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Key terms: catecholamines; employee; male employee; man; musculoskeletal pain; norepinephrine; observation; recovery; unwinding; work stressor; workplace; workplace observation

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Workplace observation of work stressors, catecholamines and musculoskeletal pain among male employees

by Achim Elfering, PhD,¹ ² Simone Grebner, PhD,¹ ³ Hans Gerber, MD,⁴ ⁵ Norbert K Semmer, PhD ¹ ²

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Objectives Prolonged activation of the sympathetic–adrenal medullary system during work and the lack of recovery after work may indicate a risk for musculoskeletal pain (ie, neck, shoulder, or low-back pain). This field study tested whether a high level of work stressors is associated with musculoskeletal pain and higher urinary norepinephrine excretion rates at work, after work, and on Sundays.

Methods Altogether 32 male employees working in adult education took part in a three-phase repeated-measures field study including workplace observation, self-report of musculoskeletal pain, and measurement of urinary catecholamine at work, in the evening after work, and at corresponding times on Sundays.

Results In univariate analyses, work stressors and workday levels of norepinephrine were significantly higher in the participants with high levels of musculoskeletal pain. Moreover, regressing workday norepinephrine excretion rates on predictor variables in the multilevel regression analysis showed a significant interaction between work stressors and musculoskeletal pain (P=0.011) with elevated excretion rates of norepinephrine in those exposed to a high level of work stressors and, at the same time, reporting higher levels of musculoskeletal pain.

Conclusions High levels of work stressors turned out to be associated with musculoskeletal pain and norepinephrine concentration in male employees. Increased activity of the sympathetic–adrenal medullary system seems to play an important role in work-related musculoskeletal pain.

Key terms norepinephrine, recovery, unwinding.

Most work-related musculoskeletal disorders have a complex and multifactorial etiology. Among other factors, biomechanical load, psychosocial stress, health behavior, and pain-related cognition are assumed to play an important role in the development of persisting musculoskeletal pain (1). Our field study tests whether catecholamine release constitutes an important link between adverse psychosocial factors at work (ie, work stressors such as time pressure and performance constraints) and musculoskeletal pain.

Reaction to work stressors leads to a catabolic state of hypervigilance, in which energy is provided so that the worker can cope with the stressors. The stress response is characterized by activation of the sympathetic–adrenal medullary system, including catecholamine secretion leading to increased heart rate and blood pressure. Moreover, activation of the sympathetic–adrenal medullary system leads to secretion of norepinephrine, which heightens muscle activity because of an increased sensitivity of the synapses and by recruiting more muscle fibers when an activity is performed (2). With regard to the activation of muscles, epinephrine amplifies the effects of norepinephrine. Studies by Lundberg and his coworker show that both mental stress and physical load elevate electromyographic (EMG) activity (3–5). Thus mental stressors can elevate the sympathetic–adrenal medullary system and EMG activity even in the absence of heavy physical load or unfavorable posture (3–5). It is noteworthy that mental stress can persist after work is over and prevent relaxation and the recovery of muscles (3). We therefore think that the effects of both physical and mental load at work and after work should be considered, which is in accordance with current models on occupational musculoskeletal pain, such as the “Brussels
versity degree, and all but one had completed vocational years. More than half of the sample (N=17) had a university organization was 60%. Altogether 35 of 39 persons agreed to participate rate among all of the employees of the organization was 43.8 (SD 8.3, range 32–58) being on medication. The mean age of the final sample was 43.8 (SD 8.3, range 32–58) man. The majority were married (N=26) and had children (N=28).

Measures
Smoking was assessed as the number of cigarettes smoked on the day of measurement, recorded in a diary in the evening of the day. The body mass index (BMI) was calculated by taking a person’s body weight and dividing it by their body height squared. Body height was reported by the participants, and body weight was measured on a weight scale.

