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Physiological Factors of Female Runners With and Without Stress Fracture Histories: A Pilot Study.

Therese E Johnston
Thomas Jefferson University

Colleen Dempsey
Thomas Jefferson University

Frances Gilman
Thomas Jefferson University

Ryan Tomlinson
Thomas Jefferson University

Ann-Katrin Jacketti
Thomas Jefferson University
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Authors

Therese E Johnston, Colleen Dempsey, Frances Gilman, Ryan Tomlinson, Ann-Katrin Jacketti, and Jeremy Close

1 Physiological Factors of Female Runners with and without Stress Fracture Histories: A Pilot
2 Study

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4 Therese E. Johnston, PT, PhD, MBA; Colleen Dempsey, RT, EdD; Frances Gilman, RT, DHSc;
5 Ryan Tomlinson, PhD; Ann-Katrin Jacketti, DPT; Jeremy Close, MD.
6 Thomas Jefferson University, Philadelphia, PA USA

7

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10

11 Corresponding author

12 Therese E. Johnston, PT, PhD, MBA

13 Professor, Department of Physical Therapy, Jefferson College of Rehabilitation Sciences

14 Jefferson (Philadelphia University + Thomas Jefferson University)

15 Jefferson – Center City Campus

16 901 Walnut Street, Room 515, Philadelphia, PA 19107

17 T 215-503-6033

18 F 215-503-3499

19 therese.johnston@jefferson.edu

20

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24 Social Media:

25 Statement: Female runners with low hip bone mineral density, menstrual changes during peak
26 training, and elevated bone turnover markers may be at increased risk of stress fracture, and thus
27 screening beyond what is commonly performed may be warranted.

28 @TJ_PTRResearch

29 #JeffersonResearch

30 @ResearchAtJeff

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47 **ABSTRACT**

48 Background: Female runners are at increased risk of stress fractures (SF) compared to men.
49 Literature is lacking in regard to best practice for preventing and treating SF in women. The
50 purpose of the study was to compare physiological measures and running related factors between
51 women with and without running-related SF histories of various ages and running abilities.
52 Hypothesis: Women with and without SF histories would differ in medical and menstrual
53 history, bone health, body composition, nutrition, and running history.
54 Study Design: Prospective cohort study
55 Level of Evidence: 2b
56 Methods: Twenty female runners with SF histories were age and running-distance matched with
57 20 women without SF histories. Data included medical, menstrual, running, injury, and
58 nutritional histories; blood histology related to nutritional, hormonal, and bone-related risk
59 factors; and bone density, fat, and lean tissue using Dual Energy X-ray Absorptiometry. Paired t-
60 tests were used to examine differences between women with and without SF histories, and
61 Spearman correlations were conducted to examine relationships between physiological factors.
62 Results: Women with SF histories had lower hip bone mineral density compared to women
63 without SF histories ($p < 0.05$). SF history was moderately correlated with menstrual changes
64 during increased training times ($r = 0.580$, $p < .0001$) but was not correlated with any other
65 physiological factor. There was a moderate correlation within the SF group ($r = 0.65$, $p = .004$) for
66 bone markers for resorption and formation both increasing, indicating increased bone turnover.
67 Conclusion: Female runners with low hip bone mineral density, menstrual changes during peak
68 training, and elevated bone turnover markers may be at increased risk of SF.

69 Clinical Relevance: Female runners need routine screening for risks associated with SF
70 occurrence. As bone mineral density and bone turnover markers are not routinely assessed in this
71 population, important risk factors may be missed.

