Original article
Scand J Work Environ Health 2002;28(3):168-175
doi:10.5271/sjweh.661

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Refers to the following texts of the Journal: 1997;23(4):243-256 1998;24(6):449-464

Key terms: back disorder; collagen metabolism; connective tissue; patient handling; physical workload; prospective cohort study; serum marker; spine; student nurse

This article in PubMed: www.ncbi.nlm.nih.gov/pubmed/12109556
Physical workload of student nurses and serum markers of collagen metabolism

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Objectives  This study explored the association between biomarkers of type I collagen metabolism and exposure to physical workload.

Methods  In a prospective cohort study, serum concentrations of markers of type I collagen synthesis and degradation were assessed monthly for student nurses who worked as nurses for a period of 6 months and compared with those of a reference group. The number of patient-handling activities was estimated from observations at the workplace. Linear generalized estimating equations were used to analyze differences in the serum concentrations of the biomarkers between the exposed group and reference group, as well as to analyze whether the number of patient-handling activities was associated with serum concentrations of the biomarkers.

Results  Serum concentrations of the biomarkers were found to differ between the groups. The biomarkers reflected a higher anabolism of type I collagen in the exposed group when compared with that of the reference group. An analysis of the effect of the number of patient-handling activities revealed that a higher exposure was associated with higher effective type I collagen synthesis within the exposed group.

Conclusions  These results indicate that serum concentrations of these biomarkers of type I collagen metabolism can reflect differences in exposure between contrasting groups, and also varying levels of exposure between persons within an occupation.

Key terms  back disorders, collagen, connective tissues, patient handling, prospective cohort study, spine.

Reviews of the epidemiologic evidence on work-related risk factors for back disorders concluded that evidence exists showing that physical demands of work (manual materials handling, bending and twisting, and heavy physical load) can be associated with the occurrence or aggravation of low-back symptoms (1–5). However, evidence on causal relationships remains inconclusive due to the lack of knowledge as to the biological relevance of different load parameters and the large variability in susceptibility between persons. Although methods and strategies to assess external exposure accurately and biomechanical models to estimate the response of the tissues to these exposures have been developed (6, 7), very little is known about biologically effective doses and early biological effects in vivo. For a better understanding of the pathomechanisms of back disorders, and a definition of adequate parameters of exposure and effect, knowledge of the effects of load is essential at the tissue level. As indicators of processes at a molecular level, biomarkers may provide an objective, quantitative measure of the response of the structures in the back to physical load. Unfortunately, no validated biomarkers are yet available for assessing the effects of back load.

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Because of the complex anatomy of the back and the multifactorial etiology of back disorders, it is complicated to identify relevant biomarkers. Physical load has been shown to cause responses in the spinal structures leading to changes in the metabolism of the extracellular matrix of connective tissues (8); therefore it is hypothesized that matrix metabolites in blood may provide potential biomarkers of the biological effects of physical back load on these structures. In connective tissues the extracellular matrix is responsible for the overall structural integrity of the tissue. When the level of load is slowly increased, the result may be an adaptive remodeling of the tissues. However, if the increase in load is sudden or large, the cells may not adapt quickly enough, and damage may result (9). Several laboratory assays have been developed to measure different matrix metabolites in blood to monitor metabolic alterations in connective tissues. In recent years, commercially available assays to measure type I collagen turnover have received considerable research attention as biomarkers for the effects of pathological processes and physical exercise on various connective tissues. Synthesis can be assessed by analyzing serum concentrations of carboxyterminal propeptide of type I collagen (PICP), whereas breakdown can be detected by analyzing carboxyterminal telopeptide region of type I collagen (CTx) (10). The ratio between PICP and CTx provides an estimate of type I collagen metabolism as a dynamic process of synchronously occurring anabolic and catabolic effects.

Type I collagen is one of the main constituents of the extracellular matrix of most connective tissues, it is tensile in nature and is found particularly in tissues subjected to tension and compression, like tendon, bone, and the anulus fibrosus of the intervertebral disc. Because of the widespread appearance of type I collagen in the human body, essentially, serum concentrations of markers of type I collagen metabolism may originate from various tissues. However, if it is assumed that collagen metabolism is not significantly altered in unaffected tissues, changes in serum concentrations of these biomarkers may provide relatively site-specific information when assessed in combination with appropriate exposure information. PICP and CTx have been used to investigate the effect of physical loading on various connective tissues induced by exercise (11–18) and active back rehabilitation (19).