Hypotheses
In our study, we expected higher musculoskeletal pain in the participants to correspond with higher levels of stressors at work (hypothesis 1). Moreover, we expected work stressors to be positively associated with the excretion rates of urinary catecholamines at work (hypothesis 2). Furthermore, we expected an interaction between stressors at work and musculoskeletal pain. Specifically, we expected comparably higher levels of norepinephrine at work (hypothesis 3a) and in the evening after work (hypothesis 3b) in the participants who reported high levels of musculoskeletal pain and, at the same time, were exposed to higher levels of stressors at work. The urinary excretion rates of norepinephrine for those who reported less musculoskeletal pain and had high levels of stressors, or reported musculoskeletal pain while having low levels of stressors at work, should have been considerably lower. Finally, we expected those who reported higher musculoskeletal pain and, at the same time, had higher levels of stressors at work to show less of a reduction in their norepinephrine levels from a workday to Sunday than the others did (hypothesis 4).

Work characteristics
The work conditions were rated by trained observers using a short version of the Instrument for Stress Oriented Task Analysis (ISTA) (11). The observers were trained for 7.5 days (12). The trainees were a research assistant, two student assistants, and five students serving a research internship for advanced psychology students. The training started with a study of the literature on task analysis, information about the occupation under study, psychological methods of observation, and bias in observation. The tasks of the trainees included self observation, interviews, and observation-based ratings. The training ended with a comprehension and memory test of the rating instrument and its dimensions.

Each participant’s work conditions were rated by two observers, and both ratings were averaged to reduce potential bias in the ratings. All of the items had a 5-point Likert format, reflecting either the intensity or the frequency of work stressors and job control. Job control was measured by four items, two of which covered method control (“employee decides which way to carry out his work” and “employee individually plans his work”), and the other two referred to time control (2 items: employee disposes his workday and work schedule” and “employee assigns his daily workhours”). Cronbach’s alpha for job control was 0.81. Task-related stressors were measured by time pressure (1 item: “employee works longer because of having too much work to do”), performance constraints (1 item: “employee has to work with inadequate devices or obsolete information”), uncertainty (1 item: “employee is in a situation in which he cannot go on with work because of a problem in another area”), and work interruptions (2 items “employee must work at several tasks simultaneously and must jump back and forth between tasks” and “employee is interrupted by telephone calls”). For a composite measure of task-related job stressors, the five-stressor items were averaged. [See the report by Grebner et al (13) for the same procedure with the questionnaire version of the

Study population and methods
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The sample consisted of 39 healthy male instructors from a public service organization in Switzerland. The participation rate among all of the employees of the organization was 60%. Altogether 35 of 39 persons agreed to the repeated measurements of urinary catecholamines. Three had to be excluded from the data analyses for being on medication. The mean age of the final sample of 32 male employees was 43.8 (SD 8.3, range 32–58) years. More than half of the sample (N=17) had a university degree, and all but one had completed vocational training. All of the participants were full-time employees and had held their present position for at least 2 (mean 5.6, SD 1.9) years. A total of 6 of the 32 employees had supervisory positions. Most of the participants were married (N=26) and had children (N=28).

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instrument.] This index of work stressors represents a heterogeneous concept, consisting of items from scales that are correlated yet do not represent a homogeneous construct. For such an index, the composite score reliability (14) is the appropriate reliability estimate. Its value was $r=0.82$.

Musculoskeletal pain
Musculoskeletal pain was assessed as pain in the neck, the shoulder region, and the lower back in the last 12 months [never (1), less than monthly (2), less than weekly (3), less than daily (4), daily (5)]. These three items are part of a scale measuring psychosomatic complaints developed by Mohr (15), based on Fahrenberg (16). The Cronbach alpha for musculoskeletal pain was 0.85, the values ranging from 3 to 15.

Domestic work
Three items addressed domestic work. The items were reported every evening at 2100, along with the time (hours and minutes) spent doing household chores, child care, and caring for relatives. A sum index of all three items indicated the total domestic workload in terms of industrial hours ($Ih = \text{hours} + \text{minutes}/60$, expressing minutes within the decimal system (ie, 3 hours 15 minutes equals 3.25). It was calculated separately for domestic work in the evening of the workdays and on Sundays.

