1	Experimental evidence that female rhesus macaques (Macaca mulatta) perceive
2	variation in male facial masculinity
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4	Kevin A. Rosenfield ^{1,2*} , Stuart Semple ¹ , Alexander V. Georgiev ^{3,4} , Dario Maestripieri ⁴ ,
5	James P. Higham ⁵ , Constance Dubuc ^{5,6}
6	
7	¹ Centre for Research in Evolutionary, Social and Interdisciplinary Anthropology,
8	University of Roehampton, Holybourne Avenue, London SW15 4JD, UK
9	
10	² Department of Anthropology, Pennsylvania State University, 409 Carpenter Building,
11	University Park, PA 16802, USA
12	
13	³ School of Natural Sciences, Bangor University, Bangor, Gwynedd, LL57 2UW, UK
14	
15	⁴ Institute for Mind and Biology, The University of Chicago, 940 East 57th St, Chicago,
16	IL 60637, USA
17	
18	⁵ Department of Anthropology, New York University, 25 Waverly Place, New York, NY
19	10003, USA
20	
21	⁶ Department of Zoology, University of Cambridge, Downing Street, Cambridge CB2
22	3EJ, UK
23	*To whom correspondence should be addressed. E-mail: <u>kar61@psu.edu</u>

24 Abstract

25 Among many primate species, face shape is sexually dimorphic, and male facial 26 masculinity has been proposed to influence female mate choice and male-male 27 competition by signalling competitive ability. However, whether conspecifics pay 28 attention to facial masculinity has only been assessed in humans. In a study of free-29 ranging rhesus macaques, *Macaca mulatta*, we used a two-alternative look-time 30 experiment to test whether females perceive male facial masculinity. We presented 107 31 females with pairs of images of male faces – one with a more masculine shape and one 32 more feminine – and recorded their looking behaviour. Females looked at the masculine 33 face longer than at the feminine face in more trials than predicted by chance. Although 34 there was no overall difference in average look-time between masculine and feminine 35 faces across all trials, females looked significantly longer at masculine faces in a subset 36 of trials for which the within-pair difference in masculinity was most pronounced. 37 Additionally, the proportion of time subjects looked toward the masculine face increased 38 as the within-pair difference in masculinity increased. This study provides evidence that 39 female macaques perceive variation in male facial shape, a necessary condition for 40 intersexual selection to operate on such a trait. It also highlights the potential impact of 41 perceptual thresholds on look-time experiments.

42

43 Keywords

44 Sexual dimorphism, mate choice, facial masculinity, look-time experiment

45 Introduction

46	Sexual selection can shape the evolution of male secondary sex characters through the
47	processes of intra- or intersexual selection, commonly associated with male-male contest
48	competition and female mate choice, respectively [1]. Although intra-sexual and
49	intersexual selection were initially believed to be independent evolutionary processes [1],
50	a growing body of evidence now indicates that traits initially shaped by intrasexual
51	selection - such as badges of dominance status, agonistic displays, large body size and
52	weapons - can sometimes be used secondarily by females as cues or signals of male
53	physical strength and competitive ability, allowing them to select optimal mating partners
54	or avoid coercive males [2–4]. As long as inter-male variation in such traits can be
55	perceived, females might be able to use them in their mating decisions.
56	In humans, there is good evidence that facial masculinity is associated with male-
57	male competition: facial masculinity has been found to be positively associated with
58	physical strength [5], testosterone levels [6,7; but see 8, in which no link was found, 9, in
59	which testosterone reactivity to competition, but not baseline testosterone levels, were
60	related to facial masculinity, and 10, in which neither reactivity nor baseline levels were
61	related to facial masculinity], aggressiveness [11,12], and unethical behaviour (propensity
62	to deceive in negotiation and cheat to increase financial gain) [13]. There is also indirect
63	evidence that facial masculinity predicts fitness, being negatively associated with the
64	probability of dying from contact aggression [14] and positively associated with number
65	of short-term mating partners [15].perceived facial masculinity and dominance are
66	closely linked [5,16], and recent research has shown that humans find viewing male faces