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73 Key Words: running, female, stress fracture, bone density

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92 **INTRODUCTION**

93 Stress fractures (SFs) are non-traumatic incomplete fractures resulting from repetitive loading on
94 normal bone or from normal loading on abnormal bone.¹¹ Running related SFs account for 69%
95 of all SFs with 95% occurring in the lower extremities and pelvis.¹¹ Women have at least 2 times
96 greater risk than men,^{13,16} and more women than men are now running. In the 2018 National
97 Runner Survey, runners were 54% female, 52% of all runners were between ages 35 and 54, and
98 60% considered themselves frequent fitness runners.²⁹

99

100 The risk factors for SFs in women are multifactorial, and include differences in anatomy, body
101 composition, metabolism, the cardiovascular system, hormonal status, and psychological status
102 as compared to men.¹⁶ Both intrinsic and extrinsic factors contribute to the occurrence of SFs.
103 Intrinsic factors are physiological¹¹ and include bone structure and density, decreased fat in
104 relation to lean tissue, and nutritional, hormonal, and bone-related health status. Menstrual
105 irregularities and energy deficiency due to an imbalance between nutritional intake and activity
106 are often present.²² Women also have greater risks due to the female athlete triad, a negative
107 energy balance between nutritional intake and activity that can lead to menstrual issues and
108 decreased bone mineral density, showing the inter-relationships of these factors.²⁰ Both pre-
109 menopausal and post-menopausal women are at risk.^{20,26} Extrinsic factors include training
110 intensity, training surfaces, diet, and footwear.¹¹

111

112 The literature is lacking in regard to best practice for preventing and treating SFs in women.
113 Surprisingly, few studies^{4,27} directly evaluate women with and without a history of SFs to assist
114 in better assessing risk and developing preventative strategies. There are several articles related

115 to risk factors,^{11,13,16,20,23} a few case reports with female runners,^{3,10,12,18} and a few observational¹⁵
116 and experimental studies.^{4,21,27,30} These studies examine various factors including bone density,
117 nutritional status, biomechanics, and menstrual status. Overall these studies show some
118 relationships between these factors. Some limitations include small sample sizes in most studies,
119 inclusion of only high level adolescent or young female runners, and mixed populations
120 (male/female or different sports). Due to these limitations and the increased SF for women, there
121 is a significant need to better understand issues related to SFs to prevent and properly treat these
122 injuries to optimize return to running, overall health, and participation. The issue is not limited to
123 women of a specific age as hormonal issues affect all women runners, thus making it important
124 to not limit studies to young elite runners. Therefore, the objective of this study is to compare
125 important physiological measures between women with and without running-related SF histories
126 of various ages and running abilities. The hypothesis was that there would be differences related
127 to medical and menstrual history, bone health, body composition, nutrition, and running history.

128

129 **METHODS**

130 Female runners, age 18-65 years, with and without running-related SF histories were recruited
131 over a 5 month time period via posted flyers and social media for this study held within an urban
132 university hospital system. A variety of social media sites were identified to decrease possible
133 selection bias. Women self-identified as runners, with no upper or lower limit set for running
134 intensity, duration, or distance. To control for differences in age and running ability, after each
135 woman with a SF history was enrolled into the study, a woman without a SF history was
136 recruited who was age-matched within 5 years and running-distance-matched within 10
137 miles/week.^{5,31} All enrolled women signed a written informed consent form approved by the

138 governing Institutional Review Board. Women with SF were included if they had a SF at any
139 time as runners. Women with and without SF histories were excluded if they had a neurologic
140 diagnosis or any systemic medical condition that would impact bone, were pregnant, or were
141 breastfeeding.

142
143 Data collection included background information and physiological measures. Participants
144 completed an online questionnaire (Qualtrics, Seattle, WA) to collect demographics as well as
145 medical, menstrual, running, injury, and nutritional histories. To examine physiological data on
146 nutritional, hormonal, and bone related risk factors,⁸ the following non-fasting serum histological
147 measures were collected and processed using standard medical laboratory procedures: complete
148 blood count, vitamin D (25-(OH)D), calcium, albumin, parathyroid hormone, estradiol,
149 testosterone, bone specific alkaline phosphatase (BALP, measure of bone formation),⁶ and N-
150 telopeptide (N-Tx, measure of bone resorption).⁶ To examine bone, fat, and lean tissue, Dual
151 Energy X-ray Absorptiometry (DXA)⁹ was used to measure areal bone mineral density (aBMD)
152 of the left hip and the lumbar spine, and full body composition using a Hologic Horizon A
153 scanner (Hologic, Marlborough, MA). The DXA machine was calibrated prior to each testing
154 session to decrease measurement error. A negative pregnancy test was required prior to
155 conducting the DXA for all participants.

