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Testing the vagal withdrawal hypothesis during light exercise under autonomic blockade: a heart rate variability study

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1 TESTING THE VAGAL WITHDRAWAL HYPOTHESIS DURING LIGHT EXERCISE

- 2 UNDER AUTONOMIC BLOCKADE: A HEART RATE VARIABILITY STUDY
- 3

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43 ABSTRACT

Introduction. We performed the first analysis of heart rate variability (HRV) at rest and exercise under full autonomic blockade on the same subjects, to test the conjecture that vagal tone withdrawal occurs at exercise onset. We hypothesized that, between rest and exercise: i) no differences in total power (PTOT) under parasympathetic blockade; ii) a PTOT fall under β 1-sympathetic blockade; iii) no differences in PTOT under blockade of both ANS branches.

50 **Methods.** 7 males (24±3 years) performed 5-min cycling (80W) supine, preceded by 51 5-min rest during control and with administration of atropine, metoprolol and 52 atropine+metoprolol (double blockade). Heart rate and arterial blood pressure were 53 continuously recorded. HRV and blood pressure variability were determined by power 54 spectral analysis, and baroreflex sensitivity (BRS) by the sequence method.

Results. At rest, PTOT and the powers of low (LF) and high (HF) frequency 55 56 components of HRV were dramatically decreased in atropine and double blockade 57 compared to control and metoprolol, with no effects on LF/HF ratio and on the normalised LF (LFnu) and HF (HFnu). At exercise, patterns were the same as at rest. 58 Comparing exercise to rest, PTOT varied as hypothesized. For SAP and DAP, resting 59 P_{TOT} was the same in all conditions. At exercise, in all conditions, P_{TOT} was lower 60 61 than in control. BRS decreased under atropine and double blockade at rest, under 62 control and metoprolol during exercise.

63 Conclusions. The results support the hypothesis that vagal suppression determined
 64 disappearance of HRV during exercise.

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69 Key words

70 Cardiovascular regulation · Arterial blood pressure · Baroreflexes · Metoprolol ·
 71 Atropine

72

73 New & Noteworthy

This study provides the first demonstration, by systematic analysis of heart rate variability (HRV) at rest and exercise under full autonomic blockade on the same subjects, that suppression of vagal activity is responsible of the disappearance of spontaneous HRV during exercise. This finding supports previous hypotheses on the role of vagal withdrawal in the control of the rapid cardiovascular response at exercise onset.

80 INTRODUCTION

81

At exercise start, the characteristics of the heart rate (HR) kinetics under vagal blockade (12) suggested that sudden withdrawal of vagal tone may occur. This hypothesis may explain the concomitant sudden increase of cardiac output (13, 25). Recently, vagal withdrawal was called upon also to explain the early changes in baroreflex sensitivity upon exercise start (4). If this is so, we should expect that the amplitude of the rapid HR and cardiac output responses would be greater, the stronger is the vagal modulation of heart activity at rest.

89 The experimental evidence, however, is not conclusive under this respect, and 90 several data seem to contradict the vagal withdrawal hypothesis. For instance, 91 although we know that resting vagal activation is greater in supine than in upright 92 position (35, 47, 49), the amplitude of the rapid cardiac output response at exercise 93 onset was found to be smaller in supine than in upright posture (27; 55). On the other 94 hand, vagal activity is reduced and sympathetic activation is increased in acute 95 hypoxia as compared to normoxia (5;18, 23, 57, 58): in spite of this, even in hypoxia 96 HR determined a large fraction of a significant cardiac output response (26). These 97 data represent a serious challenge to the vagal withdrawal hypothesis at exercise 98 onset.

99 The vagal withdrawal hypothesis at exercise onset may also be tested by 100 investigating the neural modulation of the heartbeat under pharmacological blockade 101 of either the vagal or the sympathetic or both branches of the ANS (2, 6, 15, 17, 21, 102 24, 29, 32, 33, 35, 43, 53). The analysis of spontaneous heart rate variability (HRV) 103 demonstrated that vagal blockade reduced the total power (PTOT) of HRV, acting on 104 the reduction of both its high (HF) and low frequency (LF) components. Nevertheless, 105 little attention was given so far to the analysis of HRV during exercise combined with 106 pharmacological blockade. Warren et al. (1997) reported that the powers of both the

107 LF and the HF peaks were by far lower at exercise than at rest under placebo, but they did not find differences under vagal blockade with glycopyrrolate; moreover, 108 109 esmolol administration provided similar results as placebo. The interpretation of their 110 results was undermined by the type of drug used and their study was limited by the 111 fact that they did not analyse blood pressure variability, another important indirect 112 feature of sympathetic modulation of the cardiovascular system. Polanczyk et al. (42) showed that atropine and propranolol administration did not vary the spectrum 113 114 components of HRV, contrary to their expectations.