67	rated as dominant as more rewarding, even when ratings of facial attractiveness are
68	statistically controlled [17,18]. Sexually dimorphic face shape is not merely a result of
69	ontogenetic scaling [19], suggesting that it may have been under selection independently
70	of body size. Importantly, variation in facial masculinity is perceived by the human
71	sensory system: it can be used to assess competitive ability [5,16], and more masculine
72	faces appear to be more attractive to women, at least during the fertile phase of the
73	menstrual cycle [5,20,21]. Together, this suggests that in humans, facial masculinity is
74	under either intra- or intersexual selection, or both.
75	Previous research has shown that primates pay great attention to conspecifics' faces
76	[22–24]. Facial shape is sexually dimorphic in many primate species (e.g. collared
77	mangabeys, Cercocebus torquatus: [25]; rhesus macaques, Macaca mulatta: [26]; tufted
78	capuchins, Sapajus apella: [27,28]; papionins: [29]), and, as in humans, this is not just a
79	consequence of sexual dimorphism in body size [29]. There is evidence that male facial
80	masculinity plays a role in male-male contest competition in tufted capuchins, Sapajus
81	apella: in this species, there is a positive association between male facial masculinity
82	(facial width-to height ratio) and both dominance rank [27] and assertiveness [27,30].
83	Finally, facial masculinity may be associated with greater bite strength in male primates
84	[19]. While there is evidence that other facial features are perceived and used for
85	individual recognition and social decision-making in primates [31–33], whether inter-
86	individual variation in facial masculinity is perceived by conspecifics is unknown.
87	In this study, we used an experimental approach to investigate whether free-
88	ranging female rhesus macaques perceive variation in male facial masculinity. In this

89	species, sexual dimorphism in facial features [26] may be associated with bite strength
90	[19]; under the assumption that bite strength is associated with success in contest
91	competition and may reflect overall body strength, facial masculinity thus may serve as a
92	cue of male quality or formidability to females. Therefore, we hypothesized that females
93	would show a visual preference for more masculine male faces. Previous research using
94	looking-time experiments has demonstrated that when conspecific faces are presented
95	alongside other types of stimuli, such as seashells or heterospecific faces, primates show
96	a strong conspecific bias [34-39]. To test this hypothesis, we presented adult females
97	with pairs of photographs of faces of adult males, whose facial masculinity we quantified,
98	in order to test two predictions: (1) females will have a higher overall looking time
99	towards the more masculine male face of the pair, and (2) the proportion of time spent
100	looking at the more masculine face will be positively related to the difference in
101	masculinity between the two faces presented.
102	

103 Methods

104 *Study population*

105 We studied rhesus macaques on Cayo Santiago, a 15.2-hectare island 1 km off the eastern

106 coast of Puerto Rico, managed by the Caribbean Primate Research Centre (CPRC) of the

107 University of Puerto Rico. The population of ca. 1,300 - 1,400 macaques living on the

- island at the time of the study is descended from a group of 409 individuals brought from
- 109 India in 1938 [40]. Animals are individually recognizable, with tattoos providing a

unique ID and ear notches given when they are yearlings. Dates of birth of all animals areavailable from long-term records.

112

113 Facial sexual dimorphism measurement

114 To quantify sexual dimorphism in face shape, we measured facial images of male 115 (N=69) and female (N=27) rhesus macaques, collected during the 2012 and 2013 mating 116 season following a previously described method [41]. Multiple images of males were 117 captured in RAW format from 1–3 m away from subjects using a calibrated Canon EOS 118 Rebel T2i camera with an 18-megapixel CMOS APS sensor and an EF-S 55–250 mm 119 f/4-5.6 IS lens. In order to obtain an image of the male looking straight at the camera, we 120 placed a red plastic apple immediately above the camera lens to attract their attention, 121 and collected several images in a row using the burst function, enabling us to select the 122 most forward-facing image from the series. Immediately after the capture of an image, 123 we took a photograph of a colour standard (X-rite ColourChecker passport) placed in the 124 same location and photographed under the same lighting as the subjects (*i.e.*, the 125 "sequential method": [42–45]. 126 For analysis, we chose only images of fully adult males (median age = 9 years; 127 range = 8-16 years; N=69) and females (median age = 9 years; range = 8-14 years; N=27) 128 looking directly towards the camera. For each image, we digitally measured the sizes of 129 eight facial features in GIMP 2013, as depicted in Fig. 1, and scaled the length of each 130 feature by dividing it by head height (hereafter, relative size). We then compared male

131 and female relative feature sizes using either Mann-Whitney U or independent samples t-