Catecholamine measures
The participants were instructed to empty their bladder in the morning at 0715. These specimens were not analyzed, however. The participants were asked to note the time when they voided their bladder until noon. Urinary catecholamine samples were collected at work before lunch (1200), before the end of work (1700), and in the evening (2100) by trained collectors. The measurements were repeated at home on Sunday at corresponding times. On each of three workdays and three Sundays more than 75% of the sample covered the whole time period; that is, 75% of the participants did not empty their bladder between measurements. Only two or three persons had collection periods that were shorter than 2 hours. For the afternoon period, more than 70% of the measurements covered the whole time period, and, for the evening period, approximately 80% of the measurements covered the whole time period. On Sunday, the participants were asked to avoid vigorous physical activity (such as sports, housework, or gardening) or demanding mental work in order to obtain valid physiological reference values. The participants were asked to empty their bladder at the time indicated, measure the volume of urine with a metering box, and keep a record of each voiding time and volume that occurred between the measurement points. The urine samples (100 ml) were acidified immediately by hydrochloric acid (HCl) to a pH of 3. Each sample was then divided into two aliquots of 15 ml each. The aliquots were stored in a freezer ($-20^\circ$C) until the laboratory analysis. The excretion rates of epinephrine and norepinephrine were expressed as pmol/(min x kg) [picomole, $10^{-12}$ moles per liter of urine/(minutes since last voiding x kilograms of body weight)]. The participants recorded their consumption of alcohol, nicotine and caffeine and their intake of medication in a diary each evening. The accuracy and robustness of the measurements with respect to the collection schedule, acidification, storing conditions, and comparison of the time-related index with the ratio of creatinine was tested by the authors in a previous experimental study (17).

Laboratory analysis
The aliquots were analyzed in the Chemical Laboratory of the University Hospital in Bern by high-pressure liquid chromatography. More details on the laboratory analysis, including information on accuracy, have been reported elsewhere (17).

Design
The study had a repeated measures design with respect to the catecholamine assessments. Catecholamine data were collected on a workday (Tuesday, Wednesday, or Thursday) and on the following Sunday within the same week. The data collection was repeated twice, so that 3 weeks were covered for each person. The time lag between the measurements was 6 weeks. Therefore, each participant had up to 18 urinary catecholamine measurements (3 workdays plus 3 Sunday measures for 3 weeks). In the first week, there were no missing data. One person could not take part in the second and third week. Due to situational constraints of the participants, 1 single measurement was skipped in the second week and 13 were skipped in the third week. Three further measurements (1 in the second week and 2 in the third week) were lost due to failed laboratory analyses. Altogether, 278 workday catecholamine measurements and 270 Sunday catecholamine measurements were analyzed. There was no imputation of missing values.

Data analysis
The univariate analysis included a comparison of work conditions and urinary excretion rates of catecholamine between a low (below or equal the median value of musculoskeletal pain) and a high (higher than the median...
value of musculoskeletal pain) subgroup of participants (hypothesis 1). In the univariate analyses, the day- and
time-corresponding measurements of norepinephrine and
epinephrine were averaged for each person to control for
day-specific situational influences on the urinary excretion
rates of catecholamine [ie, measures of the same
time and type of day were averaged for each person; for
instance, the three workday noon measurements from
the three workweeks were averaged, the three workday
measurements made at 1700 were averaged, etc.; for a
similar approach used for the cortisol measurements, see
the report of Pruessner et al (18)]. To test hypothesis 1,
Student’s t-test for group comparisons was calculated. In
order to test hypothesis 2, we calculated the correlations
between the work stressors and the catecholamine levels.
P-values were one-tailed with the alpha set to 5%.