115 If the vagal withdrawal hypothesis was correct, we should predict that, when comparing rest and exercise: i) no differences in PTOT, LF and HF under full vagal 116 117 blockade would be found; ii) a drastic fall in PTOT, LF and HF under selective β 1-118 sympathetic blockade would occur; iii) no differences in PTOT, LF and HF under simultaneous blockade of the two branches of the ANS would appear. Moreover, we 119 120 expected that arterial blood pressure variability would not follow the same pattern of response as HRV, because the former reflects more the peripheral sympathetic 121 122 vascular modulation than the central cardiac modulation.

123 These predictions were tested in the present study, the aim of which was to 124 investigate the effects of vagal blockade, of selective β 1-sympathetic blockade and of 125 simultaneous blockade of both branches of the ANS, at rest and during exercise, on 126 HRV and blood pressure variability.

- 127
- 128
- 129 **METHODS**

130

131 **Participants**

Seven healthy non-smoking young participants volunteered for the experiments. They were (mean \pm SD) 24.3 \pm 2.6 years old, 181.2 \pm 3.1 cm tall and weighed 78.9 \pm 6.1 kg. Exclusion criteria were: presence of history of cardiopulmonary disease and regular use of drugs at the time of the study. Participants were instructed to avoid caffeine consumption 24 hours before the visit and to refrain from performing strenuous exercise the day before testing.

All participants were preliminarily informed on the design and risks associated with the experiments and they signed a written informed consent. The study was conducted in accordance with the Declaration of Helsinki, and the protocol was approved by the local institutional ethical committee.

142

143 **Protocol and measurements**

144 The experiments were carried out in the Clinical Physiology Laboratory of the 145 University of Geneva, Switzerland. The volunteers reported to the laboratory on four 146 different days, with at least a 48-hour recovery between visits. Experiments were performed in supine posture, in order to reduce potential mechanical effects related 147 148 to the remarkable sudden increase in venous return at exercise start upright. After 149 instrumentation, a 20-gauge catheter was placed in the antecubital vein of the right 150 arm to administer drugs. A unique 5-min monitoring at rest preceded a series of three 151 5-min constant-load leg exercises, on cycle ergometer, at 80 watts, to avoid lactate threshold. Between repetitions a 5-min recovery was administered. 152

For the entire duration of the protocol, we obtained continuous recordings of the electrocardiogram (Elmed ETM 2000, Heiligenhaus, Germany), and the arterial pulse pressure profiles, obtained at a fingertip of the left arm by means of a noninvasive cuff pressure recorder (Portapres, FMS, Amsterdam, The Netherlands).

157 The R-R interval (RR) and its reciprocal, HR, were computed beat-by-beat. 158 Systolic and diastolic blood pressure (SAP and DAP, respectively) values were

159 obtained from each pulse pressure profile, using the Beatscope® software package (FMS, Amsterdam, The Netherlands). Beat-by-beat mean arterial pressure (MAP) 160 was computed as the integral mean of each pressure profile, using the same 161 162 software package. Breathing frequency also calculated from was the 163 electrocardiogram plot.

164

All the signals were digitalized in parallel by a 16-channel A/D converter (MP150, Biopac Systems, Goleta CA, USA) and stored on a computer. The acquisition rate was 400 Hz.

The protocol was performed under four experimental conditions, administered in random order: i) control, i.e. with placebo infusion, ii) parasympathetic blockade with atropine administration, ii) selective β1-adrenergic blockade with metoprolol administration, and iv) double blockade of both branches of the ANS with simultaneous atropine and metoprolol administration.

173

174 **Drug administration**

Parasympathetic blockade was achieved by administering atropine in a single 0.04 mg/kg dose (mean 3.06 ± 0.23 mg, range 2.7 - 3.4 mg), which was used in previous studies to attain full vagal blockade (14, 17, 31, 59). The half-life of a single atropine dose is 180 minutes (52) so that, blockade was maintained during the entire duration of each experiment.

