The onset of deep abdominal muscles activity during tasks with different trunk rotational torques in subjects with non-specific chronic low back pain

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ABSTRACT

Background: Although delayed onset of the deep abdominal muscles activity in subjects with non-specific chronic low back pain (CLBP) has been suggested to be related to trunk rotational torque, no study has examined the onsets associated with non-specific CLBP during a variety of tasks with different trunk rotational torque. The aim of this study is to compare the onsets of deep abdominal muscles activity among tasks with different trunk rotational torques in subjects with and without non-specific CLBP.

Methods: Twelve subjects with non-specific CLBP and 13 control subjects were included. They performed 8 types of upper limb movements. The onsets of muscular activity of bilateral internal oblique-transversus abdominis (IO-TrA) and trunk rotational torque due to the upper limb movements were measured using a surface electromyography and a three-dimensional motion analysis system.

Results: In non-specific CLBP group, right IO-TrA activities were significantly delayed during tasks with left trunk rotational torque compared with the control ($P < 0.05$), while onsets of the left IO-TrA activities were significantly later than those of the control during tasks with right rotational torque of the trunk ($P < 0.05$). There were no significant differences in onsets of both sides IO-TrA during tasks without trunk rotational torque between non-specific CLBP and control groups ($P > 0.05$).

Conclusions: The onsets of IO-TrA activities in subjects with non-specific CLBP were
delayed during tasks with rotational torque of the trunk in the opposite direction, suggesting a possibility that delayed onset of the deep abdominal muscles during rotational torque of the trunk might be etiology of chronic low back pain.
1. Introduction

Non-specific low back pain with no identifiable underlying pathology accounts for 90% of cases of low back pain [1]. Patients with non-specific acute low back pain demonstrate a favorable improvement rate in the first six weeks [2], but approximately 40% of patients will develop non-specific chronic low back pain (CLBP) [3]. One proposed mechanism for non-specific CLBP is instability of the spine due to the lack of neuromuscular control [4].

The deep abdominal muscles play an important role in lumbopelvic stability [5]. Several studies indicated that the onsets of transversus abdominis activities in response to movements with rotational torque of the trunk were prior to those of other trunk muscles [6, 7, 8]. Early activation of the deep abdominal muscles is therefore considered to contribute to the control of spinal stiffness [7, 9]. Dysfunction of the deep abdominal muscles activities could contribute to persistence of low back pain [10].

The onset latency of deep abdominal muscles in rapid movements was delayed in subjects with non-specific CLBP compared to healthy subjects [6, 8]. On the other hand, there were no significant differences in onsets of transversus fibers of the internal oblique and transversus abdominis (IO-TrA) activation during bilateral shoulder flexion between healthy individuals and subjects with non-specific CLBP [11]. Therefore, we hypothesized that the delay of the onset of the deep abdominal muscles would be related to the magnitude of trunk rotational torque during limb movement in subjects with non-specific CLBP and that the
delayed onsets of the deep abdominal muscles in response to movements with rotational
torque of the trunk might be etiology of CLBP. The purpose of the present study was to
compare the onsets of deep abdominal muscles activities among tasks with different rotational
torque in the subjects with and without non-specific CLBP.

2. Methods

2.1 Participants

Twelve subjects with non-specific CLBP and 13 healthy control individuals participated in this study. All subjects were recruited in our University thorough advertisement. Non-specific CLBP was defined as pain and discomfort localized between the 12th rib and the inferior gluteal folds for at least 3 months. Subjects with non-specific CLBP were excluded if they had neurologic symptoms, spinal and abdominal surgery in the past 12 months, pregnancy, observable spinal deformity (e.g., scoliosis or kyphosis) and suspected or diagnosed serious spine pathology. Subjects reported minimal or no pain at the time of testing. All participants provided informed consent and completed the Oswestry disability index (ODI). A visual analogue scale was used to characterize their pain. Descriptive statistics for the two groups are reported in Table 1. There were no differences in age, height, weight, and BMI between groups. This research has been approved by the Institutional Review Board of the authors’ affiliated institutions.
2.2 Experimental procedure

