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## **Effects of Anger and Trait Forgiveness on Cardiovascular Risk in Women $\leq$ 25 Years of Age**

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Running Head: Anger, trait forgiveness & cardiovascular modulation

## Abstract

High trait anger is linked to adverse cardiovascular outcomes. A potential antidote to the cardiotoxic influence of anger is trait forgiveness (TF), as it has shown associations with improved blood pressure (BP) and cardiovagal tone regulation in cardiac patients. However, it has yet to be determined if anger and forgiveness independently predict cardiovascular parameters. Trait anger (State-Trait Anger Expression Inventory-2) and TF (Tendency to Forgive Scale) were evaluated in 308 ( $M=21.11$  years  $\pm$   $SD=2.52$ ) healthy female volunteers allocated to three related, yet distinct, studies. Hierarchical multiple regressions tested the incremental contribution of TF after accounting for anger. Study 1 assessed autonomic modulation via beat-to-beat BP and spectral analysis to examine sympathovagal balance and baroreflex functioning. Study 2 used tonometry and pulse wave analysis for aortic hemodynamics. Study 3 assessed 24-hour ambulatory BP and arterial stiffness index (AASI). Hierarchical models demonstrated that anger was significantly associated with increased sympathovagal tone, increased hemodynamic indices, high ambulatory blood pressures, and attenuated BP variability and baroreflex. In contrast, TF was associated with more favorable hemodynamic effects (i.e. decreased ventricular work and myocardial oxygen consumption). In conclusion, these results demonstrate divergent cardiovascular effects of anger and forgiveness, such that anger is associated with a more cardiotoxic autonomic and hemodynamic profile, whereas TF is associated with a more cardioprotective profile. These findings suggest that interventions aimed at decreasing anger while increasing forgiveness may be clinically relevant.

**Key Words:** anger, forgiveness, cardiovascular modulation

Considerable attention has been given to the relationship between anger and increased cardiac risk<sup>1</sup>, which is relevant to both healthy and cardiac patients. For example, research indicates anger to increase the risk of coronary heart disease among initially healthy individuals and to lead to poorer prognosis for individuals with heart disease.<sup>2,3</sup> Although the mechanism linking anger to increased cardiovascular risk is not well understood, impaired cardiovascular autonomic modulation and increased ventricular workload may be implicated. A potential antidote to the cardiotoxic influence of anger and hostility may be the cardioprotective properties provided by trait forgiveness (TF). TF has been shown to lower blood pressure (BP) and improve heart rate (HR) variability.<sup>4,5</sup> There is even some evidence that forgiveness predicts mortality<sup>5</sup> suggesting that failure to forgive unconditionally may be life threatening.

We sought to investigate anger and TF and their potentially divergent relationships with cardiovascular risk factors. We employed markers of cardiovascular functioning and tested the relationship between these psychological constructs, sympathetic nervous system (SNS) activity, BP control, and non-invasive aortic hemodynamics. We tested the overall hypothesis that anger would predict markers of cardiotoxicity and that TF would be associated with cardioprotection. To this end, we carried out 3 related, yet distinct, studies to test the incremental and unique contribution of TF in comparison to anger in examining the functioning of BP, cardiac autonomic modulation, and aortic hemodynamics. Study 1 assessed autonomic modulation via beat-to-beat BP and power spectral analysis to examine the differential contribution of SNS and parasympathetic nervous system (PNS) activation on baroreflex sensitivity (BRS) and HR modulation. Study 2 assessed aortic hemodynamics via applanation tonometry and pulse wave analysis to allow the measurement of noninvasive surrogates of aortic hemodynamics. Study 3 assessed 24 hour ambulatory BP and ambulatory arterial stiffness.

## Methods

A total of 308 healthy young women ( $M=21.11$  years  $\pm$   $SD=2.52$ ) participated in this research as approved by the University's institutional review board. Subjects were allocated to one of the following studies: Study 1: cardiovascular autonomic modulation and baroreflex function; Study 2: aortic hemodynamics; and Study 3: 24-hour ambulatory blood pressure. To minimize potential cardiovascular risk confounders, participants were excluded from study participation through an online health screening assessment if they smoked, exercised regularly as defined as  $>120$  min per week in the previous 6 months, were hypertensive as defined as  $BP \geq 140/90$  mmHg, had major chronic diseases, or were taking beta blockers, antidepressants, or stimulants. Participants were asked to abstain from caffeine, alcohol, and strenuous physical activity for at least 24 hours prior to testing and were asked to not eat for 4 hours prior to testing. Participants were tested in the early follicular phase of the menstrual cycle in order to avoid potential variations in pressure wave morphology and cardiovascular functioning.