Hypotheses 3 and 4 were tested with a multilevel
analysis (19), in which the catecholamine measurements
were regressed on predictor variables with two hierarchi-
cal levels, namely, catecholamine measurements (level
1) nested within persons (level 2). We used the MLwiN
software package (20). The dependent variables were the
excretion rates of norepinephrine and epinephrine.
The predictor variables that were assessed for level 1 were
type of day (workday or Sunday), week (first, second,
or third), total time spent on household chores, child
care or care of relatives, and time of day (measurement
at 1200, 1700, or 2100). The level-2 predictor variables
were person-related (age, BMI, smoking, work stressor
index, job control, and musculoskeletal pain). All of the
predictor variables in the multilevel analyses were cen-
tered around the grand mean (19). All of the P-values in
the multilevel regression analysis were one-tailed with
the alpha set at 5%.

Results

Prevalence of musculoskeletal pain

The participants’ reports of the 1-year prevalence of
neck, shoulder, and low-back pain showed that most of
the respondents experienced musculoskeletal pain. Only
15.6% reported “never” having experienced low-back
pain (scored 1); this percentage was higher for neck
pain (37.5%) and shoulder pain (43.8%). Half of the
participants reported low-back pain “less than monthly”
(scored 2), and this percentage was a bit lower for neck
pain and shoulder pain (both 31.3%). The figures for
the three different types of pain did not differ much for
the participants who reported more frequent pain (“less
than weekly”, scored 3: 21.9% with neck pain, 12.5%
with shoulder pain, 18.8% with low-back pain; “less
than daily”, scored 4: 6.2% with neck pain, 6.2% with
shoulder pain, 9.4% with low-back pain; and “daily”,
scored 5: 3.1% with neck pain, 6.2% with shoulder pain,
6.2% with low-back pain). Neck, shoulder, and low-back
pain correlated with one another. Neck and shoulder pain
were more closely related with one another (r=0.780)
than neck pain with low-back pain (r=0.620) and should-
er pain with low-back pain (r=0.580). However, these
differences in the strength of the association were not
significant [P=0.136 and P=0.059, test of correlated cor-
relations according to Williams (21)]. The overall sum of
musculoskeletal pain ranged from 3 to 15, with half of the
participants scoring 5 or less, representing the “low
musculoskeletal pain” group. In the “high musculo-
skeletal pain” group, seven participants had a score of 6, one
had a score of 7, four scored 9, and four scored relatively
high at 11, 12, 13, and 15.

Work stressors in the groups with low and high mus-
culoskeletal pain

Table 1 shows the comparison between the groups with
low and high musculoskeletal pain with respect to the
study variables. The groups did not differ significantly
with regard to age, smoking habits, or body weight. In
line with expectations (hypothesis 1), the “high muscu-
loskeletal pain” group revealed higher scores for work
stressors (P=0.033). However, against expectations,
the “high musculoskeletal pain” group also had higher
levels of job control. The norepinephrine excretion rates
at work were higher among those with high musculo-
skeletal pain than among those with low musculoskel-
tal pain, and the group differences reached statistical
significance at noon (P=0.015) and at the end of the
workday (P=0.047).

The workday epinephrine excretion rates were also
higher in the “high musculoskeletal pain” group at the
end of the workday (P=0.050). However, there was
no corresponding pattern for group differences in the
evening of workdays and on Sundays. The mean Sun-
day urinary excretion rates of both epinephrine and
norepinephrine at 1200, 1700, and 2100 in the “high
musculoskeletal pain” group did not differ from those of
the “low musculoskeletal pain” group or were smaller.
Furthermore, the groups did not differ in the time spent
on household chores, child care, or care of relatives.

Correlation between work stressors and catechol-
amines on a workday

The correlations between workday norepinephrine and
work stressors were positive, and they reached statistical
significance for the norepinephrine measures at 1200
(r=0.297, P=0.049) and at 2100 (r=0.332, P=0.032).
The correlations between workday epinephrine and
work stressors were all positive (r=0.113 to r=0.134),
but did not reach statistical significance. There were
no significant associations between workday catecholamines and job control.