156
157 To examine differences between women with and without SF histories, paired t-tests were
158 conducted using SPSS Statistics Version 25 (IBM Corporation, Armonk, New York). Cohen's d
159 was calculated to determine effect size. To examine possible relationships between group and
160 physiological factors and among different physiological factors, Spearman correlations were

161 performed. Due to the lack of data available on medical and menstrual history, bone health, body
162 composition, nutrition, and running history that span the age ranges included, a sample of 20 per
163 group was chosen based on differences in bone turnover, body mass, and estradiol levels seen in
164 study with 37 adolescent runners.² Effect sizes were thus calculated for measures in this study.

165

166 **RESULTS**

167 Forty nine women were screened for the study. Two women with SF histories were excluded due
168 to thyroid disease, and five eligible women without SF histories were excluded as they did not
169 match with a woman with a SF. Forty two women (35.0 ± 7.4 , range 22-50 years) enrolled into
170 the study. Two participants withdrew after signing the consent form due to time constraints, and
171 data are complete for 40 participants or 20 matched pairs. Data were complete for all participants
172 except for 1 missing albumin value for the SF group and 2 missing N-Tx values for the non-SF
173 group. These data and the matched pair's values were thus excluded from data analysis.

174

175 The oldest enrolled woman was 50 years old, and she was the only participant who was post-
176 menopausal. Her match with a SF history was peri-menopausal. Women were highly educated
177 and predominately white (Table 1). Women with SF histories were 2.2 ± 2.6 years post their
178 most recent fracture (range 0.8-10 years) with 10 having fractured within the past year, 5 in the
179 last 1-3 years, and 5 more than 5 years prior. Fracture sites included tibia (n=15), metatarsal
180 (n=8), femur (n=5), cuneiform (n=1), and sesamoid (n=1) with 6 participants reporting having
181 had 2 SFs, and 2 participants reporting 3 SFs.

182

183 Tables 2 and 3 show self-reported information for running and menstrual status, respectively,
184 and there were no differences ($p=0.57-1.00$) between groups for these data. Groups were also
185 evenly distributed in regard to birth control use and type, and for the number who had ever gone
186 >3 months without a period other than during pregnancy (6 per group). However, 12 women who
187 had a SF reported that their menstrual periods changed during increased training times, while
188 only 1 reported this occurring in the non-SF group. Age when started running did not differ
189 between groups, yet 9 women with SF histories started running at 18 years or younger, while
190 only 4 without SF started this young.

191
192 In comparing physiological measures between women with and without SF histories (Table 4),
193 the only statistical difference was in hip aBMD, with lower aBMD in the women with a SF
194 history. But the effect size for this difference was low (0.19). The measure with the largest effect
195 size of 0.61 was BALP, but the difference between groups was not statically significant.

196 Correlational analysis showed that time post fracture was unrelated to bone markers (BALP, N-
197 Tx) and that hip aBMD was unrelated to any other physiological factor. SF history was
198 moderately correlated with menstrual changes during increased training times ($r=0.580$, p
199 $<.0001$) but was not correlated with any other physiological factor. While there was a low
200 correlation between BALP and N-Tx when looking at all participants together ($r=0.34$, $p=.03$),
201 there was a moderate correlation within the SF group ($r=0.65$, $p=.004$) with BALP and N-Tx
202 increasing together (Figure 1), indicating increased bone turnover.