In Study 1, after laboratory familiarization, anthropometrics were measured. Participants then completed a health questionnaire that included a health history form and an anger and TF scale. All data collection was conducted in the afternoon in a quiet, dimly lit, temperature-controlled room ( $23 \pm 1^\circ\text{C}$ ) at the same time of the day ( $\pm 2$  hours) in order to minimize potential diurnal variations in cardiovascular reactivity.<sup>6</sup> After instrument calibration and a 10 minute resting period in a seated position, beat-to-beat finger BP was recorded for 5 minutes.

In Study 2, participants were first introduced to the laboratory setting and familiarized with the study procedures. Body measurements (i.e. height and weight) were taken followed by the completion of a health questionnaire that included a health history form and an anger and TF scale. Data collection was conducted in the afternoon in a quiet, dimly lit, temperature-controlled

room ( $23 \pm 1^\circ\text{C}$ ) at the same time of the day ( $\pm 2$  hours). Participants were seated and given a 10 minute rest before any measurements were performed. Within 5 minutes after the rest period, measurements for peripheral brachial BP and applanation tonometry of the radial artery for central aortic hemodynamics were taken.

In Study 3, after completing an online health questionnaire, eligible participants were scheduled for a laboratory appointment to complete a 24 hour ambulatory BP assessment. Upon arrival, participants completed an anger scale, a TF scale, health characteristics (height and weight) were measured, and participants were fitted with an ambulatory BP device, which began between 08:00 – 11:00 hours and concluded when the recorder was returned to the laboratory the following day.

The trait subscale of the State-Trait Anger Expression Inventory-2 was used to measure trait anger.<sup>7</sup> Reliability for the sample was  $\alpha = .87$ . TF was measured using the 4-item Tendency to Forgive Scale.<sup>8</sup> Responses were summed into one overall score, with a possible range of 4 to 20. Reliability for the sample was  $\alpha = .81$ .

Beat-to-beat BP, HR, systolic BP, and diastolic BP were recorded via finger plethysmography (NIBP-100 Biopac Inc., Goleta, CA, USA). This method has been shown to provide accurate measurement of BP changes when compared with intra-arterial BP.<sup>9</sup> Mean BP was calculated as systolic blood pressure and diastolic blood pressure, where  $(1/3)$  systolic BP +  $(2/3)$  diastolic BP = mean BP. The BP peaks were used to calculate the time duration of intervals between heartbeats (RRI) and were automatically detected using commercially available software (WinCPRS, Turku, Finland). The RRIs were inspected for artifacts, premature beats and ectopic episodes in order to calculate HRV parameters. The main spectral components of the HRV that we calculated, by means of Fast Fourier transformation, were the low frequency (LF;

0.04-.15Hz) and the high frequency (HF; 0.15-0.4Hz). The use of absolute units ( $\text{ms}^2$ ) for HF and LF may be obtained in proportion to the total power (TP-VLF) which is expressed in normalized units (nu). Normalization is used to exclude the influence of other fractal components.<sup>10</sup> Since there is structural algebraic redundancy inherent in the normalized spectral HRV measures with respect to each other ( $n\text{LF}=1-n\text{HF}$ ), and also with respect to the LF/HF ratio, here we report LFnu to denote cardiac sympathovagal tone.<sup>10,11</sup>

Baroreflex functioning was evaluated via baroreflex sensitivity (BRS) calculated from the ECG and beat-by-beat BP files with the use of the cross-correlation method<sup>12</sup>, which is a time-domain sequential method for baroreflex function based on spontaneous systolic BP and R-R variability changes.