All participants stood on the floor with their arms at their sides while standing comfortably with the eyes directed straight forward. They performed 5 trials of 8 types of rapid upper limb movements: weighted left shoulder flexion and right shoulder extension (WLFRE) (Fig. 1A); left shoulder flexion and right shoulder extension (LFRE) (Fig. 1B); left shoulder flexion (LF) (Fig. 1C); bilateral shoulder extension (LERE) (Fig. 1D); bilateral shoulder flexion (LFRF) (Fig. 1E); right shoulder flexion (RF) (Fig. 1F); left shoulder extension and right shoulder flexion (LERF) (Fig. 1G); weighted left shoulder extension and right shoulder flexion (WLERF) (Fig. 1H). A weight of 1 kg was selected in order to increase the torque without substantially altering the pattern of movement [12]. The participants were instructed to flex their shoulders to approximately 60 degrees and extend them to approximately 40 degrees as fast as possible in response to a beep signal [6]. The emphasis was on the speed of limb movement rather than the accuracy of the angle. The order of each upper limb movement was randomized for each subject. Participants had 3-5 practice trials for each task prior to the measurements.

2.3 Data collection

EMG recordings of the bilateral IO-TrA, anterior deltidoid (AD), and posterior deltidoid
(PD) were acquired using a wireless surface EMG system (WEB-1000; Nihon Kohden Corporation, Tokyo, Japan) with surface-type electrode telemeters that sampled at 1000 Hz. Surface electrodes of the IO-TrA were positioned approximately 20 mm medial and inferior to the anterior superior iliac spine [13], and “surface EMG for a non-invasive assessment of muscles (SENIAM)” recommendations were followed for the AD and PD [14]. Surface electrodes were placed in parallel with the muscles fibers.

Kinematic data were collected using a motion analysis system (Cortex-64 5.5.0.1579, Motion Analysis Corporation, Santa Rosa, CA, USA) with 7 digital cameras (Hawk cameras, Motion Analysis Corporation) sampling at 200 Hz. Ten reflective markers were attached to each segment with double-sided tape bilaterally at the following locations: acromion, medial epicondyle of the humerus, lateral epicondyle of the humerus, radial styloid process, and ulnar styloid process.

2.4 Data analysis

The raw EMG signals were band-pass filtered at 20-500 Hz, full-wave rectified, and filtered using a fourth-order Butterworth low-pass filter with a cut off of 10 Hz. Following data reduction, the onsets of EMG signals were identified using MATLAB software (The MathWorks, Natick, MA, USA) as the point at which the rectified EMG signal first rose above baseline by 1 standard deviation (SD) and lasted for at least 50 msec [11]. The baseline
activity was defined as the average of the 500 msec period preceding the beep signal. In addition, each onset of EMG signal was confirmed visually [15]. Onset of IO-TrA activity was expressed relative to the onset of deltoid EMG signal (t=0) (Fig. 2). For BE and BF, the ipsilateral anterior or posterior deltoid onset was determined as t=0 [11]. For the other six tasks, the onset of activity of the anterior deltoid on the flexing side was established as t=0 [12].

Each marker trajectory was filtered using a fourth-order Butterworth low-pass filter at 12 Hz. Kinematic data were condensed to two-dimensional coordinates in the sagittal plane. Joint reaction forces at the shoulder joint were estimated using free-body diagram force analysis [16]. Trunk rotational torque around a midpoint of the bilateral acromia was calculated from transversus components of the shoulder joint reaction forces. The trunk rotational torque was normalized to body weight for each subject (Nm/kg). The peak rotational torque was determined for analysis of maximum torque during the acceleration phase, defined as the duration from when the angular acceleration at the shoulder first rose above baseline by 2 SD and stayed above this level for 20 msec, to the instant when the angular acceleration crossed the zero line [17]. In addition, peak shoulder angle and peak shoulder angular velocity were calculated during each task.

### 2.5 Statistical analysis
A priori power analysis was performed in G-power 3.1. The sample size was estimated from a pilot study we carried out with 8 subjects (4 subjects with non-specific CLBP and 4 healthy control individuals), for a calculated effect size of partial $\eta^2 = 0.472$.

