1	The longitudinal relationship between cortisol responses to mental stress and
2	leukocyte telomere attrition
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25 Abstract

26 Context: Chronic psychological stress has been associated with shorter telomeres in some studies, but 27 the underlying mechanisms are poorly understood. One possibility is that the neuroendocrine responses 28 associated with stress exposure are involved. 29 **Objective:** To testing the hypothesis that greater cortisol responsivity to acute stressors predicts more 30 rapid telomere attrition. 31 Design: We measured salivary cortisol responses to two challenging behavioral tasks. Leukocyte 32 telomere length was measured at the time of mental stress testing and 3 years later. 33 Participants: We studied 411 initially healthy men and women aged 54-76 years. 34 Main outcome measure: Leukocyte telomere length. 35 **Results:** Cortisol responses to this protocol were small, we divided participants into cortisol 36 responders (n = 156) and non-responders (n = 255) using a criterion (\geq 20%) previously shown to 37 predict increases in cardiovascular disease risk. There was no significant association between cortisol 38 responsivity and baseline telomere length, although cortisol responders tended to have somewhat 39 shorter telomeres ($\beta = -0.061$, standard error 0.049). But cortisol responders had shorter telomeres and 40 more rapid telomere attrition than non-responders on follow-up, after controlling statistically for age, gender, socioeconomic status, smoking, time of day of stress testing and baseline telomere length ($\beta = -$ 41 42 0.10, standard error 0.046, p = 0.029). The association was maintained after additional control for 43 cardiovascular risk factors ($\beta = -0.11$, p = 0.031). The difference between cortisol responders and non-44 responders was equivalent to approximately 2 years in aging. 45 **Conclusions:** These findings suggest that cortisol responsivity may mediate in part the relationship 46 between psychological stress and cellular aging.

47

49 Introduction

Telomeres are complexes of DNA and proteins situated at the ends of chromosomes that protect the genomic DNA of eukaryotic cells (1). Telomeres shorten with each cell division, and telomere length is a marker of cellular aging. Telomere function is impaired when shortening becomes critical, leading to cell senescence, genome instability and apoptosis. Leukocyte telomere length is associated with increased risk of cardiovascular disease, cancers, diabetes, dementia and all-cause mortality (2-4). These relationships have been confirmed by studies of inherited telomere syndromes (5), and by Mendelian randomization studies (6).

57 Several environmental and lifestyle factors are associated with telomere shortening, including 58 smoking, obesity and physical inactivity (7). There is growing interest in the relationship of leukocyte 59 telomere length with psychiatric conditions and psychological stress as well. Large scale investigations 60 indicate that individuals with major depressive disorder have shorter telomeres independently of 61 demographic factors and health behaviors, although findings across studies have been variable (8). 62 Anxiety disorders may also be associated with reduced telomere length (8), while a meta-analysis of 22 63 studies documented a small statistically significant relationship between greater perceived stress and 64 shorter telomeres (9). Exposure to early life adversity has been linked with reduced telomere length in some studies (10), but not in all (11). Associations with low social support (12) and hostility (13) have 65 66 also been described.

Evaluation of the importance of links between stress exposure, mental health, and telomere
dynamics would be strengthened by better understanding of potential underlying mechanisms.
Unhealthy habits such as smoking, excessive alcohol consumption and inactivity might play a role, but
many studies have observed associations with leukocyte telomere length after these factors have been
taken into account (8,9,14). The physiological responses associated with mental stressors may also be
involved. Cortisol plays a central role in the stress response because of its multiple effects on immune,