The β 1-adrenergic blockade was attained by using metoprolol tartrate (Loprésor, Novartis, Switzerland). After an initial bolus of 15 mg, metoprolol tartrate was continuously infused in an antecubital vein at a rate of 45 mg per hour, by means of an infusion pump. The efficacy of adrenergic blockade along time was evaluated on a separate session, by analysing the heart rate response following isoprenaline injection, as previously described (14). The correct metoprolol maintenance dose was identified as the dose ensuring an 80% reduction of the HR
response to isoprenaline for the entire protocol duration.

experiments with 188 For the double, simultaneous sympathetic and 189 parasympathetic blockade, the same atropine and metoprolol dose and administration procedure described here above were applied. 190

191

192Data treatment

After construction of the time series of RR. SAP and DAP from the continuous 193 194 recordings of electrocardiogram and pulse pressure profiles, Fast Fourier Transform 195 (FFT) was used to evaluate spontaneous variability of RR, SAP and DAP (35). The data length used was 5 minutes at rest and 3 minutes at exercise. In the latter case, 196 197 one repetition, that with the most stable and cleanest trace, was analysed. The total 198 power (PTOT) (0.0-0.5 Hz) of RR, SAP and DAP variabilities, corresponding to 199 variance, was initially computed. Subsequently, the powers and frequencies of LF (0.03-0.14 Hz) and HF (0.15-0.5 Hz) components of the power spectrum were 200 computed and expressed in absolute units (ms²). The very low frequency component 201 202 was neglected. The LF/HF ratio was also calculated. Normalized LF and HF (LFnu 203 and HFnu, respectively) were computed as:

$$\frac{LF \times 100}{Ptot-VLF}$$
(1)

and expressed in normalized units (28).

The spontaneous baroreflex sensitivity (BRS, expressed in ms mmHg⁻¹) was estimated from SAP and RR by means of the sequence method (3). Sequences of at least three heartbeats, corresponding to an increase or decrease in SAP and identifying a consensual change in RR interval, were selected. Linear regression analysis was applied on these sequences and the calculated slope was retained. BRS was then calculated as the mean of the slopes of all sequences per each

212 participant in each condition. Only sequences showing a coefficient of determination

of at least 0.85 were analysed.

214 Spectral analysis and BRS were performed on Matlab® environment as previously 215 described (41). Breathing Frequency was calculated with the ECG-Derived-216 Respiration method used by Moody et al. (30).

- 217
- 218

219 Statistics

Data are reported as group means \pm standard deviation. The effects of medication and exercise type on the main outcomes were analysed by 2-way ANOVA for repeated measurements. When applicable, a Tukey post-hoc test was used to locate significant differences. Differences were considered significant if p<0.05. All data were analysed with Statistica 12 © (StatSoft, Inc., Tulsa, OK).

225

226 **RESULTS**

227 All participants successfully completed the study maintaining a normal sinus 228 beat along the four experimental conditions (no arrhythmic beats were observed). 229 The mean values of measured and calculated variables at rest and during exercise for all conditions are reported in Table 1. At rest, in control condition, HR was 62.7 ± 230 8.5 min⁻¹. Under sympathetic blockade, no significant differences with respect to 231 232 control were observed. Under atropine, it was significantly higher than in control and 233 under metoprolol. Under double blockade, it was higher than in control and under 234 metoprolol, but lower than under atropine. During exercise, in control condition, HR was $105.0 \pm 12.4 \text{ min}^{-1}$, and was higher under metoprolol, atropine and double 235 blockade than in control. With respect to the corresponding values at rest, HR during 236 237 exercise increased in all conditions except double blockade.

238 At rest, in control condition SAP was 112.0±9.5 mmHg and DAP was 55.0±9.6 mmHg. With respect to control, no differences were observed for either SAP or DAP 239 240 with any investigated pharmacological treatment, although with double blockade, 241 DAP tended to be higher than in control and was significantly higher than under metoprolol. MAP was 74.0±8.6 mmHg in control and did not differ in the three 242 243 investigated pharmacological conditions, except that it was higher under double blockade than with metoprolol. Breathing frequency was 0.23±0.06 Hz in control and 244 did not change in the three conditions. At exercise, in control condition, SAP was 245 246 138.5±17.5 and DAP was 60.9±7.5 mmHg. With respect to control, SAP was significantly lower under the three pharmacological conditions. No differences were 247 observed for DAP. MAP was 86.8±9.9 mmHg in control and did not vary significantly 248 249 among conditions. With respect to the corresponding values at rest, MAP during 250 exercise was higher only in control. Breathing frequency was 0.42±0.07 Hz in control 251 and did not change in the three other conditions.