We performed the power analysis using F-test model of G*Power 3.1. Eleven subjects in each group were deemed sufficient to detect significant differences in the onset of right IO-TrA between groups with a power $(1-\beta)$ of 0.8.

Unpaired t-tests were used to examine differences in age, height, weight, body mass index (BMI), peak shoulder angle, and peak shoulder angular velocity between the non-specific CLBP and control groups. The effects of group factor (non-specific CLBP and control) and task factor (WLFRE, LFRE, LF, LERE, LFRF, RF, LERF and WLERF) on the onsets of right and left IO-TrA activity and peak rotational torque were analyzed using two-way analysis of variance (ANOVA). Where significant main effects were observed, post hoc Tukey’s HSD tests were performed. Significance was set at $\alpha = 0.05$. Statistical analyses were performed using IBM SPSS Statistics 22 software (IBM, Chicago, IL, USA).

3. Results

There was a significant task effect on the peak rotational torque ($P < 0.05$, [effect size (ES) = 0.96]), but ANOVA did not detect a significant group effect (Fig. 3). LERE and LFRF were the smallest and were not significantly different from each other (Fig. 3). WLFRE and
LFRE were not significantly different, nor were LERF and WLERF (Fig. 3). On the other
hand, LERF and WLERF produced significantly greater amounts of peak rotational torques
than RF ($P < 0.05$, [ES = 1.51]) (Fig. 3). WLFRE and LFRE caused greater peak rotational
torques than LF ($P < 0.05$, [ES = 2.09]) (Fig. 3). RF and LF produced greater absolute peak
rotational torques than LERE and LFRF ($P < 0.05$, [ES = 4.28 - 4.58]) (Fig. 3). The peak left
shoulder angle in LFRF and WLERF were significantly different between groups but there
were no significant differences of shoulder angle in other tasks (Table 2). There were no
significant differences of peak shoulder angular velocity in all tasks as well as peak rotational
torque (Table 2).”

There were significant main effects of group on the onsets of right and left IO-TrA
activity ($P < 0.05$, [ES = 0.40, 0.50]) (Fig. 4, 5). The onsets of right IO-TrA activity were
delayed in the non-specific CLBP group in comparison with the control group in WLFRE,
LFRE, and LF ($P < 0.05$, [ES = 0.89 - 1.94]) (Fig. 4). The onsets of left IO-TrA activity were
delayed in the non-specific CLBP group in comparison with the control group in RF, LERF,
and WLERF ($P < 0.05$, [ES = 1.18 - 1.48]) (Fig. 5). However, in LERE and LFRF, there are
not significant differences for the onsets of IO-TrA activity on both sides between groups ($P =
0.09 - 0.27$, [ES = 0.50 - 0.63]).

There were significant main effects of task on the onsets of right and left IO-TrA
activity ($P < 0.05$, [ES = 0.44, 0.54]). In the control group, the onsets of right IO-TrA activity
in WLFRE, LFRE and LF were significantly earlier than the other five tasks ($P < 0.05$, $[ES = 1.13 - 3.77]$) (Fig. 4). The onset of left IO-TrA activity in RF was significantly earlier than the onsets in WLFRE, LFRE, and LF ($P < 0.05$, $[ES = 1.39 - 2.25]$) (Fig. 5) and the onsets of left IO-TrA activity in LERF and WLFRE were significantly earlier than the onsets in WLFRE, LFRE, LF, LERE, and LFRF ($P < 0.05$, $[ES = 1.11 - 3.17]$) (Fig. 5). In the non-specific CLBP group, the onsets of right IO-TrA activity in WLFRE, LFRE, and LF tasks was significantly earlier than the onsets in LERF and WLERF ($P < 0.05$, $[ES = 1.15 - 1.70]$) (Fig. 4). The onsets of left IO-TrA activity in RF, LERF, and WLERF was significantly earlier than those WLFRE, LFRE, and LF ($P < 0.05$, $[ES = 1.40 - 2.34]$) (Fig. 5).