73 metabolic, and vascular processes . Animal studies indicate that embryonic exposure to corticosteroids 74 elicits increased oxidative stress and shorter telomeres in later life (15). There are large individual 75 differences in the magnitude of cortisol responses to standardized mental stress tests, and these reflect 76 variations in the capacity of neuroendocrine regulatory processes to adapt to challenge. A small number 77 of studies have shown that larger cortisol responses to mental stress are associated with shorter 78 telomeres in adults and children (16-18). For example, Tomiyama et al (19) administered a 79 standardized mental stress protocol to 28 caregivers for people with Alzheimer's disease and controls, 80 and found that telomeres were shorter in individuals who manifest greater cortisol stress responses. 81 However, these studies of telomeres and stress physiology have been cross-sectional. It is possible that 82 heightened cortisol responsivity drives telomere attrition, or conversely that greater cortisol responses 83 are characteristic of people with shorter telomeres. Null associations have also been described (20). 84 In the present study, we evaluated the relationship between cortisol responses to mental stress 85 and differences in telomere length measured at the time of mental stress testing and three years later. 86 We tested the hypothesis that cortisol stress responders would show greater telomere attrition over time 87 than non-responders. This hypothesis was examined in a sample of healthy men and women aged 54-88 76, since biological aging processes are particularly relevant to disease risk as people progress into 89 older age. We used a measure of cortisol responses to mental stress tests that has previously been 90 shown to predict the progression of subclinical coronary atherosclerosis as indexed by coronary 91 calcification (21), and the development of hypertension (22). Our analyses also took into account 92 sociodemographic and physiological factors that might also contribute to telomere shortening over 93 time.

94

95 Materials and Methods

96 **Participants**

97 We analyzed data from the Heart Scan Study, a sample of 543 men and women of white European 98 origin of the Whitehall II epidemiological cohort recruited between 2006 and 2008 to investigate 99 physiological responsivity to mental stress testing and subclinical coronary artery disease. Participants 100 were selected as having no history of coronary heart disease, and no previous diagnoses or treatment 101 for hypertension, diabetes, inflammatory diseases, or allergies. We used civil service employment 102 grade as an indicator of socioeconomic status (SES), and recruitment was stratified to include men and 103 women from higher, intermediate and lower employment grades. The women in the study were 104 postmenopausal. Participants were invited for reassessment 3 years after mental stress testing (mean 105 1087 days interval). Ethical approval was obtained from the University College London Hospital 106 Committee on the Ethics of Human Research, and all participants gave signed informed consent. All 107 procedures were carried out in accordance with approved guidelines. 108 Figure 1 shows a flow chart summarizing participant progression through the study. Telomere 109 length was measured in 501 (92.3%) respondents an average 36.2 months after stress testing. Of these, 110 411 also had telomere length measures at the time of stress testing, since assessments were not 111 introduced at the start of data collection. They constitute the sample for this study. There were no

differences on any measures between individuals included and not included in the telomere lengthanalyses.

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115 Laboratory mental stress testing

We tested participants individually in a light and temperature- controlled laboratory, with sessions beginning either in the morning at 8:30-9:30, or in the early afternoon at 13:30-14:30. Participants were instructed not to drink caffeinated beverages or smoke for at least 2h before testing and to avoid vigorous exercise and alcohol from the previous evening, and not to have taken any anti-inflammatory or anti-histamine medication for the 7 days before testing. They were rescheduled if they reported colds 121 or other infections on the day of testing. At the start of the session, we measured height, weight, waist 122 and hip circumference using standardized techniques, and body mass index (BMI) was computed. After 123 a 30 min rest period, baseline blood pressure (BP) was measured with an automated UA-779 digital 124 monitor, a blood sample was drawn, and a saliva sample was taken using salivettes (Sarstedt, Leicester, 125 UK). Two behavioral tasks designed to induce mental stress were then administered in random order 126 (21,23). Both tasks were performed for 5 min. One was a computerized version of the Stroop color-127 word interference task which involved successive presentation of target color words (e.g. red, blue) 128 printed in another color. Four names of colors printed in incongruous colors at the bottom of the 129 computer screen, and participants were requested to press the computer key that corresponded to the 130 position at the bottom of the screen of the name of the color in which the target word was printed. The 131 rate of presentation of stimuli was adjusted to the performance of the participant in order to ensure 132 sustained demands. The second task was mirror tracing, which involved tracing with a metal stylus a 133 star that could only be seen in mirror image. Each time the stylus came off the star a mistake was 134 registered and a loud beep was emitted by the apparatus (Lafayette Instruments Corp., Lafayette, IN, USA). Participants were told that the average person could complete five circuits of the star in the 135 136 available time. These tasks were selected because they have been shown to stimulate similar appraisals 137 of involvement and engagement from participants across the social gradient. A second saliva sample 138 was taken immediately after tasks, with further samples at 20, 45 and 75 min after tasks.