HRV data are shown in Table 2. Examples of HRV spectra are shown in 252 Figure 1. At rest, with respect to control, PTOT was not affected by metoprolol 253 254 administration, but it was largely and significantly decreased under atropine and 255 double blockade, due to drastically lower values of both LF and HF powers. No differences between atropine and double blockade were found. The same was the 256 257 case at exercise, although the difference were much smaller than at rest, because, when moving from rest to exercise, PTOT was drastically reduced in control and 258 259 under metoprolol. No differences for LF and HF powers between sympathetic 260 blockade and control, or between atropine and double blockade, were observed.

At rest, the LF/HF ratio at rest was unaffected by drug treatment, the only significant difference being between atropine and double blockade. The same was the case for LFnu. No differences were observed concerning HFnu. *At exercise*, the LF/HF ratio did not differ under metoprolol or atropine with respect to control, but it was lower under double blockade than in control and in the other pharmacological
conditions. The same was the case for LFnu. Coherently, HFnu was higher in double
blockade than in any other condition.

268 All data concerning spontaneous SAP and DAP variability are shown in Table 3. At rest, concerning SAP, no differences among conditions were observed for P_{TOT} . 269 270 Concerning the LF power, no differences between sympathetic blockade and control 271 were found, but it was lower under atropine and double blockade than in control and sympathetic blockade. The HF power in atropine and double blockade was lower 272 than in control and under metoprolol, although for the latter the level of significance 273 was not attained. During exercise, PTOT was lower in all three investigated 274 pharmacological conditions than in control, but no differences among conditions were 275 276 observed for both the LF and the HF powers. In control and under atropine, the LF power was higher at exercise than at rest. The LF/HF ratio was unchanged in all 277 278 conditions.

At rest, concerning DAP, no changes in P_{TOT} were found in any pharmacological condition with respect to control. The HF power did not vary among conditions, while the LF power was lower in atropine and double blockade than in control. The LF/HF ratio was lower in all conditions than in control. *During exercise*, there were no significant differences among conditions or with respect to the same condition at rest.

The BRS values at rest and exercise are shown in Figure 2. *At rest*, BRS was significantly lower under atropine and under double blockade than in control and under sympathetic blockade, which in turn did not differ between them. *During exercise*, BRS under atropine and double blockade was lower than in control and under sympathetic blockade. BRS was lower at exercise than at rest in all conditions except double blockade.

292 **DISCUSSION**

293

The analysis of spontaneous heart rate variability at rest showed that: 1. atropine administration drastically reduced PTOT, due to the fall of both LF and HF powers, with respect to control; 2. simultaneous double blockade with atropine and metoprolol provided the same results as atropine administration only; 3. metoprolol administration had no effects on heart rate variability.

When moving from rest to exercise, our results showed that: 1. no differences in PTOT, LF and HF appeared under atropine and under simultaneous atropine and metoprolol administration with respect to rest; 2. PTOT, and the LF and HF powers, were decreased by the same extent under metoprolol as in control. However, during exercise, PTOT, and the LF and HF powers were lower under atropine and double blockade than in control or with metoprolol.

These results are in line with the predictions made, and thus do not allow refutation of the vagal withdrawal hypothesis, but rather reinforce it. Although, taken separately, similar consistent results can be found in the previous literature (2, 6, 8, 10, 11, 15, 17, 21, 24, 29, 32, 33, 35, 43, 44), this is the first time that a complete picture of the role of the autonomic nervous system in determining heart rate variability in rest and exercise was obtained.

311

312 Heart rate variability

The significant increase in HR after atropine administration is in line with previous studies (9, 21, 22, 48, 50) and was opposed by the observation that, after metoprolol administration, despite a slight decrease, HR did not change significantly compared to control. These results were similar in size to those obtained in a previous study with the same drug (48). However, they are at variance with those of other studies, carried out in upright posture, showing a significant HR reduction at

rest with beta-blockade (11, 14, 15, 19). In supine posture, the predominance of
vagal modulation of HR (20, 35) may explain the lack of HR changes with metoprolol.