4. Discussion

In a present study, we found that the onsets of IO-TrA activity were significantly delayed during tasks with trunk rotational torque in the non-specific CLBP group but there were no significantly differences between groups during LERE and LFRF. In addition, the results of this study demonstrated that the onsets of IO-TrA activity were significantly related not to the magnitude of trunk rotational torque, but to direction of trunk rotational torque.

To our knowledge, this is the first study to show the onsets of the deep abdominal muscles activity are significantly delayed during tasks with trunk rotational torque but are not significantly delayed during tasks without trunk rotational torque in the non-specific CLBP
group. These results supported our first hypothesis. Previous studies reported that anticipatory
activity of the deep abdominal muscles was delayed in subjects with non-specific CLBP
during RF [6, 8]. However, the onsets of IO-TrA activity during bilateral shoulder flexion
were reported no significant difference between subjects with non-specific CLBP and healthy
individuals [11]. These different results may be explained by our study’s findings, as
neuromuscular dysfunction of the deep abdominal muscles during upper limb movement in
subjects with non-specific CLBP may be related to rotational torque of trunk. For example,
during RF, the upper limb movement generated right trunk rotational torque on the trunk in
acceleration phase of right shoulder flexion. Delayed onset of IO-TrA may lead to instability
of the lumbosacral segment for trunk rotational torque. These results suggest that feedforward
mechanism for trunk rotational torque is altered in the subjects with non-specific CLBP. The
trunk rotation has been identified as a significant risk factor for low back injury [18, 19] and
produces the onset of microdamage in the elements of the lumbar spine even if the amount of
trunk rotation is small [20, 21]. Therefore, although it is important to maintain lumbosacral
segmental stability for trunk rotation, the stability may be decreased in the non-specific CLBP
subjects due to the delayed onset of the deep abdominal muscles activity. Hodges et al. [22]
have investigated a cause-effect relationship between the delayed onset of the deep abdominal
muscles and pain. They have shown that pain produces delayed onset of transversus
abdominis activity and suggested that this delay appears to be a result rather than a cause of
pain [22]. However, these changes may increase the likelihood of chronicity of non-specific low back pain because dysfunction of trunk muscles would lead to spinal instability, repeated microtrauma and pain [4, 10]. The present findings suggest that clinicians need to pay attention to the deep abdominal muscles activity against trunk rotational torque.

Our results failed to find any significant differences in the onsets of IO-TrA activity between tasks with the same directional torque in the control group. This may indicate that the onset of IO-TrA activity is not influenced by the magnitude of trunk rotational torque. Hodges and Richardson [8] recorded the onsets of trunk muscle activity in upper limb movements at three different speeds (fast, intermediate, and slow), and found that the onset of TrA and IO activity at fast speeds was significantly different from the onset at slow speeds, but not significantly different from the onset at intermediate speeds. They explained these results by stating that the trunk muscles respond to upper limb movement above a threshold velocity or postural disturbances at fast or intermediate speeds, while TrA and IO are not needed to react to slow upper limb movements [8]. Therefore, because the three tasks with the same direction of rotational torque in our study (WLFRE- LFRE- LF, RF-LERF-WLERF) provided enough velocity or postural disturbance due to upper limb movement, it is possible that there are no significant differences between the onsets of IO-TrA activity in these tasks.

Also, in the present study, the onsets of IO-TrA activity were directionally specific not only in the control group but also in the non-specific CLBP group. Except for LERE and
LFRF, the contraction of the ipsilateral IO-TrA for the direction of trunk rotational torque lagged behind the contralateral side. The sequences of IO-TrA recruitment were consistent with previous studies indicating that left TrA and IO had earlier onset than right TrA and IO in right shoulder flexion [23]. An opposite and equal torque needs to be created to maintain the trunk position of horizontal plane during tasks with trunk rotational torque. Morris et al. [12] and Allison et al. [23] suggested that the TrA asymmetrically contracted to balance the rotational torque generated by rapid upper limb movement in healthy subjects. The present study suggests that the deep abdominal muscles activity in the subjects with non-specific CLBP is also specific to the direction.