To explore PICP and CTx as potential biomarkers of the effects of physical demands of work, we investigated the association between these biomarkers and exposure to work involving heavy manual materials handling. The objective of the study was to determine whether serum concentrations of PICP and CTx in young nurses and the PICP:CTx ratio differ from those of a sedentary reference group during a 6-month period. Furthermore, we investigated whether differences in exposure to patient-handling activities within the exposed group were associated with serum concentrations of PICP and CTx and their ratio.

**Subjects and methods**

**Design and study population**

A prospective cohort study was performed among student nurses during a 6-month work-placement period at the end of their vocational training. Apart from earlier short work-placement periods in the 4-year training program, this was the first period in which the student nurses performed nursing work for a long time at one stretch, for 4 days a week. Students were invited to participate if they had not had a job prior to the start of their vocational training. A total of 30 student nurses participated (3 men and 27 women). The participants went to different hospital departments, 10 in the internal medicine department, 7 in the surgery department, 6 in the obstetrics and gynecology department, 4 in the pediatric department, and 3 in the orthopedics department. Except for two participants, who worked in regional hospitals, all the participants worked in an academic hospital.

A reference group of 30 students (2 men and 28 women) was recruited among student nurses in the first year of their vocational training, when they had mostly theoretical education.

Once every month, starting the week before the work-placement period until 1 month after the end of the period, the participants visited our institute to donate a blood sample and to complete a questionnaire (time 0–7 months in figure 1). The reference group did not donate blood on the eighth occasion (at 7 months) because they had started to participate in a short work-placement period by then. Individual characteristics such as age, height, and weight were assessed using data from the baseline questionnaire. Furthermore, the subjects were asked about the use of medication and whether they suffered or had suffered from musculoskeletal traumas in the previous year or from joint diseases, liver or kidney diseases, blood diseases, or metabolic...
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diseases (like diabetes), because of the potentially
confounding effect on the serum concentrations
of the collagen markers. Table 1 presents the charac-
teristics of the study population. The participants were fully
informed about the procedures and filled out an in-
formed consent form. The study was approved by the
Ethics Committee of the Academic Medical Center, Uni-
versity of Amsterdam, The Netherlands.

Assessment of physical load
Exposure to patient-handling activities during the work-
placement period was estimated by means of onsite ob-
servations using TRAC (Task Recording and Analysis
on Computer) software (20). Each student nurse was
observed for a full dayshift in the third or fourth month
of the work-placement period. The frequency and dura-
tion of activities involving patient handling were ob-
served on a real-time basis. These activities were divid-
ed into lifting patients, moving patients around in bed,
transferring patients from bed to (wheel)chair and vice
versa, pushing or pulling a wheelchair or bed contain-
ing a patient.

Changes in exposure to physical load during the
work-placement period were assessed using the month-
ly questionnaire. Apart from the number of shifts they
had worked per week, the student nurses were asked to
estimate the percentage of worktime they had actually
performed tasks as opposed to observing senior nurses
and taking breaks.

To estimate exposure to physical load during leisure
time, all the subjects were asked whether they had had
side jobs, and, if so, for how many hours during the pre-
ceding month. On the basis of the job title, the jobs were
classified as either physically demanding or not (21).
Furthermore, the subjects reported whether they had
been engaged in physically demanding sports and other
leisure-time activities during the preceding month, and,
if so, for how many hours.

Exposure characteristics of the study population
The results of the workplace observations showed that
the average duration of a dayshift was 8 hours and 7
(SD 27) minutes. The student nurses lifted 5 (SD 7) pa-
tients per shift. Moving patients around in bed averaged
1 (SD 2) times per day. The average frequency of push-
ing or pulling a bed or a wheelchair with a patient was
7 (SD 7) times and 2 (SD 3) times per shift, respective-
ly, with an average duration of 4.4 (SD 5.5) minutes and
21 (SD 40) seconds per shift, respectively.