132	tests, depending on normality of the data distribution (See Table 1). The relative sizes of
133	five features (lower face height, jaw width, temporalis height, jaw height, nose length)
134	were larger in male faces, while two features (interpupil distance, face width) did not
135	differ significantly between the sexes, and one feature (eye height) was significantly
136	larger in female faces. We then ran a multiple linear regression model with each facial
137	feature as a predictor variable and sex as the independent variable. We saved the
138	unstandardised predicted variables for use as facial masculinity scores for each male and
139	female image. The derived male (mean \pm SE = 1.91 \pm 0.026) and female (mean \pm SE
140	=1.22 \pm 0.039) facial masculinity scores differed significantly (Mann-Whitney U = 10.0,
141	P < 0.001).
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4.40	
143	Insert Figure 1 approximately here
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144 145	
144 145 146	Insert Table 1 approximately here
144 145 146 147	Insert Table 1 approximately here
144 145 146 147 148	Insert Table 1 approximately here Stimulus preparation Using the masculinity scores described above, we selected as stimuli in experimental
144 145 146 147 148 149	
144 145 146 147 148 149 150	Stimulus preparation Using the masculinity scores described above, we selected as stimuli in experimental trials the 10 most masculine and 10 most feminine facial images (hereafter masculine images and feminine images, respectively) that did not contain any distracting elements,

We printed stimuli onto matte photo paper (Staples Photo Supreme) using a colour-calibrated printer (Canon Pixma Pro 100), and measured the printed face colour using a Xrite ColourMunki spectrophotometer (see [31]). Pictures were printed on letter format paper (21.5 cm × 28.9 cm), with printed images of a dimension of 18.5 cm × 18.5 cm, in such a way that face length was 17 cm.

159

160 Experimental design

161 To test for female preference for male facial masculinity, we used a look-time paradigm 162 that has been used successfully to test interest towards other facial features in this study 163 species [31,46–49]. Each test pair consisted of one masculine and one feminine image, 164 selected randomly from the set of 10 stimuli in each category. KR and one assistant 165 conducted trials on weekdays from 18 March to 29 April 2015, between 09:00 and 13:00 166 h. The stimuli were placed in frames built into an experimental apparatus, such that they 167 were 85 cm apart at their centres (the apparatus measured 50 x 120 cm; Fig. 2). The 168 relative position of the images in the frame – whether the masculine image was on the 169 right or left – was randomised. Prior to trials, the stimuli were covered by occluders. 170 Potential trial subjects available on Cayo Santiago were all females \geq 3 years old (N=476) 171 at time of study). We tested 167 of these potential subjects, each being tested only once. 172 We discarded 60 trials that lasted less than 15 s, or during which it was not possible to 173 determine which image the subject was looking toward at any point. This left 107 trials, 174 one from each of 107 subjects (median age = 8 years). Females were not retested if they 175 participated in failed trials, and females that saw stimuli when they were not being tested

were also identified and were not tested in future trials. Females who were near adultmales, sleeping, or grooming other adults were not tested.

178	For testing, we placed the experimental apparatus 2-3 m in front of a female and
179	started recording her behaviour on video (Fig. 2). To determine the location of the stimuli
180	in relation to the subject's eyes (for video coding), we directed her visual attention
181	toward the location of the covered stimuli by tapping on the occluders (in randomised
182	order). We then directed her attention away from either stimulus by tapping on the centre
183	of the apparatus, and removed the occluders to reveal the stimuli. Trials lasted for 30
184	seconds after removal of the occluders, unless the subject moved away or engaged
185	
186	Insert Figure 2 approximately here
187	
188	socially with another monkey. We used MPEG Streamclip for Mac to code only the first
189	15 seconds (following removal of occluders) of trial videos frame-by-frame, because
190	most subjects stopped looking at either stimulus before the 15 th second. During coding,
191	we assessed the total amount of time spent looking at the masculine image and at the
192	feminine image. To eliminate the possibility of coding bias, we coded all trials blind to
193	condition (<i>i.e.</i> on which side the masculine image was located).
194	
195	Potential confounds of facial masculinity
196	In order to test for possible confounding effects of other male traits, such as age