203

204

205

206 **DISCUSSION**

207 The main results from this study were that women with a SF history had lower hip aBMD than
208 their matched counterparts without a SF history, and that women with a SF history had
209 alterations in their typical menstrual cycles during more intense training times even though
210 current estradiol levels did not differ between groups. The study was conducted during the
211 months of March to June, which represented mainly off to early season training for the included
212 women. Within the SF group, there was a correlation between bone formation and resorption that
213 was not seen within the non-SF group, indicating increased bone turnover.¹⁷ Of note, DXA for
214 bone density and blood histology to examine bone resorption and formation markers are not
215 routinely performed in this population, thus important information may be missed clinically in
216 these women. As DXA is a relatively inexpensive with low radiation exposure, performing DXA
217 in this population may be cost-effective. The more expensive tests for bone resorption and
218 formation markers may then be performed based on concerning findings via DXA. Asking
219 female runners about any menstrual cycle changes during heavier training times may be an
220 important addition to a patient interview. Women who had these changes reported lighter flow,
221 shorter duration, increased spotting, irregularity, and missed cycles.

222
223 Several studies have examined menstrual dysfunction in relation to bone, but primarily in a
224 younger population. Ackerman et al.¹ reported decreased spine and whole body aBMD and
225 altered bone structure in 14-25 year old female athletes with oligoamenorrhea (6 cycles or less in
226 prior year), with greater changes seen in participants with more than 1 SF. In a study that
227 included collegiate cross-country runners, Tenforde et al.³⁰ reported that oligoamenorrhea or
228 amenorrhea and a prior SF were predictors of subsequent bone stress injuries. A small

229 percentage of participants had low aBMD, with more than half of them being runners. Nose-
230 Ogura et al.²⁴ found a relationship between amenorrhea in the teenage years and aBMD in the
231 20's for female athletes that included distance runners, suggesting the need for intervention at a
232 younger age. While these studies provide important information for female runners in these
233 younger age groups, women older than 25 years represent a large number of runners. As bone
234 mass starts to decline between 20 and 30 years of age for women,⁷ issues specific to these
235 women must also be addressed. Micklesfield et al.²² studied 613 long distance (half-marathon
236 and ultramarathon) female runners ages 16-62 years, of whom 17.3% had sustained a bone stress
237 injury, but found no differences between these women and the women without these injuries for
238 age, weight, BMI, or menstrual function. They also found that over half of all 613 women
239 reported menstrual dysfunction. Thus, further study is needed to better understand the risks.
240 These studies that relate menstrual status and aBMD as well as the results of this current study
241 indicate the need to evaluate and treat female runners for these issues early and to continue to
242 evaluate changes over time.

243

244 While there were no differences in estrogen levels between women with and without SF
245 histories, some women in the study had very low estrogen levels. The low end of the normal
246 range for estrogen levels is 24 pg/mL. Four women with SF histories and eight without had very
247 low values (<5 pg/mL), and two in each group had low values (8-23 pg/mL). The significance of
248 these low values is difficult to determine in this small sample as the women with and without SF
249 histories were equally impacted. Estrogen levels fluctuate during the menstrual cycle,²⁸ and data
250 were not collected regarding menstrual phase in this study. To gather cyclical data on female
251 runners would require measures of estrogen levels to be collected throughout the menstrual cycle

252 to identify patterns.²⁸ Assessing estrogen levels across the menstrual cycle is thus recommended
253 for future studies.

254

255 The bone turnover markers of N-Tx and BALP as measured in this study are not routinely
256 assessed in female runners but may play a role in assessing risk. While these measures were not
257 statistically significant different between groups in this study, there was a correlation between
258 increased bone formation and resorption in the SF group, indicating increased bone turnover.¹⁷ In
259 a literature review of studies of post-menopausal women by Vasikaran et al.,³² several studies
260 reported that an increase in bone turnover markers led to an additive effect on the risk for
261 fractures, and that increased bone turnover markers may predict fracture risk independently of
262 aBMD. While the population in Vasikaran et al.³² differs from the women runners in this study,
263 the use of these markers may be beneficial and more research is warranted. In a sample of
264 adolescent female cross-country runners, elevated bone markers were associated with a lower
265 BMI, menstrual irregularities, and lower estradiol and Vitamin D levels.² In contrast, Fujita et
266 al.¹⁴ measured bone resorption (urine N-Tx) twice per year in a small sample of female runners
267 ages 19-34, and found while N-Tx values were normal during training, they increased when a SF
268 occurred. These findings suggest that N-Tx may be a non-invasive way to identify SFs and
269 monitor healing. A review article by Papageorgiou et al.²⁵ reported that short term low energy
270 availability can also elevate bone markers, thus several factors need to be considered when using
271 bone markers to guide diagnosis and return to running post SF. Finally, there is mixed opinion
272 as to the effect of increased turnover. While increased formation temporarily increases bone
273 porosity and decreases stiffness, it may also induce microdamage repair following bone stress.¹⁹
274 Thus, more research is needed on the interpretation of these bone markers clinically.

