lution of 0.01  $\mu$ S (the minimum variation of the SCR [31][32]), the minimum 95 current to be amplified is of 5 nA. The maximum input current of the am-96 plifier is 15  $\mu$ A (maximum SCR and SCL), therefore the transimpedance of 97 the circuit must be of 333 k $\Omega$ . This circuit has two independent outputs that 98 allow the analysis of each component separately. The sampling frequency is 99 170 Hz. 100

# 101 2.2. Digital processing of the signal and transmission

The information coming from the analogue sensors was converted into digital form, acquired, pre-processed and transmitted via USB to a PC. The

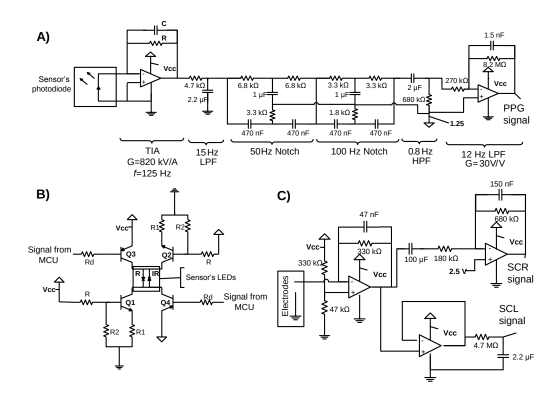


Figure 2: A) Schematic of the photoplethysmograph. B) Schematic of the LEDs driver. The signal from the microcontroller unit (MCU) controls which of the 2 LEDs is ON. C) Schematic of the skin conductance sensor. The 2 outputs are for the Skin Conductance Response (SCR) and the Skin Conductance Level (SCL).

microcontroller Atmel ATmega 16 accomplished these tasks [33][34]. Such a 104 microcontroller was chosen as it is cheap and it has a high speed, a small code 105 size and an analog multiplier (which can be used to implement digital filters). 106 Its most important characteristics are listed in Table 1. The connection of 107 the microcontroller to the system is shown in Figure 3. Notice that the 108 internal 10 bit ADC was used for sampling the skin conductance, whereas an 109 external 16 bits analog-to-digital converter (ADC) was included to sample the 110 PPG (the considered external ADC is AD7715, which is a 16-bit Sigma-Delta 111 ADC that can be interfaced with microcontrollers using the Serial Peripheral 112 Interface, SPI). 113

The red and infrared lights for the pulse oximetry sensor must be ON in 114 different cycles: the signal period was 1 ms, with a duty cycle of 0.25 and 115 the phase between the two lights was 0.5 ms. The state of the LEDs was 116 controlled using timer interrupt. An average filter (of 5 samples for pulse 117 oximetry and 20 samples for the Galvanic skin response) was implemented in 118 the microcontroller to remove experimental noise from the recorded signals 119 (the sampling frequency was about 5 kHz, so that there was sufficient time 120 to process the acquired data to produce a low pass filtered sample to be 121 acquired). The filtered samples were then transmitted to the PC with the 122 USART, which was interfaced with a USB transceiver.

D /	37.1
Parameter	Value
Maximum Speed	16 MHz
Operating voltage	2.7 V - 5.5 V
Power consumption	Active: 1.1 mA
	Idle: 0.35 mA
	Power-down: $< 1 \ \mu A$
Flash Memory	16 KB
EEPROM	512 B
Internal SRAM	1 KB
ADC Channels	8
ADC resolution	10 bits
ADC speed (max. resolution)	200 ksps
Timers max. resolution	16 bits
Serial communication	USART
	SPI
	Two-wire Serial

Table 1: General properties of the Atmega16 [33].