In the first month of the work-placement period, the
exposed subjects spent an average of 46 (SD 17)% of
their worktime performing tasks themselves; this per-
centage increased to 67 (SD 17)% in the second month.
In the last 4 months the average percentage gradually
increased from 74 (SD 15)% in the third month to 80
(SD 10)% in the final month.

At the different time points during the study period,
the average number of hours worked in physically de-
manding side jobs ranged from 13 to 19 hours per month
in the exposed group and from 13 to 16 hours per month
in the reference group. The average number of hours
spent on weight-bearing sports or leisure-time activities
ranged from 4 to 10 hours per month in the exposed group
and from 4 to 9 hours per month in the reference group.

Biomarkers
Blood samples of 7 ml were collected by means of ven-
puncture in the antecubital vein. To minimize the pos-
sible effect of circadian rhythmicity, repeated blood
samples were drawn at about the same time of day. Af-
fter coagulation, the blood was centrifuged, and serum
was stored at –20 °C until analyzed. The serum concen-
trations of PICP were measured by the Prolagen-C sand-
wich enzyme-linked immunosorbent assay (ELISA)
(Metra Biosystems, Mountain View, United States). The
CTx serum concentrations were analyzed using the se-
rum CrossLaps™ One Step ELISA (OSTEOMETER,
Herlev, Denmark) (22). All of the samples of each sub-
ject were analyzed with the same kit, with the subjects
of both study groups randomly divided over 16 kits of
each of the assays. All of the samples were analyzed in
duplicate. The intraassay precision (coefficient of vari-
ation) was 3.5% for PICP and 5.1% for CTx, whereas
the interassay coefficients of variation were 9.6% and
10.7%, respectively. To illustrate the effective change
in collagen metabolism, the PICP:CTx ratio was cal-
culated for each subject.

Table 1. Characteristics of study population.

| Group          | Age (years) | Height (meters) | Weight (kilograms) | Body mass index (kg/m²) |
|----------------|-------------|-----------------|-------------------|------------------------|
|                | Mean        | SD              | Mean              | SD                     | Mean        | SD         |
| Exposed a (N=30) | 22.4        | 1.9             | 1.73              | 0.07                   | 65.5        | 7.8        | 21.8        | 2.0        |
| Reference b (N=30) | 18.3        | 1.4             | 1.72              | 0.07                   | 66.3        | 11.2       | 22.4        | 3.3        |

*3 men, 27 women.  b2 men, 28 women.
Statistical analysis

The statistical analyses consisted of several steps. To test for baseline differences in the serum concentrations of the biomarkers between the groups, analyses of variance were performed for each of the biomarker parameters. To determine whether the serum concentrations of the biomarkers during the work-placement period differed between the exposed and reference groups, linear generalized estimating equations (GEE) were used (23). A GEE analysis accounts for the within-subject correlations between repeated measurements. Both time-dependent and time-independent covariates can be analyzed. Use of the GEE provides a pooled analysis of between-subject and within-subject associations (24). Separate analyses were performed with PICP and CTx concentrations and the PICP : CTx ratio during the work-placement period (time 1–6 months) as the time-dependent outcome variable and group and time (exact number of days) as the determinants. The interactions of “group × time” and “gender × group” were examined; if not significant, the interaction term was removed from the model.

Subanalyses were performed for the exposed group on the associations between the serum concentrations of each of the biomarkers and exposure to patient-handling activities. For each of the three biomarker parameters, a GEE analysis was conducted with the biomarker values during the work-placement period (time 1–6 months) as the time-dependent outcome variable, and the total number of patient-handling activities during one dayshift as the time-independent determinant and time (exact number of days) as an additional determinant. The interaction of “patient-handling activities × gender” was examined; if not significant, the interaction term was removed from the model. In all the analyses, the baseline serum concentration of the outcome variable, body mass index (BMI), and gender were included in the model as time-independent determinants.

The Statistical Package for Interactive Data Analysis (SPIDA, version 6.05) was used for the GEE analyses. The Statistical Package for Social Sciences (SPSS for Windows, version 9.0) was used for the analysis of variance.