and facial colouration, on stimulus image masculinity, we used Spearman's rank

198	correlations and Mann-Whitney test. Male age was taken from the long-term records of
199	CRPC. Age and masculinity were not correlated among the males used as experimental
200	stimuli ($r_s = 0.075$, $N = 20$, $P = 0.754$), and males used in masculine male images ($N =$
201	10, median age = 10) were not older than more feminine males ($N = 10$; median age = 9;
202	Mann-Whitney U = 39.5, P = 0.421). To quantify facial colour and luminance, we took
203	red (R), green (G) and blue (B) measurements from the stimuli and, based on the
204	processing of colours early in the primate visual pathway, calculated redness as the Red-
205	Green Opponency Channel, $(R - G)/(R + G)$, and darkness as the Luminance
206	(achromatic) Channel (R + G)/2 [46]. Neither facial colour ($r_s = -0.138$, N = 20, P =
207	0.559) nor facial luminance ($r_s = 0.339$, $N = 20$, $P = 0.143$) was correlated with facial
208	masculinity in the stimulus set. Furthermore, masculine ($N = 10$, median colour = 0.087,
209	median luminance = 167.75) and feminine males (N = 10, median colour = 0.097 , median
210	luminance = 145.13) did not differ in facial colour (U = 44, P = 0.684) or luminance (U =
211	71.5, $P = 0.11$). We thus concluded that any difference in the looking behaviour of our
212	subjects toward masculine and feminine stimuli would be independent of the effects of
213	male age or facial colour.
214	We also checked for confounding effects of stimulus males' familiarity to
215	females. Our operational definition of familiarity was group co-membership. Therefore,

216 using Mann-Whitney tests, we compared the proportion of trial time subjects looked at

217 masculine stimuli when they were groupmates with neither stimulus male (N=76) to trials

218 in which they were groupmates of the masculine stimulus male (N=14), the feminine

stimulus male (N=10) and both (N=7). All results were non-significant (masculine vs.

220 neither: U = 543.5, P = 0.903; feminine vs. neither: U = 378, P = 0.983; both vs. neither: 221 U = 345, P = 0.198), indicating that group co-membership did not influence subjects' 222 looking behaviour.

223

224 Data analysis

To test the prediction that females would look longer at the more masculine face of the pair, we undertook two analyses. First, we compared females' duration of looking towards the masculine and feminine images using Wilcoxon signed-rank tests. Secondly, we compared the number of trials in which females looked longer at the masculine vs. feminine stimuli to the value expected by chance (0.5) using a binomial test.

230 We also used two approaches to test the prediction that the proportion of time 231 spent looking at the more masculine face would be positively related to the difference in 232 masculinity between the two faces presented. Firstly, we calculated the relative difference 233 between the masculinity scores of the masculine and feminine image for each trial as 234 follows: ((masculine image score - feminine image score) / feminine image score) x 100, 235 with higher scores indicating larger disparities between the two images. We then ran a 236 linear model with percentage of total look-time spent looking at the masculine image 237 (relative look-time score) as the dependent variable, and relative facial masculinity score 238 as the predictor variable. As relative masculinity may be related to absolute masculinity, 239 we ran an additional linear model, this time entering both relative and absolute facial 240 masculinity score (of the more masculine image) as predictors.

241	Secondly, to further examine the salience of differences in facial masculinity, we
242	separated the dataset into two groups – one containing the 53 trials with the lowest
243	relative facial masculinity scores, the other containing the 54 trials with the highest scores
244	(results were identical if we used the lowest 54 and the highest 53 trials). We then used
245	Wilcoxon signed-rank tests to compare females' duration of look-time towards the
246	masculine and feminine images in each group. Statistical tests were two-tailed and
247	performed using IBM SPSS Statistics for Mac (21.0); α was set at 0.05.
248	
249	Results
250	Prediction 1 - females will look longer at the more masculine male face of the pair
251	Subjects' median overall look-time was 4.72 s, or 31% of the 15-s trial period.
252	When all trials were included in the analysis, median looking time for masculine images
253	(2.24 s; IQR = 1.31 - 3.45 s) did not differ from that for feminine images $(2.24 s; IQR =$
254	1.07 – 3.43s; Wilcoxon signed-rank test: Z = -0.799, N = 107, P = 0.424; Fig. 3).
255	However, subjects did look longer at the masculine than the feminine image in a
256	significantly higher proportion of trials than expected by chance (looked longer at
257	masculine image: 64 trials; looked longer at feminine image: 41 trials; 2 ties. Binomial
258	test: $p = 0.031$).
259	
260	Insert Figure 3 approximately here
261	