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<sup>124</sup> To manage the data, a software interface was developed in Visual Basic,

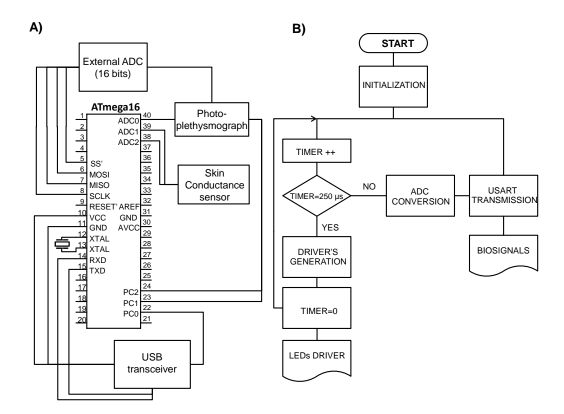


Figure 3: A) General schematic of the digital processing unit. B) Microcontroller general flow diagram.

- <sup>125</sup> with the following main functions.
- Registration of patient's information. The user should also be able to import the information from the database of pupillograms.
- Visualization of the signals. The charts update constantly to show the signals while they are being acquired.
- Storage of the signals. The information was stored in a file. This file contains the patient's information, the starting time of the acquisition, the time of each sample and the samples of the signals.

#### 133 3. Experimental tests

#### 134 3.1. Signal acquisition

#### 135 3.1.1. Instrumentation.

Images of the pupils were acquired by the Oculus system (Inventis srl, Padova, Italy), using two infrared CCD cameras (resolution 720x576 pixels, 256 grey levels) mounted on a light helmet (1.5 kg), with sampling frequency of 25 frame/s. The eyes were illuminated with an infrared diode with 880 nm of wave-length; moreover, during experiments on pupil dynamics under constant light conditions, illumination was provided by a yellow-green LED with 740 nm of wave-length.

Pulse oximetry and variations of sweat were monitored with the system described in the previous sections, considering the PPG and the GSR time series.

#### <sup>146</sup> 3.1.2. Synchronization of the signals.

The software of the considered system for pupil investigation [25] does not allow any kind of modifications or access. Therefore, the synchronization of the signals was done by registering the precise moment in which the system started recording. The signals were synchronized with a resolution of 10 ms. Given that the sampling period of the pupillogram was 40 ms, this resolution is enough.

# 153 3.1.3. Experimental set up.

The subjects were sitting in a high-back chair. The environment was kept at a constant temperature of 21°C. Visual predominance was determined. The acquisition system was then connected to the patient's non dominant hand. The helmet was applied and was maintained until the end of the recording. This phase took about 4 minutes.

The correct procedure and execution of tests was first explained to the subjects. Then, they were asked for brief tests to make sure that the instructions were well understood. This phase took about 2 minutes.

Two operators worked within the experimental set. The first took care of the subject (pretest and test instructions, helmet handling, check of the correctness of execution), the second controlled hardware and software.

#### 165 3.1.4. Experiments.

One minute long acquisitions were obtained (after approval by the Inter-166 nal Review Board of Politecnico di Torino) from 8 young, healthy subjects 167 (age  $25.1 \pm 1.1$  years; 6 females, 2 males) in different stationary conditions 168 (separated by 5 minutes rest), which require a different involvement of the 169 sympathetic and of the parasympathetic control: neutral position of the jaw 170 (rest position: RP) and habitual dental occlusion  $(HDO)^1$ , in light or dark-171 ness condition; moreover, a test was performed during a computational task, 172 which was assumed to induce a detectable stress of the subject. These 5 173 conditions (RP in light and darkness, HDO in light and darkness and com-174 putational task in darkness) were performed before, during and after the ap-175 plication of low-frequency TENS (5 minutes duration), which was expected 176 to induce relaxation. 177

#### 178 3.2. Signal processing and results

# 179 3.2.1. Pre-processing of recorded data.

Pupillometric recordings were processed through the algorithm of strongly connected components [35] to measure frame by frame the area of the pupil, expressed as the number of pixels covering it. The area of the pupil was then low pass filtered under 2 Hz (non-causal, zero-phase, Butterworth filter of order 2).