Results

Baseline values

The serum concentrations of PICP and CTx and the PICP : CTx ratio determined each month are shown for both groups in figure 2. An analysis of variance showed that the two groups did not differ with respect to the baseline serum concentrations of PICP (F_{1,56}=0.01, P=0.95) and CTx (F_{1,56}=1.80, P=0.19), and the PICP : CTx ratio (F_{1,56}=1.69, P=0.19).

Comparison of the exposed and reference groups

The results of the GEE analyses on the effect of group on the serum concentrations of PICP and CTx and the PICP : CTx ratio during the 6-month work-placement period are presented in table 2. For PICP, the time...
Table 2. Regression coefficients and 95% confidence intervals (95% CI) of the generalized estimating equations with serum concentrations of propeptide of type I collagen (PICP), carboxyterminal telopeptide region of type I collagen (CTX), and the PICP:CTX ratio measured once a month during the 6-month work-placement period as the outcome variables and group as the determinant, adjusted for time, gender, body mass index (BMI), and interaction time × group if significant (N=60).

| Biomarker       | ß      | 95% CI        |
|-----------------|--------|---------------|
| PICP            |        |               |
| Intercept       | 94.4   | 44.3–144.5    |
| Group a         | -14.0  | -24.2– -3.9   |
| Baseline PICP   | 0.7    | 0.6– 0.8      |
| Time            | -0.2   | -0.3– -0.0    |
| Gender a        | -14.4  | -29.7– 1.0    |
| BMI             | -0.3   | -1.5– 0.9     |
| Time × group    | 0.1    | 0.0– 0.2      |
| CTx             |        |               |
| Intercept       | 0.9    | 0.9– 1.6      |
| Group a         | -0.1   | -0.1– 0.0     |
| Baseline CTx    | 0.4    | 0.3– 0.6      |
| Time            | 0.0    | -0.0– 0.0     |
| Gender a        | -0.3   | -0.5– -0.1    |
| BMI             | -0.0   | -0.0– 0.0     |
| PICP/CTX        |        |               |
| Intercept       | -404.4 | -7299.9– -79.0|
| Group a         | 63.6   | 13.3–113.9    |
| Baseline ratio PICP:CTX | 0.3 | 0.2– 0.4 |
| Time (days)     | 0.2    | -0.3– 0.5     |
| Gender a        | 149.3  | 51.8–246.7    |
| BMI (kg/m²)     | 13.2   | 4.0– 22.4     |

a Gender code: reference group=1, exposed group=2.
b Gender code: men=1, women=2.

course of the serum concentrations differed for the two groups, as indicated by the significant interaction between time and group (P=0.05). When adjusted for this interaction, the regression coefficient for group was significant (P=0.01). The regression coefficient for group was negative, the negative result indicating that the serum concentrations of PICP were lower in the exposed group than in the reference group. In the CTx model, the regression coefficient for group was also negative, but not significant (P=0.07). The PICP:CTx ratio was higher in the exposed group (P<0.01).

 Associations between patient-handling activities and biomarkers

Table 3 shows the results of the GEE analyses on the associations between the serum concentrations of each of the biomarkers and the PICP:CTX ratio during the work-placement period and the observed number of patient-handling activities on a representative dayshift in that period. In the analyses of all three biomarker parameters, the interaction “gender × number of patient-handling activities” was significant (PICP, P=0.05; CTx, P<0.01; PICP:CTX ratio, P<0.01) and therefore indicated differences in the effects of the number of patient-handling activities on the serum concentrations of the biomarkers between the men and women. When adjusted for these interaction effects, the regression coefficients for patient handling in the models of PICP and CTx and the PICP:CTx ratio were significant (PICP, P=0.05; CTx, P<0.01; PICP:CTX ratio, P<0.01). For PICP and CTx the coefficient was negative, whereas for the PICP:CTx ratio a positive coefficient was found.