Indices of vascular function and aortic hemodynamics were obtained using brachial BP and applanation tonometry via pulse wave analysis (PWA) which is defined as the examination of the functioning of the arterial (central) pulse wave; allowing for accurate assessment of central hemodynamic functioning.<sup>13</sup> Brachial BP was recorded using an automated oscillometric device (HEM-705CP; Omron Healthcare, Vernon Hill, Illinois, USA). Brachial systolic and diastolic BP was used to calibrate radial waveforms obtained from a 10 second epoch using a high-fidelity tonometer (SPT-301B; Millar Instruments, Houston, TX). PWA provides a more sensitive marker of cardiovascular function than brachial BP.<sup>14,15</sup> We measured, brachial mean BP, aortic mean BP, systolic pressure time interval (STI; of left ventricular work), diastolic pressure time interval (DTI; coronary perfusion), the ratio of DTI to STI expressed as a percentage (subendocardial viability index, SVI; surrogate of myocardial perfusion and coronary flow reserve) and rate pressure product ( $\text{RPP} = \text{systolic BP} \times \text{HR}$ ; myocardial oxygen consumption).<sup>16-18</sup> All measurements were obtained in duplicates and averaged. Aortic BP waveforms were

derived using a generalized transfer function (SphygmoCor, AtCor Medical, Sydney, Australia). Only high quality measurements (> 80% operator index) were considered for analysis.

Ambulatory BP measurements were collected using validated oscillometric 90217A SpaceLabs (Spacelabs; Wokingham, Berkshire, UK) recorders and calibrated to take 4 measurements per hour for 24 hours. To calculate the ambulatory arterial stiffness index (AASI), the regression slope of ambulatory diastolic BP on ambulatory systolic BP from unedited 24 hour recordings, taken at a rate of 4 per hour, were computed for each participant. The 24 hour mean BP was calculated from the recordings. AASI was defined as 1 minus the regression slope. The stiffer the arterial tree, the closer the regression slope and AASI are to 0 and 1, respectively.<sup>19</sup> The BP dipping was defined as the degree of fall (%) in nocturnal meanarterial pressure relative to the diurnal mean BP:  $100 \times (1 - [\text{nighttime mean BP} \div \text{daytime mean BP}])$ .<sup>20</sup>

Pearson correlation coefficients evaluated univariate associations. Hierarchical multiple regression (HMR) analyses were conducted to test the relationship between anger and TF with cardiovascular parameters and to demonstrate the incremental contribution of TF from anger in accounting for variance in cardiovascular parameters. A priori alpha level of  $p < 0.05$  was considered to be significant. SPSS version 18.0 (SPSS Inc., Chicago, IL, USA) was used for all analyses.

## Results

Figure 1 shows how the participants were allocated across the 3 studies. Table 1 shows summary statistics for all continuous variables, including demographics, anger, TF, and cardiovascular parameters for Studies 1, 2, and 3.

In Study 1, one hundred thirty four participants ( $M_{\text{age}} = 21.31$  years,  $SD = 2.60$ ) qualified for study inclusion. Pearson correlations indicated no statistically significant associations ( $p <$

.05) between anger and TF scores with demographic or anthropometric characteristics. HMR analyses were performed in order to examine the unique relationship TF had with each of the beat-to-beat cardiovascular parameters while controlling for the influence of anger. HMR provided an analysis of the incremental contribution of TF scores above and beyond anger scores in accounting for variance in cardiovascular values. For each cardiovascular parameter serving as an outcome, Model 1 of the HMR contained anger as a predictor while Model 2 added TF as a predictor. Model 2 of the HMR analyses (see Table 2) indicated that anger, while controlling for TF, had significant relationships with all measured autonomic and cardiovascular outcomes with higher anger scores associated with higher HR, mean BP, nLF, and lower BRS. Model 2 of the HMR also indicated a significant relationship between TF scores and nLF (but not for HR, mean BP, or BRS) while controlling for anger. The addition of TF in Model 2 of the HMR analyses indicated that TF was able to uniquely predict 7% of the variance in nLF values; higher TF scores were associated with lower nLF.