The local maxima in the PPG were used to identify the heartbeats, from which the heart rate (HR) was estimated. The GSR was low pass filtered under 2 Hz.

The mean and standard deviation (STD) were computed for the three 188 following signals: pupillogram, HR and GSR. Moreover, the linear trend was 189 estimated as a basic indicator of the evolution in time of the signals (the trend 190 was defined as the slope of the interpolation line of the data after scaling the 191 time to range between 0 and 1 and normalizing the time series to have zero 192 mean and unit STD). These indexes were used as descriptors of the signals in 193 the different experimental conditions. The two-sided Wilcoxon signed rank 194 test (considering paired data) was then applied to investigate differences of 195 each of the indexes in specific pairs of conditions of interest, after pooling 196

<sup>&</sup>lt;sup>1</sup>During dental occlusion, the effect of muscle fatigue and the massive involvement of the autonomic system were excluded by avoiding prolonged teeth clenching. Subjects were asked to swallow and then to contact the teeth lightly without clenching. Attention was paid to check the activity of mimic muscles.

<sup>197</sup> data: RP in darkness compared to the computational task, RP compared to <sup>198</sup> HDO (in darkness), pre-TENS compared to TENS or post-TENS conditions. <sup>199</sup> The significance level was set to p < 0.05.

200 3.2.2. Results.

Examples of recorded signals are shown in Figure 4. The pupil shows an 201 irregular oscillatory behavior, as also the HR. There is a decreasing trend in 202 both HR and GSR, as if the subject was relaxing. The figure shows also the 203 spectral coherence of the pupillogram and the HR at frequencies lower than 204 1 Hz. The two signals were found to be coherent in subjects under control 205 breathing conditions, where a respiratory component was visible in both the 206 pupillogram and the HR [7]. Here, the considered normal breathing and the 207 short acquisitions resulted in significant coherence (over 0.5) only in a few 208 subjects and conditions. 209

The significance of the differences of indexes extracted from the signals recorded in different conditions is shown in Table 2. Different indexes have a greater discriminatory value comparing different conditions:

- an index from the HR showed the maximal significance (i.e., minimum p value) in discriminating computational task and RP in darkness (which can be considered as a rest state);
- indexes estimated from the pupillogram had the maximal significance
   in distinguishing light and darkness, RP and HDO, or pre-TENS and
   TENS conditions;
- indexes extracted from GSR were the most statistically different in the conditions pre-TENS versus post-TENS and TENS versus post-TENS.

Specifically, HR and mean pupil size increased when comparing computa-221 tional task with RP in darkness (due to the mental stress induced by the 222 task). The subject started sweating during the computational task, as indi-223 cated by the high positive trend of GSR. Pupil was the only system showing 224 significant differences between HDO and RP: pupil size increased as a result 225 of HDO stimulation [8][9][10]. Moreover, it was the only system showing 226 significant differences comparing light and darkness conditions (obviously, 227 increasing the diameter in darkness). 228

Pupil also indicated the relaxation induced by TENS (there is a significant reduction of pupil size during and after the application of TENS). On the

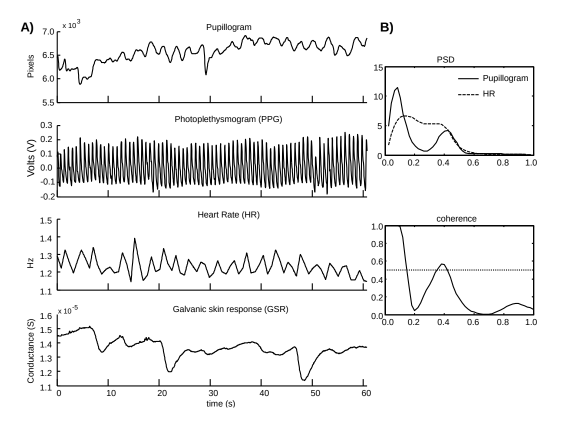


Figure 4: A) Example of data (pupillogram, PPG from which the HR is obtained, GSR), with the subject at rest position of the jaw in darkness. Power spectrum densities (PSD) of pupillogram and heart rate (using Welch's overlapped segment averaging estimator, considering 8 segments with 50% overlap) and magnitude squared coherence.

other hand, GSR increased during and after TENS, with respect to the preTENS condition. Possibly, this was due to an accumulation of sweat during
the experiment, as the sensors were kept fixed for all its duration.