Workday excretion rates of catecholamines of those who reported high musculoskeletal pain and had high levels of work stressors

Table 2 shows the regression of workday norepinephrine and epinephrine excretion rates with respect to individual characteristics and work conditions. Smoking and musculoskeletal pain were significantly related to higher excretion rates of workday norepinephrine and epinephrine. In accordance with hypothesis 3a, a significant interaction between work stressors and musculoskeletal pain indicated that increased excretion rates of norepinephrine were restricted to the participants who reported musculoskeletal pain and were also being exposed to high levels of work stressors (P=0.011). Figure 1 shows that the norepinephrine excretion rates of the other participants were lower and similar.

There was no corresponding interaction effect for the epinephrine excretion rates. The excretion rates of catecholamines slightly decreased with the time of day, but this decrease was restricted to those with a high level of musculoskeletal pain. Contrary to hypothesis 3b, the participants who reported musculoskeletal pain and were also exposed to a high level of work stressors did not show persisting higher urinary excretion rates of norepinephrine after work. The proposed three-way interaction between work stressors, musculoskeletal pain, and time of day was not statistically significant.

Table 1. Mean values and standard deviations of the study variables for the groups with low and high musculoskeletal pain.

| Variable | Musculoskeletal pain (Low N=16) | Musculoskeletal pain (High N=16) | P-value* (one tailed) |
|----------|-------------------------------|-------------------------------|----------------------|
| Age      | 43.630 8.232                  | 44.000 7.967                  | −0.126 0.800         |
| Body mass index (kg/kg²) | 24.820 3.362                  | 26.056 2.068                  | −1.252 0.111         |
| Smoking   | 0.125 0.342                   | 0.188 0.403                   | −0.473 0.320         |
| Work stressors (workplace observation) | 2.263 0.637                  | 2.681 0.601                  | −1.913 0.033         |
| Job control (workplace observation)  | 2.547 0.666                   | 3.008 0.673                   | −1.947 0.969         |
| Domestic work after regular work (housework, child care, caring for relatives) | 1.869 2.171                   | 1.864 2.164                  | 0.006 0.990         |
| Domestic work on Sundays (housework, child care, caring for relatives) | 3.536 4.235                   | 4.244 4.773                   | −0.444 0.330         |
| Norepinephrine on workdays [pmol/(min × kg)] | 3.086 1.076                  | 3.994 1.181                  | −2.273 0.015         |
| Noon      | 3.086 1.076                   | 3.994 1.181                   | −2.273 0.015         |
| End of work | 2.970 0.944                  | 3.684 1.350                   | −1.736 0.047         |
| Evening   | 3.212 1.182                   | 3.489 1.099                   | −0.688 0.249         |
| Epinephrine on workdays [pmol/(min × kg)] | 0.996 0.651                   | 1.219 0.414                   | −1.156 0.129         |
| Noon      | 0.996 0.651                   | 1.219 0.414                   | −1.156 0.129         |
| End of work | 0.754 0.345                  | 0.967 0.363                   | −1.700 0.050         |
| Evening   | 0.891 0.373                   | 0.664 0.215                   | 2.102 0.977         |
| Norepinephrine on Sunday [pmol/(min × kg)] | 2.862 1.051                   | 2.906 1.057                   | −0.119 0.453         |
| Noon      | 2.862 1.051                   | 2.906 1.057                   | −0.119 0.453         |
| Afternoon | 2.675 1.367                   | 2.106 0.756                   | 1.456 0.920         |
| Evening   | 2.833 1.080                   | 2.373 0.741                   | 1.405 0.915         |
| Epinephrine on Sunday [pmol/(min × kg)] | 0.556 0.225                   | 0.433 0.186                   | 1.826 0.961         |
| Noon      | 0.556 0.225                   | 0.433 0.186                   | 1.826 0.961         |
| Afternoon | 0.608 0.249                   | 0.585 0.302                   | 0.231 0.590         |
| Smoking   | 0.125 0.342                   | 0.188 0.403                   | −0.473 0.320         |

* P-values from one-tailed hypothesis tests.
1 Smoking indicates nonsmoking (0 = cigarettes/day) versus smoking (1 = >1 cigarette/day).
2 We expected job control to be lower in the group with a high level of musculoskeletal pain, but indeed it turned out to be higher. For job control and other marked indicators the expectation was directional, and the one-tailed P-value reflects the finding that our expectations were wrong.
3 Corrected for unequal variances.