In Study 2, eighty participants ( $M_{\text{age}} = 21.03$  years,  $SD = 2.42$ ) qualified for study inclusion. Inclusion criteria were identical to Study 1. Pearson correlations indicated no statistically significant associations ( $p < .05$ ) between anger and TF scores with demographic or anthropometric characteristics. Model 1 of the HMR contained anger as a predictor while Model 2 added TF as a predictor. Model 2 of the HMR analyses (see Table 3) indicated that anger, while controlling for TF, had significant relationships with both brachial and aortic hemodynamic pressures, STI, DTI and RPP, with higher anger scores associated with higher pressures, STI, DTI, and RPP. Model 2 of the HMR also indicated significant relationship (while controlling for anger) between TF scores and HR, STI, SVI, and RPP, with higher TF scores associated with less HR, STI, and RPP but greater SVI. The addition of TF in the Model 2 of the

HMR analyses indicated that TF was able to uniquely predict 10% of the variance in HR values, 14.2% of STI values, 12.6% in SVI values, and 9.8% in RPP values.

In Study 3, ninety four young participants ( $M_{\text{age}} = 20.99$  years,  $SD = 2.51$ ) qualified for study inclusion. Males were excluded from further analysis due to low participation rate ( $n = 6$ ). Pearson correlations indicated no statistically significant associations ( $p < .05$ ) between anger and TF scores with the demographic or anthropometric characteristics. Model 1 of the HMR contained anger and Model 2 added TF as a predictor. After controlling for anger, the addition of TF in the Model 2 of the HMR analyses (see Table 4) indicated that TF was able to uniquely predict 13% of the variance in 24 hour mean BP dipping scores with higher TF scores associated with greater mean BP dipping. Model 2 of the HMR also indicated that anger remained a significant predictor for 24 hour mean BP, mean BP dipping (mean  $BP_{dp}$ ), and AASI after controlling for TF scores with higher anger scores associated with higher 24 hour mean BP, higher AASI and less mean BP dipping.

### **Discussion**

Three studies were conducted to evaluate cardiovascular functioning underlying anger and TF in order to determine whether TF plays a unique role as a potential protective factor against the development of impaired cardiovascular functioning. The novel findings of the present studies are: (i) anger was associated with indices typical of increased hemodynamic and SNS activity, but TF was negatively related to nLF while controlling for anger, (ii) the effects of TF are unique from anger and are modulated via decreased ventricular work (STI) and ultimately decreased myocardial oxygen consumption (RPP), and (iii) anger remained a significant predictor for 24 hour mean BP, mean  $BP_{dp}$ , and AASI after controlling for TF scores, such that higher anger scores predicted higher mean BP, higher AASI and less mean BP dipping. These

results suggest that anger and TF have divergent effects on cardiovascular risk factors. To the best of our knowledge this is the first research to systemically address the impact of TF and anger on cardioprotection and cardiotoxicity, respectively.

Study 1 demonstrated that TF was associated with decreased sympathovagal tone supporting theoretical models proposed by Thoresen et al.<sup>21</sup> and Seybold et al.<sup>22</sup> to explain the relationship between TF, autonomic nervous system functioning, and health. Conversely anger was associated with cardiovascular autonomic deregulation including decreased BRS. It is worth noting that distention aortic pressure (aortic mean BP) and AASI have been linked to increased arterial stiffness and therefore may explain the anger induced decreases in BRS<sup>23</sup> Study 2 assessed aortic hemodynamics and demonstrated decreased ventricular work and decreased myocardial oxygen consumption corresponded to an increase in forgiveness. This constitutes the first empirical investigation into the cardiac mechanisms potentially responsible for the positive cardiovascular health associations with TF. Additionally, anger was associated with increased ventricular work and aortic BP suggesting that prior studies may have underestimated the impact of this psychological risk factor on cardiovascular function. Finally, Study 3 examined the relationship between anger, TF and cardiovascular functioning over a 24 hour period via ambulatory BP monitoring and found that TF may serve as a protective factor against future CVD owing to increased BP dipping. Strikingly, the cardioprotective effect sizes demonstrated by forgiveness in this research are similar to the effect sizes of known beta blockers.<sup>24,25</sup>