# 234 4. Discussion

The ANS controls many different visceral functions, including heart rate, perspiration, digestion, salivation, respiratory rate, pupillary dynamics, micturition (urination), sexual arousal, breathing and swallowing. The joint acquisition of different autonomic responses could be useful for a deeper insight into ANS physiology and pathology. For example, the study of the autonomic response is important in the following situations [36]:

- sympathetic and parasympathetic lesions after surgical procedures;
- drug's collateral effects;
- diagnosis and follow up of ANS diseases;
- poisoning;
- involuntary reactions of the patient.

The diagnosis in most of these situations is currently vague [37][38][39]. 246 The joint investigation of different autonomic responses could help in clar-247 ifying the complex dynamics of the ANS in such conditions. One of the 248 problems when designing an experimental setup is being constrained by the 249 functionalities of commercial systems, which are usually closed and allow only 250 specific protocols. Here, we were interested in investigating synchronously 251 the joint responses of the heart (i.e., the cardiac pulse), the skin (i.e., sweat 252 production) and pupil. Despite the interest in the study of pupillary dy-253 namics, at present there are no tools that are able to simultaneously obtain 254 the acquisition and processing of pupillary dynamics and those of other sys-255 tems such as the cardiovascular (e.g., through pulse oximetry) and sudorific 256 system (through skin conductivity). Being able to expand the analysis to 257 several systems in a synchronous way would allow a better knowledge of the 258 coupling pathways between somatosensory and affective / emotional needs 259 and responses of the whole of the ANS in physiological and/or pathologi-260 cal conditions, probably indicating the activation of different autonomous 261 circuits in case of different conditions. 262

In this work, we have designed an experimental setup able to record in 263 synchrony the video of pupil (done with a commercial system [25]) and dif-264 ferent responses of the ANS. A modular sensor network was designed and 265 implemented to acquire, process and transmit via USB some biomedical sig-266 nals reflecting the state of the ANS. Two sensors have been included in the 267 prototype, for pulse oximetry and skin conductivity. However, it is feasible 268 for being extended to include more body sensors, miniaturized and embedded 269 in a portable system. This could be important for future extension of the 270 work, as body sensor networks are finding many applications in the contin-271 uous monitoring of sensitive people [40][41][42][43]. Indeed, many different 272 sensors are available to monitor physiological data and could be included: 273 accelerometer, blood glucose sensor, electrodes for bioelectric signals, blood 274

<sup>275</sup> pressure sensor, gyroscope, carbon dioxide gas sensor, etc. [44].

The developed system, even if it is only a prototype, provided robust estimations of the electrodermal activity and of the pulse oximetry, which was then processed to investigate the heart inter-beat interval. Note that the PPG is immune to electrical artifacts, which could be observed on the electrocardiogram during TENS application (however, it is less precise in detecting the HR, as it also depends on the pulse wave velocity). Thus, our system is adequate for the study of the ANS response to TENS reflected in the HR.

Preliminary experimental tests are shown. Even if we considered only 283 short recordings, weak ANS stimulations (dental occlusion, light, TENS and 284 a computational task) in healthy subjects and we extracted simple indexes 285 (mean, STD and trend), we have shown statistically significant variations of 286 at least an index in each pair of conditions. This indicates that the overall 287 information provided by the joint recordings, not just that of each individual 288 signal, should be used for the discrimination of the ANS responses in the 289 different considered conditions. The results are in line with our expectations: 290 HDO and computation elicit the sympathetic response (which should result 291 in HR and GSR increasing and pupil dilation), TENS induces relaxation 292 (determining a decrease of HR, GSR and pupil size). Pupil appeared to 293 be the most sensible system, as it reflected even the small stimuli given by 294 HDO or TENS. HR showed significant variations only in a few conditions 295 (e.g., mean HR was significantly increased only by the computation task). 296 GSR showed an important trend during computation, but it was prone to 297 accumulation effects (with an average increase along the experiment, in spite 298 of the application of TENS). 299