262	Prediction 2 - the proportion of time spent looking at the more masculine face will be
263	positively related to the difference in masculinity between the two faces presented
264	Trials' relative facial masculinity scores ranged from 12.1% to 61.4%, and
265	variation in these scores explained a small but significant proportion of variation in
266	relative look time scores ($\beta = 0.29, 95\%$ CI =0.03-0.55, p = 0.03, adjusted R ₂ = 0.035;
267	Fig. 4). In other words, the greater the within-trial disparity in masculinity scores, the
268	stronger the bias toward masculine images. The relationship between relative masculinity
269	and relative look time remained significant when absolute masculinity scores were
270	included in the model (β = 0.379, 95% CI =0.06-0.70, p = 0.021), and there was no
271	additional influence of absolute masculinity on relative look times ($\beta = -0.179, 95\%$
272	CI=0.54-0.192, $p = 0.34$; total model: F [2,104] = 2.9, adjusted R ₂ = 0.034, $p = 0.06$).
273	
274	Insert Figure 4 approximately here
275	
276	Having separated trials into high (N = 54; mean \pm SE = 42.7 \pm 0.75%) and low (N
276 277	Having separated trials into high (N = 54; mean \pm SE = 42.7 \pm 0.75%) and low (N = 53; mean \pm SE = 22.7 \pm 1.14%) relative masculinity score groups, we found that in the
277	= 53; mean \pm SE = 22.7 \pm 1.14%) relative masculinity score groups, we found that in the
277 278	= 53; mean \pm SE = 22.7 \pm 1.14%) relative masculinity score groups, we found that in the low-differences group, subjects' look-times did not differ between masculine (median =
277 278 279	= 53; mean \pm SE = 22.7 \pm 1.14%) relative masculinity score groups, we found that in the low-differences group, subjects' look-times did not differ between masculine (median = 2.41 s; IQR = 1.48 – 3.66 s) and feminine images (median = 2.41 s; IQR = 1.17 – 3.97 s;

283 -3.23 s; Wilcoxon signed-rank test: Z = -2.421, N = 54, P = 0.015; Cohen's d = 0.54; Fig

- 284 5).
- 285

286 -----Insert Figure 5 approximately here-----

287

288 Discussion

289 Using a two-alternative experimental look-time paradigm, we tested the hypothesis that 290 free-ranging female rhesus macaques perceive variation in male facial masculinity. In 291 partial support of our prediction that females would look longer at the more masculine 292 male face of the pair, test subjects looked longer at masculine than feminine male faces 293 when the difference in masculinity between the two was high. Moreover, as predicted, the 294 proportion of time spent looking at the more masculine face was positively related to the 295 difference in masculinity between the two faces presented. No relationships were found 296 between male facial masculinity and either male age or facial colour, and female look-297 time did not appear to be related to familiarity to the test subject, ruling out these as 298 potential confounds of our key results. Overall, this study provides evidence from a non-299 human species that variation in male facial shape, specifically variation along a feminine-300 masculine continuum, is salient to female conspecifics.

The finding that females distributed their visual attention unevenly between masculine and feminine faces indicates that the variation in facial masculinity we measured was not only perceived by, but also salient to female rhesus macaques. It is possible that variation in facial masculinity has no reliable connection to underlying

305 physiological, behavioural or genetic factors in male rhesus macaques, in which case 306 there may be no fitness repercussions of female attention to such variation. However, as 307 male facial masculinity is related to hormone levels and behaviour in humans and other 308 primates [e.g., 7,43], it seems likely that females' ability to discriminate variation in this 309 trait is the result of evolutionary processes.

310 Work to date has indicated that the development of facial masculinity in humans 311 is under the control of testosterone [6,7, but see 10, which failed to replicate this 312 relationship] and is linked to aggressiveness and competitive ability in humans and non-313 human primates [11,27,30]. As such, it is possible that females gain from paying 314 attention to male facial masculinity because it provides information about the risks of 315 aggression males may present; this explanation has also been proposed to underlie the 316 attentional bias shown for threat grins documented in this species [46,48]. A non-317 mutually exclusive possibility is that females are attracted to facial masculinity in a 318 sexual context; a preference for males with more masculine faces as mating partners may 319 benefit females if this trait is an honest cue of male genetic quality and health. According 320 to the immuno-competence handicap hypothesis, testosterone-dependent traits can 321 provide information about male quality because androgens are immunosuppressive [51]. 322 Since the development of facial masculinity is under the control of testosterone, high 323 facial masculinity could therefore be a cue to male quality that is available to females. 324 Since visual attentional biases can be underpinned by both attraction and fear (reviewed 325 in [49]), more work is needed to establish whether female perception of variation in male 326 facial features does translate into higher reproductive output for males with more

masculine faces, such that female mate choice would play a positive role in maintainingmale facial masculinity in this species.