The novel nature of this research advances the understanding of the physiology underlying both anger and TF with analyses demonstrating that anger and TF are unique, independent predictors of autonomic and cardiovascular parameters. This novelty extends not only to the assessment techniques utilized (i.e. beat-to-beat BP, pulse wave analysis, ambulatory)

but more importantly to the discovery of new mechanisms that might account for the association between TF and health (i.e. cardiac autonomic modulation, ventricular work, myocardial oxygen consumption, nocturnal hemodynamics and BP dipping). As we view this research as an initial step into the search for physiological mechanisms associated with TF, future studies should emerge that examine stress hormones or blood catecholamine levels as additional mechanisms as they have been linked to SNS activity.<sup>26</sup> At a methodological level, future research utilizing longitudinal designs or manipulations inducing cardiovascular reactivity may replicate and greatly expand our findings. It should be noted, however, that designs examining the role that stress or anger plays in mediating the relationship between TF and cardiovascular health suggest that TF alone can uniquely account for decreased hemodynamic values.<sup>27</sup> Additionally, as there appear to be gender differences in TF<sup>28</sup> it is important for future research to examine the relationship between TF and male physiology as the current research is limited to female physiology. Furthermore, as this sample is restricted to young adults devoid of cardiovascular illness and in seemingly good health, the protective effect of TF may have been blunted or restricted.

1. Suls J. Anger and the heart: perspectives on cardiac risk, mechanisms and interventions. *Prog Cardiovasc Dis* 2013;55:538-547.
2. Chida Y, Steptoe A. The association of anger and hostility with future coronary heart disease: a meta-analytic review of prospective evidence. *J Am Coll Cardiol* 2009;53:936-946.
3. Williams JE, Paton CC, Siegler IC, Eigenbrodt ML, Nieto FJ, Tyroler HA. Anger proneness predicts coronary heart disease risk: prospective analysis from the atherosclerosis risk in communities (ARIC) study. *Circulation* 2000;101:2034-2039.
4. Friedberg JP, Suchday S, Shelov DV. The impact of forgiveness on cardiovascular reactivity and recovery. *Int J Psychophysiol* 2007;65:87-94.
5. Toussaint LL, Owen AD, Cheadle A. Forgive to live: forgiveness, health, and longevity. *J Behav Med* 2012;35:375-386.
6. Muller JE. Circadian variation in cardiovascular events. *Am J Hypertens* 1999;12:35S-42S.
7. Spielberger CD, Gorsuch RL, Lushene RE. Manual for the State-Trait Anxiety Inventory. Palo Alto, CA: Consulting Psychologists Press, 1970:1-26.
8. Brown RP. Measuring individual differences in the tendency to forgive: construct validity and links with depression. *Personality & social psychology bulletin* 2003;29:759-771.
9. Imholz BP, Wieling W, Langewouters GJ, van Montfrans GA. Continuous finger arterial pressure: utility in the cardiovascular laboratory. *Clin Auton Res* 1991;1:43-53.
10. Anon. Heart rate variability: standards of measurement, physiological interpretation and clinical use. Task Force of the European Society of Cardiology and the North American Society of Pacing and Electrophysiology. *Circulation* 1996;93:1043-1065.
11. Burr RL. Interpretation of normalized spectral heart rate variability indices in sleep research: a critical review. *Sleep* 2007;30:913-919.

- 12.** Westerhof BE, Gisolf J, Stok WJ, Wesseling KH, Karemaker JM. Time-domain cross-correlation baroreflex sensitivity: performance on the EUROBAVAR data set. *J Hypertens* 2004;22:1371-1380.
- 13.** Nichols WW, Singh BM. Augmentation index as a measure of peripheral vascular disease state. *Curr Opin Cardiol* 2002;17:543-551.
- 14.** Hashimoto J, Imai Y, O'Rourke MF. Indices of pulse wave analysis are better predictors of left ventricular mass reduction than cuff pressure. *Am J Hypertens* 2007;20:378-384.
- 15.** Roman MJ, Devereux RB, Kizer JR, Lee ET, Galloway JM, Ali T, Umans JG, Howard BV. Central pressure more strongly relates to vascular disease and outcome than does brachial pressure: the Strong Heart Study. *Hypertension* 2007;50:197-203.
- 16.** Gobel FL, Norstrom LA, Nelson RR, Jorgensen CR, Wang Y. The rate-pressure product as an index of myocardial oxygen consumption during exercise in patients with angina pectoris. *Circulation* 1978;57:549-556.
- 17.** Bunckberg GD, Fixler DE, Archie JP, Hoffman J. Experimental Subendocardial Ischemia in Dogs with Normal Coronary Arteries. *Circ Res* 1972;30:67-81.
- 18.** Tsiachris D, Tsioufis C, Syrseloudis D, Roussos D, Tatsis I, Dimitriadis K, Toutouzas K, Tsiamis E, Stefanadis C. Subendocardial viability ratio as an index of impaired coronary flow reserve in hypertensives without significant coronary artery stenoses. *J Hum Hypertens* 2012;26:64-70.
- 19.** Adiyaman A, Dechering DG, Boggia J, Li Y, Hansen TW, Kikuya M, Bjorklund-Bodegard K, Richart T, Thijs L, Torp-Pedersen C, Ohkubo T, Dolan E, Imai Y, Sandoya E, Ibsen H, Wang J, Lind L, O'Brien E, Thien T, Staessen JA. Determinants of the ambulatory arterial stiffness index in 7604 subjects from 6 populations. *Hypertension* 2008;52:1038-1044.