Table 3. Regression coefficients and 95% confidence intervals (95% CI) of the generalized estimating equations with serum concentrations of propeptide of type I collagen (PICP), carboxyterminal telopeptide region of type I collagen (CTx), and the PICP:CTX ratio measured once a month during the 6-month work-placement period in the exposed group as the outcome variables and the frequency of patient-handling activities in one shift as the determinant, adjusted for time, gender, body mass index (BMI), baseline value of the determinant and interaction gender × patient-handling activities (N=30).

| Biomarker       | ß      | 95% CI        |
|-----------------|--------|---------------|
| PICP            |        |               |
| Intercept       | 86.9   | 9.6– 164.2    |
| Patient handling | -5.1 | -10.2– 0.0    |
| Baseline PICP   | 0.7    | 0.6– 0.8      |
| Time            | -0.1   | -0.0– 0.2     |
| Gender a        | -26.3  | -45.6– 6.9    |
| BMI             | -0.3   | -3.0– 2.4     |
| Gender × patient handling | 2.6 | 0.0– 5.2 |
| CTx             |        |               |
| Intercept       | 1.3    | 0.9– 1.6      |
| Patient handling | -2.5 | -4.1– 0.9     |
| Baseline CTx    | 0.3    | 0.1– 0.6      |
| Time            | 0.0    | -0.0– 0.0     |
| Gender a        | -0.4   | -0.4– 0.3     |
| BMI             | 0.0    | -0.0– 0.0     |
| Gender × patient handling | 1.2 | 0.4– 2.1 |
| PICP/CTX        |        |               |
| Intercept       | -856.4 | -1228.7– -484.1|
| Patient handling | 20.9 | 8.3– 33.5     |
| Baseline ratio PICP:CTX | 0.3 | 0.1– 0.5 |
| Time            | 0.1    | -0.4– 0.7     |
| Gender a        | 259.6  | 200.6– 318.5  |
| BMI             | 29.4   | 16.0– 42.8    |
| Gender × patient handling | -8.9 | -16.4– 1.5  |

a Gender code: men=1, women=2.

Discussion

In this study a start was made towards exploring potential biomarkers for the effects of occupational physical back load. No valid serum markers specific for metabolic alterations in spinal structures are available. Therefore, PICP and CTx as validated assays for markers of components that are important constituents of, but not specific for, spinal tissues were considered potential
biodmers. Nurses were studied since their work, and especially their tasks involving manual handling of pa-
tients, has been characterized as hazardous with respect
to low-back disorders (4, 25, 26). However, nursing
work also involves physical loading of the whole
body. Since the serum concentrations of PICP and
CTx are not specific for spinal tissues, the results of
this study provide information on the effects of
physical workload in general, and discernment of the
relative contributions of the specific sites in the hu-
man body is not possible.

The longitudinal design of this study with monthly
repeated measurements enabled an analysis of the tem-
poral trend of the biomarkers. Biological variation of the
biomarkers is considerable. In the statistical analyses
adjustments were made for the baseline values of the
biomarkers. Despite considerable intra- and interindi-
vidual variability, serial comparison of the values of
the exposed and reference subjects indicated that the
effects of exposure exceeded the normal biological
variability. As the reference group had no occupa-
tional exposure, there was a considerable contrast in
exposure to physical load between the two study groups.
The rationale for the choice of student nurses as the
study population was the relative lack of occupational
load history. Furthermore, the population consisted of
young, healthy subjects and therefore limited the con-
 founding effects of degenerative connective tissue dis-
orders.