Concerning HRV, metoprolol failed in changing PTOT, LF and HF at rest, 321 322 indicating that a selective blockade of cardiac β -adrenergic receptors has no effects on spontaneous HR oscillations. This suggests that the sympathetic outflow to the 323 heart may not be the main determinant of HRV, although the PTOT values under 324 325 double blockade appear lower (just a tendency) than under atropine. These results 326 for PTOT, although in agreement with those of some previous studies (15, 53), are in contrast with those by Cogliati et al. (11), who showed an increase in PTOT under 327 328 atenolol, supporting the idea that the pattern was mostly due to an increase in the HF peak. This finding suggested stronger cardio-respiratory coupling under atenolol than 329 330 in control. Comparable results were obtained by others (40) using propranolol.

Spontaneous HR oscillations were almost suppressed after atropine 331 332 administration, as previously found (8, 15, 21, 29, 32, 33, 53), supporting the notion that parasympathetic outflow to the heart is the major determinant of HRV in resting 333 334 humans. This was so also under simultaneous sympathetic and vagal blockade, 335 indicating that suppression of the parasympathetic modulation of the heartbeat was 336 the most important determinant of the present results. Breathing frequency did not 337 change in the three conditions, being obviously higher at exercise than at rest. This implies that changes in HF power were not due to any change in breathing 338 339 frequency.

Coherently, in the present study, passing from rest to exercise implied a large fall in LF and HF powers in control and under metoprolol. Conversely, under atropine and double blockade, in which a suppression of the vagal modulation of HR was obtained already at rest, no changes were found at exercise with respect to rest. These results demonstrate that the well-known fall of HRV, which is usually observed during exercise (37), is essentially a consequence of the withdrawal of the vagal

outflow to the heart occurring at exercise onset (12, 25), as hypothesized. As such, 346 our results suggest that vagal withdrawal is incomplete at the investigated powers, 347 348 because the LF and HF powers during exercise were still higher in control than with 349 atropine or double blockade, the two conditions in which a full blockade of muscarinic 350 receptors was attained. On the other hand, the fact that passing from rest to exercise 351 generated comparable results with metoprolol as in control, is coherent with the 352 reported decrease of the LF peak in humans (37, 39). These data are in contrast with the generally accepted notion that, during exercise, the degree of sympathetic 353 354 activation increases (46, 54) and the modulation of the heartbeat by the sympathetic 355 efferents becomes predominant (38, 45). This may mean that HRV in exercise does not reflect the degree of ongoing sympathetic activation. 356

357 When we look at the normalized variables at rest, none of the investigated 358 drugs could change the LF/HF significantly with respect to control: this reflects the 359 finding that the effects of drug administration on the LF and HF powers at rest were 360 of the same size. In contrast, during exercise, there was a tendency toward a lower HF power than LF power. Yet this tendency, though not significant, was such as to 361 362 provide, at exercise compared to rest, significantly lower HFnu values in control and 363 under sympathetic blockade (only a tendency in A and in DB). Consequently, LF/HF ratio resulted higher at exercise than at rest, at least in these two cases. 364

365 In the context of the present hypothesis, this would suggest that the withdrawal of vagal tone at exercise onset might have had greater effects on the HF 366 367 than on the LF component of HRV. Alternatively, the relative increase of the LF component of RR variability may suggest an increase of the cardiac sympathetic 368 369 modulation. Nevertheless, LFnu in double blockade was significantly lower and HFnu 370 significantly higher than in control, despite the lack of differences in the LF/HF ratio. 371 This may be due to the non-autonomic effect of an increase in ventilation that is reflected on HRV through changes in venous return during exercise. A similar 372

condition can be observed in a neurodegenerative disease such as the pure autonomic failure. This condition is characterized by both a cardiac sympathetic and parasympathetic denervation leading to PTOT values mimicking high dosage atropine administration (16), in which a HF component of HRV, non-autonomic in origin, is present (39). These apparently contradictory results prevent us from arriving at clearcut conclusions concerning the mechanisms at the basis of relative powers in this study.

380

381 Blood pressure variability

382 Arterial blood pressure at rest was unaffected by drug administration. The fact 383 that atropine did not act on systemic blood pressure, in agreement with previous 384 studies (15, 21), is coherent with the notion that there is no cholinergic innervation in 385 most regional circulations. On the other hand, metoprolol is a selective blocker of β 1-386 adrenergic receptors that are expressed specifically in the heart, not in arterioles, so 387 that it is not expected to induce changes in blood pressure.