- 20.** Friedman O, Logan AG. Can nocturnal hypertension predict cardiovascular risk? *Integrated blood pressure control* 2009;2:25-37.
- 21.** Thoresen C, Harris A, Luskin F. Forgiveness and health: an unanswered question. *Forgiveness: Theory, research and practice*. New York: Guilford Press, 2000:254-280.
- 22.** Seybold K, Hill P, Neumann J, Chi D. Physiological and psychological correlates of forgiveness. *Journal of Psychology and Christianity* 2001;20:250-259.
- 23.** Wang MY, Huang CJ, Wu YL, Liu JC, Tsai PS. The influence of baroreflex sensitivity on ambulatory arterial stiffness index in individuals with cardiovascular risk. *Blood Press Monit* 2010;15:262-267.
- 24.** Williams B, Lacy PS, Thom SM, Cruickshank K, Stanton A, Collier D, Hughes AD, Thurston H, O'Rourke M. Differential impact of blood pressure-lowering drugs on central aortic pressure and clinical outcomes: principal results of the Conduit Artery Function Evaluation (CAFE) study. *Circulation* 2006;113:1213-1225.
- 25.** Williams B, Lacy PS. Impact of heart rate on central aortic pressures and hemodynamics: analysis from the CAFE (Conduit Artery Function Evaluation) study: CAFE-Heart Rate. *J Am Coll Cardiol* 2009;54:705-713.
- 26.** Henry JP. Biological basis of the stress response. *Integrative physiological and behavioral science : the official journal of the Pavlovian Society* 1992;27:66-83.
- 27.** Lawler-Row KA, Karremans JC, Scott C, Edlis-Matityahou M, Edwards L. Forgiveness, physiological reactivity and health: the role of anger. *Int J Psychophysiol* 2008;68:51-58.
- 28.** Miller AJ, Worthington EL, McDaniel MA. Gender and Forgiveness: A Meta-Analytic Review and Research Agenda. *Journal of Social and Clinical Psychology* 2008;27:843-876.

**Figure 1.** Participant allocation.

**Table 1.** Summarizes continuous variables, demographics statistics, and cardiovascular parameters for each study.

Study	1 (n = 134)	2 (n = 80)	3 (n = 94)
Variable (M ± SD)			
Age (years)	21.28 ± 2.61	21.01 ± 2.44	21.02 ± 2.57
Height (m)	1.67 ± 0.08	1.65 ± 0.07	1.63 ± 0.08
Weight (kg)	68.72 ± 15.85	24.57 ± 4.22	24.09 ± 4.17
BMI (kg/m <sup>2</sup> )	24.35 ± 4.26	78.41 ± 9.95	78.41 ± 9.95
State-Trait Anger	15.40 ± 1.59	15.90 ± 1.70	16.68 ± 1.05
Tendency to Forgive	12.78 ± 1.52	14.60 ± 2.67	13.77 ± 2.03
HR (bpm)	78.41 ± 9.95	74.53 ± 8.17	-
Mean BP (mmHg)	92.56 ± 7.28	83.82 ± 6.83	-
nLF	0.65 ± 0.09	-	-
BRS (ms/mmHg)	16.81 ± 10.94	-	-
Brachial systolic BP (mmHg)	-	114.38 ± 7.58	-
Brachial diastolic BP (mmHg)	-	68.86 ± 7.51	-
Aortic systolic BP (mmHg)	-	97.43 ± 6.40	-
Aortic diastolic BP (mmHg)	-	69.78 ± 7.62	-
Aortic mean BP (mmHg)	-	82.29 ± 6.68	-
STI (mmHg/s.min-1)	-	1546.90 ± 215.47	-
DT I (mmHg/s.min <sup>-1</sup> )	-	3392.82 ± 288.47	-
SVI (%)	-	223.33 ± 33.45	-
RPP (bpm x mmHg x100)	-	59.93 ± 9.62	-
AASI	-	-	0.28 ± 0.16
24-HR (bpm)	-	-	78.66 ± 8.69
24-mean BP (mmHg)	-	-	85.42 ± 6.08
Mean BP <sub>dp</sub> (%)	-	-	4.94 ± 7.37