139

140 **Biological measures**

Saliva samples were analyzed for cortisol concentration using a time resolved immunoassay with
fluorescence detection, at the Technical University Dresden, as described previously (24,25). The intraand inter-assay coefficients of variation were less than 8%. Total and high density lipoprotein (HDL)
cholesterol were measured in serum stored at 4°C within 72 h using enzymatic colometric methods.

145	Glycated hemoglobin was measured using Tosoh G7 HPLC analyzer calibrated to Diabetes Control
146	and Complications Trial (DCCT) standards. An adaptation of the method first described by Cawthon
147	(26) was used for the assessment of leukocyte telomere length. Genomic DNA was extracted from
148	peripheral blood mononuclear cells (PBMCs) in a QIAcube workstation (baseline) or manually
149	(follow-up) with the QIAamp DNA blood mini kit (Qiagen, Crawley, United Kingdom) according to
150	instructions of the manufacturer and stored in 10 mmol/L Tris-hydrochloride, 0.5 mmol/L
151	ethylenediamine tetraacetate, pH 9.0 at -20°C (baseline) or -80°C (follow-up). Relative mean TL was
152	measured by a monochrome multiplex quantitative real-time polymerase chain reaction (PCR) assay
153	with a Bio-Rad CFX96 Real-Time PCR Detection System (Bio-Rad, Hemel Hempstead, United
154	Kingdom) for samples obtained at the time of mental stress testing, and with a Roche Lightcycler 480
155	real-time PCR machine (Roche Diagnostics Corporation, Indianapolis, IN) on follow-up (27).
156	Reactions containing serial dilutions of a reference DNA standard were included in each polymerase
157	chain reaction plate to generate the telomere (T) and β -globin gene (S) standard curves required for
158	quantitation, and relative mean TL, expressed as a T/S ratio, was derived. The coefficient of variation
159	of these assays was 2.3%.

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161 **Data reduction and statistical analysis**

The mental stress protocol in this study did not generate large cortisol responses, with many respondents not showing an increase following tasks. Cortisol stress responsivity was therefore quantified by calculating differences scores between the baseline cortisol concentration and the samples obtained both immediately after tasks and 20 minutes later. Individuals who showed $a \ge 1$ nmol/L increase (equivalent to a 20% increase) between baseline and either sample were defined as cortisol responders, and the remainder as non-responders. Differences between the responder groups at baseline were analyzed using analysis of variance and χ^2 methods for continuous and categorical variables

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169 respectively. The cortisol profiles across the mental stress testing session of responder and non-170 responder groups were compared using repeated measures analysis of variance with sample as the 171 within-person factor and responder status as the between-person factor. Associations between cortisol 172 stress responsivity and telomere length at baseline were analyzed using multivariable regression, 173 including age, gender, grade of employment, smoking status and time of stress testing (morning or 174 afternoon) as covariates. A similar method was used to analyze associations between cortisol stress 175 responsivity and follow-up telomere length, except in this case baseline telomere length was included 176 as a covariate. Results are presented as standardized regression coefficients (β) with standard errors. 177 In a sensitivity analysis, we added cardiovascular risk factors (systolic BP, BMI, total and HDL 178 cholesterol, and glycated hemoglobin) to the model; these factors were not included in the main model 179 since missing data on some variables reduced the sample size. 180 Absolute measures of telomere length can vary across laboratories, but rankings of relative 181 length are highly correlated (28). In view of the different systems used at baseline and follow-up, we

182 therefore computed standardized telomere length scores for the two time points. However, repeating 183 the analyses with standardized as opposed to absolute values generated identical statistical findings, so 184 the latter are presented in the Results section.