329 Our finding that subjects' responses to experimental stimuli depended on relative 330 differences in masculinity highlights the importance of considering aspects of receiver 331 psychology in studies such as this one. We suggest two potential explanations for the 332 positive association between masculinity differences and subjects' visual bias toward 333 masculine faces. First, the differential responses may have been associated with subjects' 334 ability to perceive differences in masculinity. A critical feature of signals and cues is that 335 the information they are hypothesised to convey can only alter receiver behaviour if 336 receivers are able to perceive the differences exhibited by emitters [52]; small differences 337 may simply not be discernible. Second, subjects may effectively perceive differences 338 even when small, but such differences may not be sufficient to motivate a differential 339 response; other features, such as skin colouration or texture, may overshadow masculinity 340 differences when they are small.

The effects seen in the present study may represent responses to low-level features (i.e. more elementary features of the scenes presented in our stimuli, such as local colour, luminance or contrast) [53]. In this case, such effects might be seen as the perceptual mechanism by which rhesus macaques are stimulated by masculine facial traits. Such effects would require that low-level features are systematically linked to facial masculinity for them to result in the pattern we observed.

347 Our study did not attempt to disentangle the potential reasons for the visual biases348 we observed, but these are important avenues for future investigation. Studies of female

349	rhesus macaque mating behaviour in relation to male characteristics, like those conducted
350	by Manson [54], Dubuc et al. [41], and Georgiev et al. [55] are needed to determine
351	whether females' bias in visual attention towards more masculine faces translates into
352	differences in mating and reproductive success. Another important avenue for research is
353	to assess the potential information content of facial shape by investigating the
354	behavioural, physiological, morphological, and genetic correlates of facial masculinity.
355	Finally, as there is evidence that male facial colouration plays an important role in female
356	mate choice in this species [31,41,56], a more comprehensive analysis of the relationship
357	between facial masculinity and facial colouration is needed to better understand how
358	different facial features, and the interaction between them, may shape female preferences.
359	
360	Ethics statement
361	
362	The study was approved by the IACUC of the University of Puerto Rico, Medical
363	Sciences Campus (protocol No. A0100108). All applicable international, national, and/or
364	institutional guidelines for the care and use of animals (in particular non-human primates)
365	were followed. All procedures performed involving animals were in accordance with the
366	ethical standards of the institution(s) at which the study was conducted.
367	

368 Data accessibility

- All data and code related to this study have been: 1) Submitted as electronic
- 370 supplementary materials accompanying the manuscript, 2) Submitted to Biorxiv along
- 371 with manuscript preprint at <u>https://www.biorxiv.org/content/early/2017/11/21/222810</u>
- 372 3) Uploaded to Dryad data repository; review link:
- 373 <u>http://datadryad.org/review?doi=doi:10.5061/dryad.k79v6;</u> Temporary Dryad DOI
- 374 (pending manuscript acceptance): doi:10.5061/dryad.k79v6
- 375

376 **Competing interests**

- 377
- 378 The authors declare that they have no financial or non-financial competing interests. The
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- 380 necessarily represent the official views of any of the agencies and organizations that
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- 382
- 383 Authors' contributions
- 384
- 385 KR conceived, designed, and coordinated the study, measured facial masculinity,
- 386 performed experiments and statistical analyses, and drafted the manuscript; SS helped
- design and coordinate the study, carry out statistical analyses, and draft the manuscript;
- 388 AG and DM helped coordinate the study, and draft the manuscript; JH printed stimulus
- images, and helped coordinate the study and draft the manuscript; DC collected the

390	stimulus images, and helped design and coordinate the study, carry out statistical	
391	analyses, and draft the manuscript. All authors gave final approval for publication.	
392		
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407		
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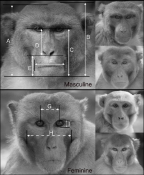
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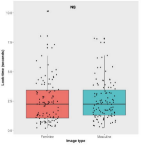
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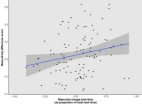
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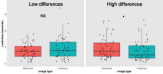
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0.633 ± 0.025	0.55
0.391 ± 0.020	0.36
1.016 ± 0.011	1.00
0.280 ± 0.024	0.26
0.354 ± 0.015	0.33
0.259 ± 0.016	0.26
0.617 ± 0.048	0.63

0.599 ± 0.022
0.366 ± 0.090
1.003 ± 0.006
0.267 ± 0.027
0.332 ± 0.017
0.261 ± 0.016
0.622 ± 0.040

т	-6.03
т	= -4.84
υ	= 288
	= -2.20
т	= -5.98
Т	= 0.624
Т	= 0.527