Abbreviations: BMI, body mass Index; HR, heart rate; BP, blood pressure; nLF, normalized low frequency; BRS, baroreflex sensitivity; STI, systolic time interval; DTI, diastolic time interval; SVI, subendocardial viability index; RPP, rate pressure product; <sub>dp</sub>, dipping value; AASI, ambulatory arterial stiffness index; 24-HR, ambulatory 24-hour HR; 24-mean BP, ambulatory 24-hour mean BP.

**Table 2.** Study 1( n = 134): Hierarchical multiple regression analyses of anger and forgiveness scores on beat-to-beat cardiovascular indices.

Variable	Model	Predictors	$\beta$	<i>sr</i>	<i>p</i>	Model $R^2$	Model $\Delta R^2$	Model <i>F</i>
HR (bpm)	1	Anger	.31	.31	<.001	.10		F(1, 132) = 13.36, p < .001
	2	Anger	.33	.32	<.001	.11	.01	$\Delta F(1, 131) = 1.53, p = .219$
		Forgive	-.11	-.10	.219			
Mean BP (mmHg)	1	Anger	.23	.23	.007	.05		F(1, 132) = 7.43, p = .007
	2	Anger	.22	.21	.015	.06	.00	$\Delta F(1, 131) = 0.54, p = .464$
		Forgive	-.06	-.06	.464			
nLF	1	Anger	.58	.58	<.001	.34		F(1, 132) = 33.89, p < .001
	2	Anger	.56	.58	<.001	.41	.07	$\Delta F(1, 131) = 22.12, p < .001$
		Forgive	-.26	-.31	.009			
BRS (ms/mmHg)	1	Anger	-.18	-.18	.041	.03		F(1, 132) = 4.26, p = .041
	2	Anger	-.18	-.18	.045	.03	.00	$\Delta F(1, 131) = 0.01, p = .917$
		Forgive	.01	-.01	.917			

Note. *df* = 1, 133. HR, heart rate; BP, blood pressure; nLF, normalized low frequency; BRS, baroreflex sensitivity.

**Table 3.** Study 2 (n=80): Hierarchical multiple regression analyses of anger, forgiveness, and aortic hemodynamic indices.