In this study, there were significant differences of left shoulder angle in LFRF and WLERF. However, there were no significant differences of left shoulder angle in WFRE, LFRE and LF which had the significant delayed onset of right IO-TrA activity. The onsets of IO-TrA activity delayed in non-specific CLBP group were observed during tasks which had no significant differences of shoulder angle between groups. Therefore, these significant differences of left shoulder angle would have little effect on the results in this study.

There were limitations to this study. First, EMG recordings were made from the surface EMG system, and this system may pick up other muscular activity. However, a previous study showed that the use of surface EMG to measure IO-TrA activity could acquire data independent of other trunk muscle activity [13]. Second, we hypothesized from a
previous study that WLERF and WLFRE had increased trunk rotational torque compared with other tasks [12]. However, there were nonsignificant differences between peak rotational torque in WLFRE and LFRE as well as in LERF and WLERF. In this study, the acceleration of the center of mass in each segment was used for calculation of trunk rotational torque. In a later analysis, we clarified that accelerations of the forearm and hand in WLFRE and WLERF were significantly decreased compared with LFRE and LERF. The nonsignificant findings of peak rotational torque could be explained by the delayed accelerations of the forearm and hand in WLFRE and WLERF with a 1 kg weight. Third, the subjects with non-specific CLBP in this study had relatively mild symptoms. The average ODI score in Japanese people in their 20s who have low back pain who were unable to engage in routine work was reported to be 15.86 [24]. In this study, which recruited subjects in a similar generation, the average ODI score was 13.0 (SD 6.6). In other words, the subjects in our non-specific CLBP group would be regarded as having mild non-specific CLBP. Therefore, the results of this study may be generalizable to subjects with mild non-specific CLBP and may be related to aggravation of non-specific CLBP.

5. Conclusion

The onsets of IO-TrA activities in subjects with non-specific CLBP were delayed during tasks with rotational torque of the trunk in the opposite direction, suggesting a
possibility that delayed onset of the deep abdominal muscles during rotational torque of the
trunk might be one of the characteristics of CLBP.

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**Figure captions**

Fig. 1. The upper limb movements of 8 types were performed by all subjects.

Fig. 2. An example of onset of right deltoid (top) and left IO-TrA (bottom) during a trial of LERF in control group. The dotted line means onset of right deltoid. The thick vertical line means onset of left IO-TrA.

Fig. 3. The peak trunk rotational torque (mean and SD) for the 8 rapid upper limb movements in the control group and non-specific CLBP group. Positive value means right trunk rotational torque and negative value means left trunk rotational torque.

Fig. 4. Onset latency (±SD) of right IO-TrA in non-specific CLBP and control group. * indicates significant differences between groups.
† indicates significantly earlier than LERE, LFRF, RF, LERF, and WLERF.
‡ indicates significantly earlier than LERF and WLERF.

Fig. 5. Onset latency (±SD) of left IO-TrA in non-specific CLBP and control group. * indicates significant differences between groups.
† indicates significantly earlier than WLFRE, LFRE, LF, LERE and LFRF.
‡ indicates significantly earlier than WLFRE, LFRE, LF.
Acknowledgements

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors. We thank Peter Mittwede, MD, PhD, from Edanz Group (www.edanzediting.com/ac) for editing a draft of this manuscript.
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↑ indicates significantly earlier than WLFRE, LFRE, LF.
# Tables

## Table 1. Subject characteristics

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<td>13</td>
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<td>7:6</td>
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<td>VAS (%)</td>
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<td>ODI (%)</td>
<td>13.0 (6.6)</td>
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Abbreviations: CLBP, chronic low back pain. BMI, body mass index. VAS, visual analog scale. ODI, Oswestry disability index. Values expressed as mean (standard deviation).
Table 2. Peak shoulder angle and peak shoulder angular velocity during each task.

<table>
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<tr>
<th>Task</th>
<th>Side</th>
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<th>Peak shoulder angular velocity (deg/sec)</th>
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Abbreviations: CLBP, chronic low back pain. Rt, right. Lt, left.

Values expressed as mean (standard deviation).

* indicates significant difference in peak shoulder angle between groups.