Serum concentrations of biomarkers of type I colla-
gen metabolism were found to be different in subjects
exposed to physically demanding work when compared
with unexposed subjects. Monthly repeated measure-
ments of healthy young student nurses exposed to phys-
ical workload for a period of 6 months showed lower
serum concentrations of the marker of type I collagen
anabolism (PICP) when compared with the results of the
reference group. The concentrations of the marker of
catabolism (CTx) also tended to be lower, but this dif-
ference was not statistically significant. The PICP : CTx
ratio, on the other hand, was significantly higher for the
exposed subjects. Thus, in the 6-month exposure peri-
od, type I collagen metabolism was more anabolic in the
exposed group. Within the exposed group, longitudinal
analyses on associations between biomarkers during the
work-placement period and number of patient-handling
activities on a workday revealed a picture similar to that
of the analyses on the effect of group (ie, higher expo-
sure was associated with lower concentrations of PICP
and CTx but with a higher PICP : CTx ratio). Consider-
ing the fact that the longitudinal design and the selec-
tion of the study population confined the effects of poten-
tial confounders, the conclusion that the differences
in biomarker concentrations can be ascribed to the ef-
fects of physical workload is a plausible one. In healthy
connective tissues, the cells respond to mechanical
load by increased synthesis of collagen. This adap-
tive remodeling serves to strengthen the tissue to
withstand higher forces (9). As already stated, the
nonspecificity of the biomarkers together with the
study design does not allow discernment of the rela-
tive contributions of connective tissues at specific sites
of the human body. Nevertheless, it can be hypothesized
that loading of the spinal structures played an important
role in the differences in biomarker concentrations found
in this study. The workplace observations provided only
crude exposure measures, but from biomechanical stud-
ies it is known that in patient-handling activities es-
pecially the spinal structures are subjected to high
compression and shear forces (26). Experiments on
cadaver spines and animal studies have shown that
exposure to these forces was associated with increased
collagen synthesis in intervertebral discs (27–29). Most
studies have focused on intervertebral discs since ma-
trix changes and structural damage to this tissue are con-
sidered main features in the association between load
and back disorders. However, adaptive remodeling
responses also occur in other spinal structures, as has
been illustrated by the finding of dense vertebrae in
weightlifters and osteophytes around the margins of
vertebral bodies in people involved in heavy manual la-
bor (8).

This study was not able to show a temporal correla-
tion between exposure and effect. Changes in exposure
were assessed crudely, but the fact that the exposed
group was unexposed at the start of the study, and spent
a lot of time observing senior nurses during the first
months of the study period, suggests an increase in ex-
posure during the first months, which stabilized in the
third month. However, in the reference as well as the
exposed groups, the biomarker concentrations did not
change significantly in time during the study period.
PICP was an exception, as the time course differed sig-
ificantly between the groups, but, when the time course
of PICP was analyzed separately for each group, the
changes in time were not significant in either group.
Blood concentrations of type I collagen metabolic mark-
ers probably reflect the actual status of type I collagen
metabolism (15). With respect to the time frame, it has
been shown that, in response to a single bout of strenu-
ous exercise (ie, running a marathon), type I collagen
synthesis increased, reaching a peak after 3 days and
returning to the baseline value 5 days after the run (16).
The current study was not designed to assess the possi-
ble effects of acute changes in physical load on collagen
metabolism. The purpose of the monthly repeated mea-
surements of the biomarkers was to provide information
on changes in the state of the tissues. The state of the
tissue and the level of adaptation determine the response
in collagen metabolism. Degradation mainly occurs

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when the load exceeds the strength of the tissue (15). The serum concentrations of the student nurses suggested that the level of physical load did not cause enhanced degradation in the 6-month study period, but merely yielded adaptive remodeling of the tissues. It would be interesting to assess follow-up measures for several years in high-risk occupations to investigate the long-term effects of physical load. In addition, a comparison of the responses in newly exposed workers with those of workers who have been exposed for years may provide useful information on the tissue responses to physical load.

The results of this exploratory study showed that serum concentrations of PICP and CTx and the PICP : CTx ratio are associated with exposure to physically demanding work. Although the relative contribution of back loading could not be determined, these results may provide a productive starting point for further research. For the potential biomarkers to be useful in identifying relevant load parameters and assessing biologically effective doses, studies are needed on aspects like the biological relevance and individual dose-response relationships. Hence future studies should, for instance, combine biomarkers of collagen synthesis and degradation with imaging studies on molecular changes in specified tissues on one hand and with biomechanical models estimating forces on the spinal structures on the other. Furthermore, ongoing advancements in biochemical technology may enhance the development and validation of valid biomarkers for the effects of back load. Recent studies reported on increased knowledge and the development of more sophisticated assays for assessing bone and cartilage metabolism (30–32). This progress enhances the possibilities of obtaining tissue-specific information. In combination with assays for proteoglycan components or type II collagen metabolites, PICP and CTx may provide a more complete picture of tissue responses to physical load.

Acknowledgments

We thank Allard van der Beek for his contribution to the statistical analyses and his comments on the text. Furthermore we are grateful to Sijmen Kuiper and Suzan van Damme for performing the laboratory analyses.

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Received for publication: 6 August 2001