Variable	Model	Predictors	$\beta$	<i>sr</i>	<i>p</i>	Model $R^2$	Model $\Delta R^2$	Model F
HR (bpm)	1	Anger	.148	.148	.229	.022		F(1, 78) = 1.47, <i>p</i> = .229
	2	Anger	.173	.173	.142	.122	.100	$\Delta F(1, 77) = 7.41, p = .008$
		Forgive	-.317	-.316	.008			
BSBP (mmHg)	1	Anger	.488	.488	<.001	.238		F(1, 78) = 20.63, <i>p</i> < .001
	2	Anger	.503	.501	<.001	.271	.033	$\Delta F(1, 77) = 2.94, p = .091$
		Forgive	-.182	-.182	.091			
BDBP (mmHg)	1	Anger	.693	.693	<.001	.480		F(1, 78) = 61.04, <i>p</i> < .001
	2	Anger	.704	.701	<.001	.497	.017	$\Delta F(1, 77) = 2.19, p = .144$
		Forgive	-.131	-.130	.144			
BMAP (mmHg)	1	Anger	.690	.690	<.001	.476		F(1, 78) = 59.87, <i>p</i> < .001
	2	Anger	.701	.702	<.001	.497	.021	$\Delta F(1, 77) = 2.72, p = .104$
		Forgive	-.145	-.200	.104			
ASBP (mmHg)	1	Anger	.613	.613	<.001	.376		F(1, 78) = 39.76, <i>p</i> < .001
	2	Anger	.620	.618	<.001	.384	.008	$\Delta F(1, 77) = .80, p = .373$
		Forgive	-.088	-.087	.373			
ADBP (mmHg)	1	Anger	3.18	.694	<.001	.482		F(1, 78) = 61.37, <i>p</i> < .001
	2	Anger	3.23	.702	<.001	.497	.015	$\Delta F(1, 77) = 1.91, p = .171$
		Forgive	-.635	-.122	.171			
AMAP (mmHg)	1	Anger	.690	.690	<.001	.476		F(1, 78) = 59.87, <i>p</i> < .001
	2	Anger	.701	.699	<.001	.497	.021	$\Delta F(1, 77) = 2.72, p = .104$
		Forgive	-.145	-.145	.104			
STI (mmHg/s.min <sup>-1</sup> )	1	Anger	.513	.513	<.001	.263		F(1, 78) = 23.56, <i>p</i> < .001
	2	Anger	.543	.542	<.001	.405	.142	$\Delta F(1, 77) = 15.54, p < .001$
		Forgive	-.378	-.377	<.001			
DTI (mmHg/s.min <sup>-1</sup> )	1	Anger	.575	.575	<.001	.330		F(1, 78) = 32.56, <i>p</i> < .001
	2	Anger	.569	.567	<.001	.335	.005	$\Delta F(1, 77) = .497, p = .483$
		Forgive	.072	.071	.483			
SVI (%)	1	Anger	-.155	-.155	.206	.024		F(1, 78) = 1.63, <i>p</i> = .206
	2	Anger	-.184	-.183	.114	.150	.126	$\Delta F(1, 77) = 9.64, p = .003$
		Forgive	.356	.355	.003			
RPP (bpm*mmHg)	1	Anger	.379	.379	.001	.144		F(1, 78) = 11.10, <i>p</i> = .001
	2	Anger	.405	.403	<.001	.242	.098	$\Delta F(1, 77) = 8.42, p = .005$
		Forgive	-.314	-.313	.005			

Abbreviations: HR, heart rate; BSBP, brachial systolic blood pressure; BDBP, brachial diastolic blood pressure; BMAP, brachial mean arterial pressure; ASBP, aortic systolic blood pressure; ADBP, aortic diastolic blood pressure; AMAP, aortic mean arterial pressure; STI, systolic time interval; DTI, diastolic time interval; SVI, subendocardial viability index; RPP, rate pressure product; *sr*, semi-partial correlation.

**Table 4.** Study 3 (n=94): Hierarchical multiple regression analyses of ambulatory hemodynamic indices.

Variable	Model	Predictors	$\beta$	<i>sr</i>	<i>p</i>	Model $R^2$	Model $\Delta R^2$	Model F
24-HR (bpm)	1	Anger	.12	.12	.250	.01		F(1, 92) = 1.34, <i>p</i> = .250
	2	Anger	.10	.10	.325	.02	.01	$\Delta F(1, 91) = 0.86, p = .357$
		Forgive	-.10	-.10	.357			
24-mean BP (mmHg)	1	Anger	.40	.40	<.001	.16		F(1, 92) = 16.97, <i>p</i> < .001
	2	Anger	.41	.41	<.001	.16	.01	$\Delta F(1, 91) = 0.94, p = .335$
		Forgive	-.09	-.09	.335			
MAP <sub>dp</sub> (%)	1	Anger	-.39	-.39	<.001	.15		F(1, 92) = 12.37, <i>p</i> < .001
	2	Anger	-.39	-.39	<.001	.28	.13	$\Delta F(1, 91) = 11.98, p < .001$
		Forgive	.36	.36	<.001			
AASI	1	Anger	.23	.23	.032	.06		F(1, 92) = 4.76, <i>p</i> = .041
	2	Anger	.24	.24	.030	.06	.00	$\Delta F(1, 11) = 0.18, p = .667$
		Forgive	-.05	-.05	.669			

Abbreviations: 24-HR, 24-hour heart rate; 24-mean BP, 24-hour mean BP; MAP<sub>dp</sub>, mean arterial pressure dipping percentage; AASI, ambulatory arterial stiffness index.