185

186 **Results**

The 411 participants included 156 cortisol responders and 255 non-responders. The characteristics of these two groups are summarized in Table 1. Participants generally had favorable risk profiles, with few smokers, blood pressure and glycated hemoglobin in the healthy range, and no marked elevation of BMI or cholesterol. There were no differences in any sociodemographic or physiological factors between the two groups. There was a non-significant tendency of cortisol responders to be more likely to have undertaken mental stress testing in the afternoon compared with non-responders (p = 0.096), so time of day was included as a covariate in the analyses.

194 Cortisol concentrations in the responders and non-responders to behavioral challenge are shown 195 in Figure 2. There was a robust interaction between responder group and trial (p < 0.001). It can be 196 seen that cortisol concentrations were similar in the two groups at baseline. But while the responder 197 group showed an average 47% increase in salivary cortisol after tasks, values declined steadily in the 198 non-responder group. Even 75 min after mental stress tests had been completed, cortisol concentration 199 remained more than 30% higher in the responder than non-responder groups.

The mean T/S ratio averaged 0.992 ± 0.07 at baseline, and 0.894 ± 0.15 at follow-up. This indicates a significant decrease in telomere length over the 3 year interval (p < 0.001). Telomere lengths at the two time points were moderately correlated (r = 0.31, p < 0.001). There was a small positive association between baseline telomere length and change over time (r = 0.20), indicating that participants with longer telomeres showed greater shortening. Telomere length on follow-up was inversely associated with age (p < 0.001), and was shorter in men than women (p < 0.001).

206 The relationship between cortisol stress responsivity and telomere length at baseline was 207 negative, though not significant (($\beta = -0.061$, SE = 0.049, p = 0.22). But we found that cortisol stress 208 responsivity was associated with shorter telomere length on follow-up after adjustment for baseline 209 telomere length, age, gender, grade of employment, smoking status and time of stress testing ($\beta = -$ 210 0.10, SE = 0.046, p = 0.029). The other independent predictors of shorter telomeres on follow-up were 211 older age, male sex, and shorter telomere length at baseline. Figure 3 illustrates the pattern of change in 212 telomere length over time in cortisol responders and non-responders to stressors, showing the greater 213 shortening over time in stress responders. There was no interaction between time of stress testing and 214 cortisol responsivity in predicting telomere length on follow-up.

The association was unchanged in the sensitivity analysis which included baseline systolic BP, BMI, total and HDL cholesterol, glycated hemoglobin, and time interval between baseline and followup; the regression coefficient for cortisol responsivity was (n = 378, β = -0.11, SE = 0.049, *p* = 0.031).

218

219 **Discussion**

220 In this study, we tested the notion that cortisol responses to mental stress would be associated with the 221 rate of telomere attrition over time. We found that healthy late middle-aged men and women who 222 responded to standardized behavioral challenges with larger increases in salivary free cortisol showed 223 greater shortening of leukocyte telomeres over a 3 year period. This association was independent of 224 baseline telomere length, age, gender, socioeconomic status (SES) defined by grade of employment, 225 smoking, cardiovascular risk factors (blood pressure, cholesterol, BMI, glycated hemoglobin) and 226 length of follow-up. The difference in telomere attrition between cortisol responders and non-227 responders corresponded to 107 base pairs on follow-up, indicating a difference of approximately two 228 years in aging (29).

229 The cortisol responses during mental stress testing in this study were small. A major purpose of 230 the study from which these data were drawn was to evaluate SES differences in stress reactivity and 231 recovery (23). Consequently, the task protocol was designed to be perceived as equally stressful across 232 the SES spectrum, and was selected after pretesting on this criterion. It did not involve socially 233 evaluative tasks such as the Trier Stress Test (TSST) that are known to elicit large cortisol responses 234 (30), since such tasks are often appraised differently by higher and lower social status individuals, 235 compromising any differences in physiological responsivity. The range of individual differences as 236 well as absolute magnitude of cortisol responses was therefore smaller than in some other 237 investigations. However, the value of the cortisol responder categorization adopted here has been 238 endorsed by evidence that individuals classified as cortisol responders show an increased risk of

incident hypertension (22) as well as more rapid progression of subclinical coronary artery disease as indexed by coronary artery calcification (21). Brief cortisol responses to short-term tasks are of little significance in themselves. However, the magnitude of acute cortisol responses is positively associated with cortisol output in everyday life (31). If these responses are representative of people's habitual profile of cortisol when confronted by the challenges of everyday life, they may contribute to chronic neuroendocrine activation that could have deleterious health consequences.