Other tests and advanced processing techniques could provide specific indications in physiology or pathology, in future joint acquisitions of different responses of the ANS. Additional measures to guarantee reproducibility of the tests need to be developed, such as careful preparation in order to stabilize hemodynamic parameters. Moreover, additional sensors or stimulation signals could also be included in the proposed device, due to its modular architecture.

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CONDITIONS	INDEXES with SIGNIFICANT DIFFERENCE
	Mean HR (p=0.0002)
	Comp.: $1.4(1.17, 2.04) - RP: 1.08(0.94, 1.57)$
Computation	STD of HR $(p=0.001)$
	Comp.: $0.28(0.17, 0.54) - \text{RP: } 0.11(0.08, 0.38)$
versus	STD of GSR (p=0.0015)
	Comp.: $1.25(0.54, 2.77) - \text{RP: } 0.2(0.1, 1.0)$
RP (D)	Trend of $GSR$ (p=0.0078)
	Comp.: $3.03(0.27, 3.28) - \text{RP:} -0.12(-1.88, 2.06)$
	Mean pupil area $(p=0.01)$
	Comp.: $6952(6180, 9050) - RP: 6565(5839, 8072)$
HDO (D)	Mean pupil area $(p=0.0029)$
	HDO: $6817(6318, 8636) - RP: 6565(5839, 8072)$
versus	Trend of pupil area (p=0.012)
	HDO: $1.05(-0.62, 1.86) - \text{RP}: 2.10(1.55, 2.63)$
RP (D)	STD of pupil area (p=0.024)
	HDO: $177(142, 279) - RP: 267(202, 308)$
pre-TENS	Mean of pupil area (p=0.009)
	pre: $6425(3806,7391) - \text{tens: } 5818(3134,7082)$
versus	$STD \ of \ GSR \ (p=0.031)$
TITING	pre: $0.11(0.05, 0.27)$ – tens: $0.21(0.72, 0.83)$
TENS	$STD \ of \ HR \ (p=0.036)$
	pre: $0.11(0.07, 0.24)$ – tens: $0.12(0.09, 0.57)$
TENS versus	Mean of GSR (p=0.0001)
post-TENS	tens: $12.91(9.09, 14.83) - \text{post: } 13.65(12.86, 16.61)$
pre-TENS	Mean of GSR (p=0.006) pre: $12.88(10.88, 13.68) - post: 13.65(12.73, 16.61)$
TONORI	Mean area of pupil (p=0.012)
versus	pre: $6425(3806, 7392) - post: 5823(3151, 6874)$
post-TENS	$STD \ of \ HR \ (p=0.022)$
	pre: $0.12(0.08, 0.25) - \text{post:} 0.16(0.08, 0.35)$
Light	$\frac{1}{Mean \ area \ of \ pupil \ (p << 0.001)}$
21810	L: $2987(2545, 3981) - D: 7312(6613, 9059)$
versus	$STD \ of \ pupil \ (p << 0.001)$
	L: $417(333,539) - D: 220(161,283)$
Darkness	Trend of pupil (p=0.014)
	L: $0.23(-0.79, 1.29) - D: 1.70(0.21, 2.32)$

Table 2: Statistical analysis of the data using Wilcoxon sign rank test. Mean, standard deviation and trend of data were considered (L: light; D: darkness). Median and quartiles of the indexes showing significant differences (p<0.05) are reported (in order of increasing p). Pupil size is indicated in pixels, HR in Hz, GSR in  $\mu$ S; their trends are measured in arbitrary units (they were computed on normalized data and time).