Table 2. Multilevel analysis of norepinephrine and epinephrine excretion rates on workdays (noon, end of work, evening). (parameter = fixed parameter estimates, SE = standard error, random effects = variance and covariance estimates of parameters that are allowed to vary on level 2, VAR = variances, IGLS = Iterative Generalized Least Squares)

| Variable | Norepinephrine | Epinephrine | Parameter | SE | P-value | Parameter | SE | P-value |
|----------|----------------|-------------|-----------|----|---------|-----------|----|---------|
| Level-2 predictor variables (person) | | | | | | | | |
| Age      | 0.034 0.017 0.025 | 0.001 0.007 0.469 | | | | | | |
| Smoking   | 0.030 0.011 0.003 | 0.004 0.005 0.176 | | | | | | |
| Work stressors | 0.244 0.398 0.269 | 0.201 0.163 0.109 | | | | | | |
| Work control | 0.286 0.262 0.138 | −0.048 0.108 0.328 | | | | | | |
| Musculoskeletal pain | 0.267 0.083≤0.001 | 0.083 0.034 0.007 | | | | | | |
| Work stressors × musculoskeletal pain | 0.440 0.191 0.011 | 0.095 0.078 0.112 | | | | | | |
| Level-1 predictor variables (measurement) | | | | | | | | |
| Domestic work | 0.005 0.110 0.484 | −0.013 0.045 0.386 | | | | | | |
| Time of day | −0.190 0.083 0.022 | −0.148 0.034≤0.001 | | | | | | |
| Workweek 1 | 0.387 0.166 0.009 | −0.034 0.068 0.308 | | | | | | |
| Workweek 2 | 0.235 0.170 0.083 | 0.070 0.069 0.156 | | | | | | |
| Work stressors × time of day | −0.077 0.138 0.287 | −0.059 0.056 0.147 | | | | | | |
| Musculoskeletal pain × time of day | 0.082 0.032 0.006 | −0.037 0.013 0.003 | | | | | | |
| Random effects | | | | | | | | |
| Intercept | 2.895 0.173 | 0.857 0.071 | | | | | | |
| VAR intercept Level 2 | 0.352 0.125 | 0.060 0.021 | | | | | | |
| VAR intercept Level 1 | 1.251 0.113 | 0.208 0.019 | | | | | | |
| Z*log likelihood (IGLS) | 881.170 | 387.146 | | | | | | |

* Sample size: N=275 (norepinephrine), N=274 (epinephrine) measurements from 32 participants.
1 One-tailed Wald Test (parameter estimates/standard error). The Wald Test is one-tailed for variances (VAR).
2 Smoking indicates the number of cigarettes smoked on the measurement day.
perception of psychosocial risk factors that is related to musculoskeletal pain, but also their work conditions as assessed by independent raters. Moreover, work stressors were related to the excretion rates of catecholamine and musculoskeletal disorders, associations that are in accordance with current models of occupational musculoskeletal pain. Work stressors—for example, time pressure—lead to a mobilization of resources. A mobilization of resources corresponds with physiological responses, including increased activation of the sympathetic–adrenal medullary system. While this activation is useful in a normal range, two deviations from “normal” may occur. The first deviation refers to excretion rates that are too high during demanding situations, that is, reactivity is too high. The second deviation refers to impaired recovery, that is, a prolonged response that takes too long to return to normal (or, possibly, never gets back to normal excretion rates). These two deviations