388 Coherently, SAP variability was much less affected by atropine and double 389 blockade than HRV. According to Zhang et al. (61), who investigated spontaneous 390 blood pressure variability under ganglionic blockade with Trimethaphan, the HF peak 391 of blood pressure variability is mediated by mechanical effects due to the breathing 392 cycle and cardiac filling: if this is so, one would not expect effects of any of the drugs used in this study on the HF power for blood pressure. In fact, the changes in HF 393 394 power due to drug administration in the present study were much smaller than for HRV, although significant under atropine and double blockade. Zhang et al. (61) also 395 396 attributed the LF power of blood pressure variability to either sympathetic activity or intrinsic vascular rhythmicity: if this is so, no changes in LF were to be found with 397 atropine, metoprolol or double blockade: in fact, we found much smaller differences 398 399 in LF power due to drug administration for blood pressure variability than for HRV.

400 Yet these changes were consensual with those of HF power, being significant under 401 atropine and double blockade. These effects might have been an indirect 402 consequence of the role that the autonomic nervous system may play in modulating 403 the dynamic relationship between HRV and blood pressure variability (7, 61), with an 404 involvement of its parasympathetic branch.

405 Most remarkable are the differences observed when passing from rest to exercise: the LF power for SAP increased in control, as expected (37, 39), and with 406 407 atropine, but not with metoprolol and in double blockade. This indicates that the increase in LF power for SAP may be a consequence of increased sympathetic 408 409 modulation during exercise. No effects were observed under any drug on the HF 410 power: this means that the HF power of SAP is independent of the activity of the two 411 branches of the ANS. The lack of exercise effects on HF power under drug stimulation explains why the PTOT did not differ significantly at exercise with respect 412 413 to rest under atropine.

DAP variability was unaffected by drug administration: this suggests that the exercise effect on the LF power of SAP, related to a selective blockade of β 1 adrenergic receptors, is mediated by a central (cardiac) rather than a peripheral (arteriolar muscle vasodilation) action of the sympathetic branch of the ANS.

418

419 **Baroreflex sensitivity**

At rest, BRS was drastically lower under atropine and double blockade than in control. This observation was consistent with what we observed for the LF peak of blood pressure variability: reduced under atropine and double blockade, unchanged under metoprolol, with respect to control. Coherently, when comparing rest with exercise in a given condition, BRS decreased in control and under metoprolol, but did not change under atropine and double blockade. These results on BRS appear in agreement with previous observations (1, 11, 56). Bringard et al. (4) postulated that

BRS is mainly modulated by the parasympathetic efferent branch on the ANS. These 427 428 data support this hypothesis. Muscarinic receptors do not modulate smooth muscle 429 tone in most arterioles, including those of skeletal muscles. Thus, the 430 parasympathetic effect on arterial blood pressure variability indexes must be indirect. Based on the present results, we speculate that baroreflexes may participate in the 431 432 modulation of the LF power of arterial blood pressure. The reduction of BRS observed during exercise (51) support the idea of alfa-index changes as previously 433 434 reported (36). In the present study, the BRS reduction at exercise was observed only 435 in control and with metoprolol, but not with atropine and double blockade. This finding reinforces the notion that withdrawal of vagal tone is responsible for the fall of BRS at 436 exercise onset (4, 34). Coherently, no differences in BRS among the four 437 438 investigated conditions were observed during exercise.

439

440 **Study limitations**

A limitation of this study may be suggested by the lack of differences between 441 442 control and metoprolol, as this may also suggest that the β 1-adrenergic blockade 443 might have been incomplete. It is of note, however, that we used the same dose and followed the same procedure of metoprolol administration as in a previous study (14) 444 in upright posture, which showed a significant resting HR decrease both in normoxia 445 446 and in acute hypoxia at rest as at exercise. Moreover, we observe that the 447 isoprenaline test provided unambiguous evidence of quasi-complete B1-adrenergic 448 blockade.

Another possible limiting factor is related to the fact that HR rate differed remarkably among conditions. This may affect the HRV indexes in time domain *per* se (59), thus possibly undermining the relation to the action of the autonomic nervous system.

453

454 CONCLUSION

The results of this study support the tested hypothesis that vagal suppression 455 is responsible of the disappearance of the spontaneous HRV during exercise. The 456 457 observed effects on arterial blood pressure variability are indirectly related to the action of the administered drugs, supporting the notion that blood pressure and HRV 458 459 are only partially-associated phenomena, possibly controlled by different physiological mechanisms 460

461

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465

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Table 1: Mean steady state values for the cardiovascular variables monitored during rest (R)
and exercise (E) in the four experimental conditions: Control, Atropine, Metoprolol, and
Double Blockade.