245 Research relating telomere length with measures of cortisol output at rest have produced mixed 246 results (32,33), suggesting that relating individual differences in cortisol responses to standardized 247 mental stress with telomere length may be a valuable strategy. Epel et al (16) found that urinary cortisol 248 concentration collected over a night following a behavioral stress battery was inversely associated with 249 telomere length in healthy women. A study of older female caregivers of partners with dementia 250 showed relationships between telomere length and cortisol responses to behavioral challenge (19), 251 while work with children as young as 5 to 6 years has demonstrated that cortisol reactivity to mildly 252 stressful tasks is inversely correlated with telomere length (17,18). By contrast, a study of older men 253 and women in Finland showed no associations between telomere length and cortisol responses to acute 254 stress exposure, but is difficult to interpret since stress testing took place an average 2.1 years after 255 telomere assays (20). Our study builds on these findings by establishing a longitudinal relationship, 256 since cortisol responsivity predicted telomere shortening over time. The results are also consistent with 257 longitudinal clinical studies indicating that telomere length is shorter during active Cushing's syndrome 258 than when patients are in remission (34).

A puzzling feature of our results is that no association was present between cortisol responsivity and telomere length at baseline. There was a negative association between cortisol responsivity and baseline telomere length, but it was not significant. It is potentially relevant is that the studies of adults that have shown associations between cortisol responsivity and telomere length have focused on individuals exposed to chronic stressors such as caregiving or having children with severe disabilities
(16,19). No association has previously been observed in general population samples of the type
involved in the present study (20). It is possible that in our sample of relatively healthy older men and
women, these associations only emerged after several years.

We found a positive correlation between baseline telomere length and the magnitude of the change in length over time. Regression to the mean has been put forward as the explanation of this phenomenon (35). However, regression to the mean is unlikely to be the explanation for the association with cortisol stress responsivity, since if anything, cortisol responders had slightly shorter telomeres at baseline. Regression to the mean would therefore operate against the effects observed here.

272 The mechanisms underlying these associations have yet to be defined in detail. Telomere length is regulated dynamically and does not decrease monotonically with advancing age (1). Faster telomere 273 274 attrition over time may result from several causes, including the expansion of leukocyte subsets that 275 occurs during inflammation and immunological responses, a decrease in telomerase activity, and 276 oxidative stress (27). Although cortisol responses might be expected to inhibit inflammation, 277 simultaneous heightened inflammation and cortisol is common in response to behavioral stress. A 278 reason for this might be because glucocorticoids have proinflammatory effects under some 279 circumstances. In vitro administration of glucocorticoids induces cytokine overexpression and NF-KB 280 activation in isolated macrophages (36), while pre-treatment with cortisol has been found to enhance 281 interleukin 6 responses to endotoxin (37). Cortisol administration in vitro also appears to reduce 282 telomerase activity (38). Frank, Watkins and Maier (39) have proposed that glucocorticoid responses to 283 stress may be neuroendocrine warning signals to the innate immune system, sensitizing 284 neuroinflammatory processes even after the corticosteroid response has dissipated. The combined 285 effect of reduced telomerase activity and oxidative stress would impinge negatively on the maintenance of telomere length, particularly in the context of chronic inflammation, thus providing a plausibleexplanation for the current findings.