275

276 In this study, a physical therapy examination was not performed as the goal was to gather
277 physiological factors rather than specific musculoskeletal impairments. Koprelainen et al.²¹
278 reported that the risks of recurrent SFs across multiple sites may include a high weekly training
279 mileage, a leg length difference, a high longitudinal arch of the foot, and forefoot varus in
280 addition to menstrual dysfunction. Thus, these factors may be important to consider in the
281 examination of runners clinically along with the measures collected in this study. As the current
282 study controlled for running distance through matching of subjects, the impact of mileage cannot
283 be determined. Other factors to consider are impact forces and kinematics, which are not easily
284 collected clinically. Popp et al.²⁷ reported that women who fractured had less bone strength and
285 greater impact forces than women without fractures,²⁷ and Becker et al.⁴ reported different
286 kinematic patterns between runners with and without navicular SFs.

287

288 **CLINICAL SIGNIFICANCE**

289 For female runners ages 20-50 years of age with varying running abilities, it is recommended
290 that screening of intrinsic and extrinsic risk factors be performed to determine potential risks for
291 SF. Based on the research of others, these factors include nutritional, hormonal,¹¹ menstrual
292 irregularities, energy deficiency,²² training intensity, training surfaces, diet, and footwear.¹¹
293 Testing of aBMD is also recommended based on this study and others,¹¹ especially for those
294 women who report menstrual changes as intensity/frequency/duration of running increase. While
295 women with these changes may be at increased risk, DXA is encouraged for all female runners
296 to better inform them about potential increased risks and educate them on prevention.
297 Histological measures of bone turnover should also be considered for those with increased risk.

298

299 **LIMITATIONS**

300 In this study, a physical examination was not performed as the goal was to gather physiological
301 factors rather than specific musculoskeletal impairments. Koprelainen et al.²¹ reported that the
302 risks of recurrent SFs across multiple sites may include a high weekly training mileage, a leg
303 length difference, a high longitudinal arch of the foot, and forefoot varus in addition to menstrual
304 dysfunction. Thus, these factors may be important to consider in the examination of runners
305 clinically along with the measures collected in this study. As the current study controlled for
306 running distance through matching of subjects, the impact of mileage cannot be determined.

307

308 Other study limitations include the small sample size, which could potentially impact the ability
309 to obtain statistical significance. Matching women based on age and running distance likely
310 reduced some of the impact of small sample size. The sample was also one of convenience and
311 thus may not represent the population of female runners as a whole. The women in this study
312 also spanned a wide age range. But despite this heterogeneity of age, differences were found
313 between groups.

314

315 **CONCLUSION**

316 Based on the results of this study, measurement of aBMD, bone turnover markers, and menstrual
317 change data during training may be important additions to the clinical examination of female
318 runners. More research is needed on the role of bone turnover markers in assessing risk of SFs
319 and return to running post SF.

320

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324

325 **CONFLICTS OF INTEREST AND SOURCE OF FUNDING**

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327 of Thomas Jefferson University.

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430 standards. *Osteoporos Int*. 2011;22:391-420. PMID: 21184054.