Measured variables		Control	Metoprolol	Atropine	Double Blockade
HR (min⁻¹)	R	62.67±8.47	59.58±7.11 ^{#§}	111.17±17.75 ^{*§}	93.71±5.48*
	Е	105.04±12.39	93.53±8.17 [#]	135.04±20.56*	103.19±8.06 [#]
RR (ms)	R	985.3 ± 185.7	1017.7 ± 104.4 [#]	548.6 ± 79.5 *§	642.1 ± 38.2 *
	E	577.9 ± 66.2	645.3 ± 51.3 [#]	455.9 ± 89.3*	584.2 ± 41.1 [#]
SAP (mmHq)	R	111.97±9.52	109.75±13.89	112.96±11.83	119.48±14.29
	Е	138.51±17.53	113.58±15.21*	108.73±15.94*	107.70±14.76*
DAP (mmHa)	R	54.95±9.64	48.96±10.81 [§]	60.95±9.10	66.16±8.43
(3)	Е	60.94±7.48	53.35±13.55	54.21±7.72	54.34±6.92
MAP (mmHq)	R	73.95±8.59	69.22±10.42 [§]	78.28±7.76	83.93±7.78
	Е	86.79±9.88	73.42±13.53	72.58±10.03	72.13±9.41
BRS (ms	R	25.74 ± 11.28	27.42 ± 8.51 [#]	2.17 ± 1.06 * [§]	3.00 ± 0.92 *
mmHg⁻¹)	Е	2.59 ± 1.76	3.17 ± 0.62 [#]	0.85 ± 0.31*	2.13 ± 0.44
BF (Hz)	R	0.23 ± 0.06	0.21 ± 0.06	0.29 ± 0.04	0.23 ± 0.07
(/	Е	0.42 ± 0.07	0.39 ± 0.04	0.40 ± 0.04	0.41 ± 0.07

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640

641 Values are means ± SD. HR: heart rate; RR: R-R interval; SAP: systolic arterial pressure;

642 DAP: diastolic arterial pressure; MAP: mean arterial pressure; BRS: spontaneous baroreflex

643 sensitivity. BF: breathing frequency.

644 N=7; 2-way ANOVA for repeated measurements; p<0.05; *: significantly different from

- 645 Control. #: significantly different from Atropine. §: significantly different from Double
- 646 Blockade. In bold: Exercise significantly different from Rest.

Table 2: Mean and standard deviations of all parameters calculated by means of heart rate
variability in the four investigated conditions: Control, Atropine, Metoprolol, and Double
Blockade.

Heart Rate variability		Control	Metoprolol	Atropine	Double blockade
ABSOLUTE					
P _{TOT} (ms2Hz ⁻¹)	R	6351.4 ± 4476.4	7883.2 ± 5965.9	22.5 ± 13.8*•	12.9 ± 4.9*•
	Е	185.4 ± 77.1	93.6 ± 30.9*	10.1 ± 3.3*•	$14.8 \pm 4.7^{*\bullet}$
LF (ms2Hz ⁻¹)	R	1717.5 ± 1290.6	2711.9 ± 2061.8	1.5 ± 1.2*•	1.1 ± 0.5*•
	Е	40.6 ± 29.3	41.3 ± 29.3	1.7 ± 1.4*•	1.6 ± 1.5*•
HF (ms2Hz ⁻¹)	R	1441.0 ± 1296.1	2552.3 ± 2245.0	$0.9 \pm 0.5^{*}$	$2.6 \pm 0.8^{*\bullet}$
	Е	10.8 ± 7.8	11.2 ± 9.2	0.3 ± 0.13*	3.1 ± 1.6*•
RELATIVE					
LF/HF	R	1.4 ± 1.0	1.4 ± 0.8	1.8 ± 0.9	$0.5 \pm 0.3^{\#}$
	Е	4.1 ± 2.0	4.0 ± 2.0	4.0 ± 2.6	$0.3 \pm 0.1^{*^{\bullet \#}}$
LFnu (%)	R	46.8 ± 19.3	46.1 ± 14.7	57.7 ± 28.2	25.9 ± 13.8 [#]
	Ε	69.6 ± 16.5	65.3 ± 21.2	61.6 ± 22.7	12.7 ± 8.0***
HFnu (%)	R	51.1 ± 18.3	51.1 ± 15.4	38.3 ± 26.9	62.6 ±15.6
	Ε	15.5 ± 8.7	17.4 ± 5.0	22.6 ± 14.4	$45.5 \pm 23.5^{*}$