288 This study has a number of limitations. The participants were middle-aged and older white 289 European men and women with no serious chronic illness, and results may not generalize to other 290 groups. Telomere length was measured in PBMCs, and values may differ in lymphocyte 291 subpopulations. Measures were also made with two different PCR machines at the two time points; 292 although this might affect comparisons of absolute values on the two occasions, it does not affect the 293 relative changes that are central to these results, so findings were the same with standardized measures 294 of telomere length. The cortisol responses were less substantial than those recorded with socially-295 evaluative stress testing, reducing the variability in responsivity profiles. We did not include a no stress 296 control group in this study, since we have previously found that the measurement protocol itself does 297 not induce physiological responses (40).

298 A strength of the study is that our findings were obtained in a well characterized longitudinal 299 population cohort, with a rather larger sample than has previously evaluated cortisol responses to acute 300 mental stress and telomere length. The results may have implications for understanding the pathways 301 through which social-environmental factors and mental ill-health impact cellular aging. If associations 302 between stress exposure and mental distress and telomere length are mediated through cortisol 303 responsivity, it is possible that the effects of mental stress on cellular aging might be reduced not only 304 by modifying stress exposure (which is not necessarily practical), but also by attenuating the 305 physiological components of the stress response.

In conclusion, the results of this study strongly suggest that heightened cortisol responsivity to psychological stress is associated with accelerated cellular aging as indexed by leukocyte telomere length. This indicates that heightened cortisol responsivity is not simply a consequence of more advanced cellular aging, but may contribute to the cellular aging process.

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434Table 1Characteristics of cortisol responders and non-responders435

Means \pm standard deviations

Variable	Non-responders (n = 255)	Responders (n = 156)	P-value	
Age (years)	63.1 ± 5.6	63.6 ± 5.7	0.36	
Men (%)	47.5	48.1	0.52	
Grade of employment (%) Higher Intermediate Lower	39.6 34.5 25.9	30.8 44.9 24.4	0.36	
Current smoker (%)	6.3	5.8	0.51	
Baseline systolic BP (mmHg)	124.8 ± 14.5	126.8 ± 15.4	0.18	
Body mass index (kg/m ²)	25.7 ± 4.3	26.1 ± 3.7	0.26	
Total cholesterol (mmol/l)	5.33 ± 0.95	5.34 ± 0.91	0.89	
HDL cholesterol (mmol/L)	1.70 ± 0.47	1.66 ± 0.47	0.72	
Glycated hemoglobin (%) mmol/mol	5.48 ± 0.39 36.3	$\begin{array}{c} 5.46 \pm 0.40\\ 36.2 \end{array}$	0.76	
Stress testing in afternoon (%)	57.3	66.0	0.096	
Follow-up interval (days)	1073 ± 62.6	1068 ± 73.3	0.48	

Table 2

Unstandardized and standardized regression coefficients (β) with standard error in parentheses

Predictor:	В	β (s.e.)	р
	0.001		0.020
Cortisol stress responsivity	-0.031	-0.10 (0.046)	0.029
Age	-0.005	-0.19 (0.047)	< 0.001
Gender	0.055	0.18 (0.046)	< 0.001
Grade of employment	0.009	0.05 (0.046)	0.32
Smoking status	0.013	0.02 (0.046)	0.66
Time of stress testing	-0.004	-0.01 (0.047)	0.77
Baseline telomere length	0.560	0.28 (0.046)	< 0.001

Predictors of follow-up leukocyte telomere length

448 **Figure Legends**

449 Figure 1 Flow chart of study participation.

450

451	Figure 2	Mean salivary cortisol concentration at baseline, immediately after behavioral tasks
452		(post-task), and 20 (+20 min), 45 (+45 min), and 75 (+75 min) minutes after tasks in
453		cortisol responders (solid line) and cortisol non-responders (dashed line). Error bars are
454		standard errors of the mean (s.e.m.).
455		
456	Figure 3	Mean telomere length (T/S ratio) in cortisol stress responders (solid line) and non-
457		responders (dashed line) at baseline and 3 year follow-up. Values are adjusted for age,
458		gender, grade of employment, smoking status and baseline telomere length. Error bars
459		are s.e.m. Telomere length is significantly different in cortisol responder and non-
460		responder groups at follow-up ($p = 0.016$).
461		

462 Figure 1463