Discussion

Many researchers in occupational health sciences recommend measuring risk factors and dependent variables with different methods (22, 23). This study included an observational assessment of work conditions and found associations between work stressors and musculoskeletal pain. There are only a few other studies that had done so earlier (24). Therefore, this study contributes to the increasing evidence that it is not only people’s

Table 3. Multilevel analysis of norepinephrine and epinephrine during work (noon, end of work) and corresponding times on Sundays. a (parameter = fixed parameter estimates, SE = standard error, random effects = variance and covariance estimates of parameters that are allowed to vary on level 2, VAR = variances, IGLS = Iterative Generalized Least Squares)

| Parameter | SE  | P-value b | Parameter | SE  | P-value b |
|-----------|-----|-----------|-----------|-----|-----------|
| Level-2 predictor variables (person) | | | | | |
| Age | 0.025 | 0.012 | 0.017 | -0.002 | 0.004 | 0.312 |
| Smoking c | 0.023 | 0.008 | 0.002 | 0.002 | 0.003 | 0.203 |
| Work stressors | 0.285 | 0.337 | 0.199 | 0.008 | 0.120 | 0.475 |
| Work control | 0.254 | 0.187 | 0.087 | -0.024 | 0.063 | 0.650 |
| Musculoskeletal pain | 0.115 | 0.073 | 0.057 | 0.031 | 0.026 | 0.475 |
| Work stressors × musculoskeletal pain | 0.209 | 0.167 | 0.106 | -0.029 | 0.060 | 0.313 |
| Level-1 predictor variables (measurement) | | | | | |
| Domestic work | 0.058 | 0.039 | 0.070 | -0.013 | 0.014 | 0.177 |
| Day (1 = workday, 2 = Sunday) | -0.864 | 0.113 | -0.001 | -0.290 | 0.041 | -0.001 |
| Time of day | -3.164 | 0.334 | -0.001 | -1.114 | 0.122 | -0.001 |
| Workweek 1 | 0.381 | 0.133 | 0.002 | -0.067 | 0.049 | 0.084 |
| Workweek 2 | 0.206 | 0.135 | 0.153 | -0.028 | 0.049 | 0.287 |
| Work stressors × day | -0.368 | 0.183 | 0.022 | 0.031 | 0.067 | 0.319 |
| Musculoskeletal pain × day | -0.047 | 0.042 | 0.134 | -0.026 | 0.015 | 0.044 |
| Work stressors × musculoskeletal pain × day | -0.077 | 0.098 | 0.218 | 0.005 | 0.036 | 0.441 |
| Random effects | | | | | |
| Intercept | 3.396 | 0.253 | 1.120 | 0.092 |
| VAR intercept level 2 | 0.033 | 0.012 | 0.003 | 0.001 |
| VAR intercept level 1 | 0.184 | 0.014 | 0.025 | 0.002 |
| 2×log likelihood (IGLS) | 1087.052 | 358.067 |

a Sample size: N=358 (norepinephrine), 357 (epinephrine) measurements from 32 participants.

b One-tailed significance level of the Wald test (parameter estimates/standard error). The Wald test is one-tailed for variances (VAR).

c Smoking indicates the number of cigarettes smoked on measurement days.

Figure 1. Interaction between work stressors and musculoskeletal pain (MSP) with respect to workday urinary norepinephrine.

Sunday excretion rates of catecholamines among those who reported a high level of musculoskeletal pain and had high levels of work stressors