- 651 Values are means ± SD. P_{TOT}: total power. LF: low frequency power. HF: high frequency
- 652 power. LF/HF: low-to-high frequency ratio; LFnu, relative low frequency power; HFnu,
- relative high frequency power. R: Rest. E: Exercise.
- 654 N=7; 2-way ANOVA for repeated measurements; p<0.05; *: significantly different from
- 655 Control. •: significantly different from Metoprolol. #: significantly different from Atropine. In
- 656 bold: Exercise significantly different from Rest

Table 3: Parameters resulting from the analysis of spontaneous variability of systolic and
 diastolic arterial pressures in the four investigated conditions: Control, Atropine, Metoprolol,
 and Double Blockade.

SAP variability		Control	Metoprolol	Atropine	Double Blockade
Ртот (ms2Hz ⁻¹)	R	25.70 ± 11.52	26.91 ± 15.50	16.99 ± 17.77	15.63 ± 8.19
	Е	70.83 ± 41.42	29.07 ± 12.24*	28.09 ± 6.77*	17.46 ± 7.00*
l F (ms2Hz ⁻¹)	R	7.03 ± 3.60	4.96 ± 1.90	1.55 ± 0.64*•	2.09 ± 1.38*•
	Е	18.68 ± 17.97	5.51 ± 1.52	10.93 ± 6.15**	5.80 ± 2.90* [#]
HF (ms2Hz ⁻¹)	R	4.04 ± 3.21	2.57 ± 1.79	1.20 ± 0.45*	1.09 ± 0.67*
	Е	5.49 ± 4.20	5.48 ± 3.96	3.29 ± 1.80	2.48 ± 0.93
LF/HF	R	2.27 ± 1.07	2.36 ± 1.09	1.46 ± 0.70	2.32 ± 1.54
	Е	2.61 ± 1.39	1.87 ± 1.32	3.46 ± 2.39	2.17 ± 0.67
DAP variability					
Ртот (ms2Hz ⁻¹)	R	9.65 ± 6.06	9.01 ± 3.47	4.64 ± 3.03	5.10 ± 2.32
	Е	7.63 ± 2.56	5.52 ± 2.56	3.92 ± 1.00	4.90 ± 3.40
LF (ms2Hz ⁻¹)	R	3.54 ± 2.57	2.56 ± 1.17	0.77 ± 0.52*	0.97 ± 0.67*
	Е	2.70 ± 1.80	1.97 ± 0.86	1.63 ± 0.48	1.11 ± 0.31
HF (ms2Hz ⁻¹)	R	2.22 ± 2.77	1.88 ± 2.10	0.40 ± 0.20	0.46 ± 0.52
, , , , ,	E	1.66 ± 1.25	1.13 ± 0.75	0.89 ± 0.52	0.92 ± 0.50
LF/HF	R	3.65 ± 1.26	2.40 ± 1.21*	2.03 ± 1.26*	3.03 ± 1.48*
	Е	3.00 ± 2.53	1.69 ± 0.98	2.05 ± 0.73	1.53 ± 0.63

- 661 Values are means ± SD. P_{TOT}: total power. LF: low frequency power. HF: high frequency
- 662 power. LF/HF: low-to-high frequency ratio. R: Rest. E: Exercise.
- 663 N=7; 2-way ANOVA for repeated measurements; p<0.05; *: significantly different from
- 664 Control. •: significantly different from Metoprolol; #: significantly different from Atropine. In
- 665 bold: Exercise significantly different from Rest

Figure 1: Heart Rate Variability (HRV) spectrum resulting from the experiments which the
shown HRV segments belong to *left column*: Rest; *right column*: Exercise; *first row*: Control; *second row*: Atropine; *third row*: Metoprolol; *fourth row*: Double blockade. N=7; X axis:
frequency (Hz). Y axis: RR power (ms2/Hz). Note: differences in Y scales. C: Control. A:
Atropine. M: Metoprolol DB: Double blockade.

Figure 2: Mean values ± SD of BRS in each investigated condition (control / atropine / metoprolol / double blockade) at rest and during exercise. BRS: Spontaneous baroreflex sensitivity. N=7; 2-way ANOVA for repeated measurements; p<0.05: *: significantly different from CTRL. #: significantly different from DB. §: significantly different from ATR. \$: significantly different from the same condition at rest.



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Figure 2