431

432

433 **TABLE 1.** Participant demographics

Item	Item Choices	Stress Fracture Group (n)	Non-fracture Group (n)
Age	Years	35.1 ± 7.2	34.4 ± 7.7
Highest Educational Degree	Bachelor's	7	7
	Master's	6	9
	Doctoral	7	4
Race	Asian	0	3
	Hispanic	1	1
	White	19	16

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451 **TABLE 2.** Running status

Item	Item Choices	Stress Fracture Group (n)	Non-Fracture Group (n)	p-value
Days per week	2	0	1	0.96
	3	11	7	
	4	4	4	
	5	2	5	
	6	2	1	
	7	1	2	
Miles per week	0-10	1	1	0.88
	11-20	6	9	
	21-30	6	6	
	31-40	4	2	
	41-50	1	1	
	>50	2	1	
Average running pace (min/mile)	<6	1	0	0.98
	6-7	0	1	
	7-8	6	2	
	8-9	2	6	
	9-10	7	4	
	10-11	4	5	
	>11	0	2	
Age when started running	<10	3	1	0.96
	11-18	6	3	
	19-25	2	9	
	26-33	5	7	
	34-40	3	0	
	>40	1	0	

452 No differences between group ($p > 0.05$) using Chi-square.

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459 **TABLE 3.** Menstrual status

Item	Item Choices	Stress Fracture Group (n)	Non-Fracture Group (n)	p-value
Age at first menstrual cycle	9-10 years	1	2	1.0
	11-12 years	9	8	
	13-14 years	6	8	
	15-16 years	4	2	
Menstrual cycle length	29 days or less	11	13	1.0
	30-35 days	2	1	
	36 days or more	1	1	
	Irregular	6	4	
	Absent	0	1	
Menstrual cycle length	N/A	0	1	0.57
	1-2 days	1	2	
	3-4 days	9	9	
	5-6 days	8	4	
	7-8 days	0	3	
	8 days or more	0	0	
	No answer	2	1	

460 No differences between group ($p > 0.05$) using Chi-square.

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472 **TABLE 4.** Blood histological, bone density, and body composition results

Measure	Normal range	Stress Fracture Group	Non-Fracture Group	p-value	Effect size
Albumin	3.2 - 4.9 g/dL	4.3 ± 0.3	4.4 ± 0.2	0.21	0.40
Vitamin D	18 - 72 pg/mL	51.0 ± 10.0	51.8 ± 21.6	0.88	0.04
Calcium	8.5 - 10.3 mg/dL	9.3 ± 0.3	9.3 ± 0.3	0.73	0.11
Estradiol	12.5 - 498 pg/mL†	76.1 ± 105	50.6 ± 67.0	0.35	0.29
Testosterone	2-45 ng/dL	18.8 ± 8.2	19.1 ± 7.8	0.90	0.03
Parathyroid Hormone	11 - 67 pg/mL	36.7 ± 14.2	34.8 ± 9.2	0.64	0.16
Bone Specific Alkaline Phosphatase	5.0 - 18.8 mcg/L	9.9 ± 2.7	8.3 ± 2.4	0.09	0.61
N-Telopeptide	6.2 - 19.0 mg/dL	11.8 ± 5.0	11.1 ± 4.9	0.67	0.15
Spine Bone Mineral Density	N/A‡ gm/cm ²	1.0 ± 0.09	1.0 ± 0.11	0.15	0.44
Hip Bone Mineral Density	N/A‡ gm/cm ²	0.9 ± 0.1	1.0 ± 0.1	0.03*	0.19
Fat percent	N/A‡ %	31.2 ± 6.1	31.0 ± 5.0	0.94	0.02
Body Mass Index	18.5-24.9 kg/m ²	22.4 ± 2.8	23.2 ± 2.9	0.36	0.28

473 * Significant p-value

474 †Pre-menopausal, influenced by menstrual cycle phase

475 ‡N/A as normal is based on age and percentiles.

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480 **Figure Caption**

481 **FIGURE 1.** Bone turnover for each group. There was a moderate correlation within the stress
482 fracture group between bone resorption (N-telopeptide) and bone formation (bone specific
483 alkaline phosphatase) but not within the non-stress fracture group. This finding indicates
484 increased bone turnover in the stress fracture group.

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