The regression of workday and Sunday catecholamines with respect to the predictor variables yielded a significant effect for day, indicating that the urinary excretion rates of catecholamine were significantly higher on workdays than on Sundays (table 3). They were also higher at 1200 than at 1700. Age and smoking were significantly positively related to higher excretion rates of norepinephrine. Contrary to hypothesis 4, the participants who reported musculoskeletal pain while being exposed to a high level of work stressors did not show persistently higher urinary excretion rates of norepinephrine on Sunday. There was no significant prediction according to musculoskeletal pain or work factors, nor was there an interaction between these variables.
Research on stress at work shows that elevated norepinephrine levels in urine predominantly characterize the mobilization of physical resources (26). When the situation mainly requires an activation of mental resources, predominantly epinephrine is activated. Depending on the remaining resources of the person, an elevation of epinephrine may correlate with feelings of activation or, in case of depleted resources, with feelings of anxiety and distress (27). With regard to the activation of muscles, however, mental load and physical load show synergistic effects (4). Indeed, synergistic effects of mental load and physical load have even been observed on the level of disc pressure (28). We, therefore, argue that the effects of physical load and mental load combine with regard to musculoskeletal pain, and their potential mediators, norepinephrine and epinephrine, combine as well. Thereby, we consider norepinephrine as more proximally linked to musculoskeletal pain through its more direct physiological effect on muscles. Nevertheless, even if norepinephrine is predominantly activated by physical load, and epinephrine by mental load, it seems likely that, in most stressful situations at work, both catecholamines will be elevated. In our study, the correlations showed a pattern that conforms to the view of norepinephrine as the more proximal link to musculoskeletal pain. Furthermore, norepinephrine was associated with work stressors. This finding is in line with the results of previous studies that showed a similar association between work conditions, musculoskeletal pain, and norepinephrine excretion (3, 10). Both sustained activation during work and lack of muscular relaxation after work may cause pain. The norepinephrine release during work was higher among those who reported pain and were exposed to a high level of work stressors at the same time. The excretion rates of norepinephrine and epinephrine were still elevated in the evening on workdays, but this difference was not significant. There was a tendency for a slightly lower norepinephrine excretion among those who were exposed to a high level of work stressors but did not report musculoskeletal pain. One might argue that these persons might be more resilient to stressors, cope more successfully with stressors, or show reactivity to stressors with a physiological system other than the sympathetic–adrenal medullary system (individual response specificity). Further studies should take into consideration these potential individual moderators. Contrary to hypothesis 4, those with musculoskeletal pain and work stressors did not show impaired recovery during their days off. One explanation for the lack of differences in musculoskeletal pain in unwinding and recovery may have been the study design itself. The participants were instructed to avoid heavy work and sports in leisure time so that the urinary excretion rates of catecholamine would reflect unwinding and recovery after work. Therefore, the comparably little workload during leisure time from household chores, child care, and caring for relatives may not be representative of the daily life of the participants. The norepinephrine levels at work were comparable with Danish reference values of urinary epinephrine and norepinephrine for 120 Danish healthy persons performing their routine work (29). For norepinephrine, geometric mean levels, expressed as micromoles per mole of creatinine, for a time window from 1530 to 1830 was 24.2 and the range was 8.5–67.1. The respective norepinephrine values in our study, expressed as micromoles per mole of creatinine, at 1700 was 26.11, and the respective range was 13.9–52.9. Therefore, the norepinephrine levels (and also epinephrine excretion) were highly comparable with the Danish reference values.

The job demand–control model (30) postulates that job control attenuates the effects of job demands on health and well-being. In this study we found no preventive effect of job control, nor a buffering effect of job control on the influence of work demands. The study even showed, unexpectedly, a higher level of observed job control among those reporting more musculoskeletal pain. Meier et al (31) recently found that job control attenuated the effects of stressors on musculoskeletal pain only for people with an internal locus of control. For people with an external locus of control, job control actually predicted more pain as stressors increased. In our study, locus of control was not assessed, but it is possible that the participants suffering from musculoskeletal pain had a more external locus of control, interfered by the effects of job control. The small sample size and the sample of healthy men with comparably well-designed jobs limit the generalizability of the study results. Therefore, there is a need for replication. On the other hand, the multimethod and repeated measures approach rules out sources of bias that weaken the conclusions that can be drawn from many other studies. This study, therefore, adds plausibility to an important possible psychobiological mechanism that links psychosocial work conditions to musculoskeletal disorders. Against the background of increasing time pressure for many employees (32), this study furthermore promotes the redesign of jobs as a way of preventing musculoskeletal disease.
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